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FACULTÉ DES SCIENCES APPLIQUÉES

Generic image classification : random and convolutional approaches

Master thesis

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Contents

C	onter	ats	1
1	Intr	oduction	2
2	Stat 2.1 2.2	Se of the art Supervised learning. Image classification.	
3	Obj	ectives	6
4	Met 4.1	4.1.3 Subwindows extraction 4.1.4 Feature descriptions 4.1.5 Classification schemes Dataset and environment Implementation 4.3.1 Software architecture 4.3.2 Technical issues	$egin{array}{c} 7 \\ 7 \\ 8 \\ 12 \\ 14 \\ 14 \\ 16 \\ 16 \\ 17 \\ 19 \\ 19 \\ 19 \\ 19 \\ 19 \\ 19 \\ 19$
5	Res 5.1	Direct classification scheme 5.1.1 Accuracy as a function of the learning set size 5.1.2 Variability 5.1.3 Accuracy as a function of the number of subwindows 5.1.4 Influence of the number of trees 5.1.5 Influence of the minimum number of samples to split 5.1.6 Influence of the number of filters 5.1.7 Influence of the filter generator 5.1.8 Influence of spatial poolings Feature learning scheme	$egin{array}{c} 21 \\ 21 \\ 22 \\ 24 \\ 25 \\ 25 \\ 25 \\ 25 \\ 25 \\ 25$
6	Con	clusion and perspective	28
Bi	bliog	raphy	29

	1			
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Chapter				
Oliapioi				

Introduction

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State of the art

The early days of computer vision have seen the development of myriads of domain specific methods, notably in the field of image classification. Image classification is the process of assigning the correct label from a predfinied set to an image. The main drawback of the domain specificity is that solutions are not necessarily transposable to other domains. The supervised machine learning framework sidestep this limitation by making no assumptions about the particular domain of application. Rather, it proposes a representation general enough for most problem to fit in, albeit with some preprocessing. Overcoming the domain specificity is achieved by letting the computer learn the discrimination scheme instead of supplying it.

This chapter is divided in two sections. In the first one, we review the supervised learning framework in general. In the second part, we focus on supervised learning image classification.

2.1 Supervised learning.

At the framework core are objects, also called individuals. The objects differ from each other by their features, also called variables. Among them is one which holds a special status: the output variable. The ultimate goal of a supervised learning algorithm is to produce a model mapping a previously unseen object's regular features to the output variable. Thus, the learning algorithm needs a space of candidate models. In order to provide the most adequate model out of the candidates, the algorithm needs two additional elements: a quality measure and an optimization strategy. It is sometimes more convenient to use the inverse of the quality measure: the loss function. We talk about *learning* because the algorithm optimizes the quality measure using a set of objects, called the learning set.

More formally, the learning set $LS = \{(\boldsymbol{x}_i, y_i) | i = 1, ..., N\}$ is composed of N objects. Each object i is described by a tuple of features: $\boldsymbol{x}_i \in \mathcal{X}$ are the regular features and $y_i \in \mathcal{Y}$ is the output variable. We represent the learning algorithm hypothesis space by \mathcal{H} and its quality measure by $\ell: \mathcal{Y} \times \mathcal{Y} \to \mathbb{R}$. The learning algorithm maps the learning set to a function $f \in \mathcal{H}: \mathcal{X} \to \mathcal{Y}$ so as to try to minimize some expectation over the loss function. Depending on the structure of \mathcal{Y} , the model either performs a regression or a classification.

The most usual quality measures are the square error $\ell(y,\hat{y}) = (y-\hat{y})^2$ for regression problems and the classification error

$$\mathbf{1}_{\neq}(y,\hat{y}) = \begin{cases} 1 \text{ if } y \neq \hat{y} \\ 0 \text{ otherwise} \end{cases}$$

for classification problems. Based on those loss functions, we can define the expected error by $E_{x,y}\{\ell(f(x),y)\}$. If we dispose of a testing set $TS = \{(x_i,y_i)|i=1,...,M\}$ with a sufficiently large M, we can assess the expected error by $\frac{1}{M}\sum_{x,y\in TS}\ell(f(x),y)$. If TS = LS, we call this

the resubstitution error. However, we are generally interested in the case where both sets are different while coming from the same distribution. We then talk about generalization error. This is the error we are trying to minimize.

The example of decision trees. Decision tree (Breiman et al. (1984)) is a good example of classification algorithm. More accurately, the decision tree is the model produced and the learning algorithm is the growing algorithm. The decision tree is a binary tree where each interior node represent a dichotomic choice regarding one regular feature and each leaf is labeled by a class. Classifying an object consist of moving from one node to another according to the node's test. Once the image reaches a leaf, its class is associated to the image.

Growing a good tree, in the sense of the quality measure, is done in a top-down fashion by a greedy heuristic. We first need to define an impurity measure which indicates how much diversity for the variable y there is in a sample. We start at the root and pick up the dichotomic choice, also called splitting criterion, which accomplishes the maximum expected reduction of impurity. Then the learning sample is split into a left and a right branch according to the test result. We can now develop both nodes recursively up till there is only objects of the same class in a node, making it a leaf. With this mechanism, the resubstition error is null, thus proving that the method minimizes the loss function. However, the generalization can be quite large. That is why other mechanism, such as other stopping criteria, are implemented.

A widely-used impurity measure is Shannon entropy. In that case, the construction algorithm has a nice interpretation. The reduction of impurity is the information gain. At each node, we choose the test which brings the most information to the classification variable.

The learning algorithm dependence on the learning set is clearer in the regression case, where we have the following bias-variance decomposition:

$$E_{LS}\{E_{x,y}\{(y - f(x)^{2})\}\} = E_{x,y}\{(y - E_{y|x}\{y\})^{2}\}$$

$$+ E_{x}\{(E_{y|x}\{y\} - E_{LS}\{f(x)\})^{2}\}$$

$$+ E_{x}\{(f(x) - E_{LS}\{f(x)\})^{2}\}$$

The first term, characterize the intrinsic difficulty of the regression task. It quantifies the average deviation of the best possible model $E_{y|x}$, the Bayes model, from the ground truth. The second term is the expected square bias. It quantifies the average error between the Bayes model and the average model, $E_{LS}\{f(x)\}$. The third term is the average variance of the model. It quantifies how much the model varies from one learning set to another. Although there is no such analytical decomposition for classification, the concepts of Bayes model, bias and model variance still abide as conceptual tools.

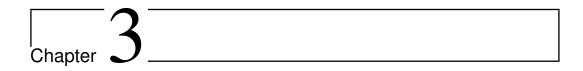
The space of candidate model \mathcal{H} is composed of elements of varying complexity. For instance, a decision tree complexity is assessed by its number of nodes. A more complex model will fit the learning set better, thus reducing the resubstitution error. However, fitting this set too well will cause overfitting: the algorithm incorporate set-specific details. This is reflected by the model variance term of the learning algorithm error: slight changes in the learning set will produce very different models. The bias decrease will, at first, surcompensate that increase, yielding a better generalization error. The compensation will work up to the point where the average is of the same order of complexity as the Bayes model. From that point on, the bias decrease will slow down while the model variability still increases. Therefore, the generalization error starts to increase, as well. The ability to control the complexity is an important characteristic of a learning algorithm. Another way around this problem is to resort to ensemble methods. Among such methods are the averaging methods which combine several models by averaging their predictions. The models are usually drawn from the same candidate space by either using different learning set

or introducing some randomization in the learning algorithm, sometimes both. Ensemble methods rely on the averaging to reduce variability and thus can enjoy more complex models. Ensemble of decision trees form classification forests.

A learning algorithm is usually dependent of some parameters, which influence the optimization strategy. They are called hyper-parameters so as not to confuse them with the other parameters on which the optimization is performed.

2.2 Image classification

Traditional learning algorithms are not able to works with "structured data" such as texts, images and graphs. Indeed, they expect the individual features to be scalar. Image classification is therefore much more about bridging the two realities than about developing brand new learning algorithm. We will now review the main such techniques.



Objectives

The hypothesis at the core of the present master thesis can be stated as follows:

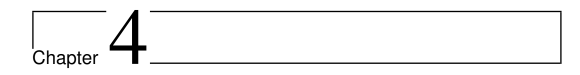
It is possible to combine the advantages of the classification forests, namely computational cost, feature importance evaluation and ease of use, with those of convolutional networks, primarily the accuracy.

The feature importance evaluation capability is one of the nicest features of the classification forests. The importance of a given feature is computed as the total reduction of impurity brought by that feature, normalize so that the feature importances sum to one. The most notable use of this measure is feature selection.

The ease of use of the forests is particularly obvious in comparison of the neural networks. With the former, the number of hyper-parameters is quite small and well understood. Therefore, tuning the method is easy and can, usually, be undertaken manually with good results. On the other hand, neural networks tuning is much more complex, as even the structure has to be adapted for each problem. Evidence of this complexity is the amount of work dedicated to this subject in the literature.

Lastly, let us mention an interesting characteristic of convolutional networks we did not pursue but which has a important impact on scalability: online learning. Indeed, classification forests require to have the total amount of data right away which will be a limitation of our method.

Validating this hypothesis constitutes our main objective. To achieve this, we developed a method based on classification forests which incorporates some convolutional networks mechanics. More specifically, random linear filters are applied to the image database, followed by one or several spatial poolings. Then, several random subwindows are extracted from each transformed image. Each subwindow is described by the row pixel values. The in-depth description of this "RandConv" method is the main subject of chapter 4. This method builds on previous works. The idea of applying predefined convolutional filters followed by several spatial poolings before extracting subwindows has already been done in . It constituted a generalization of their generic image classification scheme. The contribution of the current paper is two fold. Firstly, the RandConv framework proposes several extensions of that method, the most noticeable of which being the ability to generate the filters. This approach resembles much more the convolution networks', where the filters are actually learned. Secondly, whereas the aforementioned work was more like a proof of concept, the present study aim at analyzing more deeply this method. Indeed, proving the hypothesis is not our only goal. We also want to study closely the behavior of our classification method so as to understand its strength and limitations.



Methodology

This chapter is divided into three sections. The first one aims at fully describing our classification method. The second section details the experimental condition in which our method will be evaluated. Finally, the last one highlights implementation details and technical issues.

4.1 The RandConv framework

This section is dedicated to an in-depth description of our classification method: Rand-Conv. It stands for "Random and convolutional". The "random" part refer to both the filter generation and subwindow extraction. While the "convolutional" adjective refers to the application of the linear filters.

So as to bridge between classification forests and convolutional networks, we started from the former and added characteristics of the latter. Those characteristics are the convolutional filtering followed by spatial pooling. The RandConv method is divided into the following parts:

- 1. Generating the N linear filters
- 2. Applying the N filters to the M images of the databases
- 3. Applying the P spatial poolings to the $N \times M$ filtered images
- 4. Extracting S subwindows from each of the $N \times M \times P$ pooled and filtered images and resizing them
- 5. Describing each of the $N \times M \times P \times S$ pieces by a set of learning features

Thus, from a database of M images, we end up with a feature matrix with $M \times S$ objects described by a number of variables function of $N \times P$, among other things.

Although, the method has been designed with the use of classification forest in mind, the RandConv method, per se, is actually a feature extraction method. Its goal is to transform a set of images into a set of corresponding feature vectors. The actual classification could be carried out by any traditional learning algorithm. Nevertheless, regarding our primary objective and some other attractive properties of the trees, which will be developed in subsection 4.1.1, we will stick with classification forests in one way or another.

Algorithm 1 RandConv

```
1: procedure EXTRACT(RandConvInstance, images)
        rci \leftarrow RandConvInstance
2:
        N \leftarrow rci.nbFilters
 3:
        P \leftarrow rci.nbPoolings
 4:
        S \leftarrow rci.nbSubwindows
 5:
        F \leftarrow rci.nbFeaturesPerSubwindows
 6:
 7:
        M \leftarrow images.length
        learningMatrix[M \times S][F \times (N \times P)]
 8:
        row \leftarrow 0
9:
        colMin: colMax \leftarrow 0: F
10:
        for image \in images do
11:
            cropboxes \leftarrow rci.generateSubwindowLocations()
12:
            for filter \in rci.filters do
13:
                for pooling \in rci.poolings do
14:
                    filtered \leftarrow rci.applyFilter(filter, image)
15:
                    pooled \leftarrow rci.applyPooling(pooling, filtered)
16:
17:
                    for cropbox \in cropboxes do
                        subwindow \leftarrow rci.extractSubwindow(cropbox, pooled)
18:
                        learningMatrix[row][colMin:colMax] = rci.describe(subwindow)
19:
        \begin{array}{c} colMin: colMax \leftarrow colMax: colMax + F \\ \textbf{return} \ learningMatrix \end{array}
20:
```

4.1.1 Filter Generation and application

Mimicking the convolutional filtering is carried out by generating random linear, spatially invariant filters. More precisely, we generate the 2D finite impulse response matrices. First, the filter dimensions and then the filter coefficients are randomly drawn. This means that, contrary to the ConvNet, the coefficients are not directly learned. The coefficient learning is simulated by generating a vast number of filters and letting the learning algorithm choose the ones to emphasis.

This calls for an important remark: decision tree-based solutions are ideal classifier candidates. Firstly, their construction technique allow them to emphasis easily the interesting filters. Secondly, they deal well enough with numerous, possibly irrelevant, features. Indeed, the major impact is a reduction of the model effective complexity. The resulting accuracy drop is much less tremendous than with some other classifiers. Besides, this reduction of complexity can be balanced by the number of subwindows extracted from each image. Augmenting the dataset produces deeper trees; more complex model. Lastly, they scale well enough due to their relatively low computational cost, especially the extremely randomized tree variant.

4.1.1.1 Drawing mechanism

How to draw the filters is one of the RandConv framework cornerstone. The drawing mechanism should meet two prerequisites. Firstly, it should be able to produce unlimited, or at the least great, number of different filters. Secondly, the filters should be of some value by themselves but also together. Intuitively, a valuable filter should highlight "information" not directly accessible from the original image by the learning algorithm. We will call this characteristic the individual usefulness or simply usefulness. As for having value together, two different filters should not uncover the exact same "information". For instance, producing twice the same filter is useless. We will call this the group usefulness or co-usefulness.

Several drawing mechanisms have been developed with different characteristics in mind:

→ Custom filters

- → Discrete law generator
- → Zero-perturbation generator
- → Identity-perturbation generator
- → Maximum-distance-from-identity generator
- → Stratified perturbed generator

The first one is a special case. It consists of a set of 38 well known filter, among which the Sobel and Prewitt filters, several Laplacian filters of different sizes, the compass gradient filters, some low and high pass filters and other line detection filters. Being a small set, it violates the first prerequisite. However, this pseudo filter generator will be useful as a comparison basis: the filter are the same ones as in. Besides, these filters have practical application cases which random filters might not share. It is thus a reference point to see whether the generated filters highlight interesting "information".

The other mechanism draw randomly the filters. Before generating the coefficients of a filter, its dimensions must first be determined. The widths and heights of the impulse response matrices are drawn from an bounded set of odd, positive integers. Although we limited our tests to square matrices, this is not a strict requirement. We mainly worked with a uniform distribution of sizes, playing somewhat with the set bounds. Once again, this is not a limitation as other distribution can easily be used. For example, it is possible to create a distribution biasing towards small sizes. As for the bounds, a minimum seems to be 3. The maximum size should not be greater than twice the image size but needs probably not be greater than half this size. Indeed, greater filters might incorporate mostly non-local information. Conceptually, for a given maximum size, say $n = h \times w$, it is easy to build a bijection between the filter matrices space and \mathbb{R}^n . This representation will help us visualize the drawing mechanisms.

Discrete law generator. Once the size is fixed, every coefficient is drawn for a predefined discrete law. Even though the number of such filters is bounded for a given maximum size, this filter space is still vast enough so as to meet the first generator prerequisite. We tested the following law: -1 with a probability of .3, 0 with a probability of .4 and 1 with a probability of .3. This generator was motivated by the spatial interpretation of the convolution. It accounts for summing and substracting neighboring pixel together.

Zero-perturbation generator. Once the size is fixed, every coefficients are drawn from the same continuous probability law. Although there is no restriction on the probability law, we expect it to be symmetrical and zero-centered, hence the generator name. We used two such laws. The first one is the uniform law over reals bounded with -1 and 1. In this respect, the generator space is mappable to an hypercube centered on the origin. The second law was a Gaussian so that the probability of being outside the range [-1, 1] is equal to a given threshold. The isoprobabilities thus form hyperspheres. The points lying outside of the range can be forced to the boundary so that the generator space becomes the same hypercube as with the uniform law. Zero-centered generator were motivated by the examination of common filters which portray the same characteristic.

Identity-perturbation generator. Identity-perturbation generator work in the same fashion as its zero-perturbation counterpart. The only difference is that the hyper-structures are centered around the identity filter instead of the origin. The motivation behind this generator was to produce filtered images resembling the original while being different enough so as to be of value.

Maximum-distance-from-identity generator. This kind of generators fulfills the same purpose as the previous one. The generator space is also centered on the identity filter but its shape is different since we decided to work with the Manhattan distance. Concretely, the generator is parametrized by a maximum distance, independent of the filter sizes. The coefficients are processed in a random order. A random perturbation from the range [maximum distance, maximum distance], expectedly from a uniform distribution, is applied to the first coefficient. Before processing the next coefficient, the maximum distance is updated by substracting the absolute value of the perturbation.

Stratified perturbed generator This last class of generators are parametrized by a minimum value m, a maximum value M and a subdivision number n. For each coefficient, a value v from the set $\{m+\frac{k+1}{2}\times\frac{(M-m)}{n}|k\in\mathbb{Z},k< n\}$ is chosen randomly. This value is then randomly perturbed before being assigned to the coefficient. The perturbation is not mandatory and should stay in the range $[-\frac{(M-m)}{2n},\frac{(M-m)}{2n}]$. Expected perturbation law are Gaussian and uniform. This generator class was motivated by the idea to produce as dissimilar filters as possible so as to meet our second requirement about co-usefulness. Disregarding the perturbation, the filter space is finite but still huge. For instance, the space for a subdivision number of 10 with only the smallest filters (3x3) would still mean 10^9 filters. Whereas, 2^9 filters, i.e. a subdivision number of 2, is manageable, the other generators are able to produce filters as dissimilar. Furthermore, the following non-monotonicity property suggests that a dissimilar approach in the filter space might not be the best way to produce sets of co-useful filters. Indeed, we can use the distance to measure co-usefulness: if two filtered image are close, they probably highlight the same "information".

Non-monotonicity property. We will show that closeness in the filter space does not necessarily imply closeness of the filtering results. Closeness is to be understood as distance from a reference. Let I be an image and F, F_1 , F_2 be three linear, spatially invariant filters of possibly different sizes. Let also

$$J = I * F$$
$$J_1 = I * F_1$$
$$J_2 = I * F_2$$

We will show by counterexample that $||F - F_1|| \ge ||F - F_2|| \implies ||J - J_1|| \ge ||J - J_2||$. First let us name $e_1 = F - F_1$ and $e_2 = F - F_2$. By linearity of the convolution, we have:

$$J_1 = I * F_1 = I * (F - e_1) = (I * F) - (I * e_1) = J - (I * e_1) \iff J - J_1 = I * e_1$$

In these terms, we have to show that $||e_1|| \ge ||e_2|| \implies ||I * e_1|| \ge ||I * e_2||$. Let us take e_1 such that the coefficients sum up to zero but with a great dispersion (a Sobel filter, for example) and e_2 such that the sum of the coefficients is strictly greater than zero but with a smaller dispersion than e_1 (the 3x3 average filter, for instance). Thus, we have $||e_1|| \ge ||e_2||$. Moreover, let us consider the case of a image I with constant value c > 0. In this setting, $||I * e_1|| = 0$ while $||I * e_2|| = c \times k > ||I * e_1||$.

Therefore, playing with closeness or dissimilarities in the filter space yield no warranty about the same metrics with the filtered images. However, using the distance as measure of co-usefulness is arguably a poor choice, since close images might still highlight different aspects of the images. Considering this remark, the main shortcoming of the stratified generator is probably that, with respect to the number of generated filters we will use, it does not produce any significant advantage over other generators.

4.1.1.2 Normalization

All the generators we discussed in the previous section are able to perform a post-processing normalization of the filter. There are four normalizations:

- No normalization: the post-processing normalization is skipped.
- → Zero mean: the mean value of the filter coefficients is null.
- Unit variance: the coefficients have a unit variance.
- Yero mean and unit variance: both the previous. First the zero mean then the unit variance.
- → Unit sum: the coefficients sum to one.

The introduction of the zero mean and unit variance normalizations was primarily motivated by supplying support for learning algorithms other than classification forests. Indeed, their effect is to impose a common dynamics to all the filters. While trees can cope easily with variables of different dynamics, some classification schemes are not applicable is that setting or suffer greatly from it. As for the unit sum normalization, applied in conjunction with a generator producing positive coefficients, it produces "convex combination filters" in the following sense: for each step of the convolution, the output pixel value is a convex combination of the minimum and maximum of the neighboring original pixels (where the neighborhood is defined by the filter size). We will now look at the implication of the normalizations on the generator space and the filtering in both the spacial and frequency spaces. We will reuse the filter representation in \mathbb{R}^n and will denote by 1 the vector whose coefficients are all 1.

Zero mean normalization. In \mathbb{R}^n , the zero mean filters form the hyperplane $\{x \in \mathbb{R}^n | \mathbf{1}^T x = 0\}$. The normalization is a projection onto that hyperplane. The resulting filter y is computed as $y = x - frac1n\mathbf{1}^T x\mathbf{1}$. This operation can produce a filter which is outside of the original filter space. Since this operation is linear, the impact of the filtering are straightforwardly identifiable. Denoting I a given image, x_f a given filter, whose mean m form the constant filter m_f , y_f the normalization $y_f = x_f - m_f$ and $\mathbf{1}_f$ the constant filter with only ones as coefficients, we have:

$$I * y_f = (I * x_f) - (I * m_f) = (I * x_f) - m \times (I * \mathbf{1}_f)$$

The $(I * \mathbf{1}_f)$ correction part is independent of the filter and proportional to the mean coefficient value. The practical impact is clearer in the frequency space. Let us denote by $\rightleftharpoons_{\mathcal{F}}$ the Fourier transform:

$$I \rightleftharpoons_{\mathcal{F}} Ux_f \qquad \rightleftharpoons_{\mathcal{F}} H_x m_f \rightleftharpoons_{\mathcal{F}} H_m y_f \qquad \rightleftharpoons_{\mathcal{F}} H_y = H_x - H_m$$
$$I * y_f \rightleftharpoons W = U \times H_y = U \times (H_x - H_m) = (U \times H_x) - (U \times H_m) = Y - (U \times H_m)$$

Since m_f is a constant signal, the transfer function H_m is null everywhere except at the origin. Thus, the overall frequency response is only marginally modified and both filter achieve the same results. Therefore, the normalization does not restrict the class of filters.

Unit variance normalization. In \mathbb{R}^n , the unit variance filters form the hypersphere $\{x \in \mathbb{R}^n | x^T x = 1\}$. However, in practice, the normalization works with the current filter size and not the maximum filter size. Thus, there are several hyperspheres to consider, one per possible size. In the filter space, the normalization equals to scaling the filter so as to meet the appropriate hypersphere. The impact on filtering is immediate:

$$I * y_f = I * (\frac{1}{\sigma_x} x_f) = \frac{1}{\sigma_x} (I * x_f)$$

The whole result is scaled by the same factor. Therefor, the normalization does not restrict the class of filters.

Unit sum normalization The reasoning is identical to the zero mean normalization except for the hyperplane : $\{x \in \mathbb{R}^n | \mathbf{1}^T x = 1\}$. This normalization does not restrict the class of filters either.

4.1.1.3 Filter application

In this subsection, we cover two topics about the filter application. The first one concerns working with colors. The second one is about how the filter are actually applied.

Handling colors can be done in three ways. The first one, is to realize a 3D convolution. This results in a single output value per original pixel. However, there are two drawbacks to this approach. Firstly, in the spatial space, it means combining values of different colors together. This would work but lacks of physical interpretation. Indeed, the RGB space is only a convention. The second drawbacks has to do with the frequency space. It feels awkward to put on a same level spatial frequencies and color frequency, whatever it might mean.

The second method to handle colors is to use separate 2D filters on each channel. This produces three values per original pixels. As for the last and simplest method, is to use the same 2D filters on each color. This also produces three values per original pixels. Because the last approach seems more natural than the first one and is simpler to interpret, it is the one we adopted.

Now that we know how to handle colors, let us investigate the filter application. The convolution is carried out in the frequency space by multiplying the Fourier transform of the original image by the transfer function of the filter. The output is of the same size as the original image. The borders are handled by padding the original image with zeros.

4.1.2 Pooling goals and strategies

Now that we have fully covered the filter generation and application mechanisms, we can move on to the next part concerning the spatial pooling.

Moving windows. In the case of spatial pooling by moving windows, two elements are needed: the moving window size and the pooling function. Windows are supposed to have odd width and height. The center of the window moves to match every pixel of image. The pooling function is computed on the overlapping part of the window and the image. The resulting image as the same size as the original image.

Aggregations. In the case of spatial pooling by aggregation, the neighborhood windows do not overlap. The image is divided into several non overlapping neighborhood such that each neighborhood has the appropriate size. The pooling function is then applied on each cell of this neighborhood grid. Thus, contrary to moving windows, the resulting image is smaller and correspond to the neighborhood grid layout.

Pooling functions. The pooling function box is comprise of the minimum, maximum and average functions. In the case of the average function with moving window, we are close to defining a composition of two linear filters. The difference comes from the way the border are handled. In the case of the application of the linear filter, the outside element are replaced by zero while they are ignored in the pooling case.

4.1.3 Subwindows extraction

Once the generated filters and the spatial poolings have been applied, it is time to extract subwindows from the images. Extracting subwindows has several advantages. Firstly, it expands the number of learning objects. Depending on the feature descriptor extraction mechanism, there may be numerous features describing a given image. Using less features but more objects performs usually better. As we mentioned earlier, this is especially true with trees, at least for the "more objects" part, where a greater database means a greater tree and therefore a more complex model. Secondly, this approach is more robust towards scaling and occlusions (Marée et al. (2013)). Lastly, it can be used as a zone of interest detection

system, see for example Marée et al. (2006). This is an computationally interesting and domainfree alternative to other techniques. Although this is not the focus of the present work, the RandConv method retains this capability as well. While expanding the number of learning objects, we also need to to expand the class label accordingly. Consequently, each subwindows will be described by the label of its original image.

The number of possible subwindows is quite vast. First, let us notice that the number of subwindows of size $a \times b$ (N(a,b)) factorizes into the product of the number of subwindows along each axis: $N_v(a) \times N_h(b)$. These can be computed easily as $N_v(a) = H - a + 1$, where H represents the height of the image and similarly for $N_h(b)$, which depends on the width W. Indeed, for a column of size H, there is H origins of 1 pixel subwindows. If we take subwindows of size 2, we can take all the same origins as previously except the last one. Subwindows of size 3 cannot take the last two compare to 1 pixel subwindows, and so on. Therefore, the total number of subwindows N is:

$$N = \sum_{a=1}^{H} \sum_{b=1}^{W} N(a,b) = \sum_{a=1}^{H} \sum_{b=1}^{W} (H - a + 1)(W - b + 1) = \frac{1}{4} (H^{2}W^{2} + H^{2}W + HW^{2} + HW)$$

For 32x32 images, this yields 278,784 subwindows!

As we can see, there are numerous subwindows. Nevertheless, not all are of interests. The small size subwindows do not bring much information. For instance, taking only one pixel is not interesting. Delimiting a good threshold on the size is problem dependent, however. Despite focusing on big enough subwindows, there may still be too many of them. For instance, on 32x32 images, there are still 2025 subwindows of sizes ranging from 24x24 to 32x32. Nevertheless, this entails a redundancy level which is not needed. Since it would be difficult to establish a general heuristic to choose good candidates, we resort to drawing them randomly. At this point we have to be careful. Since we want to describe together all, i.e. within the same feature vector, the filtered images in a coherent fashion, we need to extract the same subwindows on all the filtered image belonging to the same original one. The filtering and pooling aim is to better describe a subwindow. However, for two different original images, we may choose two different sets of subwindows. By hypothesis, the chance of drawing twice the same subwindow for a given original image is small. We start by drawing the size uniformly in the affordable range. Then the upper left position of the subwindow is drawn from the possible position considering the subwindow size.

Since the subwindows are chosen uniformly, we can compute the probability of a given pixel belonging to a subwindow by counting the number of subwindows containing that pixel and dividing it by the total number of subwindows. Once again, we will use the fact that we can factorize the numbering for each axis. We will proceed by recurrence. Let us take a column of height H and index the element starting at 1 for the top element and ending at H for the bottom one. We will denote by T(i) the number of subwindows encompassing the ith element. It is immediate that T(1) = H: only one subwindow of each length can contain the first element. By symmetry this is also the case for the last element: T(H) = H. The second element is encompassed by all the subwindows of the first one but for the one-pixel subwindow. Besides this, we have to add the H-1 subwindows starting at this element. Therefore, T(2) = T(1) - 1 + (H - 1). The reasoning is similar for the next one, the only difference being that now we have to substract two previous windows from T(2): the monopixel one starting at element 2 and the bipixel one starting at element 1 (its monopixel has already been removed). Thus, T(3) = T(2) - 2 + (H - 2). Expanding the reasoning we get the general formula T(n) = T(n-1) - (n-1) + H - (n-1) = T(n-1) + H - 2(n-1). Resolving the recurrence yields $T(n) = nH - n(n+1) + 2n = nH - n^2 + n$, which verifies T(1) = T(H) = H. We just need to pay attention to the fact that we have started numbering at 1, which is not the convention. Coming back to the 2D case, we have that the number of subwindows encompassing a pixel (r+1,c+1) is $T(r,c)=(rH-r^2+r)\times(cW-c^2+c)$.

Although the subwindows can have different sizes, the feature vector describing a particular subwindows cannot. More specifically, a column of the learning matrix must correspond to a well identified variable. Thus, we need to rescale all the subwindows to a common size. Ideally, the size should be chosen so as to minimize the reinterpolation error. The interpolation algorithms are nearest neighbor, bilinear and bicubic. We will focus on the nearest neighbor because it is faster and it was found to be comparable in term of classification accuracy to the others in most cases ().

4.1.4 Feature descriptions

Starting from the original database of M images, a set of N filters and P spatial poolings, we produced $N \times P$ images for each original ones. From each of those M sets, we extracted S subwindows on each of the $N \times P$ filter images. We now have $M \times S$ learning objects described by $N \times P$ complex structures. We will describe each structure by a feature vector and concatenate all the feature vectors corresponding to the same subwindow.

Each filtered and spatial pooled image will be described by its raw pixels in a last-dimension-first fashion. Therefore, we start by the color dimension and group the three color values of the top left pixel together. Then we append the second pixel (top row, second pixel from the left) and so on for all pixels of the first row. After that, we append the second line in the same fashion and so on for all the image. Thus, there are $3(h \times w) \times (N \times P)$ features per subwindows. For instance, 100 filters with 1 pooling on 16x16 subwindows yields feature vector of $3(16 \times 16) \times 100 = 76,800$ variables.

4.1.5 Classification schemes

The previous subsections cover the preprocessing steps to go from a database of images to an actual learning matrix. Now is time to delve into the actual classification mechanism. We will use two approaches described in the following.

4.1.5.1 Direct classification

In this variant, classification will be undertaken by a special kind of classification forest: ensemble of extremely randomized trees, also known as ExtraTrees. They were introduce in Geurts et al. (2006) and resemble random forest (Breiman (2001)). In both kind of forests, only a subset of the features are examined at each node to determine the splitting criterion. This approach is called local random subspace and was introduced in Ho (1998). They differ in the following way: while random forests use bagging as an extra mechanism to introduce randomness, the ExtraTrees determine the splitting thresholds randomly. Bagging, for bootstrap aggregating, is the fact of drawing with replacement several learning samples from the original one (boostrap) and combining their predictions (aggregating). The advantage of ExtraTrees over random forests is threefold. Firstly, the bagging introduces an effective reduction of the learning size of 36% for each tree. This is not the case of the ExtraTrees, where all trees can learn from the whole learning sample. Secondly, they are much faster to build since we do not need to pick the optimal splitting thresholds of all the variables examined in a node. Lastly, they tend to perform better than their counterpart (Geurts et al. (2006)). In this variant, also called ET-DIC (ExtraTrees for direct image classification), the classification is undertaken by the ExtraTrees directly. Let us note that this scheme allows not only for individual feature importance evaluation but also for filter relevance evaluation by aggregating the importance of all the features corresponding to a given filter. This metric is surely a good candidate for filter co-usefulness evaluation.

4.1.5.2 Feature learning scheme

Also called ET-FL (ExtraTrees for feature learning) or ERC-forests (Extremely randomized clustering forests), this variant was introduce in Moosmann et al. (2008). The ExtraTrees are not used as classifier any longer. They form a preprocessing step whose output will

constitute the learning matrix of the actual classifier. The idea is to use the trees to form visual dictionary. Specifically, ExtraTrees are used in an unsupervised way by selecting randomly the splitting criterion of each node, a variant called totally randomized trees. The dictionary words are composed of T parts of different sizes, where T is the number of trees. Each part encodes the index of the leave where a given object ends up in a one-hot fashion. Thus, the part corresponding to a tree with L leaves will be a binary word with L-1 zeros and 1 one. In the learning phase, the totally randomized trees are grown from the learning sample, then the same learning sample is propagated down the tree to form the new learning matrix, which is quite sparse. Since we are using several subwindows per original image, we can aggregate by summing bitwisely the words corresponding to the original image to form a new word: an histogram word. The actual classification is then undertaken by a Support Machine Vector (SVM).

Two remarks are in order. Firstly, we have lost the ability the evaluate the feature importances. Secondly, the learning size, and consequently the number of subwindows, is now crucially important. Indeed, of all the parameters, it is one of the most influent parameters on the trees depth, and consequently the number of leaves. The second most important parameter is the pruning parameter. Pruning is the mechanism of stopping the tree creation before its completion. It can be applied either while developping the tree (pre-pruning) or artifically after the tree creation by cutting of some branches. It obviously also impact dearly the number of leaves. The main goal of pruning is to reduce the model complexity and consequently the overfitting. In the ET-DIC variant, pruning is not necessary because the voting takes care of reducing the overfitting. However, on the data fed to the SVM, there is no automatic overfitting control mechanism and the SVM will suffer from it. Therefore, pruning is also very important for this variant. Last but obviously not least, the number of trees multiply the number of leaves. Therefore, this hyper-parameter is even more sensitive than in the ET-DIC approach.

An extensive comparison of both methods was carried out in . They show that, in most cases, the ET-FL approach performs slightly better (3-4% in average). Nevertheless, ET-FL is more difficult to tune.

4.2 Dataset and environment

We worked with the CIFAR-10 database (Krizhevsky and Hinton (2009)). This dataset is composed of a learning set of 50,000 images and a testing set of 10,000 images. They are grouped into 10 mutually exclusive classes: airplane, automobile, bird, cat, deer, dog, frog, horse, ship and trucks. Both subsets contain an equal number of images from each class. The images have all the same size, 32x32, and are RGB.

This database has been chosen because it was the same one on which the precursory method from was tested. In turn, they chose this dataset because their traditional solution had difficulties with it. Their best solution with the ET-DIC method was an accuracy rate of 53.67%. The corresponding hyper-parameters were the following: 10 fully grown trees, 20 subwindows of 75-100% of the original size reshaped by nearest neighbor interpolation to 16x16 image described by raw pixel values and inspecting all the features on each node. As for the ET-FL variant, the best result was 50.07% of accuracy with 10 almost fully grown trees, 20 subwindows of 75-100% of the original size reshaped in the same manner as before and using $k = \sqrt{M}$ of inspecting features at each node, where M is the total number of features: 16x16x3. The trees are not totally grown: the minimum number of samples to split a node is fix to 10. Contrary to the majority of cases, the best ET-DIC is better than the best ET-FL. Using the aforementioned precursory method, they were able to obtain a significant raise of accuracy, attaining a new record of 74.31% with 750 trees totally randomized trees and 20 subwindows.

The best result on this database is an accuracy of 91.2% and is hold by a convolutional network (Lin et al. (2013)). The top ten best results are above 80%. Most are neural network

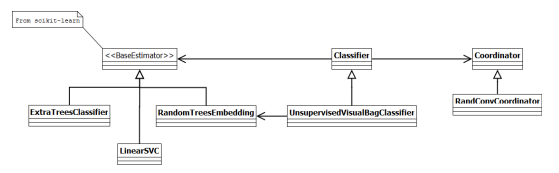


Figure 4.1: UML representation of the Classifier class and its major components

solutions and none are based on classification forests.

The learning and testing were carried out on a 64 bits 30-core 2.1 GHz computer with 288 GB of RAM.

4.3 Implementation

In this section, we dive into implementation. We will not go over all the elements in detail again but rather see how the discussions of the previous sections translate into code. In a second subsection we will also develop technical limitations.

4.3.1 Software architecture

A major concern of the design was to allow as much room as possible for flexibility, so as to be able to develop extensions and variants of the method rapidly. Consequently, the code was split into many classes. A drawback of this is that assembling all the pieces together might be difficult. Factory methods are provided to help with this issue but care must be taken to build up a coherent classifier. We will go back on this in a short while. The code is written in Python 2.7 and relies and the 0.13.3 version of Scipy (Jones et al. (01)), including Numpy 1.8.0. Not all the code is brand new. The ExtraTrees implementation comes from the scikit-learn library (Pedregosa et al. (2011)), version 0.15. We also use scikit-learn for SVM classification, although it actually consists of a wrapper to the liblinear library (). As for the subwindow extraction, it is a reorganization of the Pixit implementation (). We will examine the code in a top-down manner and focus on the important classes.

At the top, we find the Classifier class. Its aim is to supervise all the parts. The base class correspond to the ET-DIC variant. The RandConvCoordinator is responsible for all the preprocessing: filtering, pooling, subwindow extraction, feature description. After this step, the Classifier instance delegates the actual classification to a BaseEstimator instance from scikit-learn. In this case, the actual classifier is supposed to be an instance of ExtraTreesClassifier. The UnsupervisedVisualBagClassifier corresponds to ET-FL method. Between the preprocessing and the classification, we use the RandomTreesEmbedding, a totally randomized trees implementation, to build the histogram we will fed to the BaseEstimator, supposed to be a LinearSVC instance. This is summarized by figure 4.1.

We now go back to the RandConvCoordinator. Its responsibility is to transform the subwindows into feature vectors. It proceeds by subdividing the dataset to parallelize the transformation. Then, the ConvolutionalExtractor process each image. This entails filtering the image by each element of the FiniteFilter thanks to the Convolver, the applying all the spatial poolings contained in the MultiPooler and finally extracting several subwindows via the MultiSWExtractor. Once all this is done, each filtered and pooled

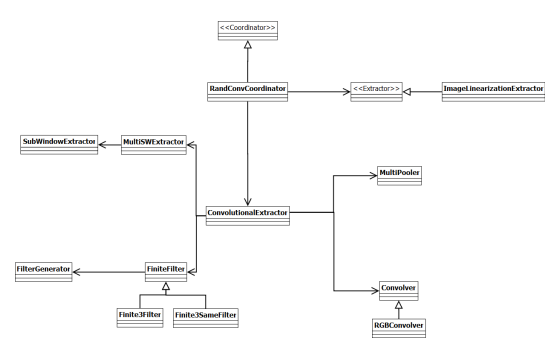


Figure 4.2: UML representation of the RandConvCoordinator and its major components

subwindow is passed through the Extractor and reassembled to form a coherent learning submatrix.

The FiniteFilter objects are containers for filters. They pre-generate a finite number of filters thanks to the FilterGenerator. We will come back to those in the next paragraph. If we are working with RGB images, we need to use either a Finite3Filter or a Finite3SameFilter. The former produces a different filter per color component while the latter uses the same filter on each color. Also, we need to use an appropriate Convolver, namely the RGBConvolver.

The subwindow extraction is carried out by the MultiSWExtractor whose sole purpose is to keep track of the subwindows to extract to the set of filtered and pooled images belonging to the same original image. The actual subwindow generation and extraction are delegated to the SubWindowExtractor.

The transformation from subwindows to feature vector is the responsibility of the Extractor instance. In this case, a ImageLinearizationExtractor object. However, other mechanism could be implemented. All this is summarized by the figure 4.2.

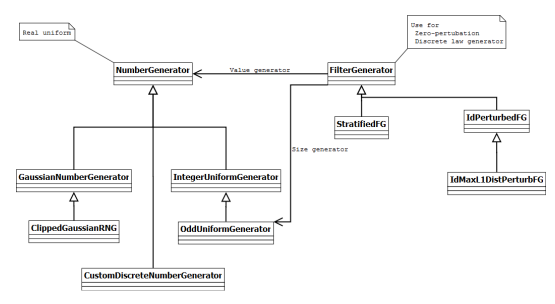
We now explore the FilterGenerator. They need two random number generators. One for drawing the values, either directly or not, and one for drawing the size. The base class is used for two of the generation methods: the discrete law generator and the zero-perturbation generator. The former is made by using a CustomDiscreteNumberGenerator while the later uses the base class of NumberGenerator. As figure 4.3 displays, there is a class dedicated to each of the other generators.

The GaussianNumberGenerator works by specifying a lower bound, an upper bound and the probability of being outside of that range. The ClippedGaussianRNG works similarly but in addition forces the values outside of the range to the appropriate bound.

The MultiPooler class involved in the RandConvCoordination is a container of spatial poolings. As the figure 4.4 depicts, our two groups of spatial poolings are presents.

4.3.2 Technical issues

The main limitation we will face is memory. The ExtraTrees implementation require 32-bits floats and the SVM, 64-bits floats. However, in the case of the ET-FL, the matrix is



 $Figure \ 4.3: \ UML \ representation \ of \ the \ {\tt FilterGenerators} \ and \ {\tt NumberGenerators}$

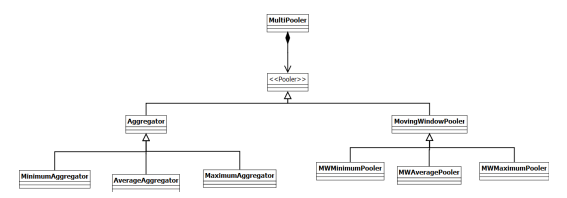


Figure 4.4: UML representaiton of the spatial poolings $\,$

mostly sparse on therefore the 64-bits floats requirement of the SVM will not be troublesome. Thus, the space cost bottleneck is the input of the ExtraTrees, which require to hold all the data in memory. Considering 100 filters, 1 spatial pooling, subwindows resized in 16x16, 3 colors, 10 subwindows and the whole learning set (50,000 images) the RandConv method will produce 153.6 GB of data. For 39 filters (the custom filters plus the original image) and 20 subwindows with 9 spatial poolings (the configuration of the best results of), 1,1 Tb would be required. Since we are limited to 288 GB, we will not be able to reproduce such a configuration.

One way of sidestepping this limitation is to build several forests with a different subset of the features, a variant which might be called "global random subset" (GRS) of features. In the case of the RandConv this can easily be done by choosing a subset of filters for each forest.

To a lesser extend, the time complexity will be an hindrance. It will not actually prevent any computation but we will have to plan carefully the experiment to carry out. For instance, our first example, which produces 153.6 GB of data, takes between 5h and 12h depending on the machine load.

4.4 Hyper-parameters summary

Before elaborating on the results, we will rapidly summarize all the hyper-parameters involved with the RandConv framework.

- → Filter generation
 - \propto Size range
 - \propto Whether or not to produce squared filters
 - \propto Value range
 - ∝ Filter generator
 - ∝ Random law
 - \propto Other filter generator specific parameters (maximum distance, number of subdivisions,...)
 - \propto Filter normalization
 - \propto whether or not to include the original image
- \hookrightarrow Spatial poolings
 - \propto Number and type of poolings
 - * Aggregation or moving window
 - * Pooling function: identity, minimum, average or maximum
 - \propto Size of the neighborhood
- → Subwindow extraction
 - \propto Number of subwindows
 - ∝ Subwindows cropping size
 - ∝ Subwindows rescaling size
 - \propto Subwindows rescaling interpolation
- → ExtraTrees
 - \propto Number of trees (default : 10)
 - \propto Number of features of the local random subspace : k (default : square root of the total number of features)

Hyper-parameter	default value				
Filter sizes	3 to 9, both included				
Square sizes	True				
Filter value range	[-1, 1]				
Filter generator	FilterGenerator (Zero-perturbed generator)				
Random law	Real uniform				
Filter normalization	None				
Include original image	True				
Number of spatial poolings	1				
Type of spatial poolings	Average moving window				
Size of the neigborhood	3x3 (moving window), 2x2 (aggregation)				
Number of subwindows	10				
Cropping size	24x24 to 32x32				
Rescaling size	16x16				
${\bf Interpolation}$	Nearest neighbor				
Number of trees	30 (ET-DIC), 750 (ET-FL)				
k	Square root of the total number of features				
Maximum depth	None				
Minimum sample to split	2 (ET-DIC), 500 (ET-FL)				
Minimum sample per leaf	1				
Bootstrap	False				

Table 4.1: Default values for hyper-parameters

 \propto Minimum sample to split : n_{min} (default : 2)

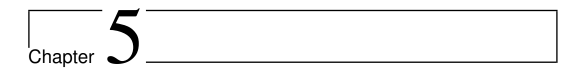
∝ Minimum sample per leaf (default: 1)

∝ Whether or not to use bootstrap (default : no bootstrap)

As we can see, the number of hyper-parameters is already large. We can divide them in two categories. On the one hand, we have structural parameters and on the other, traditional hyper-parameters. Structural parameters have a more profound impact than traditional ones. The filter generator and its random law and the number and type of spatial poolings form the structural parameters. Conceptually at least, changing one of them is closer to changing the classification method than only one of its hyper-parameters.

Three of the ExtraTrees hyper-parameters are used to control the pruning: the maximum depth, the minimum sample split and the minimum sample per leaf. The difference between the last two is that the first one does not attempt to split a node whose number of samples are under the given threshold, while the second does the split but rollback if any of the children are under the threshold.

Several of the parameters can be fixed. The value range of the filters will be set as in the filter generator descriptions. Considering the size of the images, we will mostly focus on 3x3 to 9x9 filters. Since we are working with trees, we will use no normalization of the filters. Concerning the spatial poolings, we will also restrict ourselves to small neighborhoods. For comparative purposes, we will resize subwindows to have sizes of 16x16 with nearest neighbor interpolation, as in . Moreover, we will extract subwindows of 24x24 to 32x32 pixels. For the ET-DIC variant, we will use the default trees parameters, which means no pruning. We will also use 30 trees (one per core). However, in the ET-FL approach, we will have to prune the trees and use more of them. The n_{min} parameter will be set to 500 with 750 trees. Table 4.1 displays the fix or default values of the hyper-parameters. Unless stated otherwise, these values are to be assumed in the next chapter.



Result analysis

In this chapter, we describe the experiments conducted with our new classification method and analyze their results. The chapter is divided into two sections. The first one tackles the direct classification scheme (ET-DIC), where the forest of extremely randomized trees serves as classificator. The second section describes the other variant where the extremely randomized trees are used to create a visual dictionary (ET-FL), while the actual classification is undergone by a support vector machine.

We will use the following abbreviations:

- \hookrightarrow rc: The RandConv methods without the use of the original image.
- → rci : The RandConv methods with the use of the original image. As if the first filter were the identity filter.
- → pixit: The method of , where there is no filtering or pooling. Only the subwindows extraction

When the hyper-parameters value are not explicit, the default values are to be assumed (table 4.1 of page 20).

5.1 Direct classification scheme

5.1.1 Accuracy as a function of the learning set size

Our first experiment will be to measure how the accuracies of the different methods evolve as a function of the learning set size. We will test the rci method with both types of pooling mechanisms (aggregation and moving window). The aggregation pooling uses a neighborhood of 2x2 while the moving windows are of size 3x3. Both poolings uses an averaging function. We will limit ourselves to the zero-perturbed filter generator for now. In both cases, the same 100 filter are drawn. The 10 extracted subwindows are resized to 16x16. Consequently there are $(16\times16\times100 \text{ filters}\times1 \text{ pooling})\times3 \text{ colors}=76,800 \text{ features}$. The accuracy is measured on the whole testing set composed of 10,000 images. Thepixit method will serve as reference.

We measure the accuracy for the learning set sizes of 500, 5,000, 10,000, 20,000, 30,000, 40,000 and 50,000. The result of this experiment is depicted by figure 5.1. The conclusion is clear: the aggregation pooling mechanism is not working (0.38 of accuracy). Although being of the smallest interesting size, the non-overlapping neighborhood windows already lose too much of the information. One might argue that using 3x3 moving subwindows allow for that variant to capture more of the spatial structure than 2x2 neighborhood. However, this does not seem to be the case as using 3x3 neighborhoods yields a lower accuracy of 0.35 for the whole learning set. Therefore, the bad performances of the aggregation mechanism

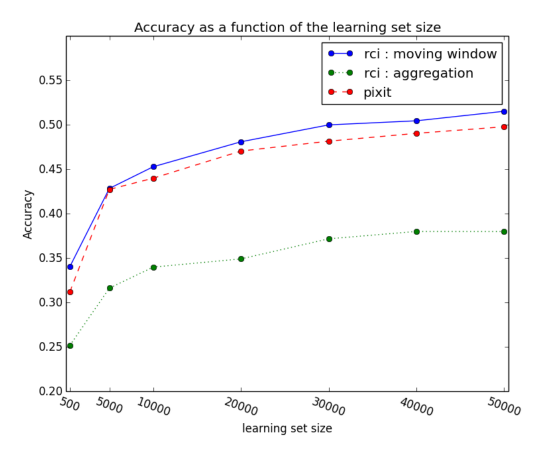


Figure 5.1: Accuracy as a function of the learning set size (30 trees, k = 277)

must indeed comes from its abusive spatial compression. This is unfortunate because those poolings suffered less from the rescaling of the extracted subwindows. Considering the gap of accuracy between both methods, the following experiments will focus on the moving window poolings.

Other observations are worth mentioning. Firstly, the "good" RandConv method performs well. It is consistently better than the pixit, although the difference might not be worth the trouble. The RandConv methods takes several hours whereas the pixit requires a couple of minutes at most. Secondly, the graph shows the impact of using only a subset of the data. This information is important in case of hyper-parameter optimization, for instance, where some of the data must be set aside as validation set.

Our best result so far is an accuracy of 0.515 due to the RandConv method with moving windows for spatial pooling. The pixit methods with the same parameters yields an accuracy of 0.50, while its best result is 0.53 for 20 subwindows, 10 trees and a minimum sample to split of 10. We will now analyze the variability of the method and how the method varies with some its hyper-parameters.

5.1.2 Variability

We now inspect the variability of the accuracy and of the filter importances. We will first look at the situation where the learning matrix is fixed and the variability can only come from the ExtraTrees. It is important to differentiate both types of variability. The one from the ExtraTrees would mainly be due to the high dimensionality of the problem. Indeed, the size of the local random subspace ($k = \sqrt{76,800} \simeq 277$) is relatively small compare to the total number of features: $\frac{277}{76,800} \simeq 3.6 \times 10^{-3}$. Therefore, different trees will end up making

Test number	Accuracy
1 Cot Hullioti	
0	0.5151
1	0.5119
2	0.5118
3	0.5136
4	0.5135
5	0.5141
6	0.5154
7	0.5114
8	0.5145
9	0.5156
10	0.5148
11	0.5118

Test number	Correlation with test 0
0	1
1	0.998
2	0.997
3	0.999
4	0.997
5	0.998
6	0.998
7	0.998
8	0.997
9	0.998
10	0.998
11	0.997

Table 5.1: Variability induced by the tree growing algorithm (same learning matrix)

different choices. This is, of course, exactly the behavior we are looking for. However, with as few trees as 30, we may expect that different forests will have a high variability on the filter importances and consequently on the accuracy.

We have run the ExtraTrees 12 times with the same hyper-parameters and the same learning matrix. Table ?? holds the accuracies and table ?? holds the first line of the correlation matrix of the filter importances between the tests. Figure ?? depicts the filter importances for the two firsts tests. The test number 0 corresponds to the results given in the previous section. As we can see, the variability is very small. More tests would be needed to establish a rigorous confidence interval but the conclusion seems clear nonetheless: ExtraTrees variability has no impact on the accuracy. This is also backed up by the evident stability of the filter importances. Such stability is due to the smoothing effect of aggregating all the feature importances of a given filter. The overall conclusion is that tree variability has no impact of the method for a given learning matrix. This stability also suggests that hyper-parameters such as the size of the local random subspace (k) and the number of trees will play little role to improving the accuracy. We will confirm that with the following experiments. The number of subwindows might, however, have more influence of the accuracy. This will also be established by one of the experiments.

Let us digress on the filter importances. From figure ??, we can see that several filters have a high importance; of the same order of magnitude than the original image (filter 0). On the other hand, some filters bring little to none information in the classification process. Either these filters are not (co-)useful or the trees are to small to incorporate their usefulness. In regard to the correlations between tests, however, the co-uselfulness and complexity hypotheses are less likely. If it were one of these problems, the low-rated filters would have developed on different trees and therefore the filter importance profiles would be less similar. Therefore, the most likely hypothesis is that those filters are simply not useful. Inspecting them in parallel of the best ones might reveal information about the classification task. In any case, removing them to make room for more interesting filters can only improve accuracy. Sadly, the gain is unlikely to be substantial. Indeed, the current accuracy gain compare to the pixit is marginal, even though we already have many filters whose importance rivals the original image.

We now turn to the variability produced by random filter generation with the other parameters fixed to their default values.

⁽a) Accuracy variability

⁽b) Correlation vector of the filter importances with test number 0

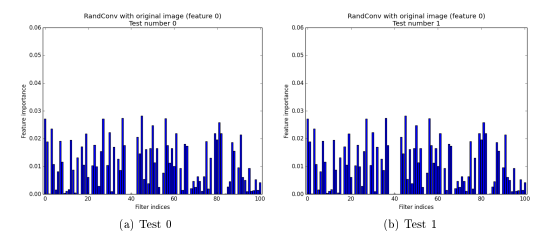


Figure 5.2: Filter importances for two different forests with the same parameters

Number of subwindows	Accuracy
1 (whole image)	0.44
1 (subwindow)	0.40
5	0.48
10	0.515
14	0.514

Table 5.2: Accuracy as a function of the number of subwindows

5.1.3 Accuracy as a function of the number of subwindows

As a reminder, the number of subwindows enlarges the learning set and therefore allows for more complex models. In this experiment, we tested how accuracy varied with the number of extracted subwindows. We used 1, 5, 10 and 14 subwindows. The memory limited us to that threshold in order to keep the other parameters to their default values (table ??, page ??). We also tested the method where no subwindows are extracted but rather the whole image is used directly. The results are reported at table ??. The filters were the same in all the tests.

As we might have suspected, using the whole image yields better accuracy than using only one random subwindows. However, a few of them already accounts for a better accuracy. Judging by the accuracy jump between 5 and 10 subwindows compare to the stagnation between 10 and 14 subwindows, we may conclude that 10 subwindows allows for complexenough models.

5.1.4 Influence of the number of trees

The accuracy is usually an increasing function of the number of trees. In this subsection, we will see how this function behaves for our method and we will look at the filter importances as well. In section 5.1.2 about the variability, we foresaw that increasing the number of trees would have little influence on the filter importances. Now is the time to validate this thesis. Once again, the other parameters will be set to the default values of table 4.1 on page 20 and the learning matrix is the same in all the tests.

In this experiment, we measure the accuracy for the following number of trees: 10, 30, 50, 100, 200, 300, 400 and 500.

The result of this experiment is depicted by figure 5.3. The general trend is indeed an increasing function of the number of trees. The local decreases are small and are due to the randomization of the tree growing algorithm. From a relatively low accuracy of 0.484 at 10 trees, the method quickly jumps to the value of 0.515 for 30 trees. The increase rate

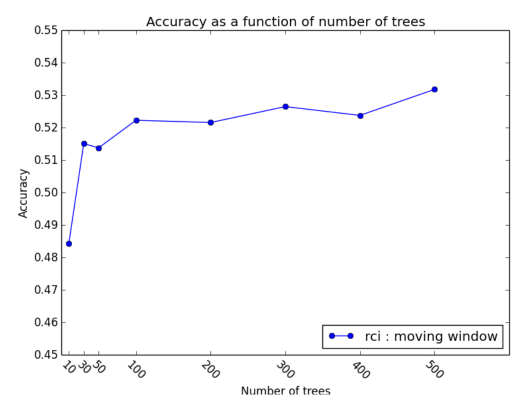


Figure 5.3: Accuracy as a function of the number of trees

is then considerably slowed down before jumping again at 500 trees, yielding 0.532. The first observation to note is that we have attained the best result of the pixit method with half the subwindows. Of course, this must be put in perspective with the variability of the RandConv method. The second conclusion we may draw is that working with 30 trees is representative of the method's accuracy. We can serenely draw conclusions with that many trees and only use more of them to grab the last few percents of accuracy.

We now turn to the filter importances in the classification process. Figure ?? shows the eight distributions and table 5.3 the corresponding correlation matrix. As we had predicted, the number of trees does not impact the filter importances. Therefore, using 30 trees is well enough to assess filter usefulness.

5.1.5 Influence of the minimum number of samples to split

This parameter control the model complexity. In the previous experiments, the trees were fully grown $(n_{min} = 2)$ and variability reduction was achieve through the smoothing effect

Number of trees	10	30	50	100	200	300	400	500
10	1							
30	0.19	1						
50	0.99	0.19	1					
100	0.05	0.06	0.05	1				
200	0.99	0.19	0.99	0.05	1			
300	0.99	0.19	0.99	0.05	0.99	1		
400	0.99	0.19	0.99	0.05	0.99	0.99	1	
500	0.14	0.15	0.15	0.12	0.14	0.15	0.15	1

Table 5.3: Correlation matrix of the filter importances with respect to the number of trees

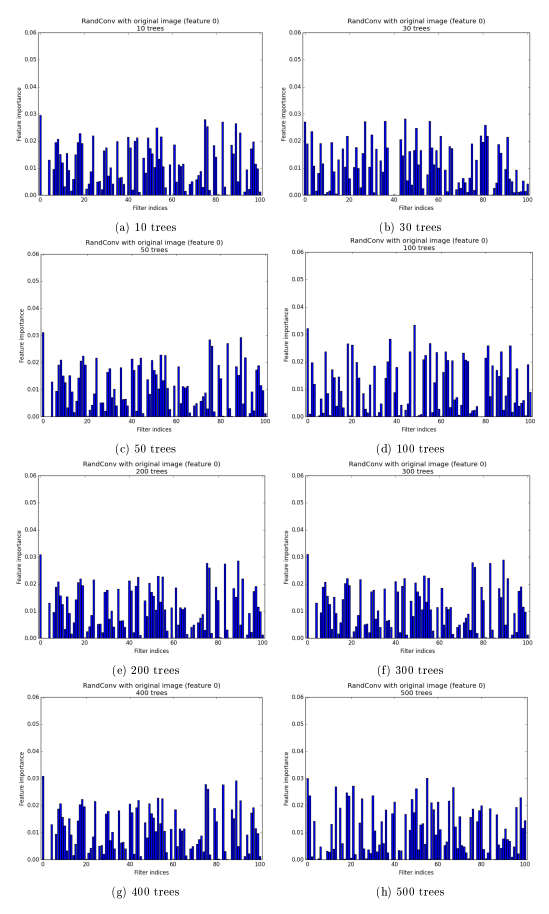


Figure 5.4: Filter importances with different numbers of trees

of the forests prediction aggregation. Still, the model might be too complex to begin with, thus requiring some pruning. We tested the following values of this hyper-parameter: 2, 10, 50, 500. The corresponding accuracies are: 0.515, 0.512, 0.498, 0.453. Once again we see that the default value performs well. Besides, pruning too much the trees is harmful: the models are not complex enough any longer.

The influence on the filter importances is more interesting. We clearly can identify two groups of filter. The first one is composed of the filters whose importance increases. This increase implies a decrease of the filter importances in the second group. It is thus possible to see where the filter is used in the trees. The first group holds the top of the tree. This was already obvious for the filters whose importance was either large or low. However, this approach sheds more light on the average filters, where we can observe divergent behaviors. Average filters of the second group are purely co-useful filters: they bring little information on their own but are helpful with the other filters. This characteristic resembles the "V-structures" of the graphical probabilistic models. On a specific classification task, these particular filters may be worth investigating. It might be difficult to identify the filters on which they depend, however.

5.1.6 Influence of the number of filters

Our expectation regarding the number of filters is that, with few of them, the accuracy might be better or worse than the pixit depending on the filter usefulnesses. With many filters, however, there should be enough usefulness altogether to be able to beat the pixit systematically. The accuracy should increase accordingly with the number of filters as long as the learning set size allows for complex-enough model. However, from the comment on the variability (subsection 5.1.2), we suspect that we will reach an upper bound quite fast.

From a more practical point of view, we will have to limit ourselves to, which should not be a problem if we were right about the upper bound.

5.1.7 Influence of the filter generator

5.1.8 Influence of spatial poolings

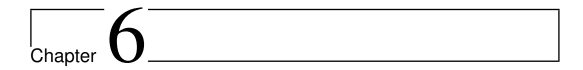
In this subsection, we tackles the influence of the spatial poolings with a couple of experiments.

5.2 Feature learning scheme

The previous section focused on the ET-DIC pros and cons. We now turn to the ET-FL mode. In this mode, we use totally randomized trees to build a visual dictionary and the actual classification is carried out by a SVM.

This mode is less prone to a deep analysis of the influence of the hyper-parameters. The parameter k is fixed to 1, and the number of subwindows, the number of trees and the minimum number of samples to split all influence the number of leaves in a well-understood fashion. An increase in the first two will produce an increase of the number of leaves and ultimately of the discriminant power. Therefore, the number of subwindows will be severely limited by the memory requirement while the number of trees limitation will be the computational time, although to a lesser extent. A good tradeoff must then be found between the number of subwindows and number of filters and spatial poolings. Since we are not averaging the results any longer, we have lost the ability to reduce the model variability and must resort to pruning in order to limit its complexity. Usually, the optimal pruning should be obtained by cross-validation. However, for our database, a good value of $n_{min} = 500$ has already been established ().

5.2.1 Accuracy



Conclusion and perspective

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