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Kevin Bailly, Maurice Milgram

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Boosting Feature Selection for Neural Network based Regression

Abstract

Head pose estimation problem is well known to be a challenging task in computer vision and is a useful tool for several applications involving human-computer interaction. This problem can be stated as a regression one where the input is an image and the output is pan and tilt angles. Finding the optimal regression is a hard problem because of the high dimensionality of the input (number of image pixels) and of the large variety of morphology and of illumination. We propose a new method combining a boosting strategy for feature selection and a neural network for the regression. Potential features are a very large set of Haar like wavelets which are well known to be adapted to face image processing. To achieve the feature selection, a new Fuzzy Functional Criterion (FFC) is introduced which is able to evaluate the link between a feature and the output without any estimation of the joint probability density function as in the Mutual Information. The boosting strategy uses this criterion at each step: features are evaluated by the FFC using weights on examples computed from the error produced by the neural network trained at the previous step. Tests are carried out on the commonly used Pointing 04 database and compared with three state-of-the-art methods. We also evaluate the accuracy of the estimation on FacePix, a database with a high angular resolution. Our method is compared positively to a Convolutional Neural Network, which is well known to incorporate feature extraction in its first layers.

Key words: Input Feature Selection, Boosting, Regression, Fuzzy Functional Criterion

1. Introduction

In a large number of regression problems, it is not easy to find relevant features from a huge set of potential input variables. In image based regression for instance, the number of pixels, which corresponds to the input dimension, is very large. Moreover, pixel level is not necessarily suitable for a particular

problem and an huge set of potential features can be extracted from an image to complete or to supply the necessary information. Just few of them are relevant, while most of them are redundant. In this paper, we present a framework to learn simultaneously relevant features and the corresponding regressor.

A large number of solutions have been proposed in feature selection literature. They can be divided into filter, wrapper and embedded approaches [5]. In filter based method, features are first selected using a specific measure of relevance such as mutual information or Pearson's correlation. The second stage consists of estimating parameters of the regression model on the selected subset. This approach is computationally efficient, but presents a major drawback [7]: the regression model is not taken into account. Thus, a suboptimal set of relevant features tends to be selected rather than the complete set of useful features.

Wrapper approaches use the prediction performance of the model to quantify the relevance of a feature subset. In this category, greedy forward selection algorithms are frequently used: they progressively integrate new variables which optimize the regression score. This strategy is really time consuming and intractable when the regression learning algorithm is too complex.

Embedded approaches incorporate the feature selection directly into the learning algorithm. In ridge regression such as LASSO [15] or Input Decay [1], a regularization term is introduced in the cost function. The aim is to train the model so that inputs which contribute poorly to the regression process are penalized. These methods are well suited when the number of potential features is quite restricted.

In section 2, we introduce BISAR (Boosted Input Selection Algorithm for Regression) which combines a filter and a wrapper approach.

BISAR uses a kind of boosting in the feature selection process. In its simplest form, boosting strategy aims to decrease the error rate of a classifier/regressor by concentrating at each iteration on examples that are particularly difficult to classify/regress. This is usually done by iteratively adapting the training samples weighting. At each iteration the weight of training examples depends on the performance of the classifier/regressor in the previous iteration.

AdaBoost [3] is a popular algorithm to iteratively build a classifier as a linear combination of so called weak classifiers. At each step, a new weak classifier is added optimizing the classification error rate with a new weighting on training samples. AdaBoost.RT, proposed by Shresta [13] for regression problems uses the so-called absolute relative error threshold ϕ to project training examples into two classes (poorly and well predicted examples) by comparing the absolute relative error with the threshold ϕ . The problem is that it is not obvious to

choose the right value for the threshold ϕ . Therefore we have selected another boosting strategy, AdaBoost.R2 [2], to compare several boosting strategies in BISAR. There are other methods like [11] and [21] that also combine these two approaches in a different way. Algorithm described in [11] is clearly dedicated to solve classification and not regression problems and aims to reduce the number of features used by each weak classifier. In [21], an approach which limits the correlations of the selected feature set for gene selection in microarray data analysis is presented. This approach seems to be specific to this kind of problem and the boosting strategy is completely different from BISAR.

The two main contributions of this paper are:

1. The Fuzzy Functional Criterion (FFC): a new filter used to select *relevant* features (section 2.1)
2. A new boosting strategy that selects incrementally new *complementary* inputs for the regressor (section 2.3).

A large number of experiments (section 3) with BISAR on a classical pattern recognition problem is covered in this article (section 4). Comparison are drawn between BISAR and state-of-the-art methods (section 5). Finally we present conclusions and prospects in section 6.

2. Boosted Input Selection Algorithm for Regression (BISAR)

The regression we propose is based on two modules working together:

1. A boosted feature selection algorithm based on a new fuzzy criterion
2. A neural network with growing input set according to the boosted incremental choice of features

In order to select useful features to perform neural network based regression, we adapt both a filtering paradigm based on our new criterion FFC (Fuzzy Functional Criterion) independent from the regression engine and a boosting paradigm based on the neural network error.

We have a set of examples $\mathbf{x}_i \in \mathbb{R}^d$. To each \mathbf{x}_i is associated a value $y_i \in \mathbb{R}$ that we want to predict. Data are divided into a training set A , a validation set V and a test set E . A set F of features H_k ($1 \leq k \leq N$) can be computed for each \mathbf{x}_i . F can be extremely large (typically more than 10 000 elements as in our case). The main objective of our method is to select a subset of features $FS \subset F$ adapted to a specific regressor. We can summarize our method as follows :

1. Initialize all examples at the same weight; FS is empty.

2. Compute the fuzzy criterion for all features in F using the training set and the current weights. The best feature according to this criterion is added to FS .
3. Train a new regressor taking as input all previously selected features.
4. Compute the new weights using the error of the regressor for each example
5. Repeat from 2 to 5 until the maximum number of input is reached
6. Select the regressor with the lowest error on the validation set V during all iterations

Hence, our method combines an iterative filtering approach for feature selection and a neural network. The selection process uses weights provided by the network error at each step. It means that, given a set of pairs (\mathbf{x}_i, y_i) of training input-output and weights on example, we determine a new score for each feature H_k . This score has to reflect the relevance of feature H_k for our problem, that is the matching between \mathbf{x}_i (a pattern) and y_i (the target value associated to \mathbf{x}_i that we want to predict).

2.1. Feature selection criterion

We have to determine a criterion to select features. This criterion must measure how much the output y depends functionally on this feature H_k . Moreover, this criterion should support a weighting function over the example set. Usual statistical measures (correlations, covariance, etc.) do not fit to our situation because of the linear hypothesis that we do not want to assume.

We are going to examine two criterions: the well-known Mutual Information (MI) and our new Fuzzy Functionnal Criterion (FFC) in next sections.

2.1.1. Mutual Information

The MI between two discrete random variables x and y is defined to be :

$$I(x; y) = - \sum_{x,y} P(x, y) \log_2 \frac{P(x, y)}{P(x) \cdot P(y)} \quad (1)$$

And:

$$I(x; y) = H(x) + H(y) - H(x, y) = H(x)H(x|y) = H(y)H(y|x) \quad (2)$$

Where $H(x)$ is the entropy of the random variable x and $H(x, y)$ is the joint entropy of these variables.

There is a continuous definition of the MI for continuous variables but it is necessary to approximate the MI before using it practically in a program. So,

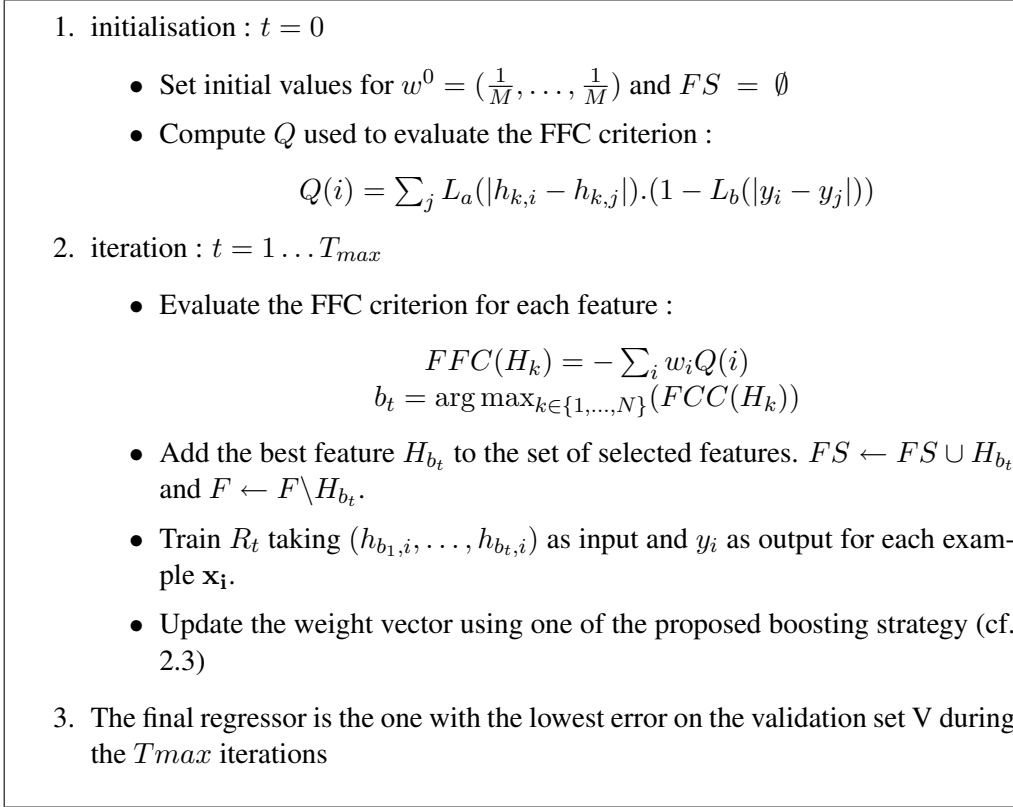


Figure 1: BISAR algorithm

the main problem is the estimation of the joint probability distribution function $P(x, y)$. We use the Parzen windows estimator to evaluate $P(x, y)$ which is a popular non-parametric approach and which is also very close to the computation performed by our FFC criterion. It is well known that the size of Parzen windows can be critical and has to be tuned finely. Unfortunately, in our case, we do not have any prior information about the probability distribution function $P(x, y)$. We will present in section 4.1 a comparison between the MI and the FFC, however, we show here some arguments against the MI, even if this measure is still a very good and broadly used tool:

1. MI computation lays on the probability distribution function (pdf) $P(x, y)$ which can be difficult to estimate, particularly if the pdf has a high dynamic
2. MI is symmetric: $I(x, y) = I(y, x)$; in our case, we want to measure the fact that y is a function of x and not the converse
3. We shall have to deal with weights on samples (see the further section on boosting) and the criterion has to incorporate these weights. Due to its

structure, we have to fully compute the MI for each feature and for each step of our algorithm

4. To transpose our method to higher dimension feature or output, it becomes exponentially harder to estimate the pdf due to the famous “curse of dimensionality”

We propose in the next paragraph a new criterion so called the Fuzzy Functional Criterion (FFC):

1. FFC does not need the joint pdf of (x, y)
2. FFC is not symmetric
3. FFC can be pre-computed independently of the weights and be very quickly computed when the weights are known
4. FFC can be easily transpose to higher dimension feature or output

2.1.2. Fuzzy Functional Criterion (FFC)

Our Fuzzy Functional Criterion is like an energy function. The basic idea is the following: we want to evaluate the relation between two variables u and v (u is a measure and v a target to be predicted). We call P the logical proposition “ u_1 and u_2 are close” and Q the logical proposition “ v_1 and v_2 are close”. If a functional smooth relation between u and v exists (i.e. $v = F(u)$ and F is a smooth function), we can say that the logical implication $P \Rightarrow Q$ is true. If the variable v takes a value in a discrete subset of the real numbers set, we have to adapt the criterion slightly. This is done using the truth table of the logical implication. We find it is equivalent to “ $\neg P$ or Q ” and also to $\neg(P \text{ and } \neg Q)$. We take the fuzzy logic formulation of P and Q based on classical triangular shaped functions [22] denoted by L and defined as:

$$L_a(e) = \begin{cases} 1 - \frac{|e|}{a} & \text{if } |e| < a \\ 0 & \text{otherwise} \end{cases} \quad (3)$$

To quantify the fact that “ u_1 and u_2 are close” is true, we just take the value $L_a(|u_1 - u_2|)$ into account, where a is the spread of the triangular function and its value is discussed later. We can do the same for v and write:

$$Z = 1 - L_a(|u_1 - u_2|)(1 - L_b(|v_1 - v_2|)) \quad (4)$$

Z is a fuzzy evaluation of our implication $P \Rightarrow Q$. To build our criterion, we have to sum up Z over all (u_1, u_2, v_1, v_2) (the constant “1” can be dropped):

$$FFC = \sum_i \sum_j -L_a(|u_i - u_j|)(1 - L_b(|v_i - v_j|)) \quad (5)$$

The sum is taken over all quadruples. In our case, we replace u_i by $h_{k,i}$ (for some feature H_k) and v_i by the target y_i . We also introduce the weighting function on examples, w_i , which is a non-negative function with $\sum w_i = 1$. These weights will be useful in the boosting process. So we can reformulate our criterion as:

$$FFC(H_k) = - \sum_i w_i \sum_j L_a(|h_{k,i} - h_{k,j}|) (1 - L_b(|y_i - y_j|)) \quad (6)$$

The criterion $FFC(H_k)$ has to be maximized. Two parameters (a and b) control the spread of the L functions and are used to normalize the criterion over all features. In many problems, the range of u and v variables are bounded and can be determined. We shall tune a and b as a fixed proportion of these ranges. Experiments have shown that the criterion is not too sensitive to this tuning. Another way to cope with the normalization issue is to replace u values by their rank. In this option, the rank can be seen as a simple normalization: if there are N examples, the number $rank(x)/N$ is always in the interval $[0, 1]$. Notice that the difference of ranks divided by the number of examples has the same distribution for all features.

The most interesting aspect is the weighting function on the examples does not imply to compute at each step a new FFC from scratch: it is possible to compute an intermediary form $Q(i)$ of FFC and to modulate it with the weighting function on examples based on the regression error. Equation (6) can be written as:

$$FFC(H_k) = - \sum_i w_i Q(i) \quad (7)$$

where :

$$Q(i) = \sum_j L_a(|h_{k,i} - h_{k,j}|) (1 - L_b(|y_i - y_j|)) \quad (8)$$

and Q can be precomputed once.

2.2. Regressor

The regressor is the second important element of our system. We have tested different regressor from the Radial Basis Function (RBF) network family. They are known to be very efficient in function interpolation. An important feature is that their output becomes close to zero when the input is very far from any training sample; this can be used, for example, for a rejection purpose. RBF

networks have three layers but are quite different from classical Multi Layer Perceptron (MLP): the hidden layer is made of RBF units (and not sigmoid) learned via a specific algorithm. Output layer is fully connected to the hidden layer and is a linear neuron (or several linear neurons). There are two parts in the training of a RBF network: during the first one, hidden cells are chosen using the EM algorithm (or in our case the K-means algorithm). In the second part, a classical gradient descent algorithm finds the optimal weights for the connections between hidden layer neurons and the output layer neuron. A hidden neuron N_i can be viewed as a couple made of a point x_i in the input space and a spread value s_i . For an input x , the output y of the neuron N_i is computed by the spherical radial function:

$$y = e^{-\frac{\|x-x_i\|}{s_i}} \quad (9)$$

It is also possible to replace the spherical function by the Gaussian function which uses the full covariance matrix S_i :

$$y = e^{-(x-x_i)^T S_i^{-1} (x-x_i)} \quad (10)$$

To learn hidden neurons values, we can use either an EM or a simple K-means algorithm. It has been shown that the latter is a more robust procedure. In EM approach, a covariance matrix of each cluster is computed at each step and it happens frequently that this matrix is poorly conditioned. This fact leads to great instabilities during the learning process. In K-means approach, covariance matrix of each cluster is computed once at the end of the process. The Euclidean distance is used in the clustering step. After the choice of the hidden neuron number N , a random initialization of centers x_i is done. Then, at each step, we select for each training sample x the nearest center x_i producing a cluster C_i of all training samples close to x_i . Then, for each cluster, we compute m_i the center of gravity of C_i and set x_i to m_i . The process runs until x_i are stable or the maximum number of iterations is reached. Once hidden neurons are learned, we determine other weights (between hidden and output layers) via a gradient descent to minimize the Mean Square Error between desired output and network output.

We also used a Generalized Regression Neural Network (GRNN) which can be considered as a simplified RBF network ([19]). Each center of the hidden unit corresponds to an example of the training set and weights between hidden and output layers are set to the target values. It means this network does not need any training step but instead just keeps all prototypes (in the hidden layer) and

all targets (in the second layer weights). This network can be viewed as a Parzen window procedure applied to an interpolation task.

2.3. Boosting strategy

At the beginning, the neural network starts with only one input cell which corresponds to the best feature given by the criterion using a uniform weighting function. In the second step, a new network is trained with 2 input cells: the new input corresponds to the second best feature which is selected among all features (except the feature already selected) using weights provided by the error of the first network. It means that examples that are poorly processed by the first network will receive higher weights than others. By doing that, we believe that the resulting weights, when fed to the criterion computation, will lead the criterion to choose a new feature better adapted to these examples. It is clear that our boosting paradigm enhances the regression system. Contrary to AdaBoost [12], iterative reweighting is only used to select the best complementary feature. Features combination is entirely handled by the regressor and weights are taken into account. It can be seen as a regularization to prevent overfitting.

Several strategies are possible to drive the selection process. One has to tune the weights according to the error and this weight adaptation has to be defined carefully.

We test three boosting mechanisms. The first one lies on a memoryless process. Weights at iteration $t + 1$ are only related to the quadratic error ϵ_k^t at the previous iteration. In our case, we adopted this simple relation:

$$w_{t+1}(i) = \frac{(\epsilon_t(i))^2}{\sum_{i=1}^M (\epsilon_t(i))^2}; \quad (11)$$

The second boosting strategy is cumulative and the weight of each example depends on the regression model error and the previous weights.

$$\tilde{w}_{t+1}(i) = \begin{cases} w_t(i) & \text{if } \epsilon_t(i) < \text{median}_i(\epsilon_t(i)) \\ \max\{\alpha w_t(i), w_{max}\} & \text{otherwise} \end{cases} \quad (12)$$

$$w_{t+1}(i) = \frac{\tilde{w}_{t+1}(i)}{\sum_{i=1}^M (\tilde{w}_{t+1}(i))} \quad (13)$$

Weights of half the examples with the highest prediction error are multiplied by a constant accumulation factor α . Here w_{max} is a constant used to avoid overfitting. Typically, α is set to 1.1 and w_{max} to 0.1.

The third reweighting procedure is the same as in AdaBoost.R2 [2]. The performance of the regressor is measured using a loss function :

$$L_t(i) = \left(\frac{\epsilon_t(i)}{\max_{i=1 \dots M} \epsilon_t(i)} \right)^2 \quad (14)$$

This function is averaged over all the weighted examples

$$\bar{L}_t = \sum_{i=1}^M w_t(i) L_t(i) \quad (15)$$

Knowing \bar{L}_t , the weight updating parameter denoted β_t is computed as follows:

$$\beta_t = \bar{L}_t / (1 - \bar{L}_t) \quad (16)$$

Finally, new weights on examples are calculated as :

$$\tilde{w}_{t+1}(i) = w_t(i) \beta_t^{(1-L_t(i))} \quad (17)$$

And normalized as :

$$w_{t+1}(i) = \frac{\tilde{w}_{t+1}(i)}{\sum_{i=1}^M (\tilde{w}_{t+1}(i))} \quad (18)$$

Contrary to other boosting algorithms for regression, no parameter has to be calibrated and tests on different data sets have shown good results in [13].

In the three reweighting procedure, weights are normalized to make their set a distribution. In the rest of this paper, the three boosting strategies will be referred as the memoryless (11), median (12) and AdaBoost.R2 (17) boosting strategy respectively.

3. Experimental setup

In order to test our approach, BISAR is applied to a head pose estimation problem. Research in head pose determination is now an important topic in pattern recognition and many face analysis applications such as face recognition systems and human computer interaction need this information. Many solutions have been suggested in [10]. One of them considers head pose estimation as a non linear regression problem. These methods try to learn a functional mapping between an image (or a set of image features) and the head direction. This section will describe the features and the data sets used in our experiments.

3.1. Image features

Faces are down-sampled to a fixed size of 32×32 pixels. Moreover, color images are converted to gray scale images and histograms are dynamically adjusted. We use four types of features. The first three descriptors correspond to the popular Haar like features [17] as depicted in Fig. 2. We choose these features because of their good performance obtained in related area such as face detection [17] and alignment [20]. Moreover they are simple and very easy to compute using the integral image [17].

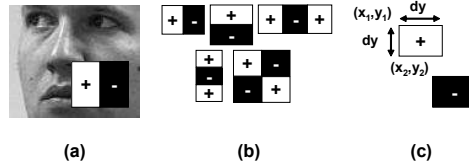


Figure 2: Image features. (a) a feature example. (b) Representation of the Haar like features used in our experiment. (c) The last type of features are the difference between the sum of the pixels within two non connected rectangular regions

Another kind of features, we propose here, are the difference between the sum of the pixels within two non connected rectangular regions. Features are parameterized by four values x_1 , y_1 , dx and dy . x and y correspond respectively to the horizontal and vertical position of the feature in the image. dx and dy are the width and height of a rectangle. Two extra parameters x_2 and y_2 are needed to describe features of the fourth category. They correspond to the position of the second rectangle in the image.

3.2. Data set 1: Pointing 04

3.2.1. Data set description

The first database used for this evaluation is the publicly available Face Pointing04 database. It was used to evaluate head pose estimation systems in the Pointing 2004 Workshop on Visual Observation of Deictic Gestures. It was also included in the International Evaluation on Classification of Events Activities and Relationships (CLEAR 2006). The corpus consists of 15 sets of images. Each set contains 2 series of 93 images of the same person at 93 different poses. The first series is used for learning. Tests are carried out on the second series.

There are 15 people in the database, male and female, wearing glasses or not and with various skin color and facial hair. The pose is determined by 2 angles,

ranging from -90° to $+90^\circ$ for the horizontal angle (pan) and from -60° to $+60^\circ$ for the vertical orientation (tilt). Fig. 3 depicts the variety of this database.



Figure 3: Pointing 04 database samples.

Ground truth was obtained by asking people to look at some specific markers in the room. In fact, this ground truth is strongly more related to the gaze direction than to the head direction. This acquisition process leads to a very challenging database with ambiguities and variations on pan angle result in very similar images.

3.2.2. Face localization

Faces in Ponting 04 are just a small area of the whole image. Thus a lot of information are not relevant and can disturb the regression process so we need somehow to crop the face area. However, because no generic multi-pose face localizer is available, we decided to manually crop the faces. We also proposed an ad hoc method similar to [4] for two main reasons:

- To reduce the bias introduced by the manual cropping and test the approach on noisy data.
- To compare results with other methods which integrate a face localization step.

Our face localization is based on skin colour. First, frontal faces are detected in the training database using the Viola-Jones face detector [17]. Pixels within the detected area are used to build a histogram on H (Hue) and S (Saturation) channels of HSV color space. Other pixels are randomly picked outside the face area in order to build a histogram of background. Face pixels are detected in the test image set with a Bayesian rule based on these skin and non skin histograms. The face bounding box size is proportional to the standard deviation of skin pixels along X and Y axis. In this method, only frontal faces of the training set are needed and no manual labelling is required. The main drawback is the weak

accuracy of this method in few cases. For example, the neck can be included in the bounding box. The lack of precision in localization can greatly degrade head pose estimation results. In result section, results are presented for both manual and automatic face localization.

3.3. Data set 2: FacePix

The second data set used for this evaluation is the publicly available FacePix database [9] from the Center for Cognitive Ubiquitous Computing (CUbiC). The FacePix database consists of three sets of face images: one set with pose angle variations, and two sets with illumination angle variations. Each of these sets are composed of a set of 181 face images (representing angles from -90° to $+90^\circ$ at 1° increments) of 30 different subjects, with a total of 5430 images. All the face images (elements) are 128 pixels wide and 128 pixels high. These images are normalized, such that the eyes are centered on the 57^{th} row of pixels from the top, and the mouth is centered on the 87^{th} row of pixels. The pose angle images appear to rotate such that the eyes, nose, and mouth features remain centered in each image.

In our experiment, we use the set with pose angle variation ranging from -90° to $+90^\circ$ with a step of 5° . 20 faces were randomly picked from the original data for training, 6 for testing and 3 for cross validation. Contrary to Pointing 04, only the pan parameter is modified. Moreover, labels on each image are reliable since the acquisition process is more accurate.

4. Experimental Results

4.1. Feature selection criterion

In this first experiment we compare the Fuzzy Functional Criterion to the Mutual Information. In order to restrict the impact of outer factors, a GRNN is used and no boosting is applied, i.e. weights on example remains constant during iterations. The optimal number of inputs is chosen using the score on the validation set. Table 1 shows FFC has the edge over MI. Moreover FFC is faster to use equation (7) when boosting is introduced. These two reasons led us to give priority to FFC criterion in the next experiments.

4.2. Boosting strategy

In this section, we compare several boosting strategies. Tests are carried out using a GRNN. Thus no parameter needs to be tuned and results do not depend on random initialization. Results are presented for both FacePix and Pointing 04

Table 1: FFC vs MI : mean absolute error in degree on the test sets. Value in brackets indicates the corresponding number of features

	Pointing 04				FacePix
	Manual cropping		Automatic cropping		Pan error
	Pan error	Tilt Error	Pan Error	Tilt error	
FFC	8.1 [°] (580)	8.4 [°] (560)	12.5 [°] (480)	15.9 [°] (100)	6.2 [°] (191)
MI	8.4 [°] (600)	8.9 [°] (560)	12.8 [°] (460)	15.7 [°] (140)	6.4 [°] (180)

databases. For this latter data set, tests are performed on manually and automatically cropped images (cf. 3.2.2). In this way, we can evaluate the robustness to noise and outliers. Results on the test set for the three boosting strategies (cf. 2.3) are reported in table 2. We also present results obtained without any boosting (denoted by 'none' in the table). We investigate the performance of BISAR with up to 600 iterations and we select the classifier with the best performance on the validation set.

Table 2: Mean absolute error in degree on the test set for different boosting strategy. Value in brackets indicates the corresponding number of features

	Pointing 04				FacePix
	Manual cropping		Automatic cropping		Pan error
	Pan error	Tilt Error	Pan Error	Tilt error	
None	8.1 [°] (582)	8.4 [°] (573)	12.5 [°] (467)	15.9 [°] (102)	6.2 [°] (191)
Memoryless	7.3 [°] (553)	8.5 [°] (267)	11.6 [°] (389)	13.5 [°] (78)	5.4 [°] (243)
Median	7.6 [°] (569)	8.9 [°] (223)	11.9 [°] (464)	16.5 [°] (55)	4.5 [°] (599)
AdaBoost.R2	8.2 [°] (382)	7.6 [°] (457)	10.9 [°] (430)	12.3 [°] (120)	6.0 [°] (265)

The effect of boosting is evident as any reweighting method outperforms results without boosting. In addition, the AdaBoost.R2 reweighting function seems slightly better since it gives better results for three of the five test sets. Figure 4 depicts the evolution of the mean absolute error against number of selected features. It shows that the error decreases faster for Adaboost.R2 reweighting method in the first few iterations. The median strategy gives better results on FacePix but it requires up to three times the number of features as compared to other methods. The memoryless procedure has good results and outperforms the

AdaBoost.R2 strategy on the pan estimation when image are manually cropped but it tends to overfit on the noisy data (cf. figure 4(b)).

The results are better on FacePix than on Pointing 04. This is most probably due to the differences underlined in 3.3 (only pan variation, bias on the cropping and the greater accuracy of the ground truth).

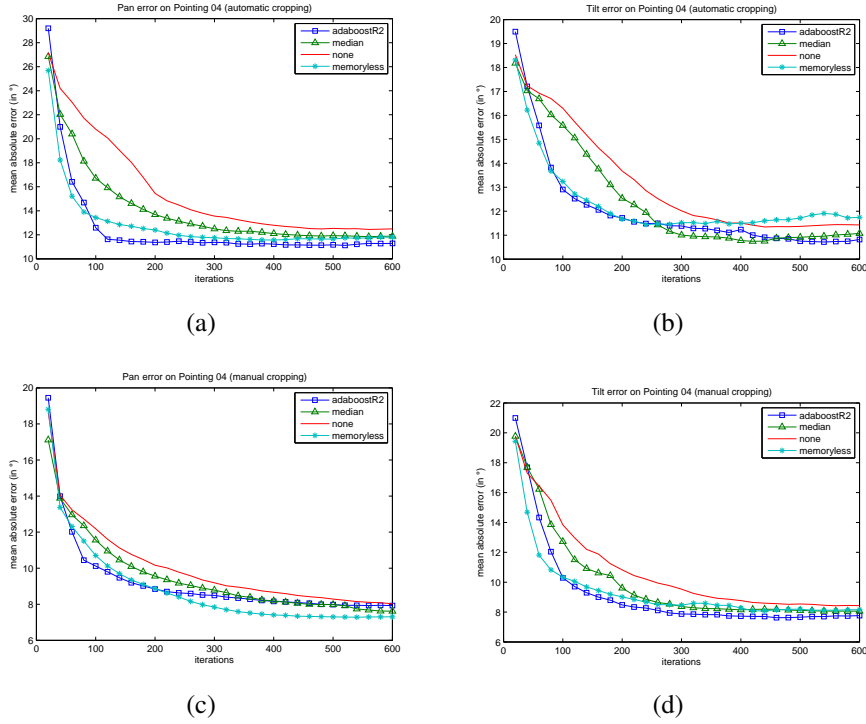


Figure 4: comparison of boosting effect against iterations

4.3. Regressor architecture

In this section, we experiment other RBF networks. We performed 300 iterations for each regressor on both Pointing and FaceFix databases. We kept AdaBoost.R2 weighting since it was found to be the best in the previous section. Test are carried out on two RBF networks which differ on the number of hidden units. RBF 1 has half the number of training example while RBF 2 has one fifth of this number. Hidden cells are chosen using a K-means algorithm and covariance matrix of each cluster is only computed at the end of the process. Table 3 shows that performance is directly related to the number of units in the hidden layer. Although the number of units is really large especially in the GRNN, no

overfitting was observed. It may be partly due to the spherical radial functions. We have chosen spherical functions because they have only d independent parameters (where d is the dimensionality of the input space) as compared with the $d(d + 3)/2$ independent parameters of a full Gaussian basis function. Moreover, training process with RBF is sometimes unstable since initial centres are randomly specified.

Table 3: Mean absolute error in degree on the test set for different regressor architecture. Value in brackets indicates the corresponding number of features

	Pointing 04		FacePix
	Pan error	Tilt Error	Pan Error
GRNN	11.4° (288)	12.3° (120)	6.0° (265)
RBF 1	14.0°(217)	15.8°(256)	7.2°(120)
RBF 2	15.2°(161)	16.0°(284)	8.5°(237)

4.4. Multiresponse regression

In this experiment, pan and tilt were jointly trained. To achieve this, we slightly modified the FFC. The output is a vector and we use the L_1 norm instead of the absolute difference of the outputs in equation (6). Learning simultaneously pan and tilt information seems, a priori, more difficult since the network has to learn two responses instead of one. However, the two pieces of information are interlinked. Tables 4 shows that the joint learning performed slightly better on pointing 04. It can also be observed that the joint learning needs to extract a smaller number of features.

Table 4: Mean absolute error in degree on the test set. Value in brackets indicates the corresponding number of features

	Manual cropping		Automatic cropping	
	Pan error	Tilt Error	Pan Error	Tilt error
joint learning	7.5° (600)	8.0°(600)	10.3° (294)	12.0° (294)
separate learning	8.2°(382)	7.6° (457)	10.9°(430)	12.3°(120)

5. Comparison with other methods

5.1. Convolutional Neural Networks

Convolutional Neural Networks (CNN) are bioinspired [6] multilayered neural networks and were first proposed by Le Cun et al. [8] for handwritten character recognition. The idea is to build an architecture using both local receptive fields with share weights and subsampling. In a typical architecture, there are convolutional layers where each unit receives input from a set of cells located in a small rectangular neighbourhood in the previous layer. With local receptive fields, cells can extract basic features (oriented edges for example). Layers extracting feature are called convolutional because of the weight sharing: all units in this layer perform the same linear operation (before applying the sigmoid function) and the process can be viewed as a simple convolution. These features are then combined by the subsequent layers in order to detect high-order features. Before this new extraction, the network reduces the resolution of the feature maps (by averaging and subsampling). This reduction has two justifications: to reduce the size of the layer and to bring some robustness against small distortions. Figure 5 describes the CNN architecture used in our experiments, following Simard recommendations [14].

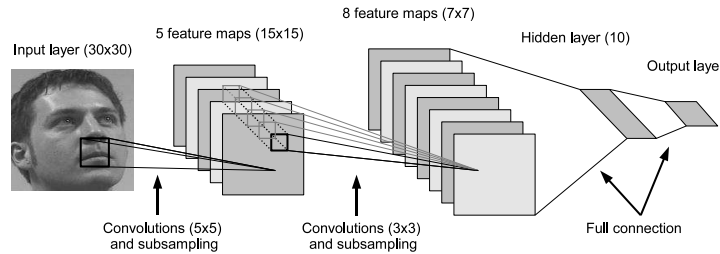


Figure 5: Architecture of the Convolutional Neural Networks

CNN can be trained with the classical backpropagation algorithm and we have used an adaptive learning rate and a momentum regularization. Table 5 summarizes comparative results between BISAR and CNN. It is worth noting that performance are quite the same on FacePix, but BISAR outperforms CNN on Pointing 04.

5.2. State-of-the-art methods on Pointing04

Performance are compared with three standard methods for this database [10]. The first one proposed by Voit et al. [18] uses a multi layer perceptron

Table 5: Comparison with convolutional neural networks

	Pointing 04		FacePix
	Pan error	Tilt Error	Pan Error
BISAR	7.3°	7.6°	4.5°
CNN	8.7°	11.5°	4.2°

(MLP) to estimate the pose. First, head is localized using a linear boundary decision classifier to detect skin color. Head area corresponds to the bounding box surrounding the biggest skin color connected component. Inputs of the MLP are the gray levels and gradient magnitudes of resized images (64pixels). Continuous outputs of the network are the estimated head angle values. In Gourier et al. [4], head poses are computed using Auto Associative Memories (one per orientation). Head pose is estimated by selecting the auto-associative network with highest likelihood score. Face was previously located using a skin color based method quite similar to [18]. The last method is proposed by Tu et al.[16]. The appearance variation caused by head pose changes is characterized by a tensor model. An image patch from the testing set is projected into the tensor subspace, and head pose can be estimated from the tensor coefficients obtained from a High Order Singular Value Decomposition (HOSVD).

Table 6: Results on tilt and pan estimation and comparison with state-of-the-art methods on Pointing04

Method	Tilt error	Classification	Pan error	Classification
Manual face localization				
BISAR	8.0°	65.4%	7.5°	60.6%
High-Order SVD [16]	8.6°	75.7%	6.2°	72.4%
Automatic face localization				
BISAR	12.0°	56.0%	10.2°	54.8%
High-order SVD [16]	17.9°	54.8%	12.9°	49.3%
Auto associative memories [4]	12.1°	53.8%	7.3°	61.3%
Multi layer perceptron [18]	12.8°	53.1%	12.3°	41.8%
Human performance [4]	9.4°	59.0%	11.8°	40.7%

Table 6 summarizes results obtained by the BISAR method and the three related approaches. Scores are obtained on the test set using the best regressor on the validation set. When faces are manually localized, BISAR results are the same on tilt estimation as the HOSVD method and less accurate on the pan estimation. But HOSVD results fall when the face is automatically cropped. This method is very sensitive to the localization accuracy. Auto associative memories outperform BISAR on pan estimation but the latter obtains a better score for tilt prediction. Moreover auto associative memories are used as generative classifier. This implies that this approach is only capable of estimating discrete pose locations and systematic problems arise as the number of detectors is increased [10].

In comparison with the multi layer perceptron, BISAR has better results on both tilt and pan estimation.

Results are also compared to human capability reported in [4]; BISAR and human performance are quite close.

6. Conclusion

In this paper, we have presented and compared a regression method by combining a new feature selection scheme with a specific RBF (Radial Basis Function) network named GRNN (Generalized Regression Neural Network). It has been shown that these tools are well adapted to problems with a very large set of potential features which is often the case in pattern recognition. Our feature selection is based on a new criterion called FFC (Fuzzy Functional Criterion). The FFC is a new measure using a fuzzy approach to evaluate the nature of the relation between two random variables x and y . If $y = F(x)$, the $FFC(y, x)$ is maximum. An important point is that our measure takes into account the fact that x and y are continuous (without sampling them) using a Parzen-window-like approach. Moreover, the FFC can be calculated once without weighting function over examples and then can be evaluated very quickly with any weighting provided later. We have selected a boosting strategy among three: memoryless, median and AdaBoost.R2 strategy. In these 3 strategies, weights are computed from the regression errors and from previous weights (except in the memoryless strategy). This selection and other benchmarking were done on the head pose determination problem. The strategy AdaBoost.R2 was found to be the best for three of the five test sets and was adopted for experiments. It also appears that GRNN is the best choice as the neural network regressor among the class of RBF networks. A comparison with CNN (Convolutional Neural Networks) has shown that BISAR is much better (Pointing Data Set) or quite equivalent (FacePix Data

Set). It is very promising since CNN are well known to be powerful regressor in Pattern Recognition community. Performances of BISAR were evaluated and found equivalent to three state-of-the-art methods on the Pointing Database. We believe that our error rate can be improved in the nearest future by incorporating new features such as gradient orientation to the potential set of features. We plan to analyze in details the relation between the boosting strategy, the cross validation and the RBF parameters. Experiments on other kinds of regression problems, for example with higher output dimension could help us to understand these issues. We think that the theory of learning could be an appropriate framework to develop and improve our approach.

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Appendix

$1 \leq k \leq N$	Number of features.
$1 \leq i \leq M$	Number of examples.
$1 \leq t \leq T$	Number of iterations.
b_t	Index of the t^{th} best feature.
A	Training set of labelled examples, $A = \{(\mathbf{x}_i, y_i) i = 1 \dots M\}$.
V	Validation set.
E	Test set.
F	Set of potential features, $F = \{H_k k = 1 \dots N\}$.
FS	Set of selected features.
\mathbf{x}_i	i^{th} example.
y_i	Target associated to \mathbf{x}_i .
H_k	k^{th} feature of F
$h_{k,i}$	Value of the k^{th} feature for the i^{th} example, $h_{k,i} = H_k(\mathbf{x}_i)$.
w_t	weighting vector at the t^{th} iteration, $w_t = \{w_t(i) i = 1, \dots, M\}$.
R_t	Regressor with t inputs.
$\epsilon_t(i)$	Absolute error of R_t on the i^{th} example.

Figure 6: Glossary of terms

Kevin Bailly is a PhD student at the Institute of Intelligent Systems and Robotics (ISIR), and a temporary assistant professor at Pierre and Marie Curie University (UPMC Paris 6), Paris, France. He holds an engineering degree from the IMAC's engineering school and a MSc in computer science from the University of Paris-Est. His main areas of research are pattern recognition, computer vision and machine learning.

Maurice Milgram is born in 1948 in Paris and obtained the Agrgation of Mathematic in 1971 and a PhD from UTC on Probabilistic Automata Networks in 1981. He starts as Assistant Professor at Technology University of Compigne (1973) then he becomes full Professor (1983) at ENSEA and joined the Robotic Laboratory of the Pierre and Marie Curie University in Paris in 1986. He founded the PARC Laboratory which is now part of the LISIF in 1992. He has supervised about 30 PhD thesis and 20 industrial contracts. His research interests concern Pattern Recognition and Image Processing.

Current work: Face localization , gaze tracking, person tracking in image sequences, gesture recognition.

Recent contracts: Person localization for a smart airbag (Faurecia), automatic directional control of the vehicles headlamp beams (Valeo), vehicle type recognition (LPR), biometry (Sagem), on-vehicle obstacle detection (PSA), baby gaze tracking (Necker Hospital).