Boosting Trees for Anti-Spam Email Filtering
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Abstract
This paper describes a set of comparative experiments for the problem of automatically filtering unwanted electronic mail messages. Several variants of the AdaBoost algorithm with confidence-rated predictions (Schapire & Singer 99) have been applied, which differ in the complexity of the base learners considered. Two main conclusions can be drawn from our experiments: a) The boosting-based methods clearly outperform the baseline learning algorithms (Naive Bayes and Induction of Decision Trees) on the PU1 corpus, achieving very high levels of the F1 measure; b) Increasing the complexity of the base learners allows to obtain better “high-precision” classifiers, which is a very important issue when misclassification costs are considered.

1 Introduction
Spam-mail filtering is the problem of automatically filtering unwanted electronic mail messages. The term “spam mail” is also commonly referred to as “junk mail” or “unsolicited commercial mail”. Nowadays, the problem has achieved a big impact since bulk emailers take advantage of the great popularity of the electronic mail communication channel for indiscriminately flooding email accounts with unwanted advertisements. The major factors that contribute to the proliferation of unsolicited spam email are the following (two): 1) bulk email is inexpensive to send, and 2) pseudonyms are inexpensive to obtain (Cranor & LaMacchia 98). On the contrary, individuals may waste a large amount of time transferring unwanted messages to their computers and sorting through those messages once transferred, to the point that they may be likely to become overwhelmed by spam.

Automatic IR methods are well suited for addressing this problem, since spam messages can be distinguished from the “legitimate” email messages because of their particular form, vocabulary and word patterns, which can be found in the header or body of the messages.

The spam filtering problem can be seen as a particular instance of the Text Categorization problem (TC), in which only two classes are possible: spam and legitimate. However, since one is the opposite of the other, it also can be seen as the problem of identifying a single class, spam. In this way, the evaluation of automatic spam filtering systems can be done by using common measures of IR (precision, recall, etc.). Another important issue is the relative importance between the two types of possible misclassifications: While an automated filter that misses a small percentage of spam may be acceptable to most users, fewer people are likely to accept a filter that incorrectly identifies a small percentage of legitimate mail as spam, especially if this implies the automatic discarding of the misclassified legitimate messages. This problem suggests the consideration of misclassification costs for the learning and evaluation of spam filter systems.

In recent years, a vast amount of techniques have been applied to TC, achieving impressive performances in some cases. Some of the top-performing methods are Ensembles of Decision Trees (Weiss et al. 99), Support Vector Machines (Joachims 98), Boosting (Schapire & Singer 00) and Instance-based Learning (Yang & Liu 99). Spam filtering has also been treated as a particular case of TC. In Cohen 96 a method based on TF-IDF weighting and the rule learning algorithm RIPPER is used to classify and filter email. Sahami et al. 98 used the Naive Bayes approach to filter spam email. Drucker et al. 99 compared Support Vector Machines (SVM), boosting of C4.5 trees, RIPPER and Rocchio, concluding that SVM’s and boosting are the top-performing methods and suggesting that SVM’s are slightly better in distinguishing the two types of missclassification. In Androutsopoulos et al. 00b Sahami’s Naive Bayes is compared against the TiMBL Memory-based learner. In Androutsopoulos et al. 00b the same authors present a new public data set which might become a standard benchmark corpus, and introduce cost-sensitive evaluation measures.

In this paper, we show that the AdaBoost algorithm with confidence-rated predictions is a very well suited algorithm for addressing the spam filtering problem. We have obtained very accurate classifiers on the PU1 corpus, and we have observed that the algorithm is very robust to overfitting. Another advantage of using AdaBoost is that no prior feature filtering is needed since it is able to efficiently manage large feature sets (of tens of thousands).

In the second part of the paper we show how increasing the expressiveness of the base learners can be exploited for obtaining the “high-precision” filters that are needed for real user applications. We have evaluated the results of such filters using the measures...
introduced in [Androutsopoulos et al. 00b], which take into account the misclassification costs. We have substantially improved the results mentioned in that work.

The paper is organized as follows: Section 2 is devoted to explain the AdaBoost learning algorithm and the variants used in the comparative experiments. In section 3 the setting is presented in detail, including the corpus and the experimental methodology used. Section 4 reports the experiments carried out and the results obtained. Finally, section 5 concludes and outlines some lines for further research.

2 The AdaBoost Algorithm

In this section the AdaBoost algorithm (Schapire & Singer 99) is described, restricting to the case of binary classification.

The purpose of boosting is to find a highly accurate classification rule by combining many weak rules (or weak hypotheses), each of which may be only moderately accurate. It is assumed the existence of a separate procedure called the WeakLearner for acquiring the weak hypotheses. The boosting algorithm finds a set of weak hypotheses by calling the weak learner repeatedly in a series of $T$ rounds. These weak hypotheses are then linearly combined into a single rule called the combined hypothesis.

Let $S = \{(x_1, y_1), \ldots, (x_m, y_m)\}$ be the set of $m$ training examples, where each instance $x_i$ belongs to a instance space $\mathcal{X}$ and $y_i \in \{-1, +1\}$ is the class or label associated to $x_i$. The goal of the learning is to produce a function of the form $f: \mathcal{X} \to \mathbb{R}$, such that, for any example $x$, the sign of $f(x)$ is interpreted as the predicted class $(-1$ or $+1)$, and the magnitude $|f(x)|$ is interpreted as a measure of confidence in the prediction. Such a function can be used either for classifying or ranking new unseen examples.

The pseudo-code of AdaBoost is presented in Figure 1. It maintains a vector of weights as a distribution over examples. The goal of the WeakLearner algorithm is to find a weak hypothesis with moderately low error with respect to these weights. Initially, the distribution $D_1$ is uniform, but the boosting algorithm exponentially updates the weights on each round to force the weak learner to concentrate on the examples which are hardest to predict by the preceding hypotheses.

More precisely, let $D_t$ be the distribution at round $t$, and $h_t: \mathcal{X} \to \mathbb{R}$ the weak rule acquired according to $D_t$. In this setting, weak hypotheses $h_t(x)$ also make real-valued confidence-rated predictions (i.e., the sign of $h_t(x)$ is the predicted class, and $|h_t(x)|$ is interpreted as a measure of confidence in the prediction). A parameter $\alpha_t$ is then chosen and the distribution $D_t$ is updated. The choice of $\alpha_t$ will be determined by the type of weak learner (see next section). In the typical case that $\alpha_t$ is positive, the updating function decreases (or increases) the weights $D_t(i)$ for which $h_t$ makes a good (or bad) prediction, and this variation is proportional to the confidence $|h_t(x_i)|$. The final hypothesis, $f$, computes its predictions using a weighted vote of the weak hypotheses.

In (Schapire & Singer 99), it is proven that the training error of the AdaBoost algorithm on the training set (i.e. the fraction of training examples $i$ for which the sign of $f(x_i)$ differs from $y_i$) is at most $\prod_{t=1}^{T} Z_t$, where $Z_t$ is the normalization factor computed on round $t$. This upper bound is used in guiding the design of both the parameter $\alpha_t$ and the WeakLearner algorithm, which attempts to find a weak hypothesis $h_t$ that minimizes $Z_t$.

2.1 Learning weak rules

In (Schapire & Singer 99) three different variants of AdaBoost MH are defined, corresponding to three different methods for choosing the $\alpha_t$ values and calculating the predictions of the weak hypotheses. In this work we concentrate on AdaBoost with real-valued predictions since it is the one that has achieved the best results in the Text Categorization domain (Schapire & Singer 00).

According to this setting, weak hypotheses are simple rules with real-valued predictions. Such simple rules test the value of a boolean predicate and make a prediction based on that value. The predicates used refer to the presence of a certain word in the text, e.g. “the word money appears in the message”. Formally, based on a given predicate $p$, our interest lies on weak hypotheses $h$ which make predictions of the form:

$$h(x) = \begin{cases} 
  c_0 & \text{if } p \text{ holds in } x \\
  c_1 & \text{otherwise}
\end{cases}$$

where the $c_0$ and $c_1$ are real numbers.
For a given predicate \( p \), the values \( c_0 \) and \( c_1 \) are calculated as follows. Let \( X_1 \) be the subset of examples for which the predicate \( p \) holds and let \( X_0 \) be the subset of examples for which the predicate \( p \) does not hold. Let \( \pi \), for any predicate \( \pi \), be 1 if \( \pi \) holds and 0 otherwise. Given the current distribution \( D_t \), the following real numbers are calculated for \( j \in \{0,1\} \), and for \( b \in \{ +1, -1 \} \):

\[
W_b^j = \sum_{i=1}^{m} D_t(i) [x_i \in X_j \land y_i = b].
\]

That is, \( W_b^j \) is the weight, with respect to the distribution \( D_t \), of the training examples in partition \( X_j \) which are of class \( b \). As it is shown in (Schapire & Singer 99) \( Z_t \) is minimized for a particular predicate by choosing:

\[
c_j = \frac{1}{2} \ln \left( \frac{W_{j+1}^j}{W_{j-1}^j} \right).
\]

and by setting \( \alpha_t = 1 \). These settings imply that:

\[
Z_t = 2 \sum_{j \in \{0,1\}} \sqrt{W_{j+1}^j W_{j-1}^j}.
\]

Thus, the predicate \( p \) chosen is that for which the value of \( Z_t \) is smallest.

Very small or zero values for the parameters \( W_b^j \) cause \( c_j \) predictions to be large or infinite in magnitude. In practice, such large predictions may cause numerical problems to the algorithm, and seem to increase the tendency to overfit. As suggested in (Schapire & Singer 00), the smoothed values for \( c_j \) have been considered.

It is important to see that the so far presented weak rules can be directly seen as decision trees with a single internal node (which tests the value of a boolean predicate) and two leaf nodes that give the real-valued predictions for the two possible outcomes of the test. These extremely simple decision trees are sometimes called decision stumps. In turn, the boolean predicates can be seen as binary features (we will use the word feature instead of predicate from now on), thus, the already described criterion for finding the best weak rule (or the best feature) can be seen as a natural splitting criterion and used for performing decision–tree induction (Schapire & Singer 99).

Following the idea suggested in (Schapire & Singer 99), we have extended the WeakLearner algorithm to induce arbitrarily deep decision trees. The splitting criterion used consists in choosing the feature that minimizes equation (2), while the predictions at the leaves of the boosted trees are given by equation (1). Note that the general AdaBoost procedure remains unchanged.

In this paper, we will denote as TreeBoost the AdaBoost.MH algorithm including the extended WeakLearner. TreeBoost[d] will stand for a learned classifier with weak rules of depth \( d \). As a special case, TreeBoost[0] will be denoted as Stumps.

3 Setting

3.1 Domain of Application

We have evaluated our system on the PU1 benchmark corpus\(^4\) for the anti-spam email filtering problem. It consists of 1,099 messages: 481 of them are spam and the remaining 618 are legitimate. The corpus is presented partitioned into 10 folds of the same size which maintain the distribution of spam messages (Androutsopoulos et al. 00). All our experiments have been performed using 10-fold cross-validation.

The feature set of the corpus is a bag-of-words model. Four versions are available: with or without stemming, and with or without stop-word removal. The experiments reported in this paper have been performed using the non-stemming non-stop-word-removal version, which consists in a set of 26,449 features.

3.2 Experimental Methodology

Evaluation Measures. Measures for evaluating the spam filtering system are introduced here. Let \( S \) and \( L \) be the number of spam and legitimate messages in the corpus, respectively; let \( S_+ \) denote the number of spam messages that are correctly classified by a system, and \( S_- \) the number of spam messages misclassified as legitimate. In the same way, let \( L_+ \) and \( L_- \) be the number of legitimate messages classified by a system as spam and legitimate, respectively. These four values form a contingency table which summarizes the behaviour of a system. The widely-used measures precision (\( p \)), recall (\( r \)) and \( F_\beta \) are defined as follows:

\[
p = \frac{S_+}{S_+ + L_+}, \quad r = \frac{S_+}{S_+ + S_-}, \quad F_\beta = \frac{(\beta^2 + 1)pr}{\beta^2p + r}.
\]

The \( F_\beta \) measure combines precision and recall, and with \( \beta = 1 \) gives equal weight to the combined measures. Additionally, some experiments in the paper will also consider the accuracy measure (\( \text{acc} = \frac{L_+ + S_+}{L_+ + S_+ + L_- + S_-} \)).

A way to distinguish the two types of misclassification is the use of utility measures (Lewis 95) used in the TREC evaluations (Robertson & Hull 01). In this general measure, positions in the contingency table are associated loss values, \( \lambda_{S+}, \lambda_{S-}, \lambda_{L+}, \lambda_{L-} \), which indicate how desirable are the outcomes, according to a user-defined scenario. The overall performance of a system in terms of the utility is \( S_+\lambda_{S+} + S_-\lambda_{S-} + L_+\lambda_{L+} + L_-\lambda_{L-} \).

Androutsopoulos et al. (Androutsopoulos et al. 00) propose particular scenarios in which misclassifying a legitimate message as spam is \( \lambda \) times more costly than the symmetric misclassification. In terms of utility, these scenarios can be translated to \( \lambda_{S+} = 0, \lambda_{S-} = -1, \lambda_{L+} = -\lambda, \lambda_{L-} = 0 \). They also introduce the weighted accuracy (WAcc) measure, a version

\(^4\)PU1 Corpus is freely available from the publications section of [http://www.iit.demokritos.gr/~ionandr](http://www.iit.demokritos.gr/~ionandr).
of accuracy sensitive to \( \lambda \)-cost:

\[
W_{\text{Acc}} = \frac{\lambda \cdot L_+ + S_-}{\lambda \cdot L + S}
\]

When evaluating filtering systems, this measure suffers from the same problems as standard accuracy [Yang 99]. Despite this fact, we will use it for comparison purposes.

### 3.3 Baseline Algorithms

In order to compare our boosting methods against other techniques, we include the following two baseline measures:

- Decision Trees. Standard TDIDT learning algorithm, using the RLM distance-based function for the feature selection. See [Marquez 99] for complete details about the particular implementation.
- Naive Bayes. We include the best results on the PU1 Corpus reported in [Androutsopoulos et al. 00], corresponding to a Naive Bayes classifier.

### 4 Experiments

This section explains the set of experiments carried out. As said in section 3, all experiments work with the PU1 Corpus.

#### 4.1 Comparing methods on the corpus

The purpose of our first experiment is to show the general performance of boosting methods in the spamfiltering domain. Six AdaBoost classifiers have been learned, setting the depth of the weak rules from 0 to 5; we denote each classifier as TreeBoost\([d]\), where \(d\) stands for the depth of the weak rules; as a particular case, we denote the TreeBoost\([0]\) classifier as Stumps. Each version of TreeBoost has been learned for up to 2,500 weak rules.

Figure 2 shows the \( F_1 \) measure of each classifier, as a function of the number of rounds used. In this plot, there are also the obtained rates of the baseline algorithms. It can be seen that TreeBoost clearly outperforms the baseline algorithms. The experiment also shows that, above a certain number of rounds, all TreeBoost versions achieve consistent good results, and that there is no overfitting in the process. After 150 rounds of boosting, all versions reach an \( F_1 \) value above 97%. It can be noticed that the deeper the weak rules, the smaller the number of rounds needed to achieve good performance. This is not surprising, since deeper weak rules handle much more information. Additionally, the figure shows that different number of rounds produce slight variations in the error rate.

A concrete value for the \( T \) parameter of the TreeBoost learning algorithm must be given, in order to obtain real classifiers and to be able to make comparisons between the different versions of TreeBoost and baseline methods. To our knowledge, it is still unclear what is the best way for choosing \( T \). We have estimated the \( T \) parameter in a validation set built for the task, with the following procedure: a) For each trial in the cross-validation experiment, 8 of the 9 training subsets are used to learn up to 2,500 rounds of boosting, and the one remaining is used as the validation set for testing the classifier with respect to the number of rounds, in steps of 25. b) The outputs of all classifiers are used to compute the \( F_1 \) measure. c) The minimum \( T \) for which the \( F_1 \) is maximum is chosen as the estimated optimal number of rounds for all classifiers.

Table 1 presents the results of all classifiers. For each one, we include the number of rounds (estimated in the validation set), recall, precision, \( F_1 \) and the maximum \( F_1 \) achieved over the 2,500 rounds learned. According to the results, boosting classifiers clearly outperform the other algorithms. Only Naive Bayes achieves a precision (95.11%) slightly lower than the obtained by boosting classifiers (the worse is 97.48%); however, its recall at this point is much lower.

![Figure 2: \( F_1 \) measure of Stumps and TreeBoost\([d]\), for increasing number of rounds](image)

**Table 1: Performance of all classifiers**

|       | \( T \) | Recall | Prec. | \( F_1 \) | \( F_{1_{\text{max}}} \) |
|-------|--------|--------|-------|-----------|-----------------|
| N. Bayes | -      | 83.98  | 95.11 | 89.19     | -               |
| D. Trees | -      | 89.81  | 88.71 | 89.25     | -               |
| Stumps  | 525    | 96.47  | 97.48 | 96.97     | 97.39           |
| TreeBoost\([1]\) | 525    | 96.88  | 97.90 | 97.39     | 97.60           |
| TreeBoost\([2]\) | 725    | 96.67  | 98.31 | 97.48     | 97.59           |
| TreeBoost\([3]\) | 675    | 96.88  | 97.90 | 97.39     | 97.81           |
| TreeBoost\([4]\) | 450    | 97.09  | 98.73 | 97.90     | 98.01           |
| TreeBoost\([5]\) | 550    | 96.88  | 98.52 | 97.69     | 98.12           |

Accuracy results have been compared using the 10-fold cross-validated paired \( t \) test. Boosting classifiers perform significantly better than Decision Trees. On the contrary, no significant differences can be observed since we do not own the Naive Bayes classifiers, no tests have been ran; but presumably boosting methods are also significantly better.
between the different versions of TreeBoost. More interestingly, it can be noticed that accuracy and precision rates slightly increase with the expressiveness of the weak rules, and that this improvement does not affect the recall rate. This fact will be exploited in the following experiments.

4.2 High-Precision classifiers

This section is devoted to evaluate TreeBoost in high-precision scenarios, where only a very low (or null) proportion of legitimate to spam misclassifications is allowed.

Rejection Curves. We start by evaluating if the confidence of a prediction, i.e., the magnitude of the prediction, is a good indicator of the quality of the prediction. For this purpose, rejection curves are computed for each classifier. The procedure to compute a rejection curve is the following: For several points \( p \) between 0 and 100, reject the \( p\% \) of the predictions whose confidences score lowest, both positive or negative, and compute the accuracy of the remaining \((100 - p)\%\) predictions. This results in higher accuracy values as long as \( p \) increases. Figure 3 plots the rejection curves computed for the six learned classifiers. The following conclusions can be drawn:

- The confidence of a prediction is a good indicator of its quality.
- Depth of weak rules greatly improves the quality of the predictions. Whereas Stumps needs to reject the 73% of the less confident examples to achieve a 100% of accuracy, TreeBoost[5] only needs 23%. In other words, deeper TreeBoost filters concentrate the misclassified examples closer to the decision threshold.
- The previous fact has important consequences for a potential final email filtering application, with the following specification: Messages whose prediction confidence is greater than a threshold \( \tau \) are automatically classified: spam messages are blocked and legitimate messages are delivered to the user. Messages whose prediction confidence is lower than \( \tau \) are stored in a special fold for dubious messages. The user has to verify if these are legitimate messages. This specification is suitable for having automatic filters with different degrees of strictness (i.e., different values for the \( \tau \) parameter). \( \tau \) values could be tuned using a validation set.

Cost–Sensitive Evaluation. In this section, TreeBoost classifiers are evaluated using the \( \lambda \)-cost matrices introduced in section 3. Three scenarios of strictness are presented in (Androutsopoulos et al. 00b): a) No cost considered, corresponding to \( \lambda = 1 \); b) Semi-automatic scenario, for a moderately accurate filter, giving \( \lambda = 9 \); and c) Completely automatic scenario, for a very high accurate filter, assigning \( \lambda = 999 \). As noted in section 3, we will consider these scenarios as particular utility matrices.

In [Schapire et al. 98] a modification of the AdaBoost algorithm for handling general utility matrices is presented. The idea is to initialize the weight distribution of examples according to the given utility matrix, and then run the learning algorithm as usual. We have performed experiments with this setting, but the results are not convincing: only the initial rounds of boosting are affected by the initialization based on utility; after a number of rounds, the performance seems to be like if no utility had been considered. Since our procedure for tuning the number of rounds can not determine when the initial stage ends, we have rejected this approach. We think that the modification of the AdaBoost algorithm should also consider the weight update.

Another approach consists in adjusting the decision threshold \( \theta \). In a default scenario, corresponding to \( \lambda = 1 \), an example is classified as spam if its prediction is greater than 0; in this case, \( \theta = 0 \). Increasing the value of \( \theta \) results in a higher precision classifier. (Lewis 97) presented a procedure for calculating the optimal decision threshold for a system, given an arbitrary utility matrix. The procedure is valid only when the system outputs probabilities, so the predic-
tion scores resulting from the boosting classifications should be mapped to probabilities. A method for estimating probabilities given the output of AdaBoost is suggested in [Friedman et al.], using a logistic function. Initial experiments with this function have not worked properly, because relatively low predictions are sent to extreme probability values. A possible solution would be to scale down the predictions before applying the probability estimate; however, it can be observed that prediction scores grow with both the number and the depth of the used weak rules. Since many parameters are involved in this scaling, we have rejected the probability estimation of predictions.

Alternatively, we make our classification scheme sensitive to $\lambda$ factor by tuning the $\theta$ parameter to the value which maximizes the weighted accuracy measure. Once more, the concrete value for $\theta$ is obtained using a validation set, in which several values for the parameter are tested. Table 2 summarizes the results obtained from such procedures, giving $\lambda$ factor values of 9 and 999. Results obtained in [Androutsopoulos et al. 10] are also reported.

Again, TreeBoost clearly outperforms the baseline methods. With $\lambda = 9$, very high-precision rates are achieved, maintaining considerably high recall rates; it seems that the depth of TreeBoost slightly improves the performance, although no significant differences can be achieved. For $\lambda = 999$, precision rates of 100% (which is the implicit goal in this scenario) are achieved, except for Stumps, maintaining fair levels of recall. However, recall rates are slightly unstable with respect to the depth of TreeBoost varying from 64.45% to 76.30%. Our impression is that high values in the $\lambda$ factor seem to introduce instability in the evaluation, which becomes oversensitive to outliers. In this particular corpus (which contains 1,099 examples), weighted accuracy does not seem to work properly when giving $\lambda$ values of 999, since the misclassification of only one legitimate message leads to score worse than if any email had been filtered (this would give $W\text{Acc} = 99.92\%$). Moreover, for 100% precision values, the recall variation from 0% to 100% only affects the measure in 0.08 units.

In order to give a clearer picture of the behaviour of classifiers when moving the decision threshold, we include in Figure 4 the precision-recall curves of each classifier. These curves are built giving $\theta$ a wide range of values, and computing for each value the recall and precision rates. In these curves, high-precision rates of 100%, 99%, 98% and 95% have been fixed so as to obtain the recall rate at these points. Table 3 summarizes these samples. All the variants are indistinguishable at level of 95% of precision. However, when moving to higher values of precision ($\geq 95\%$) a significant difference seems to occur between Stumps and the rest of variants using deeper weak rules. This fact proves that increasing the expressiveness of the weak rules can improve the performance when requiring very high precision filters. Unfortunately, no clear conclusions can be drawn about the most appropriate depth. Parenthetically, it can be noted that TreeBoost[4] achieves the best recall rates in this particular corpus.

| Method       | 100%  | 99%   | 98%   | 95%   |
|--------------|-------|-------|-------|-------|
| Stumps       | 62.37 | 87.94 | 94.17 | 98.75 |
| TreeBoost[1] | 81.91 | 91.26 | 96.88 | 98.75 |
| TreeBoost[2] | 81.49 | 90.64 | 97.08 | 98.54 |
| TreeBoost[3] | 77.54 | 93.13 | 96.88 | 98.54 |
| TreeBoost[4] | 80.24 | 96.25 | 97.71 | 98.75 |
| TreeBoost[5] | 77.75 | 93.55 | 97.29 | 98.75 |

Table 3: Recall rate of filtered spam messages with respect to fixed points of precision rate

![Figure 4: Precision-Recall curves and recall values for the fixed precision rates at 100%, 99%, 98% and 95%. x axis: recall; y axis: precision.](image)

5 Conclusions

The presented experiments show that AdaBoost learning algorithm clearly outperforms Decision Trees and Naive Bayes methods on the public benchmark PU1 Corpus. In this data set, the method is resistant to overfitting and $F_1$ rates above 97% are achieved. Procedures for automatically tune the classifier parameters, such as the number of boosting rounds, are provided.
In scenarios where high-precision classifiers are required, AdaBoost classifiers have been proved to work properly. Experiments have exploited the expressiveness of the weak rules when increasing their depth. It can be concluded that deeper weak rules tend to be more suitable when looking for a very high precision classifier. In this situation, the achieved results on the PU1 Corpus are fairly satisfactory.

Two AdaBoost classifiers capabilities have been shown to be useful in final email filtering applications: a) The confidence of the predictions suggests a filter which only blocks the more confident messages, delivering the remaining messages to the final user. b) The classification threshold can be tuned to obtain a very high precision classifier.

As a future research line, we would like to study the presented techniques in a larger corpus. We think the PU1 corpus is too small and also too easy: default parameters produce very good results, and the tuning procedures result only in slight improvements. Moreover, some experiments not reported here (which study the effect of the number of rounds, the use of richer feature spaces, etc.) have shown us that the confidence of classifiers depends on several parameters. Using a larger corpus, the effectiveness of the tuning procedures would be more explicit and, hopefully, clear conclusions about the optimal parameter settings of AdaBoost could be drawn.

Another line for future research is the introduction of misclassification costs inside the AdaBoost learning algorithm. Initial experiments with the method proposed in [Schapire et al. 98] have not worked properly, although we believe that learning directly classifiers according to some utility settings will perform better than tuning a classifier once learned.

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Table 2: Cost–sensitive evaluation results

| Method      | $\lambda = 9$ | $\lambda = 999$ |
|-------------|---------------|-----------------|
|             | $\theta$     | Recall | Prec. | WAcc  | $\theta$     | Recall | Prec. | WAcc  |
| Naive Bayes | -             | 78.77  | 96.65 | 96.38 | -             | 46.96  | 98.80 | 99.47 |
| Stumps      | 9.162         | 89.60  | 99.08 | 98.58 | 16.298        | 84.20  | 99.51 | 99.66 |
| TreeBoost[1]| 10.2          | 93.55  | 98.74 | 98.59 | 46.87         | 74.43  | 100   | 99.98 |
| TreeBoost[2]| 24.156        | 93.76  | 98.90 | 98.76 | 96.717        | 76.30  | 100   | 99.98 |
| TreeBoost[3]| 45.165        | 91.48  | 99.32 | 98.87 | 126.388       | 74.01  | 100   | 99.98 |
| TreeBoost[4]| 19.063        | 94.80  | 99.35 | 99.14 | 123.097       | 64.45  | 100   | 99.97 |
| TreeBoost[5]| 37.379        | 93.97  | 99.12 | 98.92 | 177.909       | 66.53  | 100   | 99.97 |

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