

Learning to Predict One or More Ranks in Ordinal Regression Tasks

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Abstract. We present nondeterministic hypotheses learned from an ordinal regression task. They try to predict the true rank for an entry, but when the classification is uncertain the hypotheses predict a set of consecutive ranks (an interval). The aim is to keep the set of ranks as small as possible, while still containing the true rank. The justification for learning such a hypothesis is based on a real world problem arisen in breeding beef cattle. After defining a family of loss functions inspired in Information Retrieval, we derive an algorithm for minimizing them. The algorithm is based on posterior probabilities of ranks given an entry. A couple of implementations are compared: one based on a multiclass *SVM* and other based on Gaussian processes designed to minimize the linear loss in ordinal regression tasks.

1 Introduction

In the last few years, ordinal regression has become an important issue in Machine Learning research. See [1] and [2] for a state of the art introduction. The aim of ordinal regression is to find hypotheses able to predict classes or *ranks* that belong to a finite ordered set. Applications include Information Retrieval [3], Natural Language Processing [4], collaborative filtering [5], finances [6], and user preferences [7].

The approach presented in this paper explores a new kind of predictions in ordinal regression. We shall build hypotheses that try to predict the true rank for an entry, but when the classification is uncertain the hypotheses predict an interval of ranks. The aim is to return a set of consecutive ranks, such that the set is as small as possible, while still containing the true rank. As we shall learn hypotheses for ordinal regression tasks with multiple outcomes, like nondeterministic automata, we shall call them *nondeterministic ordinal regressors*. From another point of view, these hypothesis could be called set-valued predictors.

Predictors of more than one class are not completely new. Given an error ϵ , the so called confidence machines, make *conformal predictions* [8]: they produce a set of labels containing the true class with probability greater than $1 - \epsilon$. Other approaches arose in the context of hierarchical organization of biological objects: predicting gene functions [9], or mapping biological entities into ontologies [10].

In the next Section we shall show the usefulness of these nondeterministic hypotheses in a real world application context: the assessment of muscle proportion in carcasses of beef cattle. This is an important question in cattle breeding since this proportion determines, on the one hand, the prices to be obtained by carcasses, and on the other hand, the genetic value of animals to select studs for the next generation.

We formalize the problem of nondeterministic predictions as a special kind of *Information Retrieval*. Thus, we define a family of loss functions F_β and derive an algorithm for minimizing this loss. The algorithm needs the estimation of posterior probabilities of ranks given the entries. Then, we compare a couple of implementations built on the estimations provided by a *SVM* [11], and by the Gaussian approach of [1] devised for ordinal regression tasks.

The last Section of the paper presents an exhaustive set of experiments carried out in order to test the performance of the nondeterministic approach. Thus, in addition to the beef cattle learning task, we shall use 24 datasets publicly available that were previously used in ordinal regression tasks [1,12].

2 The Round Profile of Bovines

The problem that motivated the research reported in this paper arose when we were trying to make reliable predictions of the value of the carcasses of beef cattle. This learning task was proposed by ASEAVA, the Association of Breeders of a beef breed of the North of Spain, *Asturiana de los Valles*. This is a specialized breed with many double-muscled individuals; their carcass have dressing percentages over 60%, with muscle content over 75%, and with a low (8%) percentage of fat [13]. The market target of these carcasses is made up of those consumers that prefer lean meat without any marbling [7,14,15].

Even if the animals are not going to be slaughtered, the prediction of carcass value of a beef cattle is interesting since it can be considered as a kind of assessment that is useful for breeders to select the progenitors of the next generation. Thus, the ICAR (International Committee for Animal Recording) acknowledges as a good practice the recording of live animal assessments; since these assessments are a description of an animal's morphology that reveals part of its economic value.

The records so obtained can be used for the evaluation of programs of genetic selection of dairy, dual purpose and specialized beef breeds. The growth of the scores over years of selection for specific goals can be seen as a measure of the success of the selection policy. On the other hand, when the assessed traits are heritable, the scores can be directly used for selection purposes given that they are capturing part of animal's genetic value.

Traditionally, the assessment procedures were based on *visual* appreciations of well trained technicians that had to rank a number of morphological characteristics that include linear lengths of significant parts of animals' bodies. Although this process has been used successfully, it is clear that there is a problem with the repeatability of the assessments; not only between assessors, but



Fig. 1. The assessment of the round profile of a beef cattle is a measurement of the roundness of the lines drawn in the pictures. Thus, the leftmost cow in the top row is a paradigm of the animals which have rank 1, while the following are representative examples of ranks 2, 3 and 4 respectively.

even within assessors scoring the same animal in different times. In order to overcome these difficulties, we developed a new assessment method described in [16] that is almost completely repeatable and can be carried out using just 3 lengths (in centimeters) plus the appreciation of the curvature of the round profile (see the curves in Figure 1). For this learning task we used a kernel based method described in [17].

The aim of the assessment of round profiles is to rank the muscularity of animals. Therefore, it is a very important attribute for describing beef cattle. However, the curvature of the round profile is assessed by *visual* appreciations of experts. But visual appreciations is a source of problems. Thus, for instance, in [18], the authors describe an experiment in which a set of expert graders were asked to rank a collection of mushrooms into three major and eight subclasses of commercial quality. Grader consistency was assessed by repeated classification (four repetitions) of two 100-mushroom sets. Grader repeatability ranged from 6% to 15% misclassification.

Therefore, returning to beef cattle, to ensure the repeatability of the whole process, we should skip the subjective appreciation of the rank of round profiles. Thus, a new learning task arises: to estimate this rank from repeatable live animal descriptions. For this purpose, we built a dataset with 891 pairs of animal descriptions (6 lengths in centimeters of their bodies, live weight, and sex) and ranks (in a scale of 1-4). To ensure a uniform criterion, the first author of this paper measured and ranked the round profile of the 891 live animals.

But this is a difficult learning task. The classification accuracy achieved by a multiclass SVM was 77%; the implementation used was a probabilistic *libsvm* [11]. These results are not improved when using a learner specially devised for ordinal regression tasks. Thus, using the MAP approach of [1], the accuracy was 76%. The details about how we obtained these scores are included in the last section devoted to report a number of experimental results. On the other hand, we shall prove that a nondeterministic hypothesis contains the true class more than 84% of cases, while the average number of ranks predicted is just 1.21 or 1.30 depending of the learner used.

Moreover, the nondeterministic approach is more useful than the plain deterministic one in this problem for several reasons. First, the reliability of hypothesis predictions is higher than in the deterministic case. Therefore, when the hypothesis predicts only one rank, the estimation of the rank is very probably the true one. Second, when the prediction is an interval of more than one rank, we can appeal to a more expensive procedure to finally decide the true class. In this case, we may turn to an *actual* expert, or we can wait until the natural growth of the animal make the classification more clear.

However, sometimes even a nondeterministic prediction may be useful to discard an animal as stud for the next generation: a prediction of $[1, 2]$ must imply a poor genetic value as meat producer, provided that the hypothesis is sufficiently reliable.

3 Formal Framework

Let \mathcal{X} be an input space, and \mathcal{Y} a finite set of ordered ranks. Without any loss of generality, we can assume that $\mathcal{Y} = \{1, \dots, k\}$ for a given k . We shall consider an ordinal regression task given by a training set $S = \{(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_n, y_n)\}$ drawn from an unknown distribution $Pr(X, Y)$ from the product $\mathcal{X} \times \mathcal{Y}$. Within this context, we propose the following.

Definition 1. A nondeterministic hypothesis is a function h from the input space to the set of non-empty intervals (subsets of consecutive ranks) of \mathcal{Y} ; in symbols,

$$h : \mathcal{X} \longrightarrow \text{Intervals}(\mathcal{Y}). \quad (1)$$

The aim of such a learning task is to find a nondeterministic hypothesis h from a space \mathcal{H} that optimizes the *expected prediction performance (or risk)* on samples S' independently and identically distributed (i.i.d.) according to the distribution $Pr(X, Y)$:

$$R^\Delta(h) = \int \Delta(h(\mathbf{x}), y) d(Pr(\mathbf{x}, y)), \quad (2)$$

where $\Delta(h(\mathbf{x}), y)$ is a loss function that measures the penalty due to the prediction $h(\mathbf{x})$ when the true value is y .

In nondeterministic ordinal regression, we would like to favor those decisions of h that contain the true ranks, and a smaller rather than a larger number of

ranks. In other words, we interpret the output $h(\mathbf{x})$ as an imprecise answer to a query about the right rank of an entry $\mathbf{x} \in \mathcal{X}$. Thus, the nondeterministic ordinal regression can be seen as a kind of *Information Retrieval* task for each entry.

In Information Retrieval, performance is compared using different measures in order to consider different perspectives. The most frequently used are the *Recall* (proportion of all relevant documents that are found by a search) and *Precision* (proportion of retrieved documents that are relevant). The harmonic average of the two amounts is used to capture the goodness of a hypothesis in a single measure. In the weighted case, the measure is called F_β . The idea is to measure a tradeoff between *Recall* and *Precision*.

For further references, let us recall the formal definitions of these Information Retrieval measures. Thus, for a prediction of a nondeterministic hypothesis $h(\mathbf{x})$ with $\mathbf{x} \in \mathcal{X}$, and a rank $y \in \mathcal{Y}$, we can compute the following contingency matrix, where $z \in \mathcal{Y}$,

$$\begin{array}{c|cc} & y = z & y \neq z \\ \hline z \in h(\mathbf{x}) & a & b \\ z \notin h(\mathbf{x}) & c & d \\ \hline \end{array} \quad (3)$$

where each entry (a, b, c, d) is the number of times that happens the corresponding combination of memberships. Thus, notice that a can only be 1 or 0, depending on whether the rank y is in the prediction $h(\mathbf{x})$ or not; b is the number of ranks different from y included in $h(\mathbf{x})$; $c = 1 - a$; and d is the number of ranks different from y that are not in $h(\mathbf{x})$.

According to the matrix (Eq. 3), if h is a nondeterministic hypothesis, and $(\mathbf{x}, y) \in \mathcal{X} \times \mathcal{Y}$, we have the next definitions.

Definition 2. *The Recall in a query (i.e. an entry \mathbf{x}) is defined as the proportion of relevant ranks (y) included in $h(\mathbf{x})$:*

$$R(h(\mathbf{x}), y) = \frac{a}{a + c} = a = 1_{y \in h(\mathbf{x})}. \quad (4)$$

Definition 3. *The Precision is defined as the proportion of retrieved ranks in $h(\mathbf{x})$ that are relevant (y):*

$$P(h(\mathbf{x}), y) = \frac{a}{a + b} = \frac{1_{y \in h(\mathbf{x})}}{|h(\mathbf{x})|}. \quad (5)$$

In other words, given an hypothesis h , the *Precision* for an entry \mathbf{x} , that is $P(h(\mathbf{x}), y)$, is the probability of finding the true rank (y) of the entry (\mathbf{x}) by randomly choosing one of the ranks of $h(\mathbf{x})$.

Finally, the tradeoff is formalized by

Definition 4. *The F_β , in general is defined by*

$$F_\beta(h(\mathbf{x}), y) = \frac{(1 + \beta^2)a}{(1 + \beta^2)a + b + \beta^2c}. \quad (6)$$

Table 1. For an entry \mathbf{x} with rank 1, ($y = 1$), *Precision*, *Recall*, F_1 , and F_2 for different predictions of a nondeterministic classifier h

$h(\mathbf{x})$	<i>Precision</i>	<i>Recall</i>	F_1	F_2
[1, 2, 3]	0.33	1	0.50	0.71
[1, 2]	0.50	1	0.67	0.83
[1]	1	1	1	1
[2, 3, 4]	0	0	0	0

Thus, for a nondeterministic classifier h and a pair (\mathbf{x}, y) ,

$$F_\beta(h(\mathbf{x}), y) = \begin{cases} \frac{1+\beta^2}{\beta^2+|h(\mathbf{x})|} & \text{if } y \in h(\mathbf{x}) \\ 0 & \text{otherwise.} \end{cases} \quad (7)$$

The most frequently used F-measure is F_1 . For ease of reference, let us state that

$$F_1(h(\mathbf{x}), y) = \frac{2_{y \in h(\mathbf{x})}}{1 + |h(\mathbf{x})|}. \quad (8)$$

To illustrate the use of F-measures of an entry, let us consider an example. If we assume that the true rank of an entry \mathbf{x} is 1, ($y = 1$), then, depending on the value of $h(\mathbf{x})$, Table 1 reports the *Recall*, *Precision*, F_1 , and F_2 . We observe that the reward attached to a prediction containing the true rank with another extra rank ranges from 0.667 for F_1 to 0.833 for F_2 ; while the amounts are lower when the prediction includes 2 extra ranks.

Once we have the definition of F_β for individual entries, it is straightforward to extend it to a test set. So, when S' is a test set of size n , the average loss on it will be computed by

$$\begin{aligned} R^\Delta(h, S') &= \frac{1}{n} \sum_{j=1}^n \Delta(h(\mathbf{x}'_j), y'_j) = \frac{1}{n} \sum_{j=1}^n \left(1 - F_\beta(h(\mathbf{x}'_j), y'_j)\right) \\ &= \frac{1}{n} \sum_{j=1}^n \left(1 - \frac{1 + \beta^2}{\beta^2 + |h(\mathbf{x}'_j)|} 1_{y'_j \in h(\mathbf{x}'_j)}\right). \end{aligned} \quad (9)$$

It is important to realize that for a deterministic hypothesis h this amount is the average "0/1" loss, since all predictions are singletons, $|h(\mathbf{x})| = 1$. Thus, the nondeterministic loss used here is a generalization of the error rate of deterministic classifiers. Furthermore, the average *Recall* and *Precision* on test sets can be similarly defined. In this case, the *Recall* on a test set is the proportion of times that $h(\mathbf{x}')$ includes y' , and is thus a generalization of the deterministic *accuracy*.

Algorithm 1. The nondeterministic ordinal regressor **nd_•**, an algorithm for computing the prediction with one or more ranks for an entry \mathbf{x} provided that the posterior probabilities of ranks are given

Input: object description \mathbf{x}
Input: $\{Pr(j|\mathbf{x}) : j = 1, \dots, k\}$
for $i=1$ **to** k **do**
 $[Start^+(i), Pr_Inter^+(i)] = \max \left\{ \sum_{t=j}^{j+i-1} Pr(t|\mathbf{x}) : j = 1, \dots, k - i + 1 \right\}$
 /* $Pr_Inter^+(i)$ is the highest probability of the intervals of length i */
 /* This interval starts at class $Start^+(i)$ */
end for
 $Min = \operatorname{argmin} \left\{ 1 - \frac{1+\beta^2}{\beta^2+i} Pr_Inter^+(i) : i = 1, \dots, k \right\}$
return $[Start^+(Min), Start^+(Min) + Min - 1]$

4 How to Learn Intervals of Ranks with Posterior Probabilities

In the general ordinal regression setting presented in Section 3, let \mathbf{x} be an entry of the input space \mathcal{X} , and let us now assume that we know the conditional probabilities of ranks given the entry, $Pr(rank = j|\mathbf{x})$ for $j \in \{1, \dots, k\}$. In this context, we wish to define

$$h(\mathbf{x}) = Z \in Intervals\{1, \dots, k\} \quad (10)$$

that minimizes the risk defined in (Eq. 2) when we use the nondeterministic loss given by F_β (Eqs. 6, 7, and 9). We shall prove that such $h(\mathbf{x})$ can be computed by Algorithm 1.

Proposition 1. (*Correctness*) *If the conditional probabilities $Pr(j|\mathbf{x})$ are known, Algorithm 1 returns the nondeterministic prediction $h(\mathbf{x})$ that minimizes the risk given by the loss $1 - F_\beta$.*

Proof. To minimize the risk (Eq. 2), it suffices to compute

$$\Delta_{\mathbf{x}}(Z) = \sum_{y \in \mathcal{Y}} \Delta(Z, y) Pr(y|\mathbf{x}) = \sum_{y \in \mathcal{Y}} (1 - F_\beta(Z, y)) Pr(y|\mathbf{x}), \quad (11)$$

with $Z \in Intervals\{1, \dots, k\}$. Then, we only have to define

$$h(\mathbf{x}) = \operatorname{argmin}\{\Delta_{\mathbf{x}}(Z) : Z \in Intervals\{1, \dots, k\}\}. \quad (12)$$

First we shall prove that when Z is an interval of length i , say $Z = [s, s + i - 1]$, given \mathbf{x} , the value of Equation (11) can be expressed in function of i and the probability of the interval. In fact, with a probability of $1 - Pr(Z|\mathbf{x})$, we expect a loss of 1: the *true* rank will not be one of the interval Z . On the other hand,

with the probability of Z , the *true* rank will be in $h(\mathbf{x})$, and therefore the loss will be 1 minus the F_β of the prediction $h(\mathbf{x}) = Z = [s, s + i - 1]$. In symbols,

$$\begin{aligned} \Delta_{\mathbf{x}}(Z) &= \Delta_{\mathbf{x}}([s, s + i - 1]) \\ &= \left(1 - \sum_{j=s}^{s+i-1} Pr(j|\mathbf{x})\right) 1 + \left(\sum_{j=s}^{s+i-1} Pr(j|\mathbf{x})\right) \left(1 - \frac{1 + \beta^2}{\beta^2 + i}\right) \\ &= 1 - \frac{1 + \beta^2}{\beta^2 + i} \sum_{j=s}^{s+i-1} Pr(j|\mathbf{x}). \end{aligned} \tag{13}$$

Therefore, the interval of length i with lower loss starts at $Start^+(i)$ according to the Algorithm 1, moreover, its loss is

$$1 - \frac{1 + \beta^2}{\beta^2 + i} Pr_Inter^+(i). \tag{14}$$

Thus if Min is the length that gives rise to the lowest loss, the output of the Algorithm is the value of Equation 12 as we wanted to prove.

In practice, posterior probabilities are not known: they are estimated by algorithms that frequently try to optimize the classification accuracy of a hypothesis that returns the class with the highest probability. In other words, probabilities are discriminant values instead of thorough descriptions of the distribution of classes in a learning task. Therefore, the actual role of β in Algorithm 1 is that of a parameter that fixes the thresholds to decide the number of ranks to predict. Hence, like other parameters, β should be tuned in order to achieve optimal results. Thus, depending of the learning task and the probabilistic learner, to reach the highest F_1 scores, it might be necessary to use in Algorithm 1 a value of β different from 1.

5 Experimental Results

In this section we report the results of a set of experiments designed to evaluate the nondeterministic learners proposed in this paper. The aim is to compare, on the one hand, the F_1 scores of well known deterministic learners and their nondeterministic counterparts. It may be argued that these comparisons are not completely fair since the F_1 score tolerates predictions of more than one rank, where it is easier to include the true one. In any case, we report these comparisons in order to test the capabilities of nondeterministic versions to achieve slightly better F_1 scores than their deterministic counterparts. On the other hand, we shall compare the *Recall* and size of predictions attained by nondeterministic learners.

Additionally, since we are dealing with ordinal regression tasks, we check the performance of nondeterministic algorithms in linear loss (sometimes called *MAD*, mean absolute deviation, or *MAE*, mean absolute error). For this purpose,

Table 2. Description of the datasets used in the experiments. The classes are real numbers, and they were discretized in 5 or 10 equal-frequency bins. The splits in train/test were suggested by the experiments reported in [1].

DATASET	#ATTRIBUTES	#TRAIN	#TEST
PYRIMIDINES	27	50	24
TRIAZINES	60	100	86
WISCONSIN BC	32	150	44
MACHINE CPU	6	150	59
AUTO MPG	7	200	192
STOCK	9	300	650
BOSTON	13	300	206
ABALONE	8	300	3877
BANK	32	300	7892
COMPUTER	21	300	7892
CALIFORNIA	8	300	20340
CENSUS	16	300	22484

we must assume singleton predictions; thus, we shall consider the center of each interval as the prediction attached to every interval $h(\mathbf{x})$. The idea is to consider that each rank r can be interpreted as the interval $[r - 0.5, r + 0.5]$ in the real line; thus, a prediction of, say, $[3, 4]$ represents the real interval $[2.5, 4.5]$, and the center point is 3.5.

We used two kinds of learning tasks. In addition to the dataset of beef cattle profiles explained in Section 2, we used a collection of 12 benchmarks (Table 2) that were originally used for metric regression learning tasks. They are publicly available at Luís Torgo’s repository¹. When they were used for ordinal regression in papers like [1,12], the continuous class values were discretized. We used versions with five and ten bins with the same frequency of training examples. The resulting rank values were ordered according to the original metric classes.

To compare the performance of different approaches, we randomly split each data set into training/test partitions. Table 2 reports the characteristics of these datasets and the sizes of splits. The partition was repeated 20 times independently.

Since the nondeterministic approach proposed in this paper is based on the estimation of posterior probabilities of ranks, we used two alternative methods for this stage. First, we used a multiclass *SVM* that estimates the probability of each class given an entry; the implementation used was *libsvm* [11]. The nondeterministic version built from it, following Algorithm 1, was called *nd_SVM*. Second, we used the *MAP* approach of [1] that was devised for ordinal regression tasks. It provides estimations of posterior probabilities using Gaussian processes. The nondeterministic counterpart was called *nd_MAP*. The use of *MAP* in the experiments required reduced sizes of training sets (Table 2) similar to those used in [1]. Nevertheless, the computational requirements of *SVM* would allow us to use *nd_SVM* in tasks of bigger sizes.

¹ <http://www.liaad.up.pt/~ltorgo/Regression/DataSets.html>

Parameter setting. With the *SVM* we used a *rbf* kernel. To set the regularization parameter C and the *rbf* kernel parameter σ , we performed a grid search using a 2-fold cross validation repeated 5 times. The initial search was done with $C \in \{10^{-3}, \dots, 10^3\}$ (respectively $\sigma \in \{10^{-3}, \dots, 10^2\}$) varying the exponent in steps of 1. Let C' and σ' be the best parameters found; then followed a fine search from $C' - 0.8$ (respectively $\sigma' - 0.8$) to $C' + 0.8$ (respectively $\sigma' + 0.8$) with a step of 0.2. Additionally, for *nd_SVM* we searched within $\beta \in \{0.5, 1, 1.5\}$, while the fine search explored the best $\beta - 0.2$, and the best $\beta + 0.2$. We looked for a β , instead of simply using $\beta = 1$, since we wished to compensate any possible inaccuracy in the estimation of probabilities.

The *MAP* learner was used with its default parameters, and no additional tuning was required. The nondeterministic version *nd_MAP* used the search for β of the *nd_SVM*.

The scores achieved in F_1 are shown in Table 3. The nondeterministic learner based on *MAP* bears favorable comparison with the learner based on *SVM*. Thus, *nd_MAP* wins in 18 out of 24 datasets, while *nd_SVM* only wins 3 times out of 24; most of these victories are statistically significant using a Wilcoxon rank sum test of 1-tail over the 20 trials. Comparing the performance over the 24 datasets, we also appreciate significant differences (using a Wilcoxon test with $p < 0.01$) in favor of *nd_MAP*. Therefore, the nondeterministic version of *MAP* outperforms the version based on *SVM* in F_1 when we are using sizes of training sets similar to those showed in Table 2. In the comparison of deterministic versus nondeterministic, in all cases the nondeterministic version outperforms its deterministic counterpart; all but one cases are statistically significant with $p < 0.01$.

The scores in *Recall* are reported in Table 4. Again *nd_MAP* wins in 17 out of 24 datasets, while *nd_SVM* only wins 4 times out of 24; however, now the differences are not so frequently significant. To compare *Recall* scores with those achieved by the deterministic versions, let us remember that for deterministic algorithms, the proportion of successful predictions (accuracy) is also the F_1 and the *Recall*. Therefore, comparing the last two columns of Table 3 and the *Recall* columns of Table 4, we appreciate that the nondeterministic learners outperform the deterministic versions. Thus, in 5 bins datasets, the differences are about 0.24, while in 10 bins datasets the differences are even higher: about 0.31. These results are logical since nondeterministic predictions have more opportunities to include the true ranks.

The average sizes of predictions are shown in the last two columns of Table 4. Here we observe that in the learning tasks of 5 bins these sizes in average are below 2, while with 10 bins, the predictions used more than 3 ranks in average.

The explanation for these facts is straightforward. The nondeterministic algorithms tend to accumulate as many ranks as they are allowed by the F_1 ; thus, in tasks in which the deterministic learners have a poor performance, the corresponding nondeterministic learner may include more ranks in their predictions than in easier tasks. And it is clear that the learning tasks with 5 bins are easier than versions with 10 bins.

Table 3. F_1 scores of the two nondeterministic algorithms and their deterministic counterparts. The results are the averages over 20 trials. In bold face we emphasize the highest score of each dataset. Additionally we test the statistical significance of some interesting differences: between nd_MAP and nd_SVM (see the first column labeled by *si.*), nd_MAP versus MAP (second *si.* column), and nd_SVM versus SVM (last *si.* column). The symbols † (respectively ‡) show that differences are statistically *significant* using a threshold of 0.05 (respectively 0.01) in a Wilcoxon rank sum test.

# BINS	DATASET	nd_MAP (si.)	nd_SVM	MAP (si.)	SVM (si.)
5	PYRIMIDINES	0.58 ‡	0.52	0.57	0.45 †
	TRIAZINES	0.40 †	0.39	0.34 †	0.30 †
	WISCONSIN BC	0.38 †	0.34	0.29 †	0.26 †
	MACHINE CPU	0.66 †	0.64	0.60 †	0.59 †
	AUTO MPG	0.73 †	0.69	0.72 †	0.67 †
	STOCK	0.86 †	0.87 †	0.86 †	0.86 †
	BOSTON	0.71 †	0.68	0.68 †	0.66 †
	ABALONE	0.53 †	0.53 †	0.47 †	0.47 †
	BANK	0.50 †	0.47	0.44 †	0.40 †
	COMPUTER	0.71 †	0.71	0.69 †	0.68 †
	CALIFORNIA	0.57 †	0.57 †	0.52 †	0.52 †
	CENSUS	0.53 †	0.51	0.48 †	0.46 †
AVERAGE (5 B)		0.597	0.576	0.555	0.527
10	PYRIMIDINES	0.35 †	0.27	0.28 †	0.19 †
	TRIAZINES	0.23 †	0.23 †	0.16 †	0.16 †
	WISCONSIN BC	0.21 †	0.19	0.15 †	0.13 †
	MACHINE CPU	0.46 †	0.45	0.36 †	0.37 †
	AUTO MPG	0.51 †	0.47	0.44 †	0.35 †
	STOCK	0.73 †	0.76 †	0.70 †	0.74 †
	BOSTON	0.47 †	0.48 †	0.41 †	0.42 †
	ABALONE	0.35 †	0.34	0.28 †	0.27 †
	BANK	0.31 †	0.28	0.24 †	0.20 †
	COMPUTER	0.53 †	0.51	0.48 †	0.45 †
	CALIFORNIA	0.39 †	0.37	0.32 †	0.30 †
	CENSUS	0.34 †	0.32	0.27 †	0.25 †
AVERAGE (10 B)		0.407	0.388	0.342	0.320
AVERAGE ALL		0.502	0.482	0.449	0.423

Considering the performance over all datasets, we can only find significant differences in *Recall* with $p < 0.06$; while the differences in size of predictions are definitively not significant.

Finally, Table 5 shows the scores achieved in linear loss. This is a relevant measure since we are dealing with ordinal regression learning tasks. Although the nondeterministic algorithms were not designed to improve the linear loss, we observe a good performance. Let us recall here that MAP is a state of the art learner for these tasks. In datasets of 5 bins, MAP wins nd_MAP 8 times out of 12, with only 2 times out of 12 victories for nd_MAP . While nd_MAP wins 7 times out of 12, against 4 wins out of 12 for MAP . The result is that differences over the 24 datasets are not statistically significant.

On the other hand, in linear loss, nd_MAP outperform nd_SVM in most of the datasets with differences statistically significant, see Table 5. The performance

Table 4. Scores of *Recall* and average *size* of predictions ($|h(\mathbf{x})|$) for nondeterministic algorithms. Notice that for deterministic algorithms, the proportion of successful predictions (accuracy) is also the F_1 and the *Recall* (see Table 3). The best scores for each dataset are in bold. When the differences are statistically significant in a Wilcoxon rank sum test, they are marked with † (threshold of 0.05) or ‡ (0.01).

# BINS	DATASET	<i>Recall</i>		AVER. $ h(\mathbf{x}) $			
		<i>nd_MAP</i> (SI.)	<i>nd_SVM</i>	<i>nd_MAP</i> (SI.)	<i>nd_SVM</i>		
5	PYRIMIDINES	0.71	0.73	1.53	‡	1.95	
	TRIAZINES	0.78	†	0.71	3.04	2.75	
	WISCONSIN BC	0.79	†	0.83	3.21	‡	3.84
	MACHINE CPU	0.84	†	0.80	1.67	1.61	
	AUTO MPG	0.80	0.80	1.22	‡	1.34	
	STOCK	0.92	†	0.91	1.16	‡	1.12
	BOSTON	0.80		0.78	1.30		1.35
	ABALONE	0.75	‡	0.70	2.02	‡	1.80
	BANK	0.74	†	0.77	2.08	‡	2.45
	COMPUTER	0.82		0.81	1.35		1.34
	CALIFORNIA	0.78		0.75	1.83	†	1.74
	CENSUS	0.74		0.73	1.93		1.89
	AVERAGE (5 B)		0.788	0.777	1.862		1.931
10	PYRIMIDINES	0.73		0.71	3.32	‡	4.74
	TRIAZINES	0.73		0.68	5.80		5.17
	WISCONSIN BC	0.68	‡	0.86	5.42	‡	8.18
	MACHINE CPU	0.80		0.77	2.75		2.69
	AUTO MPG	0.76		0.76	2.04	‡	2.29
	STOCK	0.83		0.82	1.33	‡	1.21
	BOSTON	0.66		0.72	1.92	‡	2.20
	ABALONE	0.68	†	0.64	3.35	‡	3.06
	BANK	0.70		0.69	3.85	‡	4.33
	COMPUTER	0.74	†	0.72	1.95		1.97
	CALIFORNIA	0.69		0.67	2.92		2.84
	CENSUS	0.66		0.66	3.17		3.23
	AVERAGE (10 B)		0.721	0.725	3.151		3.492
AVERAGE ALL		0.755	0.751	2.507		2.712	

over all datasets is again statistically significant with $p < 0.01$. Finally, let us point out that the nondeterministic *nd_SVM* outperforms (significantly with $p < 0.01$) *SVM*.

Profiles. Tables 6 summarize the scores achieved in the learning task described in Section 2. We used two datasets of sizes 300 and 500. The scores are quite similar for both sizes. Almost always, *nd_SVM* outperforms *nd_MAP* significantly ($p < 0.01$) in F_1 and linear loss, although the scores are quite similar. On the other hand, *nd_MAP* is superior to *nd_SVM* in *Recall*, but again the scores are similar and the significance is only achieved with $p < 0.1$ in one of the datasets. The differences are clearly significant ($p < 0.01$) in the size of the predictions; *nd_SVM* only requires an average of 1.21 or 1.22 ranks to reach a proportion of 85% of predictions that contain the true rank.

Table 5. Linear loss scores of the two nondeterministic algorithms and their deterministic counterparts. The lowest scores for each dataset are highlighted in bold. When the differences are statistically significant in a Wilcoxon rank sum test, they are marked with † (threshold of 0.05) or ‡ (0.01).

# BINS	DATASET	nd_MAP (SI.)	nd_SVM	MAP (SI.)	SVM (SI.)
5	PYRIMIDINES	0.55 ‡	0.65	0.52 †	0.77 †
	TRIAZINES	1.07	1.10	1.18 ‡	1.34 ‡
	WISCONSIN BC	1.12 †	1.19	1.36 †	1.44 †
	MACHINE CPU	0.48	0.48	0.45 †	0.46 †
	AUTO MPG	0.32 †	0.37	0.30 †	0.35 †
	STOCK	0.16 †	0.15	0.14 †	0.14 †
	BOSTON	0.36 †	0.41	0.34 †	0.40 †
	ABALONE	0.72 †	0.73	0.72	0.75 †
	BANK	0.76 †	0.83	0.76	0.90 †
	COMPUTER	0.37	0.38	0.34 †	0.36 †
	CALIFORNIA	0.63 †	0.62	0.59 †	0.60 †
	CENSUS	0.73 †	0.77	0.71 †	0.79 †
	AVERAGE (5 B)	0.607	0.640	0.618	0.692
	10	PYRIMIDINES	1.24 †	1.88	1.31 †
TRIAZINES		2.23	2.21	2.79 †	2.58 †
WISCONSIN BC		2.44	2.48	3.08 †	3.14 †
MACHINE CPU		0.95	0.96	0.95	1.04 †
AUTO MPG		0.74 †	0.80	0.69 †	1.38 †
STOCK		0.34 †	0.30	0.31 †	0.28 †
BOSTON		1.06 †	0.87	1.04 †	0.89 †
ABALONE		1.47 †	1.53	1.57 †	1.71 †
BANK		1.55 †	1.76	1.65 †	2.11 †
COMPUTER		0.75 †	0.76	0.72 †	0.80 †
CALIFORNIA		1.22 †	1.26	1.26 †	1.36 †
CENSUS		1.49 †	1.58	1.58 †	1.71 †
AVERAGE (10 B)		1.289	1.365	1.413	1.603
AVERAGE ALL		0.948	1.002	1.015	1.147

6 Conclusions

We have presented a new kind of ordinal regressors: they are able to predict a variable number of consecutive ranks (an interval of ranks) for each entry. We call such set-valued hypotheses nondeterministic regressors. Roughly speaking, the approach presented in this paper addresses the problem of deciding what to predict when it is possible to envision that the label returned by a learning algorithm is uncertain. The utility of these predictions was illustrated in the context of a real world application: the assessment of muscle proportion in beef cattle carcasses.

After presenting the formal framework as a kind of Information Retrieval, we proposed a family of loss functions for nondeterministic ordinal regression: the complementary of F_β measures. Next we derived an algorithm to minimize such loss functions provided we know the posterior probabilities of each rank given the entry to be ranked. To check the influence of the estimation of conditional

Table 6. Profiles (see Section 2). Dataset characterizations, and scores achieved by deterministic and nondeterministic algorithms. All differences are statically significant ($p < 0.01$) but those achieved in *Recall* ($p < 0.1$).

DATASET	#ATTRIBUTES	#TRAIN	#TEST
PROFILES 500	8	500	391
PROFILES 300	8	300	591

DATASET		<i>nd_MAP</i> (SI.)		<i>nd_SVM</i>		<i>MAP</i> (SI.)		<i>SVM</i> (SI.)	
F_1	PROFILES 500	0.78	‡	0.79		0.76	‡	0.77	‡
	PROFILES 300	0.77	‡	0.78		0.76	‡	0.77	‡
<i>Linear loss</i>	PROFILES 500	0.28	†	0.28		0.29	†	0.27	
	PROFILES 300	0.29	‡	0.28		0.30	‡	0.27	‡

DATASET	<i>Recall</i>		AVER. $ h(\mathbf{x}) $		
	<i>nd_MAP</i> (SI.)	<i>nd_SVM</i>	<i>nd_MAP</i> (SI.)	<i>nd_SVM</i>	
PROFILES 500	0.85	0.84	1.28	‡	1.21
PROFILES 300	0.85	0.84	1.30	‡	1.22

probabilities we compared two implementations. The first one, *nd_SVM* is based on a probabilistic *SVM*, while the second (*nd_MAP*) is built on *MAP*, a learner specialized in ordinal regression learning tasks that uses Gaussian processes to estimate posterior probabilities.

The experiments reported in the previous Section show that *nd_MAP* outperforms *nd_SVM* in almost all measures of performance. Therefore, it is clear the importance of having good probability estimations. However, *MAP* is slower than *SVM*, and it is not possible to handle datasets of medium or large size with the approach based on Gaussian processes.

We think that the main goal of nondeterministic ordinal regressors is not to achieve similar (in fact better) F_β than their deterministic counterpart. We would like to emphasize the dramatic improvement in the proportion of predictions that include the true rank, when the price to be paid for that increase is usually a tiny proportion of predictions with more than one rank.

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