Ordered Correlation Forest*

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Abstract

Empirical studies in various social sciences often involve categorical outcomes with inherent ordering, such as self-evaluations of subjective well-being and self-assessments in health domains. While ordered choice models, such as the ordered logit and ordered probit, are popular tools for analyzing such outcomes, they may impose restrictive parametric and distributional assumptions. This paper proposes a novel estimator, the ordered correlation forest, which overcomes these limitations. The proposed estimator modifies a standard random forest splitting criterion to build a collection of forests, each estimating the conditional probability of a single class. Under an "honesty" condition, predictions are consistent and asymptotically normal. The weights induced by each forest are used to estimate the variance of the predictions and obtain estimation and inference about the covariates' marginal effects. Evidence from synthetic and real data sets shows that the proposed estimator features a superior prediction performance than alternative estimators.

Keywords: Ordered non-numeric outcomes, choice probabilities, machine learning.

JEL Codes: C14, C25, C55

^{*}I especially would like to thank Franco Peracchi for feedback and suggestions. I am also grateful to Michael Lechner, Jana Mareckova, and seminar participants at University of Rome Tor Vergata and SEW-HSG research seminars for comments and discussions. Gabriel Okasa generously shared the code for implementing the DGPs in the simulation. The R package for implementing the methodology developed in this paper is available at https://github.com/riccardo-df/ocf. The associated vignette is at https://riccardo-df.github.io/ocf/.

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1 Introduction

Categorical outcomes with a natural order are commonly observed in empirical studies across social sciences. For example, happiness research typically employs large surveys to collect self-evaluations of subjective well-being (Frey & Stutzer, 2002), and health economics is heavily based on self-assessments in several health domains (see, e.g., Peracchi & Rossetti, 2012, 2013). These outcomes are usually measured on a discrete scale with five or ten classes, where the classes can be arranged in a natural order without any knowledge about their relative magnitude.

Ordered choice models are a popular class of statistical models used to analyze the relationship between this kind of outcome and a set of covariates (see e.g., Greene & Hensher, 2010). These models target the estimation of the conditional choice probabilities, which represent the probability that the outcome belongs to a certain class given the values of the covariates. Common examples of ordered choice models include ordered probit and ordered logit models. However, these models are limited by their dependence on parametric and distributional assumptions that are often based on analytical convenience rather than knowledge about the underlying data generating process. As a result, econometricians may need to consider alternative techniques to produce more accurate and reliable predictions.

This paper proposes a novel estimator that addresses these limitations, the ordered correlation forest. The proposed estimator modifies a standard random forest splitting criterion to build a collection of forests, each estimating the conditional probability of a single class. Evidence from synthetic and real data sets shows that the ordered correlation forest features a superior prediction performance than alternative estimators. Furthermore, the new estimator inherits the asymptotic properties of random forests proven by Wager and Athey (2018), namely the consistency and asymptotic normality of their predictions. This allows valid inference about conditional probabilities to be made using conventional methods, although requires the individual trees to satisfy a fairly strong condition called honesty (Athey & Imbens, 2016). Honesty is a subsample-splitting technique that ensures that dif-

ferent observations are used to place the splits and compute leaf predictions and is crucial to achieving consistency of the predictions. The particular honesty implementation used by the ordered correlation forest estimator allows for a weight-based estimation of the variance of the predicted probabilities. This is achieved by rewriting the random forest predictions as a weighted average of the outcomes (Athey et al., 2019). The weights, which are obtained for the predicted probabilities, can be properly transformed to obtain estimation and inference about the covariates' marginal effects (for a similar approach, see Lechner & Okasa, 2019; Lechner & Mareckova, 2022).

The rest of the paper unfolds as follows. Section 2 provides a brief overview of ordered choice models and discusses some alternative estimation strategies. Section 3 presents the ordered correlation forest estimator, explaining estimation and inference about the statistical targets of interest. Section 4 uses synthetic and real data sets to compare the ordered correlation forest with alternative estimators. Section 5 concludes.

2 Ordered Choice Models

Ordered choice models are a class of statistical models used to analyze the relationship between an ordered non-numeric outcome Y_i and a set of covariates X_i (McCullagh, 1980). These models are typically motivated by postulating the existence of a latent and continuous outcome variable of interest Y_i^* , assumed to obey the following regression model:

$$Y_i^* = g(W_i) + \epsilon_i, \quad \mathbb{E}\left[\epsilon_i | W_i\right] = 0 \tag{2.1}$$

where W_i consists of a set of raw covariates. We may approximate the regression function $g(W_i)$ by a linear-in-parameter model:

$$g(W_i) = \sum_{j=1}^k \beta_j X_{i,j} + V_{i,k} = X_i^T \beta + V_{i,k}$$
 (2.2)

where $X_i = (X_{i,1}, \dots, X_{i,k})$ contains k constructed covariates (generally the basic covariates

 W_i plus interactions and polynomials thereof) and $V_{i,k}$ is an approximation error that is assumed to be independent of X_i .¹ Substituting the linear approximation (2.2) into (2.1) gives:

$$Y_i^* = X_i^T \beta + U_i \tag{2.3}$$

where the random error $U_i = \epsilon_i + V_{i,k}$ depends on k through the approximation error $V_{i,k}$.

However, we generally observe only a discretized version Y_i of Y_i^* , which takes on integer values $m=1,\ldots,M$ corresponding to different categories or classes. Unknown threshold parameters $-\infty=\zeta_0<\zeta_1<\cdots<\zeta_{M-1}<\zeta_M=\infty$ define intervals on the support of Y_i^* , each corresponding to one of the M categories of the observed variable Y_i :

$$\zeta_{m-1} < Y_i^* \le \zeta_m \implies Y_i = m, \quad m = 1, \dots, M$$
 (2.4)

Although the M classes have a natural ordering, their relative magnitude is unknown, thus limiting our ability to make precise quantitative comparisons.

Researchers are typically interested in the estimation of the conditional choice probabilities, defined as:

$$p_m(X_i) := \mathbb{P}(Y_i = m | X_i) \tag{2.5}$$

However, the marginal effect of the j-th covariate on $p_m(\cdot)$ is a more interpretable measure for ordered choice models. The marginal effect is defined based on the continuous or discrete nature of the covariate:

$$p'_{m,j}(x) := \frac{\partial p_m(x)}{\partial x_j} \tag{2.6}$$

$$p'_{m,i}(x) := p_m(\lceil x_i \rceil) - p_m(\lfloor x_i \rfloor) \tag{2.7}$$

where x_j is the j-th element of the vector x and $\lceil x_j \rceil$ and $\lfloor x_j \rfloor$ correspond to x with its j-th

 $^{^{1}}$ We allow for k > n.

element rounded up and down to the closest integer. We can summarize the marginal effects in various ways, such as computing the marginal effect at the mean $p'_{m,j}(\bar{x})$, with \bar{x} denoting a vector of means. Alternatively, we can compute the marginal effect at the median, the mean marginal effect, and the median marginal effect.

Assumptions on the distribution of U_i are generally imposed to derive a closed-form expression of the conditional probabilities and marginal effects. Popular choices are the standard normal distribution function and the standard logistic distribution function, which produce the ordered probit and ordered logit models. Estimation is generally performed using standard maximum likelihood methods.

Although easy to interpret and computationally efficient, these models feature several limitations. First, they impose strong distributional and functional form assumptions generally derived from analytical convenience rather than knowledge about the underlying data generating process. Second, the definition and estimation of the marginal effects have the restrictive property of single-crossing, meaning that these effects can change sign only once when moving from the smallest class to the largest. Third, if the number of covariates is larger than the number of observations, estimation breaks down.

Several alternatives have been proposed in the literature to overcome these limitations. For example, Boes and Winkelmann (2006) discuss generalizations of the standard ordered choice models. However, they still rely on parametric and distributional assumptions which limit the increase in flexibility they provide.

Recent developments in statistical learning (see, e.g., Hastie et al., 2009; Efron & Hastie, 2016) offer ways to relax these assumptions. However, classification algorithms do not leverage the ordering information embedded in the structure of the outcome, and regression algorithms treat the outcome as if it is measured on a metric scale. Thus, applying these algorithms "off-the-shelf" can lead to biased and inefficient estimation of conditional probabilities.

To overcome these limitations, one approach is to transform ordered non-numeric out-

comes into a metric scale using scores based on the classes of the observed outcome, thus allowing us to use any regression algorithm on the transformed outcome. For example, Hothorn et al. (2006) propose using the midpoint values of the intervals defined on the support of the latent outcome as score values. In the cases where Y_i^* is not observed, this translates into setting the scores equal to the class labels of Y_i . However, this assumes that the intervals are of equal length, which may not be accurate in practice. To address this issue, Hornung (2020) proposes the ordinal forest estimator, which optimizes the class intervals and uses score values corresponding to these optimized intervals in a standard regression forest. The optimization process involves growing multiple forests using randomly generated candidate score sets, and constructing the final score values by summarizing the score sets with the smallest out-of-bag error. Hornung (2020) shows that the ordinal forest estimator outperforms a standard regression forest that uses class labels as score values using both real and synthetic data sets. However, the optimization process can be computationally expensive, which may limit its practical use for large data sets or real-time applications.

Another approach involves expressing conditional probabilities as conditional expectations of binary variables, which can be estimated by any regression algorithm. One first strategy, which we label *multinomial machine learning*, is to express conditional probabilities as follows:

$$p_{m}(X_{i}) = \mathbb{E}\left[\mathbb{1}\left(Y_{i} = m\right) | X_{i}\right]$$

$$= \psi_{m}(X_{i})$$
(2.8)

This allows us to estimate each $\psi_m(X_i)$ separately using any regression algorithm to get an estimate of conditional probabilities:

$$\hat{p}_{m}^{MML}\left(X_{i}\right) = \hat{\psi}_{m}\left(X_{i}\right) \tag{2.9}$$

However, $\hat{p}_m^{MML}(\cdot)$ does not leverage the information embedded in the ordered structure of the outcome. To overcome this limitation, an alternative strategy that we label *ordered*

machine learning expresses conditional choice probabilities as the difference between the cumulative probabilities of two adjacent classes:

$$p_{m}(X_{i}) = \mathbb{P}(Y_{i} \leq m|X_{i}) - \mathbb{P}(Y_{i} \leq m - 1|X_{i})$$

$$= \mathbb{E}\left[\mathbb{1}(Y_{i} \leq m)|X_{i}] - \mathbb{E}\left[\mathbb{1}(Y_{i} \leq m - 1)|X_{i}\right]$$

$$= \mu_{m}(X_{i}) - \mu_{m-1}(X_{i})$$

$$(2.10)$$

Then we can separately estimate $\mu_m(\cdot)$ for all $m=1,\ldots,M-1$ using any regression algorithm and pick the difference between the cumulative probabilities of two adjacent classes to estimate $p_m(\cdot)$:²

$$\hat{p}_{m}^{OML}(X_{i}) = \hat{\mu}_{m}(X_{i}) - \hat{\mu}_{m-1}(X_{i})$$
(2.11)

However, $\hat{p}_m^{OML}(\cdot)$ can potentially produce negative predictions, thereby contradicting the definition of probabilities.³

3 Estimation and Inference

In this section, I discuss the implementation of the ordered correlation forest (OCF) estimator. First, I illustrate the estimation of conditional choice probabilities and marginal effects. Second, I discuss the conditions required for the asymptotic normality and consistency of OCF predictions. Finally, I show how to conduct approximate inference about the statistical targets of interest.

²Lechner and Okasa (2019) combine ordered machine learning with random forests (Breiman, 2001) and discuss how to estimate and conduct inference about marginal effects.

³Although we might resolve this issue by setting negative predictions to zero, such a solution is suboptimal, and an alternative estimator that does not require truncation may perform better. Therefore, the choice between multinomial and ordered machine learning remains an open question that requires careful empirical investigation.

3.1 Estimation

Similar to the ordered machine learning approach (see equation 2.10), the OCF estimator computes the prediction of conditional choice probabilities as the difference between the cumulative probabilities of two adjacent classes. However, instead of estimating $\mu_m(\cdot)$ and $\mu_{m-1}(\cdot)$ separately, OCF internally performs this computation in a single random forest. This allows us to tie the estimation of $\mu_m(\cdot)$ and $\mu_{m-1}(\cdot)$ to correlate the errors made in estimating these two expectations. Additionally, it avoids negative predictions.

To see the importance of correlating the estimation errors, we can decompose the mean squared error of a prediction $\hat{p}_{m}^{OML}(\cdot)$ at x as follows:⁴

$$MSE\left(\hat{p}_{m}^{OML}(x)\right) = \mathbb{E}\left[\left\{\hat{p}_{m}^{OML}(x) - p_{m}(x)\right\}^{2}\right]$$

$$= \mathbb{E}\left[\left\{\hat{\mu}_{m}(x) - \hat{\mu}_{m-1}(x) - \mu_{m}(x) + \mu_{m-1}(x)\right\}^{2}\right]$$

$$= MSE\left(\hat{\mu}_{m}(x)\right) + MSE\left(\hat{\mu}_{m-1}(x)\right) - 2EC\left(\hat{\mu}_{m}(x), \hat{\mu}_{m-1}(x)\right)$$
(3.1)

where the last term is the error correlation and captures the degree to which the errors made in estimating $\mu_m(\cdot)$ and $\mu_{m-1}(\cdot)$ are correlated:

$$EC(\hat{\mu}_{m}(x), \hat{\mu}_{m-1}(x)) = \mathbb{E}\left[\left\{\hat{\mu}_{m}(x) - \mu_{m}(x)\right\} \left\{\hat{\mu}_{m-1}(x) - \mu_{m-1}(x)\right\}\right]$$
(3.2)

Equation (3.1) shows that $\hat{p}_{m}^{OML}(\cdot)$ is a suboptimal estimator. Estimating $\mu_{m}(\cdot)$ and $\mu_{m-1}(\cdot)$ separately minimizes only the mean squared error terms and ignores the error correlation. Correlating the errors could improve estimation performance since errors that move in the same direction cancel out when taking the difference $\hat{\mu}_{m}(\cdot) - \hat{\mu}_{m-1}(\cdot)$.

To address this limitation, OCF constructs a collection of forests, one for each of the M classes of Y_i . However, rather than the standard criterion, OCF uses equation (3.1) as the splitting rule to build the individual trees in the m-th forest. This allows the estimator to account for the error correlation that $\hat{p}_m^{OML}(\cdot)$ ignores. Intuitively, OCF seeks splits that

⁴This decomposition can be applied to any estimation strategy that involves calculating the difference between two surfaces. For example, Lechner and Mareckova (2022) leverage this decomposition to estimate heterogeneous causal effects.

not only provide good estimates of $\mu_m(\cdot)$ and $\mu_{m-1}(\cdot)$, but also correlate the errors made in estimating these expectations.

To use (3.1) as the splitting rule, we need to estimate its components. This, in turn, requires an estimator of $\mu_m(\cdot)$ in each node. An unbiased estimator of $\mu_m(\cdot)$ in a child node $C_j \subset X$ consists of the proportion of observations in C_j whose outcome is lower than or equal to m:

$$\check{\mu}_m(X_i) = \frac{1}{|C_j|} \sum_{i: X_i \in C_j} \mathbb{1}(Y_i \le m)$$
(3.3)

This leads us to estimating $MSE\left(\check{\mu}_{m}\left(\cdot\right)\right)$ and $EC\left(\check{\mu}_{m}\left(\cdot\right),\check{\mu}_{m-1}\left(\cdot\right)\right)$ in each node by their sample analogs:

$$\widehat{MSE}_{C_{j}}(\check{\mu}_{m}(X_{i})) = \frac{1}{|C_{j}|} \sum_{i:X_{i} \in C_{i}} \left[\mathbb{1} \left(Y_{i} \leq m \right) - \check{\mu}_{m}(X_{i}) \right]^{2}$$
(3.4)

$$\widehat{EC}_{C_{j}}\left(\check{\mu}_{m}\left(X_{i}\right),\check{\mu}_{m-1}\left(X_{i}\right)\right) = \frac{1}{|C_{j}|} \sum_{i:X_{i} \in C_{j}} \mathbb{1}\left(Y_{i} \leq m\right) \mathbb{1}\left(Y_{i} \leq m-1\right) - \check{\mu}_{m}\left(X_{i}\right) \check{\mu}_{m-1}\left(X_{i}\right) \quad (3.5)$$

Then, in the m-th forest, OCF constructs individual trees by recursively partitioning each parent node $\mathcal{P} \subseteq X$ into two child nodes $C_1, C_2 \subset \mathcal{P}$ such that the following minimization problem is solved:

$$\min_{C_1, C_2} \sum_{j=1}^{2} \widehat{MSE}_{C_j} \left(\check{\mu}_m \left(X_i \right) \right) + \widehat{MSE}_{C_j} \left(\check{\mu}_{m-1} \left(X_i \right) \right) - 2\widehat{EC}_{C_j} \left(\check{\mu}_m \left(X_i \right), \check{\mu}_{m-1} \left(X_i \right) \right)$$
(3.6)

Once the recursive partitioning stops, each tree in the m-th forest unbiasedly estimates $p_m(\cdot)$ at x by computing the proportion of observations in the same leaf as x whose outcome equals m:

$$\hat{p}_{m,b}^{OCF}(x) = \check{\mu}_{m}(x) - \check{\mu}_{m-1}(x)$$

$$= \frac{1}{|L_{m,b}(x)|} \sum_{i \in L_{m,b}(x)} \mathbb{1}(Y_{i} = m)$$
(3.7)

where $L_{m,b}\left(x\right)$ is the set of observations falling in the same leaf of the b-th tree as the

prediction point x. The predictions from each tree are then averaged to obtain the forest predictions:⁵

$$\hat{p}_{m}^{OCF}(x) = \frac{1}{B_{m}} \sum_{b=1}^{B_{m}} \hat{p}_{m,b}^{OCF}(x)$$
(3.8)

where $b = 1, ..., B_m$ indexes the trees in the m-th forest. In contrast to ordered machine learning, OCF ensures model consistency, as the predictions $\hat{p}_m^{OCF}(\cdot)$ always lie in the unit interval by construction.

Estimation of marginal effects proceeds as proposed by Lechner and Okasa (2019). For discrete covariates, we can plug an estimate $\hat{p}_{m}^{OCF}(\cdot)$ of $p_{m}(\cdot)$ into equation (2.7) to have a straightforward estimator of $p'_{m,j}(\cdot)$. For continuous covariates, we use a nonparametric approximation of the infinitesimal change in x_{j} :

$$\hat{p}_{m,b}^{\prime OCF}(x) = \frac{\hat{p}_{m}^{OCF}(\widehat{|x_{j}|}) - \hat{p}_{m}^{OCF}(\widehat{|x_{j}|})}{\bar{x}_{j} - \underline{x}_{j}}$$
(3.9)

where $\widehat{[x_j]}$ and $\widehat{[x_j]}$ correspond to x with its j-th element set to $\overline{x}_j = x_j + \omega \sigma_j$ and $\underline{x}_j = x_j - \omega \sigma_j$, with σ_j the standard deviation of x_j and $\omega > 0$ a tuning parameter.

3.2 Asymptotic Properties

Wager and Athey (2018) establish the consistency and asymptotic normality of random forest predictions. However, besides some regularity and technical assumptions, there are certain conditions regarding the construction of individual trees that must be satisfied. In the following, I define these conditions.

The first condition requires that the trees use different observations to place the splits and compute the leaf predictions. This condition is called *honesty* and is crucial to bounding the bias of forest predictions.

This is true also for $\hat{p}_{m}^{MML}(\cdot)$ and $\hat{p}_{m}^{OML}(\cdot)$.

Definition 1 (Honesty). A tree is honest if it uses the outcome Y_i to either place the splits or compute the leaf predictions, but not both.

Wager and Athey (2018) implement honesty by first drawing a subsample from the original sample S for each tree and then splitting the subsample into two halves, using one half to grow the tree and the other half to compute leaf predictions (see also Athey et al., 2019). Alternatively, Lechner and Mareckova (2022) suggest a different implementation (also used by Lechner & Okasa, 2019). First, they split the original sample S into a training sample S^{tr} and an honest sample S^{hon} . Then, they use random subsamples from S^{tr} to construct trees and compute leaf predictions using only S^{hon} . This strategy enables weight-based inference about leaf predictions and their transformations, such as marginal effects. OCF adopts this strategy as well (see Section 3.3).

The second condition is that the leaves of the trees must become small in all dimensions of the covariate space as the sample size increases. This is necessary for achieving consistency of the predictions and is accomplished by introducing randomness in the tree-growing process and enforcing a regularity condition on how quickly the leaves get small.

Definition 2 (Random-split). A tree is random-split if, at every step of the tree-growing procedure, the probability that the next split occurs along the j-th covariate is bounded below by π/k , for some $0 < \pi \le 1$, for all j = 1, ..., k.

Definition 3 (α -regularity). A tree is α -regular if each split leaves at least a fraction α of the observations in the parent node on each side of the split and the trees are fully grown to depth d for some $d \in N$, that is, there are between d and 2d-1 observations in each terminal node of the tree.

To achieve α -regularity, OCF ignores splits that do not satisfy this condition. The algorithm always selects the best split from among the candidate splits that would maintain at least a fraction α of the parent node's observations on both sides of the split. This way, we can rule out any influence of the splitting rule on the shape of the final leaves.

Third, trees must be constructed using subsamples drawn without replacement, rather than bootstrap samples, as originally proposed by Breiman (2001).

Lastly, to derive the asymptotic normality, trees must be symmetric.

Definition 4 (Symmetry). A predictor is symmetric if the (possibly randomized) output of the predictor does not depend on the order in which observations are indexed in the training and honest samples.

Under these conditions, we can establish consistency and asymptotic normality of the predictions. For completeness, I here report the main theorem by Wager and Athey (2018).

Theorem 3.1. Suppose that we have n independent and identically distributed training examples $(X_i, Y_i) \in [0, 1]^k \times \mathcal{R}$. Suppose moreover that the covariates are independently and uniformly distributed $X_i \sim U([0, 1]^k)$, that $\theta(x) = \mathbb{E}[Y|X=x]$ and $\theta_2(x) = \mathbb{E}[Y^2|X=x]$ are Lipschitz-continuous, and finally that Var[Y|X=x] > 0 and $\mathbb{E}[|Y-\mathbb{E}[Y|X=x]|^{2+\delta}|X=x] \le M$ for some constants $\delta, M > 0$, uniformly over all $x \in [0, 1]^k$. Given this data-generating process, let \mathcal{T} be an honest, α -regular with $\alpha \le 0.2$, and symmetric random-split tree in the sense of Definitions 1-4, and let $\hat{\theta}_n(x)$ be the estimate for $\theta(x)$ given by a random forest with base learner \mathcal{T} and a subsample size s_n . Finally, suppose that the subsample size s_n scales as:

$$s_n \approx n^{\beta} \text{ for some } \beta_{min} := 1 - \left(1 + \frac{k}{\pi} \frac{\log(\alpha^{-1})}{\log((1 - \alpha)^{-1})}\right) < \beta < 1$$

Then, random forest predictions are asymptotically Gaussian:

$$\frac{\hat{\theta}_n(x) - \theta(x)}{\sigma_n(x)} \implies \mathcal{N}(0, 1) \text{ for a sequence } \sigma_n(x) \to 0$$

3.3 Inference

In addition to Theorem 3.1, Wager and Athey (2018) show that the asymptotic variance of random forest predictions $\sigma_n(\cdot)$ can be consistently estimated by adapting the infinitesimal

jackknife estimator proposed by Wager et al. (2014) to the case of subsampling without replacement. This approach can be used to estimate the variance of a prediction $\hat{p}_{m}^{OCF}(\cdot)$ at x. However, generalizing this method to estimate the variance of marginal effects $\hat{p}_{m,j}^{\prime OCF}(\cdot)$ is not straightforward.

To overcome this limitation, OCF employs an alternative approach that leverages the weight-based representation of random forest predictions (Athey et al., 2019) and adapts the weight-based inference proposed by Lechner and Mareckova (2022) (see also Lechner & Okasa, 2019). In particular, OCF implements honesty in a way that guarantees that the weight assigned to the i-th unit is independent of the outcomes of other units. This allows for the derivation of a straightforward formula for the variance of honest predicted probabilities and marginal effects.

First, we express OCF predictions as weighted averages of the outcomes. Let S denote the observed sample. The following provides an expression for a prediction $\hat{p}_m^{OCF}(\cdot)$ at x numerically equivalent to that in (3.8):

$$\hat{p}_{m}^{OCF}(x) = \sum_{i \in \mathcal{S}} \hat{\alpha}_{m,i}(x) \, \mathbb{1}(Y_{i} = m)$$

$$\hat{\alpha}_{m,b,i}(x) = \frac{\mathbb{1}(X_{i} \in L_{m,b}(x))}{|L_{m,b}(x)|}, \quad \hat{\alpha}_{m,i}(x) = \frac{1}{B_{m}} \sum_{b=1}^{B_{m}} \hat{\alpha}_{m,b,i}(x)$$
(3.10)

where the weights $\hat{\alpha}_{m,1}(x), \ldots, \hat{\alpha}_{m,|\mathcal{S}|}(x)$ determine the forest-based adaptive neighborhood of x. They represent how often the i-th observation in \mathcal{S} shares a leaf with x in the m-th forest. This measures how important the i-th observation is for fitting $p_m(\cdot)$ at x. Notice that $\sum_{i \in \mathcal{S}} \hat{\alpha}_{m,i}(x) = 1$ for all x.

Calculating the variance of a prediction $\hat{p}_{m}^{OCF}(x)$ in (3.10) is challenging because the weight assigned to the *i*-th unit $\hat{\alpha}_{m,i}(x)$ is a function of both S and X_{i} . Thus, this weight depends on the outcomes of all other units in S, which complicates the formula for the variance.

However, the formula for the variance simplifies under the particular honesty implemen-

tation of OCF. Let \mathcal{S}^{tr} and \mathcal{S}^{hon} be a training sample and an honest sample obtained by randomly splitting the observed sample \mathcal{S} . Also, let $\hat{\alpha}_{m,i}^{tr}(\cdot)$ be the weights induced by a forest constructed using only \mathcal{S}^{tr} . Then, an honest prediction $\hat{p}_{m}^{OCF_{H}}(\cdot)$ at x is obtained by the following weighted average of observations in \mathcal{S}^{hon} :

$$\hat{p}_{m}^{OCF_{H}}\left(x\right) = \sum_{i \in S^{hon}} \hat{\alpha}_{m,i}^{tr}\left(x\right) \mathbb{1}\left(Y_{i} = m\right) \tag{3.11}$$

The new weight assigned to the *i*-th unit $\hat{\alpha}_{m,i}^{tr}(x)$ is a function of \mathcal{S}^{tr} and of X_i . Thus, under i.i.d. sampling this weight is independent of the outcomes of other units in \mathcal{S}^{hon} . This allows us to derive a simple formula for the variance of an honest prediction $\hat{p}_m^{OCF_H}(x)$:

$$\mathbb{V}\left(\hat{p}_{m}^{OCF_{H}}\left(x\right)\right) = |\mathcal{S}^{hon}| \,\mathbb{V}\left(\hat{\alpha}_{m,i}^{tr}\left(x\right) \,\mathbb{1}\left(Y_{i} = m\right)\right) \tag{3.12}$$

We can estimate this variance by its sample analog.

By plugging (3.11) into (3.9), we obtain the following estimator of honest marginal effects:

$$\hat{p}_{m,j}^{\prime OCF_{H}}(x) = \frac{1}{\bar{x}_{j} - \underline{x}_{j}} \left\{ \sum_{i \in S^{hon}} \hat{\alpha}_{m,i}^{tr}(\lceil \widehat{x_{j}} \rceil) \mathbb{1} (Y_{i} = m) - \sum_{i \in S^{hon}} \hat{\alpha}_{m,i}^{tr}(\lceil \widehat{x_{j}} \rfloor) \mathbb{1} (Y_{i} = m) \right\}$$

$$= \frac{1}{\bar{x}_{j} - \underline{x}_{j}} \sum_{i \in S^{hon}} \tilde{\alpha}_{m,i}^{tr}(\lceil \widehat{x_{j}} \rceil, \lceil \widehat{x_{j}} \rceil) \mathbb{1} (Y_{i} = m)$$

$$(3.13)$$

with $\tilde{\alpha}_{m,i}^{tr}(\widehat{|x_j|},\widehat{|x_j|}) = \hat{\alpha}_{m,i}^{tr}(\widehat{|x_j|}) - \hat{\alpha}_{m,i}^{tr}(\widehat{|x_j|})$ a transformation of the original weights. Using the same argument as before, under i.i.d. sampling the weight assigned to the *i*-th unit $\tilde{\alpha}_{m,i}^{tr}(\widehat{|x_j|},\widehat{|x_j|})$ is independent of the outcomes of other units in S^{hon} . Thus the variance of an honest marginal effect $\hat{p}_{m,j}^{'OCF_H}(x)$ can be expressed as follows:

$$\mathbb{V}\left(\hat{p}_{m,j}^{\prime OCF_{H}}\left(x\right)\right) = \frac{|\mathcal{S}^{hon}|}{\left(\overline{x}_{j} - \mathbf{x}_{j}\right)^{2}} \,\mathbb{V}\left(\widetilde{\alpha}_{m,i}^{tr}(\left\lceil\widehat{x}_{j}\right\rceil,\left\lceil\widehat{x}_{j}\right\rfloor)\mathbb{1}\left(Y_{i} = m\right)\right) \tag{3.14}$$

As before, we can estimate this variance by its sample analog.

According to Theorem 3.1, the honest predicted probabilities in (3.11) are consistent and asymptotically normal, provided that the weights $\hat{\alpha}_{m,i}^{tr}(\cdot)$ are induced by a forest composed of

 α -regular with $\alpha \leq 0.2$ and symmetric random-split trees grown using subsampling without replacement. With these conditions met, we can use the estimated standard errors of honest predicted probabilities $\hat{p}_m^{OCF_H}(\cdot)$ to conduct valid inference as usual, e.g., by constructing conventional confidence intervals.

Furthermore, under the same conditions the honest marginal effects in (3.13) are a linear combination of normally distributed predictions, and thus have a normal distribution as well. Therefore, we can also construct conventional confidence intervals for honest marginal effects $\hat{p}'_{m,j}^{OCF_H}$ (·) using their estimated standard errors.

4 Comparison with Alternative Estimators

This section compares the performance of the ordered correlation forest (OCF) estimator with several alternative methods, using both synthetic and real data sets. To provide a comprehensive evaluation, I employ both multinomial and ordered machine learning techniques, combining them with random forests (Breiman, 2001) and penalized logistic regressions with an L1 penalty (Tibshirani, 1996). I refer to the resulting estimators as multinomial random forest (MRF), multinomial L1 regression (ML1), ordered random forest (ORF), and ordered L1 regression (OL1). Additionally, I include the ordered logit (LOGIT) model as a parametric benchmark for comparison.

4.1 Simulation Results

I choose the DGPs to replicate part of the simulation study in Lechner and Okasa (2019). I consider three different designs that differ in the model for the latent outcome variable:

 $^{^6}ORF$ has been extensively discussed in Lechner and Okasa (2019).

Design 1.
$$Y_i^* = X_i^T \beta + U_i$$
Design 2.
$$Y_i^* = \sum_{j=1}^k X_{ij} \mathbb{1} (X_{ij} > 0) \beta_j + U_i$$
Design 3.
$$Y_i^* = \sin (2X_i)^T \beta + U_i$$

with $U_i \sim logistic$ (0, 1) in all designs. The three designs share all the other settings described below. For each design, I consider four sample sizes, $|S| \in \{500, 1000, 2000, 4000\}$. Thus, I consider overall twelve different scenarios.

I obtain the observed outcomes Y_i by discretizing Y_i^* into nine classes:

$$\zeta_{m-1} < Y_i^* \le \zeta_m \implies Y_i = m, \quad m = 1, \dots 9$$

I construct the threshold parameters ζ_1, \ldots, ζ_8 as follows. First, I draw eight values $\zeta_m^q \sim U(0.09, 0.91)$ and sort them in ascending order, so that $\zeta_m^q \leq \zeta_{m+1}^q$. Then, I generate a sample of 1,000,000 Y_i^* and set $\zeta_m = Q(\zeta_m^q)$, with $Q(\cdot)$ the empirical quantile function of Y_i^* . This way, the threshold parameters are unevenly spaced, and the class widths are randomized and unequal.

I generate k = 30 covariates $X_i \sim \mathcal{N}(0, \Sigma)$. The components of the coefficient vector $\boldsymbol{\beta}$ are $\beta_1, \ldots, \beta_5 = 1, \beta_6, \ldots, \beta_{10} = 0.75$, and $\beta_{11}, \ldots, \beta_{15} = 0.5$. The remaining covariates have null coefficients, that is, they are "noise" covariates. The variance-covariance matrix Σ is block diagonal and induces correlation among signal covariates as well as among noise covariates, but there is zero correlation between signal and noise covariates:

$$\Sigma = \begin{pmatrix} \mathbf{A}_{signal} & 0 \\ 0 & \mathbf{A}_{noise} \end{pmatrix} \qquad a_{i,j}^{signal} = a_{i,j}^{noise} = \begin{cases} 1, & i = j \\ 0.8, & i \neq j \cap \{i, j \text{ are odd}\} \\ 0, & otherwise \end{cases}$$

After drawing a sample S, I estimate the conditional choice probabilities using LOGIT, MRF, ML1, ORF, OL1, and two versions of OCF, the "adaptive" version OCF_A and the

⁷If the distance between two adjacent values ζ_m^q and ζ_{m+1}^q is not sufficient, I redraw another set of values.

"honest" version OCF_H . This way, we can quantify the loss in the precision derived from using fewer observations to build the forests, representing the price to pay for valid inference. I feed the estimators with all covariates without adding any polynomials, interaction terms, or other transformations of the covariates. Thus, LOGIT and OL1 are correctly specified in $Design\ 1$ and misspecified in the other designs. To implement OCF_H , I randomly split S into a training sample S^{tr} used to construct the trees and an honest sample S^{hon} used to compute the leaf predictions. I choose $|S^{tr}| = |S^{hon}| = |S|/2$.

I rely on an external validation sample S^{val} of size $|S^{val}| = 10,000$ to assess the prediction performance of the estimators. This large number of observations helps minimize the sampling variance. For each replication r = 1, ..., R, I calculate the mean squared error, mean absolute error, and ranked probability score for each estimator:

$$MSE_r = \frac{1}{|S^{val}|} \sum_{i \in S^{val}} \sum_{m=1}^{M} \left[p_m(X_i) - \hat{p}_{m,r}(X_i) \right]^2$$
 (4.1)

$$MAE_{r} = \frac{1}{|\mathcal{S}^{val}|} \sum_{i \in \mathcal{S}^{val}} \sum_{m=1}^{M} \left| p_{m} \left(X_{i} \right) - \hat{p}_{m,r} \left(X_{i} \right) \right|$$

$$(4.2)$$

$$RPS_{r} = \frac{1}{|S^{val}|} \sum_{i \in S^{val}} \frac{1}{M-1} \sum_{m=1}^{M} \left[\mu_{m} (X_{i}) - \hat{\mu}_{m,r} (X_{i}) \right]^{2}$$
(4.3)

with $\hat{p}_{m,r}(\cdot)$ the estimated conditional probabilities in the r-th replication, and $\hat{\mu}_{m,r}(x) = \sum_{j=1}^{m} \hat{p}_{j,r}(x)$ the estimated cumulative distribution function. Notice that, by simulation design, we can compute the true probabilities. I summarize these performance measures by averaging over the replications.

Table 4.1 displays the results obtained with R = 1,000 replications. The simulation shows that OCF_A outperforms all other estimators uniformly in all the considered scenarios, except for LOGIT and OL1 in Design~1. Unsurprisingly, LOGIT and OL1 perform best in Design~1, since they correctly specify the parametric model and the distributional assumption of the error term. However, their performance deteriorates when the model is misspecified, and in Design~3 even OCF_H demonstrates superior performance despite using half of the

observations to train the model.

Ordered machine learning emerges as the superior choice over multinomial machine learning, with notable performance advantages observed in terms of RPS. Specifically, ORF always features lower MSE, MAE, and RPS than MRF, and OL1 outperforms ML1 in $Design\ 1$ and $Design\ 2$. In $Design\ 3$, the MSE of OL1 is around 2–6% larger than that of ML1, and minimal disparities are observed in the other performance measures.

However, OCF_A always outperforms OL1 when the latter is misspecified, with an advantage that ranges between 13–87% in terms of MSE, 7–50% in terms of MAE, and 10–212% in terms of RPS, with larger advantages observed in Design~3. Moreover, while OCF_A and ORF achieve the same ranked probability scores, the former consistently exhibits lower mean

	Design 1				Design 2				Design 3			
	500	1,000	2,000	4,000	500	1,000	2,000	4,000	500	1,000	2,000	4,000
Panel 1:	\overline{MSE}											
LOGIT	0.070	0.057	0.052	0.050	0.093	0.086	0.083	0.081	0.185	0.179	0.177	0.176
MRF	0.171	0.154	0.141	0.131	0.092	0.079	0.068	0.061	0.140	0.127	0.116	0.107
ML1	0.192	0.182	0.175	0.171	0.099	0.091	0.085	0.081	0.184	0.180	0.177	0.174
ORF	0.158	0.145	0.135	0.126	0.085	0.076	0.067	0.061	0.138	0.125	0.113	0.104
OL1	0.123	0.088	0.069	0.059	0.098	0.079	0.067	0.060	0.195	0.188	0.182	0.178
$OCF_{_A}$	0.150	0.137	0.128	0.119	0.074	0.066	0.058	0.053	0.124	0.113	0.103	0.095
$OCF_{\scriptscriptstyle H}$	0.186	0.169	0.157	0.147	0.094	0.082	0.073	0.066	0.153	0.141	0.129	0.118
Panel 2: \overline{MAE}												
LOGIT	0.390	0.351	0.337	0.329	0.564	0.544	0.534	0.529	0.923	0.915	0.912	0.910
MRF	0.803	0.749	0.708	0.675	0.610	0.558	0.515	0.480	0.779	0.735	0.697	0.663
ML1	0.910	0.881	0.862	0.850	0.654	0.621	0.597	0.581	0.963	0.950	0.940	0.933
ORF	0.734	0.697	0.668	0.641	0.582	0.542	0.506	0.477	0.748	0.707	0.670	0.639
OL1	0.547	0.452	0.395	0.362	0.614	0.548	0.502	0.474	0.969	0.950	0.935	0.925
$OCF_{\scriptscriptstyle A}$	0.732	0.692	0.660	0.633	0.538	0.499	0.466	0.439	0.718	0.679	0.645	0.616
OCF_{H}	0.886	0.832	0.790	0.755	0.638	0.591	0.553	0.521	0.850	0.806	0.760	0.721
Panel 3: RPS												
LOGIT	0.009	0.007	0.006	0.006	0.029	0.027	0.026	0.025	0.094	0.091	0.089	0.089
MRF	0.034	0.030	0.027	0.024	0.027	0.023	0.020	0.017	0.046	0.041	0.036	0.033
ML1	0.046	0.043	0.041	0.039	0.035	0.032	0.030	0.028	0.094	0.092	0.090	0.089
ORF	0.029	0.026	0.024	0.022	0.021	0.018	0.016	0.014	0.038	0.034	0.030	0.028
OL1	0.012	0.009	0.008	0.007	0.024	0.021	0.020	0.019	0.093	0.091	0.090	0.089
$OCF_{\scriptscriptstyle A}$	0.030	0.026	0.024	0.022	0.022	0.019	0.016	0.014	0.040	0.035	0.031	0.028
OCF_H	0.044	0.038	0.034	0.031	0.034	0.029	0.026	0.022	0.065	0.056	0.048	0.041

Table 4.1: Comparison with alternative estimators. The first three panels report the average over the replications of MSE_r (\overline{MSE}), MAE_r (\overline{MAE}), and RPS_r (\overline{RPS}).

squared error and mean absolute error across all considered scenarios. In $Design\ 1$, the advantage of OCF_A over ORF is relatively small, with the MSE of ORF being approximately 5% larger than that of OCF_A , and slight differences in MAE observed. The superiority of OCF_A becomes more pronounced in $Design\ 2$ (around 15% in terms of MSE and 8% in terms of MAE) and $Design\ 3$ (around 10% in terms of MSE and 4% in terms of MAE).

Finally, we compare the prediction performance of OCF_A and OCF_H to quantify the cost of honesty resulting from using fewer observations to construct the forests. The results indicate that in terms of MSE, the cost ranges between 23%–26%, while in terms of MAE, the cost ranges between 17%–21%. The cost is larger in terms of RPS, ranging between 41%–64%. Despite this cost, the MSE of OCF_H is lower than that of LOGIT when the latter is misspecified.

4.2 Empirical Results

I utilize the same data sets considered by Janitza et al. (2016), Hornung (2020), and Lechner and Okasa (2019). These data sets differ in terms of the number of covariates, observations, and classes of the observed outcome. Table 4.2 provides a summary of the data sets. For further details on the background of each data set, the reader is referred to Janitza et al. (2016).

To assess the prediction accuracy of each estimator, I employ a ten-fold cross-validation procedure. Specifically, I randomly divide each data set into ten folds S^1, \ldots, S^{10} with roughly equal sizes. For each fold $f = 1, \ldots, 10$, I fit all the estimators using the observations

Data Sets											
Data set	Sample Size	Outcome	Class	N. Covariates							
vlbw	218	Apgar score	1 (Life-threatening)	_	9 (Optimal)	10					
mammography	412	Last mammography	1 (Never)	_	3 (Over a year)	5					
support	798	Functional disability	1 (None)	_	5 (Fatal)	15					
nhanes	1,914	Health status	1 (Excellent)	_	5 (Poor)	26					
wines	4,893	Quality	1 (Moderate)	_	6 (High)	11					

Table 4.2: Summary of data sets, sorted in increasing order of sample size.

from all the other folds except for S^f . Then, I calculate the same performance measures of Section 4.1 using the held-out S^f :

$$MSE_{f} = \frac{1}{|S^{f}|} \sum_{i \in Sf} \sum_{m=1}^{M} \left[\mathbb{1} \left(Y_{i} = m \right) - \hat{p}_{m,f} \left(X_{i} \right) \right]^{2}$$
(4.4)

$$MAE_{f} = \frac{1}{|S^{f}|} \sum_{i \in S^{f}} \sum_{m=1}^{M} \left| \mathbb{1} \left(Y_{i} = m \right) - \hat{p}_{m,f} \left(X_{i} \right) \right|$$
(4.5)

$$RPS_{f} = \frac{1}{|\mathcal{S}^{f}|} \sum_{i \in \mathcal{S}^{f}} \frac{1}{M-1} \sum_{m=1}^{M} \left[\mathbb{1} \left(Y_{i} \le m \right) - \hat{\mu}_{m,f} \left(X_{i} \right) \right]^{2}$$
(4.6)

with $\hat{p}_{m,f}(\cdot)$ the estimated conditional probabilities using all the other folds except for S^f , and $\hat{\mu}_{m,f}(x) = \sum_{j=1}^m \hat{p}_{j,f}(x)$ the estimated cumulative distribution function. Finally, I repeat this process ten times. This approach eliminates the dependence of the results on a particular training-validation sample split.

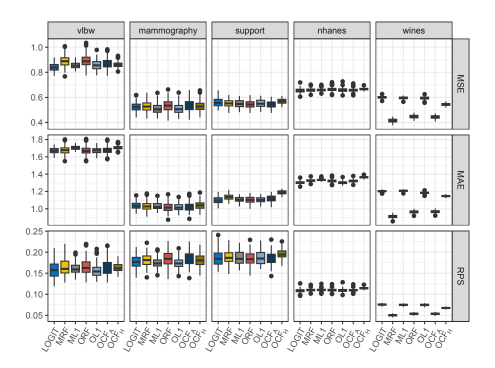


Figure 4.1: Prediction performance on real data sets. Each row contains boxplots showing the median and interquartile range of the estimated mean squared error (upper row), mean absolute error (mid row), and ranked probability score (lower row). Each column refers to a different data set, with the data set name displayed at the top of each column. Data sets are sorted according to their sample size.

Figure 4.1 reports the results. It displays boxplots showing the median and interquartile range of the estimated mean squared error (upper row), mean absolute error (mid row), and ranked probability score (lower row), together with their minima and maxima. Overall, the results indicate that OCF_A performs competitively compared to the other estimators. Contrary to the simulation results, there is no clear superiority between ordered and multinomial machine learning. The performance of the estimators does not exhibit significant differences, except in the case of the wines data set, where OCF_A emerges as one of the best estimators together with MRF and ORF. This result highlights the advantage of forest-based methods over those based on penalized regressions.

The cost of honesty resulting from constructing the forests with fewer observations is relatively small and varies with sample size. Across the three largest data sets, the cost ranges between 1%–22% in terms of median MSE, between 3%–18% in terms of median MAE, and between 4%–25% in terms of median RPS. In the two smallest data sets, minimal differences in performance between OCF_A and OCF_H are observed.

5 Conclusion

This paper proposes a novel estimator for ordered non-numeric outcomes, the *ordered* correlation forest. The estimator modifies a standard random forest splitting criterion to build a collection of forests, each estimating the conditional probability of a single class. Evidence from synthetic and real data sets shows that the proposed estimator features a superior prediction performance than alternative estimators.

Under an honesty condition, the proposed estimator inherits the asymptotic properties of random forests proven by Wager and Athey (2018), namely the consistency and asymptotic normality of its predictions. This allows valid inference about conditional probabilities

⁸The cross-validation exercise yields a smaller sample size compared to the simulation results presented in the previous section. Consequently, estimates of expected MSE, MAE, and RPS can be more imprecise and influenced by outliers. I report the distribution of the estimated MSE, MAE, and RPS using boxplots to provide a more robust assessment of the prediction performance of each estimator.

to be made using conventional methods. Moreover, transforming the weights induced by each forest provides a methodology to obtain estimation and inference about the covariates' marginal effects.

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