Monitoring Model Deterioration with Explainable Uncertainty Estimation via Non-parametric Bootstrap

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Abstract

Monitoring machine learning models once they are deployed is challenging. It is even more challenging to decide when to retrain models in real-case scenarios when labeled data is beyond reach, and monitoring performance metrics becomes unfeasible. In this work, we use non-parametric bootstrapped uncertainty estimation, the SHAP values provided by explainable uncertainty estimation, as a way to monitor the deterioration of machine learning models in deployment environments, as well as determine the source of model deterioration when target labels are not available. Classical methods are purely aimed at detecting distribution shift, which can lead to false positives, whereas our method achieves better performance than the current state-of-the-art. We show that both our model deterioration detection system and our uncertainty estimation method achieve better performance than the current state-of-the-art. Finally, we use explainable AI techniques to gain an understanding of the drivers of model deterioration. We release an open source Python package, doubt, which implements our proposed methods, as well as the code used to reproduce our experiments.

Introduction

Monitoring machine learning models in production is not an easy task. There are situations when the true label of the deployment data is available, and performance metrics can be monitored. But there are cases where it is not, and performance metrics are not so trivial to calculate once the model has been deployed. Model monitoring aims to ensure that a machine learning application in a production environment displays consistent behavior over time.

Being able to explain or remain accountable for the performance or the deterioration of a deployed model is crucial, as a drop in model performance can affect the whole business process (Mougan, Kanellos, and Gottron 2021), potentially having catastrophic consequences1. Once a deployed model has deteriorated, models are retrained using previous and new input data in order to maintain high performance. This process is called continual learning (Diethe et al. 2018) and it can be computationally expensive and put high demands on the software engineering system. Deciding when to retrain machine learning models is paramount in many situations.

Traditional machine learning systems assume that training data has been generated from a stationary source, but data is not static, it evolves. This problem can be seen as a distribution shift, where the data distributions of the training set and the test set differ. Detecting distribution shifts has been a longstanding problem in the machine learning (ML) research community (Shimodaira 2000; Sugiyama, Krauledat, and Müller 2007; Sugiyama and Müller 2005; Tasse 2017; Zadrozny 2004; Stolzenberg and Relles 1997; Heckman 1990; Cortes et al. 2008; Huang et al. 2006; He et al. 2014), as it is one of the main sources of model performance deterioration (Candela et al. 2009). Furthermore, data scientists in machine learning competitions claim that finding the train/validation split that better resembles the test (evaluation) distribution is paramount to winning a Kaggle competition (Guschin et al. 2018).

However, despite the fact that a shift in data distribution can be a source of model deterioration, the two are not identical. Indeed, if we shift a random noise feature we have caused a change in the data distribution, but we should not expect the performance of a model to decline when evaluated on this shifted dataset. Thus, we emphasize here that our focus is on model deterioration and not distribution shift, despite the correlation between the two.

Established ways of monitoring distribution shift when the real target distribution is not available are based on statistical changes either the input data (Diethe et al. 2018; Rabanser, Günnemann, and Lipton 2019) or on the model output (Garg et al. 2021). These statistical tests correctly detect univariate changes in the distribution but are completely independent of the model performance and can therefore be too sensitive, indicating a change in the covariates but without any degradation in the model performance. This can result in false positives, leading to unnecessary model retraining. It is worth noting that several authors have stated the clear need to identify how non-stationary environments affect the behavior of models (Diethe et al. 2018).

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Aside from merely indicating that a model has deteriorated, it can in some circumstances be beneficial to identify the cause of the model deterioration by detecting and explaining the lack of knowledge in the prediction of a model. Such explainability techniques can provide algorithmic transparency to stakeholders and to the ML engineering team (Mougan, Kanellos, and Gottron 2021; Bhatt et al. 2021; Koh and Liang 2017; Ribeiro, Singh, and Guestrin 2016; Sundararajan, Taly, and Yan 2017).

This paper’s primary focus is on non-deep learning models and small to medium-sized tabular datasets, a size of data that is very common in the average industry, where, non-deep learning-based models achieve state-of-the-art results (Grinsztajn, Oyallon, and Varoquaux 2022; Borisov et al. 2022; Elsayed et al. 2021).

Our contributions are the following:
1. We develop a novel method that produces prediction intervals using bootstrapping with theoretical guarantees, which achieves better coverage than previous methods on eight real-life regression datasets from the UCI repository (Dua and Graff 2017).
2. We use this non-parametric uncertainty estimation method to develop a machine learning monitoring system for regression models, which outperforms previous monitoring methods in terms of detecting deterioration of model performance.
3. We use explainable AI techniques to identify the source of model deterioration for both entire distributions as a whole as well as for individual samples, where classical statistical indicators can only determine distribution differences.
4. We release an open source Python package, doubt, which implements our uncertainty estimation method and is compatible with all scikit-learn models (Pedregosa et al. 2011).

Related Work

Model Monitoring

Model monitoring techniques help to detect unwanted changes in the behavior of a machine learning application in a production environment. One of the biggest challenges in model monitoring is distribution shift, which is also one of the main sources of model degradation (Candela et al. 2009; Diethe et al. 2018).

Diverse types of model monitoring scenarios require different supervision techniques. We can distinguish two main groups: Supervised learning and unsupervised learning. Supervised learning is the appealing one from a monitoring perspective, where performance metrics can easily be tracked. Whilst attractive, these techniques are often unfeasible as they rely either on having ground truth labeled data available or maintaining a hold-out set, which leaves the challenge of how to monitor ML models to the realm of unsupervised learning (Diethe et al. 2018). Popular unsupervised methods that are used in this respect are the Population Stability Index (PSI) and the Kolmogorov-Smirnov test (K-S), all of which measure how much the distribution of the covariates in the new samples differs from the covariate distribution within the training samples. These methods are often limited to real-valued data, low dimensions, and require certain probabilistic assumptions (Diethe et al. 2018; Malinin et al. 2021).

Another approach suggested by Lundberg et al. (2020b) is to monitor the SHAP value contribution of input features over time together with decomposing the loss function across input features in order to identify possible bugs in the pipeline as well as distribution shift. This technique can account for previously unaccounted bugs in the machine learning production pipeline but fails to monitor the model degradation.

It is worth noting that prior work (Garg et al. 2021; Jiang et al. .) has focused on monitoring models either on out-of-distribution data or in-distribution data (Neyshabur et al. 2017, 2019). Such a task, even if challenging, does not accurately represent the different types of data a model encounters in the wild. In a production environment, a model can encounter previously seen data (training data), unseen data with the same distribution (test data), and statistically new and unseen data (out-of-distribution data). That is why we focus our work on finding an unsupervised estimator that replicates the behavior of the model performance.

The idea of mixing uncertainty with dataset shift was introduced by Ovadia et al. (2019). Our work differs from theirs, in that they evaluate uncertainty by shifting the distributions of their dataset, where we aim to detect model deterioration under dataset shift using uncertainty estimation. Their work is also focused on deep learning classification problems, while we estimate uncertainty using model agnostic regression techniques. Further, our contribution allows us to pinpoint the features/dimensions that are main causes of the model degradation.

Garg et al. (2021) introduces a monitoring system for classification models, based on imposing thresholds on the softmax values of the model. Our method differs from theirs in that we work with regression models and not classification models, and that our method utilizes external uncertainty estimation methods, rather than relying on the model’s own “confidence” (i.e., the outputted logits and associated softmax values).

Rabanser, Günnemann, and Lipton (2019), presents a comprehensive empirical investigation of dataset shift, examining how dimensionality reduction and two-sample testing might be combined to produce a practical pipeline for detecting distribution shift in a real-life machine learning system. They show that the two-sample-testing-based approach performs best. This serves as a baseline comparison within our models, even if their idea is more focused on binary classification, whereas our works focus on building a regression indicator.

Uncertainty

Uncertainty estimation is being developed at a fast pace. Model averaging (Kumar and Srivastava 2012; Gal and Ghahramani 2016; Lakshminarayanan, Pritzel, and Blundell 2017; Arnez et al. 2020) has emerged as the most common approach to uncertainty estimation. Ensemble and sampling-based uncertainty estimates have been successfully applied to many use cases such as detecting misclassifications (Ren et al. 2019), out-of-distribution inputs (D’Angelo and Hen-
ning 2021), adversarial attacks (Carlini and Wagner 2017; Smith and Gal 2018), automatic language assessments (Mal- linin 2019) and active learning (Kirsch, Van Amersfoort, and Gal 2019). In our work, we apply uncertainty to detect and explain model performance for seen data (train), unseen and identically distributed data (test), and statistically new and unseen data (out-of-distribution).

Kumar and Srivastava (2012) introduced a non-parametric method to compute prediction intervals for any ML model using a bootstrap estimate, with theoretical guarantees. Our work is an extension of their work, where we take into account the model’s variance in the construction of the prediction intervals. The result, as we will see in the experiments section, is that such intervals have better coverage in such high-variance scenarios.

Barber et al. (2021) recently introduced a new non-parametric method of creating prediction intervals, using the Jackknife+. Our method differs from theirs in that we are using general bootstrapped samples for our estimates, rather than leave-one-out estimates. In the experimental, we will see that the two methods perform similarly, but that our method is again more accurate in a high-variance scenario.

Methodology

Evaluation of Deterioration Detection Systems

The problem we are tackling in this paper is evaluating and accounting for model predictive performance deterioration. To do this, we simulate a distribution shift scenario in which we have access to the true labels, which we can use to measure the model deterioration and thus evaluate the monitoring system. A naive simulation in which we simply manually shift a chosen feature of a dataset would not be representative, as the associated true labels could have changed if such a shift happened “in the wild”.

Therefore, we propose the following alternative approach. Starting from a real-life dataset \( D \) and a numerical feature \( F \) of \( D \), we sort the data samples of \( D \) by the value of \( F \), and split the sorted \( D \) in three equally sized sections: \( \{D_{\text{below}}, D_{\text{tr}}, D_{\text{upper}}\} \subseteq D \). The model is then fitted to the middle section \( D_{\text{tr}} \) and evaluated on all of \( D \). The goal of the monitoring system is to input the model, the labeled data segment \( D_{\text{tr}} \), and a sample of unlabelled data \( S \subseteq D \), and output a “monitoring value” which behaves like the model’s performance on \( S \). Such a prediction will thus have to take into account the training performance, generalization performance as well as the out-of-distribution performance of the model.

In the experimental section, we compare our monitoring technique to several other such systems. To enable comparison between the different monitoring systems, we standardize all monitoring values as well as the performance metrics of the model. From these standardized values, we can now directly measure the goodness-of-fit of the model monitoring system by computing the absolute difference between its (standardized) monitoring values and the (standardized) ground truth model performance metrics. Our chosen evaluation method is very similar to the one used by Garg et al. (2021). They focus on classification models and their systems output estimates of the model’s accuracy on the dataset. They evaluate these systems by computing the absolute difference between the system’s accuracy estimate and the actual accuracy that the model achieves on the dataset.

As we are working with regression models in this paper, we will only operate with a single model performance metric: mean squared error. We will introduce our monitoring system, which is based on an uncertainty measure, and will compare our monitoring system against statistical tests based on input data or prediction data. In that section, we will also compare our uncertainty estimation method to current state-of-art uncertainty estimation methods.

Uncertainty Estimation

In order to estimate uncertainty in a general way for all machine learning models, we use a non-parametric regression technique, which is an improvement of the technique introduced by (Kumar and Srivastava 2012). This method aims at determining prediction intervals for outputs of general non-parametric regression models using bootstrap methods.

Setting \( d \in \mathbb{N} \) to be the dimension of the feature space, we assume that the true model \( y : \mathbb{R}^d \rightarrow \mathbb{R} \) is of the form \( y(x) = \delta(x) + \varepsilon(x) \), where \( \delta : \mathbb{R}^d \rightarrow \mathbb{R} \) is a deterministic and continuously differentiable function, and the observation noise \( \varepsilon : \mathbb{R}^d \rightarrow \mathbb{R} \) is a uniform random field such that \( \varepsilon(x_1), \ldots, \varepsilon(x_t) \) are iid for any \( x_1, \ldots, x_t \in \mathbb{R}^d \), have zero mean and finite variance. We will assume that we have a data sample \( X \) of size \( N \), as well as a convergent estimator \( \hat{\delta}^{(n)} \) of \( \delta \), meaning the following:

**Definition 1** Let \( \hat{\delta}^{(n)} : \mathbb{R}^d \rightarrow \mathbb{R} \) be a function for every \( n \in \mathbb{N} \). We then say that \( \hat{\delta}^{(n)} \) is a convergent estimator of a function \( \delta : \mathbb{R}^d \rightarrow \mathbb{R} \) if:

1. \( \hat{\delta}^{(n)} \) is deterministic and continuous, for all \( n \in \mathbb{N} \).
2. There is a function \( \delta : \mathbb{R}^d \rightarrow \mathbb{R} \) such that \( \hat{\delta}^{(n)} \) converges pointwise to \( \delta \) as \( n \rightarrow \infty \).

We define an associated bias function \( \beta(x) := \delta(x) - \hat{\delta}(x) \). Note that in Kumar and Srivastava (2012) they assumed that \( \mathbb{E}[\hat{\delta}^{(n)}(x) - \delta(x)]^2 \rightarrow 0 \) for \( n \rightarrow \infty \), effectively meaning that the candidate model would be able to perfectly model the underlying distribution given enough data. It turns out that their method does not require this assumption, as we will see below. Aside from removing this assumption, the primary difference between our approach and Kumar and Srivastava (2012) is that our approach extends the latter by maintaining good coverage in a high-variance situation, as we will also see below. We start by rewriting the equation for the true model as follows:

\[
y(x) = \delta(x) + \varepsilon(x) = \hat{\delta}^{(N)}(x) + \beta(x) + \varepsilon_v^{(N)}(x) + \varepsilon(x),
\]

where \( \varepsilon_v^{(N)}(x) := \delta(x) - \beta(x) - \hat{\delta}^{(N)}(x) \) is the model variance noise. Note that

\[
\varepsilon_v^{(n)}(x) = \hat{\delta}^{(n)}(x) - \hat{\delta}^{(n)}(x) \rightarrow 0 \text{ as } n \rightarrow \infty.
\]

To produce correct prediction intervals we thus need to estimate the distribution of the observation noise, bias and model variance noise.
Estimating Model Variance Noise  To estimate the model variance noise term $\varepsilon_b^{(N)}(x)$ we adapt the technique in Kumar and Srivastava (2012) to our scenario, using a bootstrap estimate. Concretely, we bootstrap our dataset $B > 0$ times, fitting our model on each of the bootstrapped samples $X_b$ and generating bootstrapped estimates $\hat{\delta}_b^{(N)}(x)$ for every $b < B$. Centering the bootstrapped predictions as $m_b^{(N)}(x) := \mathbb{E}_b[\hat{\delta}_b^{(N)}(x)] - \hat{\delta}_b^{(N)}(x)$, we have that

\[ D_b[\hat{m}_b^{(N)}(x)] = D_b[\mathbb{E}_b[\hat{\delta}_b^{(N)}(x)] - \hat{\delta}_b^{(N)}(x)] \]
\[ \rightarrow D_X[\hat{\delta}(x) - \hat{\delta}(N)(x)] \]
\[ = D_X[\varepsilon_b^{(N)}(x)] \]

as $B \to \infty$, giving us our estimate of the model variance noise.

Estimating Bias and Observation Noise  We next have to estimate the bias $\beta(x)$ and the observation noise $\varepsilon(x)$. By rewriting (1) we get that that

\[ \beta(x) + \varepsilon(x) = y(x) - \hat{\delta}(N)(x) - \varepsilon_b^{(N)}(x), \]

so since we already have an estimate for $\varepsilon_b^{(N)}(x)$, it remains to estimate the residual $y(x) - \hat{\delta}(N)(x)$. In Kumar and Srivastava (2012) this was estimated purely using the training residuals without using any bootstrapping, whereas our approach will estimate the expected value of this residual via a bootstrap estimate, by using bootstrapped validation residuals $y(x) - \hat{\delta}_b(N)(x)$, where $x$ is not in the $b$’th bootstrap sample $X_b$. Concretely, we have that

\[ D_{(b, x \in X \setminus X_b)}[y(x) - \hat{\delta}_b(N)(x)] \]
\[ \rightarrow D_{(X, x \notin X)}[y(x) - \hat{\delta}(N)(x)] \]

as $B \to \infty$. An initial estimate is thus

\[ D_{(x, x \notin X)}[\hat{\beta}(x) + \varepsilon(x)] \]
\[ \approx D_{(b, x \in X \setminus X_b)}[y(x) - \hat{\delta}_b(N)(x) - m_b(N)(x)] \]

Denote (11) by $\text{valError}_b^{(N)}$. The problem with this approach is that the resulting prediction intervals arising from these validation errors are going to be too wide, as the bootstrap samples only contain on average 63.2% of the samples in the original dataset (Friedman et al. 2001), causing the model to have artificially large validation residuals. To fix this, we follow the approach in Friedman et al. (2001) and use the 0.632+ bootstrap estimate instead, defined as follows. We start by defining the no-information error

\[ \text{noInfoError}^{(N)} := \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} (y(x_i) - \hat{\delta}(x_j))^2, \]

corresponding to the mean-squared error if the inputs and outputs were independent. Next, define the associated training residuals $\text{trainError}_b^{(N)}$ as:

\[ D_{(b, x \in X_b)}[y(x) - \hat{\delta}_b^{(N)}(x) - (\mathbb{E}_b[\hat{\delta}_b^{(N)}(x)] - \hat{\delta}_b^{(N)}(x))] \]

Combining these two, we set the relative overfitting rate $\text{overfittingRate}_b^{(N)}$ to be:

\[ \text{valError}_b^{(N)} - \text{trainError}_b^{(N)} \]
\[ \text{noInfoError} - \text{trainError}_b^{(N)}, \]

which denotes how much we should weigh the validation error over the training error. In case of no overfitting, we get that $\text{valWeight}_b^{(N)} = 0.632$ and this reduces to the standard 0.632 bootstrap estimate (Friedman et al. 2001), whereas in case of severe overfitting the weight becomes 1 and thus only prioritizes the validation error.

Our final estimate of $\beta(x) + \varepsilon(x)$ is thus

\[ D_X[\hat{\beta}(x) + \varepsilon(x)] \approx D_b[\hat{\delta}_b^{(N)}], \]

where

\[ \hat{\delta}_b^{(N)} := (1 - \text{valWeight}_b^{(N)}) \times \text{trainError}_b^{(N)} + \text{valWeight}_b^{(N)} \times \text{valError}_b^{(N)}. \]

Note that this estimate is only an aggregate and is not specific to any specific value of $x$, as opposed to the model variance estimate in equation (4).

Prediction Interval Construction  Calculating the estimate of the prediction interval is then a matter of joining the results from the section of model variance noise and bias observation noise, in the same way as in Kumar and Srivastava (2012). As the estimate of $\beta(x) + \varepsilon(x)$ does not depend on any new sample, we can pre-compute this in advance by bootstrapping $B$ samples $X_b$, fit our model to each and calculate the $\hat{\delta}_b^{(N)}$ using Equation (16). Now, given a new data point $x_0$ and $\alpha \in (0, 1)$, we can estimate an $\alpha$ prediction interval around $x_0$ as follows. We again bootstrap $B$ samples $X_b$, fit our model to each and calculate the $\hat{\delta}_b^{(N)}(x_0)$ values. Next, we form the set $C^{(N)}(x_0) := \{ m_b(N)(x_0) + \hat{\delta}_b^{(N)} | b < B \}$, and our interval is then (start, end), where

\[ \text{start} := \hat{\delta}(x_0) - q_0 \left( C^{(N)}(x_0) \right) \]
\[ \text{end} := \hat{\delta}(x_0) + q_{1-\frac{\alpha}{2}} \left( C^{(N)}(x_0) \right), \]

with $q_0(C^{(N)}(x_0))$ being the $\alpha$th quantile of $C^{(N)}(x_0)$.

Detecting the Source of Uncertainty/Model Deterioration  Using uncertainty as a method to monitor the performance of an ML model does not provide any information on what features are the cause of the model degradation, only a goodness-of-fit to the model performance. We propose to solve this issue with the use of Shapley values.
We start by fitting a model $f_0$ to the training data, $X^{\text{train}}$. We next shift the test data by five standard deviations (call the shifted data $X^{\text{ood}}$) and compute uncertainty estimates $Z$ of $f_0$ on $X^{\text{ood}}$. We next fit a second model $g_0$ on $(X^{\text{ood}})$ to predict the uncertainty estimate $Z$, and compute the associated Shapley values (Lundberg et al. 2020b) of $g_0$. These Shapley values thus signify which features are the ones contributing the most to the uncertainty values. With the correlation between uncertainty values and model deterioration that we hope to conclude from the experiment described in the experimental section, this also provides us with a plausible cause of the model deterioration, if deterioration has taken place. Particularly, this methodology can be extended to large-scale datasets and deep learning-based models.

**Experiments**

Our experiments have been organized into three main groups: Firstly, we compare our non-parametric bootstrapped estimation method with the previous state-of-the-art, Kumar and Srivastava (2012) and Barber et al. (2021). Secondly, we assess the performance of our proposed uncertainty method for monitoring the performance of a machine learning model. And then, we evaluate the usability of the explainable uncertainty for identifying the features that are driving model degradation in local and global scenarios. In the main body of the paper, we present the results over several real-world datasets in the appendix we provide the experiments on synthetic datasets that exhibits, non-linear and linear behavior.

**Uncertainty Method Comparison**

To demonstrate the accuracy of our prediction intervals introduced in the uncertainty estimation section, we compare the coverage of the intervals with the NASA method from Kumar and Srivastava (2012) on eight regression datasets from the UCI repository (Dua and Graff 2017). The statistics of these datasets can be seen in Table 1.

| Dataset                | # Samples | # Features |
|------------------------|-----------|------------|
| Airfoil Self-Noise     | 1,503     | 5          |
| Bike Sharing           | 17,379    | 16         |
| Concrete Strength      | 1,030     | 8          |
| QSAR Fish Toxicity     | 908       | 6          |
| Forest Fires           | 517       | 12         |
| Parkinsons             | 5,875     | 22         |
| Power Plant            | 9,568     | 4          |
| Protein                | 45,730    | 9          |

Table 1: Statistics of the regression datasets used in this paper.

We split each of the eight datasets into a 90/10 train/test split, uniformly at random. Next, we fit a linear regression, a decision tree, and a gradient boosting decision tree on the training split. We chose these three models to have an example of a model with large bias (the linear regression model), a model with large variance (the decision tree model), and an intermediate model that achieves state-of-the-art performance in many tasks, the gradient boosting model. We will use the xgboost (Chen and Guestrin 2016) implementation of the gradient boosting model. After fitting the three models we compute $\alpha$-prediction intervals for $\alpha \in \{0.75, 0.76, \ldots, 0.99\}$, using our “Doubt” prediction intervals, the “NASA” prediction intervals from (Kumar and Srivastava 2012) as well as the “MAPIE” prediction intervals from (Barber et al. 2021), the latter implemented with the MAPIE package\(^3\). We can then compare the coverage of the three methods on the eight test sets.

As the goal of an $\alpha$ prediction interval is to have a coverage of $\alpha$, we can measure the performance of a prediction interval system by reporting the absolute difference between the actual coverage of the interval and this ideal coverage $\alpha$. In Table 2 we report the mean and standard deviations of these absolute differences, for each of the three model architectures. We have performed pairwise two-tailed paired t-tests on all absolute differences, and the best-performing prediction interval methods are marked in bold for each model architecture.

We see (cf. Table 2) that there is no significant difference between the three methods in the high bias case with the linear regression model. In the case of the XGBoost model, a model with higher variance, both the MAPIE and Doubt methods outperform the NASA method, but there is no significant difference between the MAPIE method and the Doubt method in this case. In the high-variance scenario with the decision tree, however, the Doubt intervals achieve significantly better coverage than both of the other two methods.

**Evaluating Model Deterioration**

The scenario we are addressing is characterized by regression data sets that have statistically seen data (train data), iid statistically unseen data (test data), and out-of-distribution data. Following the open data for reproducible research guidelines described in Arnold et al. (2019) and for measuring the performance of the proposed methods, we have used eight open-source datasets (cf. Table 1) for an empirical comparison coming from the UCI repository (Dua and Graff 2017). As described in the methodology, in order to benchmark our algorithm we, for each feature $F$ in each dataset $D$, sort $D$ according to $F$ and split $D$ into three equally sized sections $\{D_{\text{below}}, D_{\text{tr}}, D_{\text{upper}}\} \subseteq D$. We then train the model on $D_{\text{tr}}$ and test the performance of all of $D$. In this way we obtain a mixture of train, test, and out-of-distribution data, allowing us to evaluate our monitoring techniques in all three scenarios.

In evaluating a monitoring system we need to make a concrete choice of the sampling method to get the unlabelled data $S \subseteq D$. We are here using a rolling window of fifty samples, which has the added benefit of giving insight into the performance of the monitoring system on each of the three sections $D_{\text{lower}}, D_{\text{tr}}$ and $D_{\text{upper}}$ (cf. Figure 1).

We compare our monitoring system using the uncertainty estimation method against: (i) two classical statistical methods on input data: the Kolmogorov-Smirnov test statistic (K-S) and the Population-Stability Index (PSI) (Diethe et al. 2018), (ii) a Kolmogorov-Smirnov statistical test on the predictions between train and test (Garg et al. 2021) that we

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\(^3\)https://github.com/scikit-learn-contrib/MAPIE
Table 2: A comparison of different prediction interval methods, where the metric used is the mean absolute deviation from the ideal coverage (lower is better), with its associated standard deviation. Here NASA is the method described in Kumar and Srivastava (2012), MAPIE is the method from Barber et al. (2021) and Doubt is our method. The best results for each model architecture are shown in bold.

| Model     | Linear Regression | XGBoost       | Decision Tree |
|-----------|-------------------|---------------|---------------|
| NASA      | 3.854 ± 5.380     | 20.216 ± 12.405 | 20.669 ± 9.771 |
| MAPIE     | 4.207 ± 4.755     | 5.264 ± 4.258 | 7.788 ± 4.782 |
| Doubt     | 3.917 ± 4.870     | 4.861 ± 3.938 | 5.137 ± 3.984 |

denominate prediction shift and (iii) the previous state-of-the-art uncertainty estimate MAPIE. We evaluate the monitoring systems on a variety of model architectures: generalized linear models, tree-based models as well as neural networks.

The average performance across all datasets can be found in Table 3. From these we can see that our methods outperform K-S and PSI in all cases except for the Random Forest case, where our method is still on par with the best method, in that case, K-S. We have included a table with each dataset and all the estimators in the appendix, where it can be seen that both K-S and PSI easily identify a shift in the distribution but fail to detect when the model performance degrades, giving too many false positives.

### Detecting the Source of Uncertainty

For this experiment, we make use of two datasets: a synthetic one (see the appendix) and the popular House Prices regression dataset 4, where the goal is to predict the selling price of a given property. We select two of the features that are the most correlated with the target, GrLivArea and TotalBsmtSF, and also create a new feature of random noise, to have an example of a feature with minimum correlation with the target. A model deterioration system should therefore highlight the GrLivArea and TotalBsmtSF features, and not highlight the random features.

Concretely, we compute an estimation of the Shapley values using TreeSHAP (Lundberg et al. 2020b), which is an efficient estimation approach values for tree-based models, that allows for this second model to identify the features that are the source of the uncertainty, and thus also provide an indicator for what features might be causing the model deterioration.

We fitted an MLP on the training dataset, which achieved a $R^2$ value of 0.79 on the validation set. We then shifted all three features by five standard deviations and trained a gradient boosting model on the uncertainty values of the MLP on the validation set, which achieves a good fit (an $R^2$ value of 0.94 on the hold-out set of the validation). We then compare the SHAP values of the gradient boosting model with the PSI and K-S statistics for the individual features.

In Figure 2, classical statistics and SHAP values to detect the source of the model deterioration are compared. We see that the PSI and K-S value correctly capture the shift in each of the three features (including the random noise). On the other hand, our SHAP method highlights the two substantial features (GrLivArea and TotalBsmtSF) and correctly does not assign a large value to the random feature, despite the distribution shift.

Figure 3 shows features contributing to pushing the model output from the base value to the model output. Features pushing the uncertainty prediction higher are shown in red, and those pushing the uncertainty prediction lower are in red.

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4See the appendix for a more detailed table.

4https://www.kaggle.com/c/house-prices-advanced-regression-techniques
In this work, we have provided methods and experiments to monitor and identify machine learning model deterioration via non-parametric bootstrapped uncertainty estimation methods, and use explainability techniques to explain the source of the model deterioration.

Our monitoring system is based on a novel uncertainty estimation method, which produces prediction intervals with theoretical guarantees and which achieves better coverage than the current state-of-the-art. The resulting monitoring system more accurately detects model deterioration than methods using classical statistics. Finally, we used SHAP values in conjunction with these uncertainty estimates to identify the features that are driving the model deterioration at both a global and local level, and qualitatively showed that these more accurately detect the source of the model deterioration compared to classical statistical methods.

**Limitations:** We emphasize here that due to computationally limitations, we have only benchmarked on datasets of relatively small to medium size (cf. Table 1), and further work needs to be done to see if these results are also valid for datasets of significantly larger size. This work also focused on tabular data and non-deep learning models.

**Reproducibility Statement**

To ensure reproducibility of our results, we make the data, data preparation routines, code repositories, and methods publicly available. Our novel uncertainty methods are included in the open-source Python package doubt. We describe the system requirements and software dependencies of our experiments. Our experiments were run on an 8 vCPU server with 60 GB RAM.

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Table 3: Performance of model monitoring systems for model deterioration for a variety of model architectures on eight regression datasets from the UCI repository (Dua and Graff 2017). The scores are the means and standard deviations of the absolute deviation from the true labels on \( D_{lower} \) and \( D_{upper} \) (lower is better). K-S and PSI are the monitoring systems obtained by computing the Kolmogorov-Smirnov test values and the Population Stability Index, respectively, Prediction Shift is the statistical comparison of the model prediction, and Doubt is our method. The best results for each model architecture are shown in bold. See the Appendix for all the raw scores.

| Method        | Linear Reg. | Poisson | Decision Tree | Random Forest | Gradient Boosting | MLP   |
|---------------|-------------|---------|---------------|---------------|-------------------|-------|
| PSI           | 0.87 ± 0.08 | 0.93 ± 0.08 | 0.97 ± 0.10 | 0.95 ± 0.08   | 0.95 ± 0.08   | 0.84 ± 0.16 |
| K-S           | 0.81 ± 0.10 | 0.94 ± 0.20 | 0.52 ± 0.12 | 0.50 ± 0.12   | 0.61 ± 0.19   | 0.72 ± 0.22 |
| PredictionShift | 0.86 ± 0.13 | 1.00 ± 0.15 | 0.80 ± 0.14 | 0.73 ± 0.18   | 0.75 ± 0.20   | 0.74 ± 0.22 |
| MAPE          | 0.77 ± 0.10 | 0.83 ± 0.18 | 0.60 ± 0.16 | 0.86 ± 0.15   | 0.73 ± 0.18   | 0.74 ± 0.38 |
| Doubt         | 0.71 ± 0.14 | 0.79 ± 0.14 | 0.49 ± 0.10 | 0.74 ± 0.18   | 0.58 ± 0.23   | 0.68 ± 0.38 |

Figure 3: Individual explanation that displays the source of uncertainty for one instance. The previous method allowed only for comparison between distributions, now with explainable uncertainty, we are able to account for individual instances. In red, features pushing the uncertainty prediction higher are shown; in blue, those pushing the uncertainty prediction lower.

See the Appendix for all the raw scores.
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