Boosted Conformal Prediction Intervals

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Abstract

This paper introduces a *boosted conformal procedure* designed to tailor conformalized prediction intervals toward specific desired properties, such as enhanced conditional coverage or reduced interval length. We employ machine learning techniques, notably gradient boosting, to systematically improve upon a predefined conformity score function. This process is guided by carefully constructed loss functions that measure the deviation of prediction intervals from the targeted properties. The procedure operates post-training, relying solely on model predictions and without modifying the trained model (e.g., the deep network). Systematic experiments demonstrate that starting from conventional conformal methods, our boosted procedure achieves substantial improvements in reducing interval length and decreasing deviation from target conditional coverage.

1 Introduction

Black-box machine learning algorithms have been increasingly employed to inform decision-making in sensitive applications. For instance, deep convolutional neural networks have been applied to diagnose skin cancer [14], and AlphaFold has been utilized in the development of malaria vaccines [24, 25]; here, scientists have employed AlphaFold to predict the structure of a key protein in the malaria parasite, facilitating the identification of potential binding sites for antibodies that could prevent the transmission of the parasite [25]. These instances highlight the critical need for understanding prediction accuracy, and one popular approach to quantify the uncertainty associated with general predictions relies on the construction of prediction sets guaranteed to contain the target label or response with high probability. Ideally, we would like the coverage to be valid conditional on the values taken by the features of the predictive model (e.g., patient demographics).

Conformal prediction [3] stands out as a flexible calibration procedure that provides a wrapper around any black-box prediction model to produce valid prediction intervals. Imagine we have a data set $\{(X_i,Y_i)\}_{i=1}^n$ and a test point (X_{n+1},Y_{n+1}) drawn exchangeably from an unknown, arbitrary distribution P (e.g. the pairs (X_i,Y_i) may be i.i.d.). Taking the data set and the observed features X_{n+1} as inputs, conformal prediction forms a prediction interval $C_n(X_{n+1})$ for Y_{n+1} with valid marginal coverage, i.e. such that $\mathbb{P}(Y_{n+1} \in C_n(X_{n+1})) = 0.95$ or any nominal level specified by the user ahead of time. This is achieved by means of a conformity score E(x,y;f), where (x,y) represents a data point while f represents any aspects of the distribution that we have estimated. For instance, the score may be given by the magnitude of the prediction error $|y - \hat{\mu}(x)|$, where $\hat{\mu}(x)$ represents the model prediction of the expected outcome, in which case f is simply $\hat{\mu}$. Roughly, we

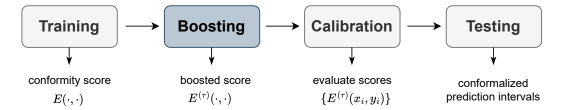


Figure 1: Illustration of the boosted conformal prediction procedure. We introduce a boosting stage between training and calibration, where we boost τ rounds on the conformity score function $E(\cdot,\cdot)$ and obtain the boosted score $E^{(\tau)}(\cdot,\cdot)$. The number of boosting rounds τ is selected via cross validation. A detailed description of the procedure is presented in Algorithm 1.

would include y in the prediction interval if $E(X_{n+1}, y; f)$ does not take on an atypical value when compared with $\{E(X_i, Y_i; f)\}$, $i = 1, \ldots, n$. Selecting an appropriate conformity score is akin to choosing a test statistic in statistical testing, where two statistics may yield the same Type I error rate yet differ substantially in other aspects of performance.

One central issue is that while the conformal procedure guarantees marginal coverage, it does not extend similar guarantees to other desirable inferential properties without additional assumptions. In response, researchers have introduced a variety of conformity scores, including the locally adaptive (Local) conformity score [16], the conformalized quantile regression (CQR) conformity score [18], and its variants, CQR-m [22] and CQR-r [23]. Among these, CQR has often demonstrated superior empirical performance in terms of both interval length and conditional coverage [18].

This paper introduces a boosting procedure aimed at enhancing an arbitrary score function.¹ By employing machine learning techniques, namely, gradient boosting, our objective is to modify the Local or CQR score functions (or other baselines) to reduce the average length of prediction intervals or improve conditional coverage while maintaining marginal coverage. While this paper focuses primarily on length and conditional coverage, our methods can be tuned to optimize other criteria; we elaborate on this in Section 7.

Our boosted conformal procedure searches within a family of generalized scores for a score achieving a low value of a loss function adapted to the task at hand. Specifically, to evaluate the conditional coverage of prediction intervals, we build a loss function that maximizes deviation from the target coverage rate in the leaves of a shallow contrast tree [21]. Searching within a strategically designed family of score functions, rather than directly retraining or fine-tuning the fitted model under the task-specific loss function, yields greater flexibility and avoids the costs associated with retraining or fine-tuning. Further, this boosting process is executed post-model training, requiring only the model predictions and no direct access to the training algorithm.

Source code for implementing the boosted conformal procedure is available online at https://github.com/ran-xie/boosted-conformal. Details regarding the acquisition and preprocessing of the real datasets are also provided in the GitHub repository.

2 The split conformal procedure

We begin by outlining the key steps of the split conformal procedure applied to a family $\{(X_i, Y_i)\}_{i=1}^n$ of exchangeable samples (e.g., i.i.d.).

- Training. Randomly partition [n] into a training set I_1 and a calibration set I_2 . On the training set, train a model by means of an algorithm \mathbb{A} to produce a conformity score function $E(\cdot,\cdot;f)$. The structure of this score function is predetermined, whereas the model f is learned from \mathbb{A} . An example of a conformity score is $E(x,y;f)=|y-\hat{\mu}(x)|$, where $\hat{\mu}(x)$ is a learned regression function so that f is here simply $\hat{\mu}$.
- Calibration. Evaluate the function $E(\cdot,\cdot;f)$ on each instance in the calibration set and obtain scores $\{E_i\}_{i\in I_2}$, with each $E_i=E(X_i,Y_i;f)$. The $(1-\alpha)$ th empirical quantile

¹An implementation of the boosted conformal procedure (BoostedCP) is available online at https://github.com/ran-xie/boosted-conformal.

²The term 'score' will henceforth refer to the conformity score unless stated otherwise.

of the score, $Q_{1-\alpha}(E, I_2)$, is calculated as

$$Q_{1-\alpha}(E, I_2) = \inf\{z : \mathbb{P}\left(Z \le z\right) \ge 1 - \alpha\},\$$

where Z follows the distribution $\frac{1}{|I_2|+1}(\delta_{\infty}+\sum \delta_{E_i})$, and δ_a is a point mass at a.

• Testing. For a new observation X_{n+1} , output the conformalized prediction interval

$$C_n(X_{n+1}) = \{ y \in \mathbb{R} : E(X_{n+1}, y; f) \le Q_{1-\alpha}(E, I_2) \}.$$
 (1)

If ties between $\{E_i\}_{i\in I_2}$ occur with probability zero, it holds that

$$1 - \alpha \le \mathbb{P}(Y_{n+1} \in C_n(X_{n+1})) \le 1 - \alpha + \frac{1}{|I_2| + 1},\tag{2}$$

see [16]. By introducing additional randomization during the calibration step, the prediction interval can be tuned to obey $\mathbb{P}(Y_{n+1} \in C_n(X_{n+1})) = 1 - \alpha$, see [4]. This adjustment is not critical here and we omit the details.

Locally adaptive conformal prediction (Local for short) [16] introduces a score function that aims to make conformal prediction adapt to situations where the spread of the distribution of Y varies significantly with the observed features X. On the training set, run an algorithm A to fit two functions $\mu_0(\cdot)$ and $\sigma_0(\cdot)$, where $\mu_0(X)$ estimates the conditional mean $\mathbb{E}[Y\mid X]$, and $\sigma_0(X)$ the dispersion around the conditional mean, frequently chosen as the conditional mean absolute deviation (MAD), $\mathbb{E}[|Y-\mu_0(X)|\mid X]$. With $f=(\mu_0,\sigma_0)$, the locally adaptive (Local) score function is:

$$E(x, y; f) = |y - \mu_0(x)| / \sigma_0(x). \tag{3}$$

For a new observation X_{n+1} , the conformalized prediction interval (1) takes on the simplified expression $[\mu_0(X_{n+1}) - Q_{1-\alpha}(E, I_2)\sigma_0(X_{n+1}), \mu_0(X_{n+1}) + Q_{1-\alpha}(E, I_2)\sigma_0(X_{n+1})].$

Conformalized quantile regression (CQR) [17] also aims to adapt to heteroskedasticity by calibrating conditional quantiles, which often results in shorter prediction intervals. Apply quantile regression to produce a pair of estimated quantiles $(\hat{q}_{\alpha/2}(x),\hat{q}_{1-\alpha/2}(x))$, where $\hat{q}_{\beta}(X)$ is the estimated β th quantile of the conditional distribution of Y. The CQR score function is defined as

$$E(x, y; f) = \max\{\hat{q}_{\alpha/2}(x) - y, y - \hat{q}_{1-\alpha/2}(x)\},\tag{4}$$

where $f = (\hat{q}_{\alpha/2}, \hat{q}_{1-\alpha/2})$. For a new observation X_{n+1} , following (1) yields the prediction interval

$$\left[\hat{q}_{\alpha/2}(X_{n+1}) - Q_{1-\alpha}(E, I_2), \hat{q}_{1-\alpha/2}(X_{n+1}) + Q_{1-\alpha}(E, I_2)\right]. \tag{5}$$

Generalized conformity score families. To construct a Local conformity score, we estimate two functions $\mu_0(\cdot)$ and $\sigma_0(\cdot)$ to plug into (3). Since these components are constructed without looking at performance downstream, it is reasonable to imagine that other choices may enjoy enhanced properties. How then should we systematically select $\mu(\cdot)$ and $\sigma(\cdot)$? To address this, we define a generalized Local score family $\mathcal F$ containing all potential score functions of the form

$$\mathcal{F} := \{ E(\cdot, \cdot; f) : E(x, y; f) = |y - \mu(x)| / \sigma(x), \sigma(\cdot) > 0 \}, \tag{6}$$

where $f = (\mu, \sigma)$. For each $E(\cdot, \cdot; f) \in \mathcal{F}$, the conformalized prediction interval is given by

$$[\mu(X) - Q_{1-\alpha}(E, I_2)\sigma(X), \mu(X) + Q_{1-\alpha}(E, I_2)\sigma(X)]. \tag{7}$$

Turning to CQR, one notable limitation is the uniform adjustment of prediction intervals by the constant factor $Q_{1-\alpha}(E,I_2)$, as shown in (5). This approach is suboptimal in the presence of heteroskedasticity, as it applies an identical correction to prediction intervals of varying widths for each X=x. Thus, simply updating the fitted quantiles $(\hat{q}_{\alpha},\hat{q}_{1-\alpha/2})$ and plugging them into the original score function would be inadequate, as the structure of the original score imposes significant limitations on the effectiveness of conformalized prediction intervals. To address this, several variants including CQR-m [22] and CQR-r [23] have been proposed. Focusing on CQR-r, it employs a flexible score function, defined as $E(x,y;f) = \max\{\hat{q}_{\alpha/2}(x)-y,y-\hat{q}_{1-\alpha/2}(x)\}/(\hat{q}_{1-\alpha/2}(x)-\hat{q}_{\alpha/2}(x))$, with $f=(\hat{q}_{\alpha/2},\hat{q}_{1-\alpha/2},\hat{q}_{1-\alpha/2}-\hat{q}_{\alpha/2})$. Following (1), conformalized prediction intervals become

$$\left[\hat{q}_{\alpha/2}(X) - \hat{\sigma}(X)Q_{1-\alpha}(E, I_2), \hat{q}_{1-\alpha/2}(X) + \hat{\sigma}(X)Q_{1-\alpha}(E, I_2)\right],\tag{8}$$

where $\hat{\sigma}=\hat{q}_{1-\alpha/2}-\hat{q}_{\alpha/2}$. Intuitively, the adjusted score function allows prediction bands to adjust in proportion to their width, instead of adding a constant shift as in CQR. However, despite the intuitive appeal of adjusted scores as a seemingly more reasonable "allocation" of the conformal correction, empirical studies reveal that they do not result in narrower prediction intervals when compared to CQR [23]. This phenomenon is largely due to the uniform direction of the conformal adjustment, represented by $Q_{1-\alpha}(E,I_2)$, across all observations. In particular, if $Q_{1-\alpha}(E,I_2)<0$, indicating that the true target y predominantly lies within the estimated quantile range $[\hat{q}_{\alpha/2},\hat{q}_{1-\alpha/2}]$, there is a uniform narrowing of the predicted interval across all samples.

In light of these insights, we propose a novel score family, \mathcal{H} , designed to augment the flexibility of the conformity score functions:

$$\mathcal{H} := \left\{ E(\cdot, \cdot; f) : E(x, y; f) = \max \left\{ \mu_1(x) - y, y - \mu_2(x) \right\} / \sigma(x), \mu_1(\cdot) \le \mu_2(\cdot), \sigma(\cdot) > 0 \right\}, (9)$$

where $f = (\mu_1, \mu_2, \sigma)$, which leads to conformalized prediction intervals of the form

$$[\mu_1(X) - \sigma(X)Q_{1-\alpha}(E, I_2), \mu_2(X) + \sigma(X)Q_{1-\alpha}(E, I_2)]. \tag{10}$$

Notably, $\ensuremath{\mathcal{H}}$ includes the Local, CQR, and CQR-r scores as special cases.

3 Boosted conformal procedure

It is clear from above that a model is trained to produce a conformity score $E(\cdot,\cdot;f)$; e.g., we may learn a regression function $\hat{\mu}(\cdot)$ to plug it into a score function $|y-\hat{\mu}(x)|$. To overcome the limitation of working with an arbitrarily selected score function, we introduce a boosting step before calibration, see Figure 1. In a nutshell, we use gradient boosting to iteratively improve upon a predefined score $E(\cdot,\cdot;f)$ now denoted as $E^{(0)}(\cdot,\cdot)$, where the superscript indicates the 0th iteration.

To achieve this, we construct a task-specific loss function ℓ , which takes a dataset $\mathcal D$ and a score function $E(\cdot,\cdot;f)$ as inputs, and outputs $\ell(E(\cdot,\cdot;f);\mathcal D)$ measuring how closely the conformalized prediction interval aligns with the analyst's objective. This loss function ℓ is designed to be differentiable with respect to each of the model components produced by the training algorithm. Importantly, it does not require knowledge of the gradient of f(x) with respect to x. In the example above, taking the labels as fixed, this means that for each feature $x_i \in \mathcal D$, $i=1,\ldots,n$, if we set $\hat y_i = \hat \mu(x_i)$, then the loss $\ell(E(\cdot,\cdot);\mathcal D)$ is a function of $\{\hat y_i\}_{i=1}^n$, and the derivative $\partial \ell(E(\cdot,\cdot);\mathcal D)/\partial \hat y_i$ is well defined. In Sections 5.1 and 6.1, we present examples of such derivatives.

Each boosting iteration updates the score function sequentially, employing a gradient boosting algorithm such as XGBoost [12] or LightGBM [15]. These algorithms accept as input a dataset \mathcal{D} , a base score function $E(\cdot,\cdot;f)$, a custom loss function ℓ , gradients of ℓ with respect to f (denoted $\nabla_f \ell$), and a number of boosting rounds τ . We may write the boosting procedure as

$$(E^{(0)}(\cdot,\cdot),\ldots,E^{(\tau)}(\cdot,\cdot)) = \text{GradientBoosting}(\mathcal{D}, E(\cdot,\cdot;f), \ell, \nabla_f \ell, \tau). \tag{11}$$

This yields a boosted score function $E^{(\tau)}(\cdot,\cdot)$, which is then used for calibration and for constructing prediction intervals. The number τ is calculated using k-fold cross-validation on the training dataset, selecting τ from potential values up to a predefined maximum T (e.g., 500). We partition the dataset into k folds and for each $j=1,\ldots,k$, we hold out fold j for sub-calibration and the remaining k-1 folds for sub-training. We apply T rounds of gradient boosting (11) on the sub-training data, generating T+1 candidate score functions $E_j^{(0)}(\cdot,\cdot),\ldots,E_j^{(T)}(\cdot,\cdot)$. Each score function is then evaluated on sub-calibration data, using the loss function ℓ to compute losses at all epochs, i.e., for each fold $j=1,\ldots k$,

$$\{L_i^{(t)}\}_{t=0}^T = \{\ell(E_i^{(t)}; \text{fold}_i)\}_{t=0}^T.$$

Last, τ is selected as the round that minimizes the average loss across all k folds:

$$\tau = \arg\min_{0 \le t \le T} \sum_{i=1}^{k} L_i^{(t)},\tag{12}$$

see Figure 2. This cross-validation step simulates the calibration step in conformal prediction and effectively prevents the overfitting of the score function.

Since boosting is conducted on the training data, the boosted procedure satisfies the same marginal coverage guarantee as the split conformal procedure, as formalized below.

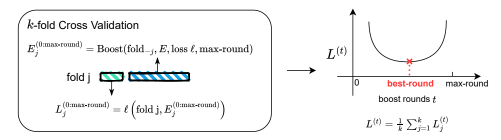


Figure 2: Schematic drawing showing the selection of the number of boosting rounds via cross-validation. Left: we hold out fold j, and use the remaining k-1 folds to generate candidate scores $E_j^{(t)}$, $t=0,\ldots,$ max-round. The performance of each score is evaluated on fold j using the loss function ℓ . Right: best-round minimizes the average loss across all k folds. A detailed description of the procedure is presented in Algorithm 1.

Proposition 3.1. Let $\{(X_i, Y_i)\}_{i=1}^m$ be the held out calibration set, and (X_{m+1}, Y_{m+1}) be a pair of new observation. If the m+1 samples are exchangeable, and ties between $\{E^{(\tau)}(X_i, Y_i)\}_{i=1}^m$ occur with probability zero, the confromalized prediction interval (1) computed from score function $E^{(\tau)}(\cdot, \cdot)$ satisfies the coverage guarantee (2).

Searching within generalized conformity score families. To update the Local score function (3), we search within the generalized score family \mathcal{F} (6). First, we initialize $\mu^{(0)} = \mu_0$ and $\sigma^{(0)} = \sigma_0$. After completing τ iterations of boosting on the training set, we obtain the boosted score function $E^{(\tau)}(x,y) = |y-\mu^{(\tau)}(x)|/\sigma^{(\tau)}(x)$. Notably, we can update any score function within \mathcal{F} . For instance, to update $E(x,y;f) = |y-\hat{\mu}(x)|$, we simply initialize $\mu^{(0)} = \hat{\mu}$, and take $\sigma^{(0)}$ to be the constant function equal to one. Similarly, to update the CQR score function (4), we search within the score family \mathcal{H} (9). First, we initialize a triple $\mu_1^{(0)} = \hat{q}_{\alpha/2}$, $\mu_2^{(0)} = \hat{q}_{1-\alpha/2}$, $\sigma^{(0)} = \hat{q}_{1-\alpha/2} - \hat{q}_{\alpha/2}$. After τ boosting rounds, we obtain the boosted score function $E^{(\tau)}(x,y) = \max\{\mu_1^{(\tau)}(x) - y, y - \mu_2^{(\tau)}(x)\}/\sigma^{(\tau)}(x)$.

Algorithm 1 Boosting stage

Input:

Training data $(X_i, Y_i) \in \mathbb{R}^p \times \mathbb{R}$, i = 1, ..., n; base conformity score function $E^{(0)}(\cdot, \cdot)$ Loss function ℓ ; target mis-coverage level $\alpha \in (0, 1)$

Number k of cross-validation folds; maximum boosting rounds T

Procedure:

Randomly divide $\{1, ..., n\}$ into k folds

for $j \leftarrow 1$ to k do

Set fold j as sub-calibration set, and the remaining k-1 folds as sub-training set

On the sub-training set, call GradientBoosting (11) to obtain candidate scores $\{E_j^{(t)}\}_{t=0}^T$ On the sub-calibration set, evaluate $L_i^{(t)} = \ell(E_i^{(t)}), t = 0, \dots, T$

end for

Set boosting rounds $\tau \leftarrow \arg\min_{t} \frac{1}{k} \sum_{j=1}^{k} L_{j}^{(t)}$ as in (12)

On the training set, call GradientBoosting (11) to obtain boosted functions $\{E^{(t)}\}_{t=0}^{\tau}$

Output:

Boosted conformity score function $E^{(\tau)}(\cdot,\cdot)$

4 Related Works

Adapting the classical conformal procedure to improve properties of the conformalized intervals has been one of the primary focuses of recent literature. Noteworthy contributions—including CF-GNN [28] and ConTr [26]—approach this problem by introducing modifications to the training stage

of the procedure. As outlined in Section 2, a model is trained to produce a score function $E(\cdot, \cdot; f)$. The model f usually depends on a set of model parameters, e.g., neural network parameters θ . Denote the trained model f by f_{θ} . CF-GNN and ConTr retrain or fine-tune the model by using a carefully constructed loss function, which may aim to produce narrower prediction intervals or prediction sets of reduced cardinality in classification problems. This process generates a new set of model parameters θ' . The new model $f_{\theta'}$ is then plugged into the *same* predefined conformity score function—namely CQR [28] or the adaptive prediction set score (APS) [26]—to produce $E(\cdot, \cdot, f_{\theta'})$.

There are two primary limitations. First, the score function imposes constraints on the properties of conformalized intervals as explained in Section 2. Our approach introduces more flexibility by constructing a family of generalized score functions that is a superset of $\{E(\cdot,\cdot;f_{\theta}):\theta\in\Theta\}$, where Θ is the parameter space of the training model. This family is strategically designed to contain an oracle conformity score ideally suited to the task at hand, e.g., achieving exact conditional coverage. Second, current methodologies necessitate fine-tuning or retraining models from scratch, requiring both access to the training model and significant computational resources. In contrast, our boosted conformal method operates directly on model predictions and circumvents these issues.

Conditional coverage of conformalized prediction intervals has also attracted significant interest, characterized by efforts to establish theoretical guarantees and achieve numerical improvements. Prior work established an impossibility result [8, 20], which states that exact conditional coverage in finite samples cannot be guaranteed without making assumptions about the data distribution. Subsequently, Gibbs et al. [27] developed a modified conformal procedure that guarantees conditional coverage for predefined protected sub-groups, i.e. subsets of the feature space. Our approach differs from the previous works by introducing a numerical method directly aimed at improving the conditional coverage, $\mathbb{P}(Y \in C_n(X)|X=x)$, across all potential values of x.

5 Boosting for conditional coverage

Maintaining valid marginal coverage, our goal is to produce a prediction interval C_n obeying

$$\mathbb{P}(Y \in C_n(X_{n+1})|X_{n+1} = x) \approx 1 - \alpha \tag{13}$$

for all possible values of x. To this end, we present a loss function that quantifies the conditional coverage rate of any prediction interval. Requiring merely a dataset \mathcal{D} and a prediction interval $C_n(\cdot)$ as inputs, it also serves as an effective evaluation metric, which may be of independent interest.

5.1 A measure for deviation from target conditional coverage

From now on, we let E be the score function $E(\cdot,\cdot;f)$. Set $\mathcal{D}=\{(X_i,Y_i)\}_{i=1}^n$ and denote by $C_n(\cdot)$ the conformalized prediction interval constructed from E. We shall assess the deviation of $C_n(\cdot)$ from the target conditional coverage by means of Contrast Trees [21]. As background, a contrast tree iteratively identifies splits within the feature space \mathcal{X} in a greedy fashion, aiming to maximize absolute within-group deviations from the target conditional coverage rate $(1-\alpha)$. For a subset R of the data point indices [n], let $\mathcal{D}_R=\{X_j,Y_j\}_{j\in R}$. The absolute within-group deviation is computed as

$$d(C_n(\cdot); \mathcal{D}_R) = \left| |R|^{-1} \sum_{j \in R} \mathbb{1}(Y_j \in C_n(X_j)) - (1 - \alpha) \right|.$$
 (14)

The overall empirical maximum deviation is then defined as

$$\ell_M(E; \mathcal{D}) = \max_{1 \le m \le M} d\left(C_n(\cdot); \mathcal{D}_{\hat{R}_m}\right), \tag{15}$$

where $\hat{R}_1 \cup \cdots \cup \hat{R}_M$ is a partition of [n], which itself depends on E and \mathcal{D} . Specifically, it is computed by running a contrast tree for M iterations. At each iteration, the algorithm not only seeks to isolate regions with large deviations but also discourages splits where any subset \hat{R}_m is too small.

To update score functions via gradient boosting as described in (11), we would need a differentiable approximation of the maximum deviation. To this end, we construct approximations for the following three components of the loss function. With an abuse of notation, in subsequent discussions, we shall employ the same notations to denote these differentiable approximations.

1. Approximation for the prediction interval $C_n(\cdot)$ in (14): the prediction interval is formulated as (7) for the generalized Local score, and as (10) for the generalized CQR score.

Denote the upper and lower limits of $C_n(\cdot)$ by $u(\cdot)$ and $l(\cdot)$. We approximate the empirical quantile $Q_{1-\alpha}(E,I_2)$ in $u(\cdot)$ and $l(\cdot)$ with a smooth quantile estimator $Q_{1-\alpha}^s$. Given r scalars $\{z_i\}_{i=1}^r$, $Q_{1-\alpha}^s$ is constructed as:

$$Q_{1-\alpha}^{s}(\{z_i\}_{i=1}^r) := \langle HD(r), s(\mathbf{z}) \rangle, \tag{16}$$

where $\langle \cdot, \cdot \rangle$ represents the dot product. Here, $\mathrm{HD}(r) = [W_{r,1},...,W_{r,r}]$ is the weight vector corresponding to the Harrel-Davis distribution-free empirical quantile estimator [1], and $s(\mathbf{z})$ is a differentiable ordering $\{\tilde{z}_{(i)}\}_{i=1}^r$, arranged in the ascending order. In practice, the derivative of $s(\mathbf{z})$ with respect to each z_i is given by the package developed in [19]. This approach is a smooth approximation of the Harrel-Davis quantile estimator $Q_{1-\alpha}^{\mathrm{HD}}$, constructed as a linear combination of the order statistics, $Q_{1-\alpha}^{\mathrm{HD}} = \langle \mathrm{HD}(r), \{z_{(i)}\}\rangle\rangle = \sum_{i=1}^r W_{r,i} z_{(i)}$, where $W_{r,i}$ takes the value $I_{(1-\alpha)(r+1),\alpha(r+1)}(i/r) - I_{(1-\alpha)(r+1),\alpha(r+1)}((i-1)/r)$ and $I_{a,b}(x)$ represents the incomplete beta function.

2. Approximation for absolute deviation d_i (14): the indicator function in (14) can be approximated by the product of two sigmoid functions,

$$\mathbb{1}(Y_j \in C_n(X_j)) = \mathbb{1}(u(X_j) - Y_j \ge 0)\mathbb{1}(Y_j - l(X_j) \ge 0)$$

$$\approx S_{\tau_1}(u(X_j) - Y_j)S_{\tau_1}(Y_j - l(X_j)),$$

where τ_1 is a parameter, trading off smoothness and quality of the approximation. The sigmoid function $S_{\tau_1}(x)$ is defined as $S_{\tau_1}(x) = (1+e^{-\tau_1 x})^{-1}$.

3. Approximation for maximum deviation: we employ a log-sum-exp function [2] to derive the differentiable approximation of ℓ_M as

$$\ell_M(E; \mathcal{D}) := \tau_2^{-1} \log \sum_{m=1}^M \exp(\tau_2 d_m(C_n(\cdot); \mathcal{D}_m)),$$
 (17)

where τ_2 is a parameter, serving the same purpose as τ_1 .

Here, we demonstrate calculating the derivative of the smooth approximation (17) with respect to each component of the generalized Local score, expanding it as follows:

$$\ell_{M}(E; \mathcal{D}) = \tau_{2}^{-1} \log \sum_{m=1}^{M} \exp\left(\tau_{2} \left| \left| R_{m} \right|^{-1} \sum_{j \in R_{m}} S_{\tau_{1}}(u(X_{j}) - Y_{j}) S_{\tau_{1}}(Y_{j} - l(X_{j})) - (1 - \alpha) \right| \right),$$

where

$$S_{\tau_1}(u(X_j) - Y_j) = \left(1 + \exp\left[-\tau_1(\mu_j + Q_{1-\alpha}^s(\{E_i\}_{i=1}^n)\sigma_j - Y_j)\right]\right)^{-1},$$

$$S_{\tau_1}(Y_j - l(X_j)) = \left(1 + \exp\left[-\tau_1(Y_j - \mu_j + Q_{1-\alpha}^s(\{E_i\}_{i=1}^n)\sigma_j)\right]\right)^{-1},$$

with $\mu_i = \mu(X_i)$, $\sigma_i = \sigma(X_i)$, $E_i = |Y_i - \mu_i|/\sigma_i$. As a result, for each feature X_i within \mathcal{D} , we can evaluate $\partial \ell_M(E; \mathcal{D})/\partial \mu_i$ and $\partial \ell_M(E; \mathcal{D})/\partial \sigma_i$ via the chain rule.

5.2 Boosting score functions for conditional coverage

Since the empirical maximum deviation ℓ_M (15) is non-differentiable, we opt for the differentiable approximation during the gradient boosting step (11). Nonetheless, we utilize the original ℓ_M to select the number of boosting rounds as in step (12) and to evaluate the conditional coverage of the conformalized prediction interval on the test set.

5.2.1 Theoretical guarantees

The oracle score function achieving conditional coverage as defined in (13) belongs to both proposed generalized score families.

Proposition 5.1 (Asymptotic expressiveness). Let $\{X_i, Y_i\}_{i=1}^n$ be i.i.d. with continuous joint probability density distribution. Under the split conformal procedure, for any target coverage rate $1-\alpha$, as $n \to \infty$, there exists (μ^*, σ^*) and $(\mu_1^*, \mu_2^*, \sigma^*)$ such that the corresponding generalized Local (6) and CQR (9) score functions recover conditional coverage at rate $1-\alpha$, as defined in (13).

It goes without saying that there is no reason to assume that the optimal μ^* corresponds to the conditional mean, median or any quantile of Y given X, or that the optimal σ^* corresponds to the standard deviation or the mean absolute deviation of Y given X, as in the original Local score (3). That said, our greedy strategy has no guarantee on global optimality and this is why the choice of the starting point—whether it is the Local or CQR score function—plays a role in the performance.

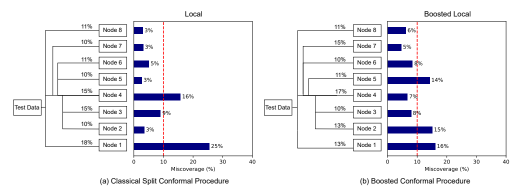


Figure 3: Comparison of test set conditional coverage evaluated on the dataset meps-19: (a) shows the classical Local-type conformal procedure and (b) our boosted Local-type conformal procedure. The target miscoverage rate is set to $\alpha=10\%$ (red). Miscoverage rate is computed at each leaf of the contrast tree, constructed to detect deviation from the target rate. Each leaf node is labeled with its size, namely, the fraction of the test set it represents.

Table 1: Test set maximum deviation loss ℓ_M evaluated on various conformalized intervals. The best result achieved for each dataset is highlighted in bold.

Max. Conditional Coverage Deviation (%), target miscoverage $\alpha=10\%$										
Dataset	Met	hod	Improvement	Met	hod	Improvement				
	Local	Boosted	improvement	CQR	Boosted	improvement				
bike	10.979	5.638	-48.65%	4.934	4.925	-0.17%				
bio	5.303	4.862	-8.31%	5.069	4.700	-7.29%				
community	25.755	13.466	-47.71%	12.688	12.105	-4.59%				
concrete	10.740	8.763	-18.40%	9.039	8.265	-8.56%				
meps-19	15.357	5.656	-63.17%	5.507	5.507	-0.00%				
meps-20	16.939	6.998	-58.69%	7.614	7.184	-5.65%				
meps-21	17.627	7.832	-55.57%	8.165	8.067	-1.20%				

5.2.2 Empirical results on real data

We apply our boosted conformal procedure to the 11 datasets previously analyzed in [23, 18, 22]. Details on the datasets are provided in Section A.6 in the Appendix. In each dataset, we randomly hold out 20% as test data. All experiments are repeated 10 times, starting from the data splitting. We refer to Section A.7 for details on the models and hyper-parameters we employ for the training and boosting stages.

We evaluate the conditional coverage of the prediction intervals as the maximum within-group deviations across a partitioned test set (15). This partition is obtained through a contrast tree algorithm described in Section 5.1. Figure 3 illustrates the comparison between miscoverage rates of prediction intervals at each leaf of the contrast tree. These intervals are derived under the classical Local conformal procedure and our boosted conformal procedure. Notably, the conditional coverage of the boosted prediction interval more closely aligns with the target rate $1-\alpha$.

The experiment results summarized in Table 1 indicate that applying boosting significantly enhances the performance of the baseline Local procedure. In contrast, boosting on CQR does not yield significant improvements—a sign that CQR already targets conditional coverage. (Before boosting, the prediction intervals generated by the baseline Local procedure exhibit conditional coverage deviations up to three times greater than those of the baseline CQR procedure.) It is noteworthy, however, that after boosting, the conditional coverage of the Local procedure improves to a level comparable to that of the boosted CQR procedure. While generally slightly less effective, nevertheless surpasses the performance of the boosted CQR procedure in two cases. Results on the remaining datasets are deferred to Tables A2 and A3.

6 Boosting for length

We begin by specifying the oracle prediction interval with minimum length. For a random variable Z, the High Density Region (HDR) at a specified significance level α , denoted as $\mathrm{HDR}_{\alpha}(Z)$, is defined as the shortest deterministic interval that covers Z with probability at least $1-\alpha$. The boundaries of $\mathrm{HDR}_{\alpha}(Z)$, the lower limit $Q_{l(\alpha)}$ and the upper limit $Q_{u(\alpha)}$, obey the condition $\mathbb{P}(Z \in [Q_{l(\alpha)}, Q_{u(\alpha)}]) \geq 1-\alpha$. For a pair of (X,Y) drawn from P, for every value of $x \in \mathbb{R}^p$, the oracle prediction interval at that point is expressed as

$$HDR_{\alpha}(Y|X=x) = \left[Q_{l(\alpha)}(Y|X=x), Q_{u(\alpha)}(Y|X=x)\right]. \tag{18}$$

Before introducing the boosting strategy, we present a word of caution against optimizing exclusively for this objective. Importantly, to maintain valid marginal coverage, the shortest prediction interval is prone to overcover when the spread of Y|X (the conditional distribution of Y given X) is small, and undercover when the spread of Y|X is large. This may be undesirable.

Similar to Proposition 5.1, we can show that the generalized score families exhibit the necessary expressiveness to contain the oracle conformity score, achieving optimal length while ensuring valid marginal coverage. The formal proof is deferred to Section A.3.

6.1 A measure for length

Consider a dataset $\mathcal{D} = \{(X_i, Y_i)\}_{i=1}^n$ and a score function E. Denote the corresponding conformalized prediction interval by $C_n(\cdot)$, with its quality measured by the average length:

$$\ell_L(E; \mathcal{D}) = n^{-1} \sum_{i=1}^n |C_n(X_i)|. \tag{19}$$

To derive a differentiable approximation of ℓ_L , we approximate the empirical quantile $Q_{1-\alpha}$ in the conformalized intervals (7) and (10) with the smooth quantile estimator $Q_{1-\alpha}^s$ constructed in (16). Here, we demonstrate calculating the derivative of the smooth approximation of ℓ_L with respect to each component of the generalized Local score, expanding it as follows based on the previously outlined approximation steps:

$$\ell_L(E; \mathcal{D}) = 2\bar{\sigma}Q_{1-\alpha}^s(\{E_i\}_{i=1}^n), \qquad E_i = |Y_i - \mu_i|/\sigma_i,$$

with $\mu_i = \mu(X_i)$, $\sigma_i = \sigma(X_i)$, $\bar{\sigma} = n^{-1} \sum_{i=1}^n \sigma_i$. As a result, for each feature X_i within \mathcal{D} , we can evaluate $\partial \ell_L(E;\mathcal{D})/\partial \mu_i$ and $\partial \ell_L(E;\mathcal{D})/\partial \sigma_i$ via the chain rule. For instance,

$$\frac{\partial \ell_L\left(E;\mathcal{D}\right)}{\partial \mu_i} = -2\bar{\sigma} \frac{\partial Q_{1-\alpha}^s(\{E_j\}_{j=1}^n)}{\partial E_i} \frac{\mathrm{sign}(Y_i - \mu_i)}{\sigma_i}.$$

6.2 Empirical results on real data

We apply our boosted conformal procedure to the same datasets described in Section 5.2.2. Detailed information on the models and hyperparameters used during the training and boosting stages can be found in Section A.7. Partial experiment results are summarized in Table 2. Notably, the boosting performance highlighted in bold exhibits significant improvement compared to previously documented results [17, 23]. We see a pronounced enhancement with the blog dataset; before boosting, the Local prediction intervals are on average 42% longer than those generated by CQR. After boosting, these intervals outperform the boosted CQR intervals by 32%. Using CQR as the baseline also yields substantial improvements, a decrease in averaged length exceeding 10% in six out of the eleven datasets. The meps-21 dataset, in particular, shows an improvement of up to 18% relative to the baseline. Results on the remaining datasets can be found in Tables A4 and A5. Figure 4 compares the conformalized prediction intervals derived from baseline Local and CQR scores with those obtained from the boosted scores. To effectively visualize the impact of boosting, we conduct a regression tree analysis on the training set to predict the label Y, setting the maximum number of tree nodes to four. This regression tree is then applied to the test set, allowing for a detailed comparison of the prediction intervals across each of the four distinct leaves.

7 Discussion

We introduced a post-training conformity score boosting scheme aiming to optimize for conditional coverage or length of the conformalized prediction interval. An intriguing avenue for future exploration involves simultaneously optimizing both length and conditional coverage, potentially trading

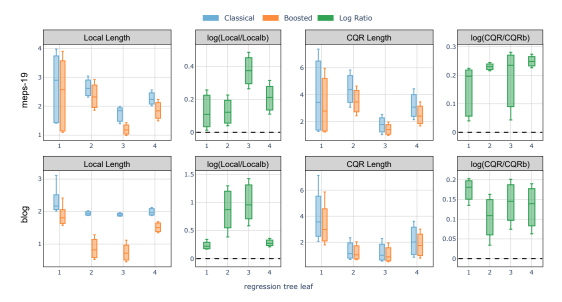


Figure 4: Comparison of test set average interval length evaluated on the meps-19 and blog datasets: classical Local and CQR conformal procedure versus the boosted procedures (abbreviated as 'Localb' and 'CQRb') compared in each of the 4 leaves of a regression tree trained on the training set to predict the label Y. A positive log ratio value between the regular and boosted interval lengths indicates improvement from boosting. The target miscoverage rate is set at $\alpha=10\%$.

Table 2: Test set average interval length ℓ_L evaluated on various conformalized prediction intervals. The best result achieved for each dataset is highlighted in bold.

Average Length, target miscoverage $\alpha=10\%$											
Dataset	Met	hod	Improvement	Met	hod	Immuorramant					
Dataset	Local	Boosted	mprovement	CQR	Boosted	Improvement					
blog	2.056	0.972	-52.74%	1.445	1.434	-0.71%					
facebook-1	1.896	1.383	-27.03%	1.198	1.072	-10.47%					
facebook-2	1.854	1.363	-26.51%	1.200	1.075	-10.41%					
meps-19	2.070	1.685	-18.60%	2.554	2.136	-16.35%					
meps-20	2.081	1.836	-11.80%	2.667	2.357	-11.62%					
meps-21	2.063	1.795	-12.99%	2.585	2.105	-18.55%					

off these objectives by incorporating user-specified weights [29]. Additionally, we can readily adapt our procedure to meet various application-specific objectives. For instance, we can optimize for conditional coverage on predefined feature groups, a common task in enhancing fairness in distributing social resources across different demographic groups [27]. Similarly, we can modify our procedure to reduce the length of prediction intervals for predefined label groups, which can be seen as reallocating resources to decrease uncertainty for certain groups at the expense of higher uncertainty for other groups [26]. Candidate loss functions tailored to these objectives are detailed in Section A.1. Lastly, the primary emphasis of this paper centers on the design of the conformity score boosting scheme and formalizing the optimization of conditional coverage in mathematical terms, leaving room for computational optimization to enhance performance and runtime efficiency. In essence, the gradient boosting algorithm in our procedure can be replaced with any gradient-based machine learning model. Thus, another interesting future direction would be to explore whether alternative algorithms could enhance performance.

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A Appendix

A.1 Candidate loss functions for additional application-specific objectives

Conditional coverage on predefined feature groups: this task can be viewed as a specialized application within our broader strategy of boosting for conditional coverage, as detailed in Section 5. There, the primary challenge was to develop a loss function that accurately measures deviations from the target conditional coverage rate. We achieved this by using contrast trees to identify partitions in the feature space that maximize these deviations, effectively identifying subgroups in need of protection. This process is simplified when the partitions correspond to prespecified groups, allowing us to continue using the empirical maximum deviation as a candidate loss function.

Consider a dataset $\mathcal{D}=\{(X_i,Y_i)\}_{i=1}^n$ and a score function E. Denote by $C_n(\cdot)$ the conformalized prediction interval constructed from E. Let G_1,\ldots,G_M be prespecified feature index groups. Within each set $\mathcal{D}_i=\{(X_j,Y_j)\}_{j\in G_i}$, compute the absolute deviation d_i as

$$d_i\left(C_n(\cdot); \mathcal{D}_i\right) = \left|\frac{1}{|G_i|} \sum_{j \in G_i} \mathbb{1}\left(Y_j \in C_n(X_j)\right) - (1 - \alpha)\right|. \tag{20}$$

The overall empirical maximum deviation is then defined as

$$\ell(E; \mathcal{D}) = \max_{1 \le i \le M} d_i(C_n(\cdot); \mathcal{D}_i). \tag{21}$$

Interval length conditional on predefined label groups: for a dataset $\mathcal{D} = \{(X_i, Y_i)\}_{i=1}^n$ and a score function E, let $\mathcal{Y}_1, \dots, \mathcal{Y}_M$ be the prespecified label groups. A natural minimization objective for balancing uncertainty among these groups is defined as:

$$\ell(E; \mathcal{D}) = \sum_{i=1}^{M} w_i \frac{1}{\sum_{j=1}^{n} \mathbb{1}(Y_j \in \mathcal{Y}_i)} \sum_{j=1}^{n} \mathbb{1}(Y_j \in \mathcal{Y}_i) |C_n(X_j)|,$$

where (w_1, \ldots, w_M) represents a set of user-specified weights.

A.2 Proof of Proposition 5.1

Our proof relies on the following lemma.

Lemma A.1 (Expressiveness). Given any sample pair X and Y with a continuous joint probability density distribution, and a prediction interval $[c_l(\cdot), c_u(\cdot)]$ with marginal coverage equal to $1 - \alpha$, there exist specific function sets: $(\mu(\cdot), \sigma(\cdot))$ for the Local type, and $(\mu_1(\cdot), \mu_2(\cdot), \tilde{\sigma}(\cdot))$ for the CQR type, such that asymptotically:

- 1. The conformalized prediction interval (7), derived using the generalized Local type conformity score $f_{\mu,\sigma}$, accurately recovers $[c_l(\cdot),c_u(\cdot)]$.
- 2. Similarly, the conformalized prediction interval (10), based on the generalized CQR type conformity score $E_{\mu_1,\mu_2,\tilde{\sigma}}$, also recovers $[c_l(\cdot),c_u(\cdot)]$.

Proof of Lemma A.1. Recall that the generalized Local score (6) characterized by (μ, σ) takes the form

$$E_{\mu,\sigma}(x,y) = \frac{|y - \mu(x)|}{\sigma(x)}.$$
(22)

Asymptotically, the conformalized prediction interval is given by

$$[\mu(X) - Q_{1-\alpha}(E_{\mu,\sigma})\sigma(X), \mu(X) + Q_{1-\alpha}(E_{\mu,\sigma})\sigma(X)]. \tag{23}$$

Here, $Q_{1-\alpha}$ represents the population quantile. Set

$$\mu(x) = \frac{c_l(x) + c_u(x)}{2}, \quad \sigma(x) = \frac{c_u(x) - c_l(x)}{2}.$$

By assumption, we have

$$\mathbb{P}(Y \in [c_l(X), c_u(X)]) = 1 - \alpha.$$

With a simple change of variables, the above inequality is equivalent to

$$\mathbb{P}\left(\left|\frac{Y - \mu(X)}{\sigma(X)}\right| \le 1\right) = 1 - \alpha.$$

In other words, this is equivalent to

$$Q_{1-\alpha}(E_{\mu,\sigma}) = 1.$$

We have thus proved the result for the generalized Local type conformity score $E_{\mu,\sigma}$. In the same spirit, we can prove the result for the generalized CQR type conformity score $E_{\mu_1,\mu_2,\tilde{\sigma}}$ by taking

$$\mu_1 = c_l + \frac{c_u(x) - c_l(x)}{2}, \quad \mu_2 = c_u - \frac{c_u(x) - c_l(x)}{2}, \quad \tilde{\sigma} = \frac{c_u(x) - c_l(x)}{2}.$$

Recall that a generalized CQR score function (9) characterized by (μ_1, μ_2, σ) is defined as:

$$E_{\mu_1,\mu_2,\sigma}(x,y) = \max \{\mu_1(x) - y, y - \mu_2(x)\} / \sigma(x), \tag{24}$$

which leads to the asymptotic conformalized prediction intervals of the form

$$[\mu_1(X) - \sigma(X)Q_{1-\alpha}(E_{\mu_1,\mu_2,\sigma}), \mu_2(X) + \sigma(X)Q_{1-\alpha}(E_{\mu_1,\mu_2,\sigma})]. \tag{25}$$

Plugging in μ_1 , μ_2 , $\tilde{\sigma}$ defined above, we immediately have

$$\mathbb{P}(Y \in [c_l(X), c_u(X)]) = 1 - \alpha$$

$$\iff \mathbb{P}(c_l(X) - Y \le 0, Y - c_u(X) \le 0) = 1 - \alpha$$

$$\iff \mathbb{P}\left(\frac{c_l(X) - Y + \tilde{\sigma}(X)}{\tilde{\sigma}(X)} \le 1, \frac{Y - c_u(X) + \tilde{\sigma}(X)}{\tilde{\sigma}(X)} \le 1\right) = 1 - \alpha$$

$$\iff Q_{1-\alpha}(E_{\mu_1, \mu_2, \sigma}(x, y)) = 1.$$

Proof of Proposition 5.1. It suffices to take $c_l(x) = Q_{\alpha/2}(Y|X=x), c_u(x) = Q_{1-\alpha/2}(Y|X=x)$ and apply Lemma A.1.

A.3 Boosting for length: theoretical guarantees

Similar to Proposition 5.1, we show in Proposition A.2 below that the generalized Local and CQR score families exhibit the necessary expressiveness to contain the oracle score, achieving optimal length while ensuring valid marginal coverage.

Proposition A.2 (Asymptotic expressiveness). *Under the assumptions of Proposition 5.1, for any target coverage rate* $1 - \alpha$, *as* $n \to \infty$, *the following statements hold true:*

- 1. There exists (μ^*, σ^*) such that the corresponding generalized Local score function (6) recovers the shortest oracle prediction interval (18).
- 2. There exists $(\mu_1^*, \mu_2^*, \sigma^*)$ such that the corresponding generalized CQR score function (9) recovers the shortest oracle prediction interval (18).

Proof of Proposition A.2. It suffices to take $c_l(x) = Q_{l(\alpha)}(Y|X=x)$, $c_u(x) = Q_{u(\alpha)}(Y|X=x)$ and apply Lemma A.1, where $Q_{l(\alpha)}$ and $Q_{u(\alpha)}$ are the lower and upper limits of the High Density Region defined in (18).

A.4 COR type conformity score boosting

A generalized CQR score function (9) is uniquely defined by a triple $(\mu_1(\cdot), \mu_2(\cdot), \sigma(\cdot))$. We will show how searching for a generalized CQR score can be reduced to searching for a Local generalized score. To begin with, we shall say that score functions are equivalent if they recover identical conformalized prediction intervals.

Definition A.3. Let $\{X_i, Y_i\}_{i=1}^n$, (X_{n+1}, Y_{n+1}) be i.i.d. with continuous joint probability density distribution, and let [n] be partitioned into a training set I_1 and a calibration set I_2 . Consider two conformity score functions, E_1 and E_2 , which produce conformalized prediction intervals $C_1(\cdot)$ and $C_2(\cdot)$, respectively. For any target coverage rate $1-\alpha$, E_1 and E_2 are equivalent if $C_1(\cdot)=C_2(\cdot)$ when marginal coverage rates $\mathbb{P}(Y_{n+1}\in C_1(X_{n+1}))$ and $\mathbb{P}(Y_{n+1}\in C_2(X_{n+1}))$ match.

Building on this definition, we are now equipped to establish the following equivalences: **Lemma A.4.** *Under the assumptions of Definition A.3, the following statements hold:*

- 1. For the CQR-r score function defined in Section 2, there is an equivalent generalized Local score function characterized by a pair $(\mu(\cdot), \sigma(\cdot))$, where $\mu = (\mu_1 + \mu_2)/2$, $\sigma = (\mu_2 \mu_1)/2$.
- 2. For any generalized Local score function characterized by the pair $(\mu(\cdot), \sigma(\cdot))$, there is an equivalent generalized CQR score function characterized by a triple $(\mu(\cdot), \mu(\cdot), \sigma(\cdot))$.

The proof of the above Lemma is deferred to Section A.5. Leveraging these equivalences, we carry out the boosted conformal procedure as follows: first, we initialize a triple $\mu_1^{(0)} = \hat{q}_{\alpha/2}$, $\mu_2^{(0)} = \hat{q}_{1-\alpha/2}$, $\sigma_1^{(0)} = \hat{q}_{1-\alpha/2} - \hat{q}_{\alpha/2}$, which characterizes the CQR-r score function. Next, we find an equivalent generalized Local score function characterized by a pair $(\mu^{(0)}, \sigma^{(0)})$ chosen according to Lemma A.4. After τ boosting rounds, we obtain the boosted pair $(\mu^{(\tau)}, \sigma^{(\tau)})$ and the corresponding score function. Finally, we recover an equivalent generalized CQR score function

$$E^{(\tau)}(x,y) = \max \left\{ \mu_1^{(\tau)}(x) - y, y - \mu_2^{(\tau)}(x) \right\} / \sigma_1^{(\tau)}(x),$$

characterized by the triple $(\mu_1^{(\tau)}, \mu_2^{(\tau)}, \sigma_1^{(\tau)})$ chosen according to Lemma A.4.

A.5 Proof of Lemma A.4

Recall that the generalized Local score (6) characterized by (μ, σ) takes the form

$$E_{\mu,\sigma}(x,y) = \frac{|y - \mu(x)|}{\sigma(x)}.$$
(26)

The conformalized prediction interval is given by

$$[\mu(X) - Q_{1-\alpha}(E_{\mu,\sigma}, I_2)\sigma(X), \mu(X) + Q_{1-\alpha}(E_{\mu,\sigma}, I_2)\sigma(X)]. \tag{27}$$

A generalized CQR score function (9) characterized by (μ_1, μ_2, σ) is defined as:

$$E_{\mu_1,\mu_2,\sigma}(x,y) = \max \{\mu_1(x) - y, y - \mu_2(x)\} / \sigma(x),$$

which leads to conformalized prediction intervals of the form

$$[\mu_1(X) - \sigma(X)Q_{1-\alpha}(E_{\mu_1,\mu_2,\sigma},I_2),\mu_2(X) + \sigma(X)Q_{1-\alpha}(E_{\mu_1,\mu_2,\sigma},I_2)].$$

1. Plugging in the triple $\mu_1(x)=\hat{q}_{\alpha/2},\,\mu_2(x)=\hat{q}_{1-\alpha/2},\,\sigma_1(x)=\hat{q}_{1-\alpha/2}-\hat{q}_{\alpha/2},$ which characterize the CQR-r score function, we have the conformalized prediction interval

$$[\mu_1(X) - \sigma_1(X)Q_{1-\alpha}(E_{\mu_1,\mu_2,\sigma_1},I_2),\mu_2(X) + \sigma_1(X)Q_{1-\alpha}(E_{\mu_1,\mu_2,\sigma_1},I_2)].$$

Set

$$\mu(X) = \frac{\mu_1(X) + \mu_2(X)}{2}, \sigma(X) = \frac{\mu_2(X) - \mu_1(X)}{2},$$

then the generalized Local conformity score $E_{\mu,\sigma}(x,y) = |y - \mu(x)|/\sigma(x)$ recovers conformalized prediction intervals of the form

$$\begin{split} \left[\mu(X) - \sigma(X)Q(E_{\mu,\sigma},I_2), \mu(X) + \sigma(X)Q(E_{\mu,\sigma},I_2)\right] \\ = & \left[\frac{\mu_1(X) + \mu_2(X)}{2} - \frac{\mu_2(X) - \mu_1(X)}{2}Q(E_{\mu,\sigma},I_2), \\ & \frac{\mu_1(X) + \mu_2(X)}{2} + \frac{\mu_2(X) - \mu_1(X)}{2}Q(E_{\mu,\sigma},I_2)\right] \\ = & \left[\mu_2(X) - (\mu_2(X) - \mu_1(X))\frac{Q(E_{\mu,\sigma},I_2) - 1}{2}, \\ & \mu_1(X) + (\mu_2(X) - \mu_1(X))\frac{Q(E_{\mu,\sigma},I_2) - 1}{2}\right] \\ = & \left[\mu_1(X) - \sigma_1(X)\frac{Q(E_{\mu,\sigma},I_2) - 1}{2}, \mu_2(X) + \sigma_1(X)\frac{Q(E_{\mu,\sigma},I_2) - 1}{2}\right]. \end{split}$$

From the monotonicity of the interval lengths with respect to the empirical quantiles, we have that the two score functions are equivalent by Definition A.3.

2. Let a generalized Local score function be $E_{\mu,\sigma}(x,y) = |y - \mu(x)|/\sigma(x)$. Then it suffices to observe that

$$|y - \mu(x)| = \max\{y - \mu(z), \mu(x) - y\}.$$

A.6 Additional information on real datasets

In Table A1, we provide the predicted label, dimensions, and source for each dataset. Data cleaning and preprocessing are in accordance with the methods described by Romano et al. [17].

Table A1: Datasets for our empirical analyses, with the predicted label, number of samples (n), and features (d).

Name	Label	n	d	Source
bike	bike rental counts	10886	18	[9]
bio	deviation of predicted from native protein structure	45730	9	[10]
blog	number of comments in the next 24 hours	52397	280	[11]
community	crime rate per community	1994	100	[7]
concrete	concrete compressive strength	1030	8	[5]
facebook-1	Facebook comment volume	40948	53	[5]
facebook-2	Facebook comment volume	81311	53	[13]
meps-19	utilization of medical services	15785	139	[30]
meps-20	utilization of medical services	17541	139	[31]
meps-21	utilization of medical services	15656	139	[32]
star	total student test scores up to the third grade	2161	39	[6]

All datasets, except for the meps and star data sets, are licensed under CC-BY 4.0. The Medical Expenditure Panel Survey (meps) data is subject to copyright and usage rules. The licensing status of the star dataset could not be determined.

A.7 Experimental Setup

In each dataset, we randomly hold out 20% as test data. The remaining data is divided into a training set and a calibration set, each taking up a proportion of γ and $1-\gamma$. We explore training ratios γ ranging from 10% to 90%. Results corresponding to the optimal value of the hyperparameter γ are recorded in Table 1, following the practice of Sesia er al. [23].

In the training stage, we employ the random forest regressor from Python's scikit-learn package to learn the baseline Local score function. The hyperparameters are the package defaults, except for the total number of trees, which we set to 1000, and the minimum number of samples required at a leaf node, which we set to 40, as recommended by Romano et al. [17]. For the baseline CQR score function, we adopt a black-box neural network quantile regressor with three fully connected layers and ReLU non-linearities, following the practice of Sesia et al. [23]. In the boosting stage, we set the hyper-parameters τ_1 , τ_2 in the approximated loss (17) to 50. The approximated loss is then passed to the Gradient Boosting Machine from Python's XGBoost package along with a base conformity score. We set the maximum tree depth to 1 to avoid overfitting and perform cross-validation for the number of boosting rounds, as outlined in Section 3. All other hyperparameters are set to package defaults.

All experiments were conducted on a dual-socket AMD EPYC 7502 32-Core Processor system, utilizing 8 of its 128 CPUs each time. The runtime for each dataset and random seed varies by dataset size, ranging from 10 minutes to 5 hours.

A.8 Additional results and error bars

In Tables A2 to A5, we present additional results on marginal coverage, maximum conditional coverage deviation (ℓ_M), and average interval length (ℓ_P) for each real dataset (including those not reported in Tables 1 and 2), both before and after boosting. Notably, in each case, boosting is applied to optimize either conditional coverage or average interval length. As a result, the non-targeted characteristic may or may not improve after boosting.

Table A2: Additional information on conformalized intervals obtained before and after boosting for conditional coverage with the Local conformity score as baseline. The target miscoverage rate is set to $\alpha=10\%$.

Dataset		ℓ_L	ℓ_M	(%)	Marginal Cov.(%)		
Dataset	Local	Boosted	Local	Boosted	Local	Boosted	
bike	1.775	2.201	10.979	5.638	89.927	89.646	
bio	1.602	1.614	5.303	4.862	90.024	90.093	
blog	2.080	3.403	51.353	48.591	89.995	90.040	
community	1.824	10.759	25.755	13.466	90.376	89.549	
concrete	1.058	1.062	10.740	8.763	90.583	91.359	
facebook-1	1.896	4.790	26.020	25.917	90.201	90.056	
facebook-2	1.881	2.273	42.807	42.437	89.966	89.989	
meps-19	2.074	2.926	15.357	5.656	90.120	90.497	
meps-20	2.102	2.778	16.939	6.998	89.963	90.031	
meps-21	2.069	2.537	17.627	7.832	90.064	90.054	
star	0.189	0.179	9.658	9.348	90.808	90.831	

Table A3: Additional information on conformalized intervals obtained before and after boosting for conditional coverage with the CQR conformity score as baseline. The target miscoverage rate is set to $\alpha=10\%$.

Dataset	ℓ_L		ℓ_M	$\ell_M(\%)$		Marginal Cov.(%)		
Dataset	CQR	Boosted	CQR	Boosted	CQR	Boosted		
bike	0.555	0.540	4.934	4.925	90.073	90.184		
bio	1.518	1.515	5.069	4.700	89.841	89.853		
blog	1.761	1.766	27.760	26.836	90.222	90.244		
community	1.718	1.740	12.688	12.105	90.340	90.194		
concrete	0.484	0.489	9.039	8.265	90.451	90.652		
facebook-1	1.374	1.371	13.407	13.255	90.465	90.247		
facebook-2	1.465	1.409	18.257	18.002	89.763	90.001		
meps-19	2.784	2.784	5.507	5.507	90.257	90.257		
meps-20	2.769	2.743	7.614	7.184	89.991	90.006		
meps-21	2.834	2.815	8.16	8.067	90.169	90.067		
star	0.199	0.209	9.728	9.630	91.085	91.339		

Table A4: Additional information on conformalized intervals obtained before and after boosting for length with the Local conformity score as baseline. The target miscoverage rate is set to $\alpha = 10\%$.

Dataset		ℓ_L	ℓ_M	$\ell_M(\%)$		Marginal Cov.(%)	
Dataset	Local	Boosted	Local	Boosted	Local	Boosted	
bike	1.775	1.360	22.590	19.616	89.927	89.862	
bio	1.562	1.514	5.995	5.791	89.937	89.962	
blog	2.056	0.972	52.440	54.858	89.988	89.978	
community	1.728	1.678	26.066	24.822	89.499	89.323	
concrete	1.010	0.698	11.029	10.518	90.631	90.728	
facebook-1	1.896	1.384	26.020	34.259	90.201	89.944	
facebook-2	1.854	1.363	42.624	50.697	90.020	89.972	
meps-19	2.070	1.685	18.626	14.623	90.054	90.070	
meps-20	2.081	1.836	17.897	14.643	89.869	89.849	
meps-21	2.063	1.795	18.795	13.324	89.914	89.920	
star	0.179	0.179	9.976	9.407	90.901	90.577	

Table A5: Additional information on conformalized intervals obtained before and after boosting for length with the CQR conformity score as baseline. The target miscoverage rate is set to $\alpha = 10\%$.

Dataset		ℓ_L	ℓ_M	$\ell_M(\%)$		Marginal Cov.(%)	
Dataset	CQR	Boosted	CQR	Boosted	CQR	Boosted	
bike	0.553	0.489	9.530	10.092	90.041	90.418	
bio	1.516	1.468	5.408	5.265	89.670	89.880	
blog	1.445	1.434	29.875	34.011	90.149	90.260	
community	1.693	1.699	14.006	13.536	89.499	89.699	
concrete	0.391	0.393	10.342	10.700	88.932	89.223	
facebook-1	1.198	1.073	20.132	30.901	89.937	90.013	
facebook-2	1.200	1.075	29.233	34.475	90.035	90.010	
meps-19	2.554	2.136	10.357	11.285	90.228	90.399	
meps-20	2.667	2.357	10.826	10.720	89.875	89.838	
meps-21	2.585	2.105	10.968	10.565	89.946	89.863	
star	0.195	0.194	9.982	10.142	91.455	91.432	

We have previously reported the evaluated losses ℓ_M and ℓ_L for each dataset, averaged over ten random seeds. Tables A6 and A7 below detail the distribution of these evaluations, providing the mean, 10% quantile, and 90% quantile for the test set deviations in conditional coverage (ℓ_M) and average interval length (ℓ_L) . These statistics are derived from 110 test set evaluations across 11 datasets and 10 random training-test splits. We opt to report empirical quantiles instead of standard deviations due to the asymmetric and non-Gaussian nature of the data.

Table A6: Distribution of the test set conditional coverage deviation ℓ_M evaluated on various conformalized prediction intervals across 11 datasets and 10 random training-test splits.

Max. Conditional Coverage Deviation (%), target miscoverage $\alpha = 10\%$									
Statistics	Meth	nod	Method						
	Local	Boosted	CQR	Boosted					
mean	21.140%	16.319%	11.106%	10.771%					
10% quantile	7.267%	4.604%	4.890%	4.910%					
90% quantile	47.832%	44.712%	22.585%	18.697%					

Table A7: Distribution of the test set average interval length ℓ_L evaluated on various conformalized prediction intervals across 11 datasets and 10 random training-test splits.

Average Length, target miscoverage $\alpha = 10\%$									
Statistics	Met	hod	Met	hod					
Statistics	Local	Boosted	CQR	Boosted					
mean	1.677	1.317	1.483	1.319					
10% quantile	0.950	0.513	0.346	0.351					
90% quantile	2.082	1.829	2.655	2.210					

A.9 Experiments under different miscoverage rates

In Tables A8 and A9, we illustrate the performance of boosting for length with the Local conformity score as baseline with miscoverage rates set to 5% and 20%, respectively.

Table A8: Additional information on conformalized intervals obtained before and after boosting for length with the Local conformity score as baseline. The target miscoverage rate is set to $\alpha = 5\%$.

Dataset		ℓ_L		ℓ_M	(%)	Margina	l Cov.(%)
Dataset	Local	Boosted	Improvement	Local	Boosted	Local	Boosted
bike	2.523	1.828	-27.54%	11.553	11.626	94.927	94.972
bio	1.881	1.801	-4.25%	5.013	4.694	94.857	94.802
blog	4.217	1.923	-54.39%	37.976	34.505	95.027	95.048
community	2.578	2.217	-13.99%	16.406	14.554	95.414	94.812
concrete	1.129	0.838	-25.80%	8.55	8.086	94.709	94.515
facebook-1	2.667	2.262	-15.18%	20.85	20.052	95.104	95.049
facebook-2	2.441	2.092	-14.30%	41.645	39.943	94.941	95.016
meps-19	3.618	2.825	-21.92%	10.515	9.178	95.005	95.008
meps-20	3.951	3.049	-22.84%	11.043	9.693	94.907	94.922
meps-21	4.030	3.033	-24.74%	8.918	8.910	95.061	95.026
star	0.207	0.207	-0.11%	6.57	6.906	95.358	95.289

Table A9: Additional information on conformalized intervals obtained before and after boosting for length with the Local conformity score as baseline. The target miscoverage rate is set to $\alpha=20\%$.

Dataset		ℓ_L		ℓ_M	(%)	Marginal	Cov.(%)
Dataset	Local	Boosted	Improvement	Local	Boosted	Local	Boosted
bike	1.312	0.983	-25.07%	24.649	28.561	79.403	80.152
bio	1.248	1.213	-2.80%	10.838	10.540	79.729	79.594
blog	1.904	0.513	-73.07%	45.813	64.360	79.756	79.920
community	1.337	1.234	-7.71%	25.047	28.831	79.674	79.674
concrete	0.833	0.549	-34.10%	13.831	20.177	80.728	80.825
facebook-1	1.624	0.747	-54.02%	29.384	47.994	80.190	79.731
facebook-2	1.580	0.749	-52.60%	36.488	62.070	79.890	79.930
meps-19	1.821	1.020	-44.03%	20.561	24.763	80.326	80.013
meps-20	1.843	1.095	-40.60%	19.137	24.194	79.684	79.823
meps-21	1.831	1.064	-41.86%	18.227	23.172	80.674	80.057
star	0.142	0.141	-0.71%	15.420	14.517	80.647	80.370

A.10 Training a gradient boosting algorithm with our custom loss functions

Our proposed boosted conformal procedure serves as a post-training step designed to refine the conformity score $E(\cdot,\cdot;f)$ obtained during model training. This procedure can leverage pre-trained models when available. In the absence of pre-trained models, we can alternatively train a gradient boosting algorithm directly using our custom loss functions. For example, in the context of the local score, we may initialize with $\mu^{(0)}=0$, $\sigma^{(0)}=1$ and then apply our boosted conformal procedure. As discussed in Section 7, this approach is flexible enough to replace gradient boosting with any gradient-based algorithm, such as neural networks, trained under our custom loss functions. This framework aligns with that of [26], which introduces a neural network trained with a custom loss function to minimize the average prediction set size in classification tasks.

In Figure A1, we compare the performance of the two approaches optimizing for average interval length, searching within the generalized Local score family $\mathcal F$ defined in (6). The primary distinction between the two procedures lies in the initialization: the first approach employs $\mu^{(0)}=0$, $\sigma^{(0)}=1$, while the second derives $\mu^{(0)}$ and $\sigma^{(0)}$ from a trained random forests model. We run the experiments on the meps-19 dataset and compare the performance across different splits of the training and calibration data. In this context, the percentage of training data refers to the proportion of training data within the combined training and calibration datasets. Our results indicate that cross-validation selects a greater number of boosting iterations when we directly train the gradient boosting algorithm, resulting in longer runtime. However, the average interval length and maximum conditional coverage deviation after boosting are notably smaller for the boosted conformal procedure we introduced in this paper.

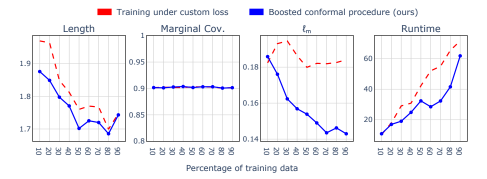


Figure A1: Comparison of boosted interval length, marginal coverage, maximum conditional coverage deviation (ℓ_M), and runtime between direct training of a gradient-based algorithm (red) and boosting on a pre-trained conformity score (blue).

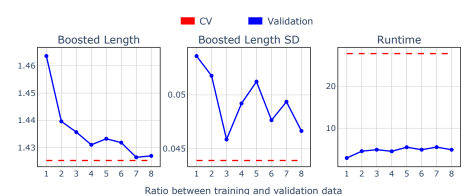


Figure A2: Comparison of the average boosted length, standard deviation of boosted length, and average runtime of the boosting procedure when selecting the optimal number of boosting rounds using 5-fold cross-validation versus a hold-out validation set of varying sizes. Experiments are conducted on the bike dataset, with a target miscoverage rate of $\alpha=10\%$.

A.11 Selecting optimal boosting rounds via hold-out validation set

In our boosted conformal procedure, we use cross-validation on the training set to determine the optimal number of boosting rounds, a process that can be time-consuming. An alternative approach is to hold out a fraction of the training set for validation. While more computationally efficient, this method introduces a trade-off: a smaller validation set can lead to greater variability in prediction intervals and model performance, whereas a larger validation set may reduce the effective training set size, potentially limiting the model's performance. To explore this trade-off, we conduct experiments on the bike dataset, optimizing for prediction interval length. We compare performance across two settings: 5-fold cross-validation and a hold-out validation set, with the training-to-validation set ratio ranging from 1:1 to 8:1. For each setting, we run 100 experiments, recording the average boosted length, the standard deviation of boosted lengths, and the average runtime. The results are shown in Figure A2.

A.12 Additional figures on individual datasets

In this section, we present a series of supplementary figures. First, we showcase the improvements in conditional coverage achieved through the boosted procedure for each benchmark dataset. Figure A3 details results for datasets meps-20 and meps-21. Figure A4 details results for datasets community, bike, and concrete.

Next, we illustrate enhanced interval lengths. Figure A5 details results for datasets meps-20, meps-21, and bike. Figure A6 details results for datasets facebook-1, facebook-2, and concrete. Finally, we demonstrate in Figure A7 how cross-validating the number of boosting rounds effectively prevents the gradient boosting algorithm from overfitting.

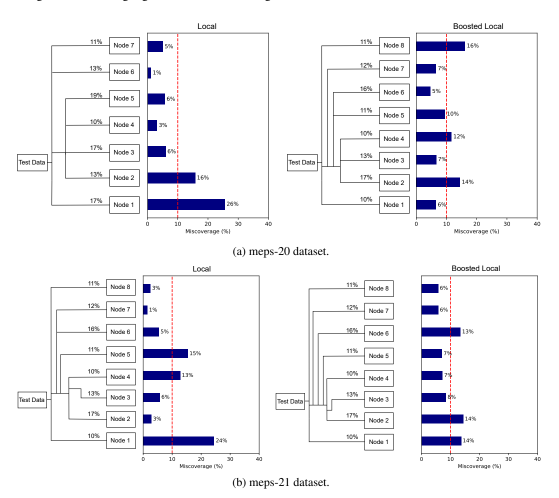


Figure A3: See the caption of Figure 3 for details.

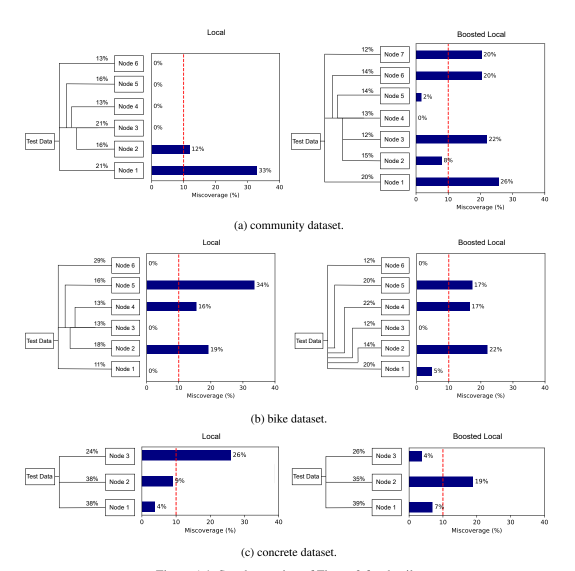


Figure A4: See the caption of Figure 3 for details.

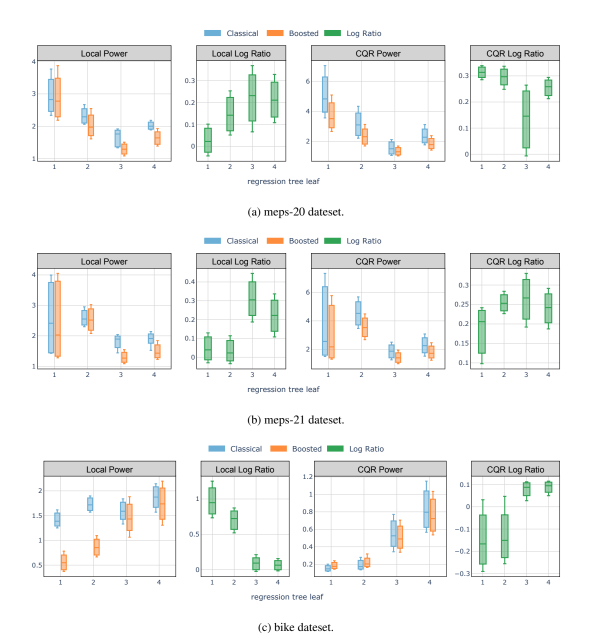


Figure A5: See the caption of Figure 4 for details.

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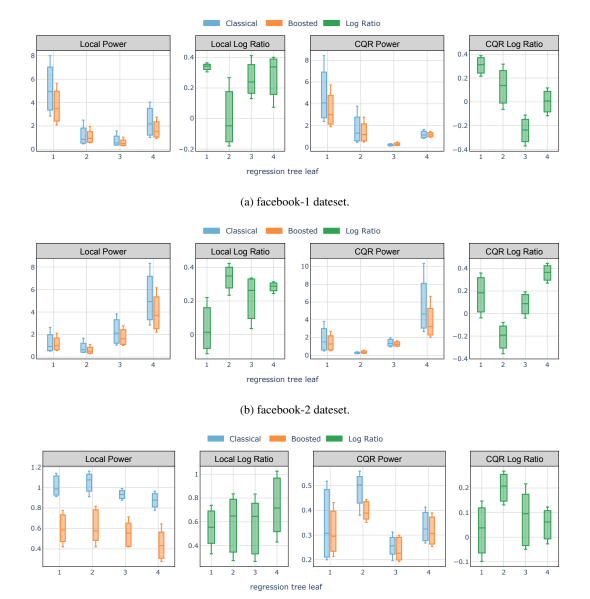


Figure A6: See the caption of Figure 4 for details.

(c) concrete dateset.

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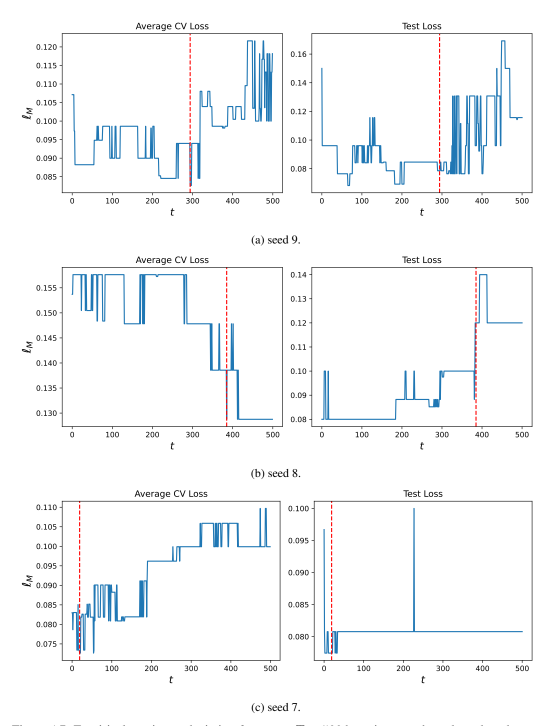


Figure A7: Empirical maximum deviation ℓ_M across T=500 boosting rounds evaluated on dataset concrete under random seeds 7, 8, 9, train-calibration ratio 60%: The left panel illustrates the cross-validated loss, computed as the average across k=3 sub-calibration folds. The right panel displays the test loss. The optimal number of boosting rounds τ , determined through cross-validation as specified in (12), is highlighted in red.

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Answer: [No]

Justification: Real data sets for our empirical analyses are publicly accessible and described in Section A.6. Currently, the code is accessible upon request. Although the code has not been released yet, we have provided detailed documentation for reproducibility in Sections 5.2.2 and A.7.

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