Approximating Full Conformal Prediction at Scale via Influence Functions

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Abstract
Conformal prediction (CP) is a wrapper around traditional machine learning models, giving coverage guarantees under the sole assumption of exchangeability; in classification problems, a CP guarantees that the error rate is at most a chosen significance level \( \varepsilon \), irrespective of whether the underlying model is misspecified. However, the prohibitive computational costs of full CP led researchers to design scalable alternatives, which alas do not attain the same guarantees or statistical power of full CP. In this paper, we use influence functions to efficiently approximate full CP. We prove that our method is a consistent approximation of full CP, and empirically show that the approximation error becomes smaller as the training set increases; e.g., for 1,000 training points the two methods output p-values that are < 0.001 apart: a negligible error for any practical application. Our methods enable scaling full CP to large real-world datasets. We compare our full CP approximation (ACP) to mainstream CP alternatives, and observe that our method is computationally competitive whilst enjoying the statistical predictive power of full CP.

1 Introduction
Conformal prediction (CP) is a post-hoc approach to providing validity guarantees on the outcomes of machine learning (ML) models; in classification, an ML model wrapped with “full” CP outputs prediction sets that contain the true label with (chosen) probability \( 1 - \varepsilon \), under mild distribution assumptions. Unfortunately, full CP is notoriously computationally expensive. Many have proposed alternative methods to avoid the full CP objective; these include: split (or “inductive”) CP (Papadopoulos et al. 2002), cross-CP (Vovk 2015), jackknife+ (Barber et al. 2021), RAPS (Angelopoulos et al. 2020), CV+ (Romano, Sesia, and Candes 2020). While these methods have shown practical promise, they do not attain the same validity guarantee as full CP or its statistical power (e.g., prediction set size). Recent work optimized full CP for ML models that support incremental and decremental learning by speeding up the leave-one-out (LOO) procedure required for the prediction set calculation (Cherubin, Chatzikokolakis, and Jaggi 2021); however, this approach may not scale to complex models such as neural networks.

Herein, we first discuss how to approximate the full CP objective. We focus on full CP for classification, and optimize it for ML models trained via ERM (e.g., logistic regression, neural networks). The key insight we leverage is that, for each test point, full CP: (i) retrains the underlying ML model on the additional test point, and (ii) performs a LOO procedure for each training point. We observe we can approximate both steps, and avoid retraining each time, by using first order influence functions (Hampel 1974). We term our method Approximate full Conformal Prediction (ACP), and we prove finite-sample error guarantees: as the training set grows, ACP approaches full CP. We then show that a stronger regularization parameter for training the underlying ML model improves the approximation quality.

We empirically demonstrate that ACP is competitive with existing methods on MNIST (LeCun 1998), CIFAR-10 (Krizhevsky, Nair, and Hinton 2009), and US Census (Ding et al. 2021). Unlike full CP, ACP scales to large datasets for real-world ML models (logistic regression, multilayer perceptrons, and convolutional neural networks). Performance-wise, ACP is consistently better than existing alternatives in terms of statistical power: it attains the desired error rate \( \varepsilon \) with tighter prediction sets; Figure 1 shows on CIFAR-10 examples where, unlike other methods, ACP learns smaller prediction sets that still contain the true label.

2 Preliminaries
We describe full CP, and then introduce influence functions, our main optimization tool.

2.1 Notation and Full CP
Consider a training set \( Z = (X, Y) \in (\mathcal{X} \times \mathcal{Y})^N \). For a test object \( x \in \mathcal{X} \) and a chosen significance level \( \varepsilon \in [0, 1] \), a CP returns a set \( \Gamma_x^\varepsilon \subseteq \mathcal{Y} \) containing \( x \)'s true label with probability at least \( 1 - \varepsilon \). This guarantee (validity) holds for any exchangeable distribution on \( Z \cup \{(x, y)\} \). Since the error rate of a CP is guaranteed by validity, a data analyst only needs to control the tightness (efficiency) of its prediction set; average \( |\Gamma_x^\varepsilon| \) is a common efficiency criterion (Vovk et al. 2016). Efficiency is controlled by improving the underlying model that CP wraps.

Underlying model. A CP can be built around virtually any ML model \( \theta \). We assume the underlying model is trained via
...and hence, CP) on the basis of a model: the deleted scheme computes the loss at a point after removing it from the training set \( Z \). Crucially, for each possible label \( \epsilon \), \( z, \theta \) sume to be convex and twice differentiable in \( \theta \). This assumption holds for many popular loss functions. Let \( \theta_Z \equiv \arg\min_{\theta \in \Theta} R(Z, \theta) \) be the ERM solution; we assume \( \theta_Z \) to be unique, and discuss relaxations in Section 7.

Nonconformity measure. The underlying model is used to construct a nonconformity measure, which defines a CP. A nonconformity measure is a function \( A : (X \times Y) \times (X \times \hat{Y})^N \rightarrow R \), which scores how conforming (or similar) an example \( (x, y) \) is to a bag of examples \( \hat{Z} \). We focus on the two most common ways of defining nonconformity measures (and, hence, CP) on the basis of a model: the deleted and the ordinary scheme (Vovk, Gammerman, and Shafer 2005).

Full CP (deleted). Consider example \( \hat{z} \) and a training set \( Z \). The nonconformity measure can be defined from the deleted (LOO) prediction: \( A(z_i, Z) = \ell(z_i, \theta_{Z \cup \{\hat{z}\} \setminus \{z_i\}}) \), for all \( z_i \in Z \cup \{ \hat{z} \} \). Computed this nonconformity measure requires training the model on \( Z \cup \{ \hat{z} \} \setminus \{ z_i \} \). This scheme computes the loss at a point after removing it from the model’s training data.

Algorithm 1 shows how the nonconformity measure is used in full CP. For a test point \( x \), CP runs a statistical test for each possible label \( y \in Y \) to decide if it should be included in the prediction set \( \Gamma^x \). The statistical test requires computing a nonconformity score \( \alpha_i \) by running \( A \) for each point in the augmented training set \( Z \cup \{ (x, \hat{y}) \} \); then, a p-value is computed, and a decision is taken based on the threshold \( \varepsilon \). This algorithm is particularly expensive. Crucially, for each test point, and for every candidate label, one needs to retrain the underlying ML model \( N + 1 \) times.

Full CP (ordinary). A computationally faster scheme is achieved by taking the loss at the point: \( A(z_i, Z) = \ell(z_i, \theta_{Z \cup \{\hat{z}\}}) \). We refer to this as the ordinary scheme (Algorithm 3). This method is inherently faster than the deleted approach, as it only requires training one model per test example and candidate label. However, the ordinary scheme generally leads to less efficient predictions (Section 5).

Optimizing CP. The complexity of full CP depends on: (i) the number of training points \( N \), and (ii) the number of labels \( |Y| \). Optimizing w.r.t. \( |Y| \) is necessary for regression settings, where full CP is not applicable directly; this was done, for specific choices of nonconformity measures, by Papadopoulos, Vovk, and Gammerman (2011); Nouretdinov, Melluish, and Vovk (2001); Lei (2019); Ndiaye and Takeuchi (2019); Ndiaye (2022). Our work focuses on optimizing w.r.t. \( N \); this enables applying full CP classification to large datasets. Future work may combine our optimizations and CP regression strategies to obtain faster regressors on large training sets (e.g., Cherubin, Chatzikokolakis, and Jaggi 2021).

2.2 Influence Functions

Influence functions (IF) are at the core of our proposal. For a training example \( z_i \in Z \), let \( I_\theta(z_i) = -\frac{1}{N} \nabla_\theta \ell(z_i, \theta) \) be the influence of \( z_i \) on model \( \theta \), where \( \nabla_\theta \ell \) is the Hessian; by assumption, \( \nabla_\theta \) exists and is invertible. A standard result by Hampel (1974) shows that:

\[
\theta_{Z \setminus \{z_i\}} - \theta \approx -I_\theta(z_i). \tag{1}
\]

\( I_\theta \) says how much \( z_i \) affects \( \theta \) during training. We can apply influence functions for computing the influence of a point \( z_i \) on any functional. In our work, we are interested in the influence on the loss function at a point \( z \). Let \( I_r(z, z_i) = \nabla_\theta \ell(z, \theta) I_\theta(z_i) \). Then, similarly to above, we have

\[
\ell(z, \theta_{Z \setminus \{z_i\}}) - \ell(z, \theta) \approx -I_r(z, z_i). \tag{2}
\]

3 Approximate Full Conformal Prediction

Our proposal (ACP) hinges on approximating the nonconformity scores via IF. We describe our approach, and prove theoretical results on its consistency and approximation error.

3.1 Approach

The bottleneck of running full CP is the computation of the nonconformity scores \( \alpha_i = \ell(z_i, \theta_{Z \cup \{\hat{z}\} \setminus \{z_i\}}) \). Each score is determined by computing the loss of the model at point \( z_i \in Z \cup \{ \hat{z} \} \) after adding point \( \hat{z} \) and removing point \( z_i \) from the model’s training data \( Z \). There are two ways to...
approximate $\alpha_i$ via IF: we can approximate the contribution of adding and removing the points to the learned model $\theta_Z$, and then evaluate its loss at $z_i$ (indirect approach), or we can directly approximate the contribution of the points on the loss function (direct approach). We describe both below.

Indirect approach. We can use Equation 1 to approximate model $\theta_{Z\cup\{\hat{z}\}\setminus\{z_i\}}$ and then compute its loss. That is, let $\tilde{\theta}_{Z\cup\{\hat{z}\}\setminus\{z_i\}} = \theta_Z + I_{\theta_Z}(\hat{z}) - I_{\theta_Z}(z_i)$. Then:

$$\alpha_i \approx \ell(z_i, \tilde{\theta}_{Z\cup\{\hat{z}\}\setminus\{z_i\}}).$$

(Direct approach.) We can directly compute the influence on the loss. Let $\theta_Z$ be a model trained via ERM on the entire training set $Z$. The direct approximation for the score is:

$$\alpha_i \approx \ell(z_i, \tilde{\theta}_{Z\cup\{\hat{z}\}\setminus\{z_i\}}) \equiv \ell(z_i, \theta_Z) + I_{\ell}(z_i, \hat{z}) - I_{\ell}(z_i, z_i).$$

$I_{\ell}(z_i, \hat{z})$ and $-I_{\ell}(z_i, z_i)$ are the influence of including point $\hat{z}$ and excluding $z_i$ (Equation 2). Algorithm 2 (ACP) shows how both approaches enable approximating full CP.

ACP gives a substantial speed-up over full CP. In contrast to full CP, ACP has a training phase, in which we: compute the Hessian, the gradient for each point $z_i$, and provisional scores $\ell(z_i, \theta_Z)$ for $z_i \in Z$. For predicting a test point $\hat{z}$, it suffices to compute its influence by using the Hessian and gradients at $z_i$ and $\hat{z}$, which is cheap, and update the provisional scores (see time complexities in Table 4). This enables ACP to scale to large real-world datasets such as CIFAR-10 (Section 5).

3.2 Theoretical Analysis

In this section, we establish the consistency of ACP: its approximation error gets smaller as the training set grows. Further, we study its finite-sample validity, and how the underlying model’s regularization parameter affects its approximation error. The consistency of ACP for the indirect approach comes from a result by Giordano et al. (2019). Proving consistency for the direct approach requires a condition, which we state in the next part as a conjecture.

Direct approximation is better than indirect We conjecture that the direct approach approximates better than the indirect one. Intuitively, it is much easier to approximate the loss at a point (direct) than to estimate the effect of a training point on the model weights, which lay in a high-dimensional space (indirect). We observed this conjecture to hold consistently across a number of simulations (Section 4). Formally:

**Condition 1.** Assume that the loss $\ell$ is convex and differentiable. Then the direct method (Equation 4) is a better approximation than the indirect one (Equation 3). That is, let

$\alpha_i = \ell(z_i, \theta_{Z\cup\{\hat{z}\}\setminus\{z_i\}})$;

$$|\ell(z_i, \theta_{Z\cup\{\hat{z}\}\setminus\{z_i\}}) - \alpha_i| \leq |\ell(z_i, \theta_{Z\cup\{\hat{z}\}\setminus\{z_i\}}) - \alpha_i|.$$

Without Condition 1, we can prove consistency for the indirect approach, but not for the direct approach.

Consistency of ACP We show that ACP is a consistent estimator of full CP. We establish this equivalence in the most generic form possible: we demonstrate that nonconformity scores produced by Algorithm 2 approximate those produced by full CP. In turn, the p-values (and, consequently, error rates) of the two methods get increasingly closer.

Our result is an extension of the work by Giordano et al. (2019), who showed that IF consistently estimate a model’s parameters in a LOO setting. This result holds under a set of assumptions (Assumption 1 in Appendix B), which Giordano et al. (2019) showed to hold for a variety of settings; e.g., they hold when $Z$ are well-behaved IID data and $\ell(\cdot, \theta)$ is an appropriately smooth function. Note that Assumption 1 limits the set of applicable nonconformity measures, e.g., by assuming them to be continuously differentiable.

**Theorem 2** (Consistency of approximate full CP). Under Assumption 1 and Condition 1, let $\alpha_i = \ell(z_i, \theta_{Z\cup\{\hat{z}\}\setminus\{z_i\}})$, and suppose $\ell$ is K-Lipschitz. For every $N$ there is a constant $C$ such that for every $z_i \in Z \cup \{\hat{z}\}$:

$$|\hat{\ell}(z_i, \theta_{Z\cup\{\hat{z}\}\setminus\{z_i\}}) - \alpha_i| \leq KC \frac{\max\{C_g, C_h\}^2}{N},$$

for finite constants $C_g, C_h$ s.t. $\sup_{\theta \in \Theta} \frac{1}{\sqrt{N}} ||\nabla_\theta \ell(z, \theta)||_2 \leq C_g$ and $\sup_{\theta \in \Theta} \frac{1}{\sqrt{N}} ||\nabla_\theta^2 \ell(z, \theta)||_2 \leq C_h$.

This result gives a finite-sample bound for the error of the direct approach for ACP; the error of the indirect approach is
also bounded as a byproduct of the same proof. We conclude that, as \( N \) grows, ACP’s scores get increasingly closer to those produced by full CP. We evaluate this in Section 4.

**Validity of ACP** Lin, Trivedi, and Sun (2021) state that finite-sample validity is not guaranteed when the LOO is estimated with IF since they cannot be exactly computed. They exemplify this issue in the Discriminative Jackknife (Alaa and Van Der Schaar 2020), which approximates the IF using Hessian-Vector-Products (Pearlmutter 1994).

Although we alleviate part of the issue by computing the exact Hessian, we cannot guarantee that our LOO estimation is exact. Basu, Pope, and Feizi (2021) also summarize several issues with using IF in deep learning. Nevertheless, ACP still inherits the high efficiency of full CP, and we observe that validity holds in practice (Section 5). Future work can prove if the approximate scores follow the same distribution as the true ones and, consequently, if exchangeability still holds.

**Relation to regularization parameter** By extending a result by Koh et al. (2019), we investigate the effect of the ERM regularization parameter on ACP’s approximation error. This result makes fairly simplistic assumptions (Appendix B.2).

**Theorem 3 (Approximation goodness w.r.t. regularizer).** Suppose the model is trained via ERM with regularization parameter \( \lambda \). Under the assumptions of Proposition 5, Assumption 2, and neglecting \( O(\lambda^{-3}) \) terms, we have the following cone constraint between the true nonconformity measure \( \alpha_i = \ell(z_i, \theta_{Z^i}) \) and its direct approximation \( \tilde{\ell}(z_i, \theta_{Z^i}) \equiv \ell(z_i, \tilde{\theta}_Z) + I_\ell(z_i, \tilde{\theta}) - I_\ell(z_i, \tilde{\theta}_Z) \), where \( \ell(\lambda) = (1 + 3\sigma_{\max}^2/2\lambda + \sigma_{\max}^2/2\lambda^2) \), and \( \sigma_{\max} \) is the maximum eigenvalue of the Hessian \( H \):

\[
\ell(z_i, \theta_Z) + I_\ell(z_i, \tilde{\theta}) - g(\lambda) I_\ell(z_i, z_i) \leq \alpha_i \leq \tilde{\ell}(z_i, \theta_{Z^i}).
\]

4 Experiments on Synthetic Data
We study the properties of ACP outlined in Section 3.2 on synthetic data (Appendix C); the underlying model is logistic regression with cross-entropy loss. Results are averaged across 100 test points.

**Direct and indirect approximation.** We empirically evaluate Condition 1, which claims that the direct method (Equation 4) is never worse than indirect (Equation 3). Figure 2 shows the absolute distance between full CP and ACP’s nonconformity scores as a function of the training set size. Results confirm that direct is always better than indirect, although the two get close for large \( N \). Importantly, the nonconformity scores produced by ACP get increasingly better at approximating those of full CP as the training set grows (cf. Theorem 2). We shall now focus on the direct approach.

**Approximation goodness.** We evaluate how well ACP approximates full CP, under various parameter choices, as the training set grows. Figure 3a shows the difference between the nonconformity scores of full CP and ACP as the number of features ranges in 5-100. The number of features does impact the IF approximation, although the error becomes negligible as the training set increases. Theorem 3 shows that, unsurprisingly, a larger \( \lambda \) (i.e., stronger regularization) implies better approximation. We confirm this in Figure 3b. Our analysis focuses on the approximation error between nonconformity scores; yet, we remark that a small error between scores implies a more fundamental equivalence between full CP and ACP: their p-values should also have a small distance.

Figure 3c compares full CP and ACP’s p-values. The difference is smaller than \( 10^{-3} \) with a training set of 600, and it becomes negligible with \( N = 10k \) training examples. Observe that in CP the p-value is thresholded by the significance value \( \varepsilon \) to obtain a prediction (Algorithm 1). As practitioners are generally interested in values \( \varepsilon \) with no more than 2 decimals of precision (e.g., \( \varepsilon = 0.15 \)), we argue that an approximation error smaller than \( 10^{-3} \) between p-values is more than sufficient for any practical application. Figure 3d compares the error rate (for \( \varepsilon = 0.1 \)) between full CP and ACP. We observe that, after 500 training points, the two methods have the same error.

5 Experiments with Real Data
We compare mainstream CP alternatives with ACP on the basis of their predictive power (efficiency). Because of its computational complexity, it is infeasible to include full CP in these experiments. Nevertheless, given the size of the training data, the consistency of ACP (Theorem 2), and the results in Section 4, we expect ACP to perform similarly to full CP.

5.1 Existing Alternatives to Full CP
There are several alternative approaches to CP for classification. In this work, we compare ACP with:

- Split (or “inductive”) Conformal Prediction (SCP) (Padopoulos et al. 2002) works by dividing the training set into proper training set and calibration set. The model is fit on the proper training set, and the calibration set is used to compute the nonconformity scores.

![Figure 2: Comparison between direct and indirect approximations](image-url)
We show similar instances in Appendix E.2. In the next part, we run an illustrative experiment for a CIFAR-10 test point. We observe this behavior generalizes to larger test sets. As a way of comparing the curves, we include the AUC w.r.t. $\varepsilon$; we observe that, while all the methods output the true label, the method would still be accurate with an $\varepsilon$ larger than $\varepsilon$. Finally, for each method we report the highest $\varepsilon$ have the smallest AUC. For the interval $[0, 0.2]$, ACP (deleted and ordinary) have the largest values.

Datasets. We select datasets to illustrate the performance of ACP in various scenarios: a simple classification problem with images (MNIST (LeCun 1998)), a more complex setting (CIFAR-10 (Krizhevsky, Nair, and Hinton 2009)), and a binary classification with tabular data (US Census (Ding et al. 2021)); details in Appendix C.2.

5.2 A Warm-Up Example

We run an illustrative experiment for a CIFAR-10 test point picked uniformly at random. We consider a neural network with 3 layers of 100, 50, and 20 neurons; we refer to this network as MLP. Appendix C.3 gives implementation details.

Figure 4a shows the prediction set for a fixed $\varepsilon = 0.05$; we observe that, while all the methods output the true label, the prediction set of ACP (deleted) is the tightest (i.e., more efficient). We also report how the prediction set size changes w.r.t. $\varepsilon$, for ACP (Figure 4b) and for all methods (Figure 4c). As a way of comparing the curves, we include the AUC for the interval $\varepsilon \in [0, 0.2]$. ACP (deleted and ordinary) have the smallest AUC. Finally, for each method we report the highest $\varepsilon$ for which the prediction set contains the true label. A higher value indicates that, for this test example, the method would still be accurate with an $\varepsilon$ larger than 0.05, which would correspond to tighter prediction sets. Once again, ACP (deleted and ordinary) have the largest values. We show similar instances in Appendix E.2. In the next part, we observe this behavior generalizes to larger test sets.

We observe an unstable behavior in the predictions of RAPS and CV+: their prediction set size considerably oscillates as $\varepsilon$ increases. The reason is that their prediction sets are not guaranteed to be nested; that is, $\varepsilon > \varepsilon'$ does not imply that the prediction sets $\Gamma^\varepsilon \subseteq \Gamma^{\varepsilon'}$. Specifically, because RAPS and CV+ use randomness to decide whether to include a label in the set, the true label may appear and then disappear for a smaller significance level. This may not be desirable in some practical applications. The prediction set for ACP and SCP monotonically decreases with $\varepsilon$, by construction.

5.3 Experimental Setup

We evaluate the methods for five underlying models: three multilayer perceptrons with architectures (neurons per layer): 20-10 (MLP$_A$), 100 (MLP$_B$) and 100-50-20 (MLP$_C$); logistic regression (LR); and a convolutional neural network (CNN). In experiments with MNIST and CIFAR-10, the dimensionality is first reduced with an autoencoder (AE) in all settings except the CNN. We defer implementation details to Appendix C.3-C.4.

Considering these five settings enables comparing the methods both for underparametrized regimes (e.g., LR) and for better performing models (e.g., CNN). Note that CP’s guarantees hold regardless of whether the underlying model is misspecified. Further, observe that most of these models are non-convex, where the ERM optimization problem does not have a unique solution; this contradicts the IF assumption (Section 2). Nonetheless, our empirical results show that ACP works well – it performs better than the other proposals; in Section 7 we discuss relaxations of this assumption.

5.4 Results

For each experiment and method, we report averaged metrics over 100 test points; we also run statistical tests to check if differences are (statistically) significant.

Prediction set size. We measure efficiency as the average prediction set size. We report this as a function of $\varepsilon$, discretized with a step $\Delta \varepsilon = 0.01$. Figures 5a and 5b show the average prediction set size in MNIST and CIFAR-10 for MLP$_C$. ACP (deleted and ordinary) consistently outperform

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1RAPS allows a non-randomized version, although with a more conservative behavior and considerably larger prediction sets.
all other methods; deleted is better than ordinary. ACP is significantly more efficient than RAPS and CV+. SCP and ACP (O) perform similarly on MNIST, but their difference is remarked on CIFAR-10; this suggests that SCP is a cheap alternative to ACP on relatively easier tasks. Appendix E.3 reports results for the rest of the models and for the US Census, showing similar behavior.

Efficiency AUC. We use an $\varepsilon$-independent metric to aid this comparison: the area under the curve (AUC) of the prediction set size in the interval $\varepsilon \in [0, 0.2]$; a smaller AUC means better efficiency. ACP (deleted or ordinary) prevails on all methods, datasets, and model combinations (Table 1). An exception is LR on US Census, where RAPS has a better efficiency than ACP and SCP; simpler tasks and models may be well served by the computationally efficient RAPS.

We observe that both deleted and ordinary ACP are better than RAPS and SCP; simpler tasks and models may be well served by the computationally efficient RAPS. Welch one-sided tests (reject with p-value < 0.1) confirm that both deleted and ordinary ACP are better than RAPS and CV+; they further show that either deleted or ordinary ACP are better than SCP on most tasks. We refer to Section 7 for directions to improve ACP’s performance.

Validity. As a way of interpreting why ACP performed better than the other methods, we measure their empirical error rate with a fixed $\varepsilon = 0.2$. Indeed, whilst all methods guarantee a probability of error of at most $\varepsilon$, a more conservative (i.e., smaller) empirical error may correspond to larger significance level in most cases; this indicates this method fully exploits its error margin to optimize efficiency.

Table 1: Efficiency AUC ($\varepsilon \in [0, 0.2]$). The table indicates that differences in the AUC are statistically significant as compared to ACP (D) ($\dagger$) and ACP (O) ($\ddagger$).
Table 2: Difference between the expected error ($\varepsilon$) and the empirical error rate on CIFAR-10. A positive value indicates a conservative prediction set; a negative value is a validity violation, which may be due to statistical fluctuation.

<table>
<thead>
<tr>
<th>Model</th>
<th>ACP (D)</th>
<th>ACP (O)</th>
<th>SCP</th>
<th>RAPS</th>
<th>CV+</th>
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<tr>
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<td>-0.01</td>
<td>0.02</td>
<td>0.04</td>
</tr>
<tr>
<td>CNN</td>
<td>-0.01</td>
<td>0.01</td>
<td>-0.01</td>
<td>0.04</td>
<td>0.10</td>
</tr>
</tbody>
</table>

Table 3: Fuzziness of ACP and SCP on MNIST and CIFAR-10. A smaller fuzziness corresponds to a higher efficiency. The table indicates that differences are statistically significant as compared to ACP (D) (*) and ACP (O) (†).

Table 6: Related Work

Full CP (Vovk, Gammerman, and Shafer 2005) is notoriously expensive. Many alternatives have been proposed. Arguably the most prominent are SCP (Vovk, Gammerman, and Shafer 2005), CV+ (Romano, Sesia, and Candes 2020), RAPS (Angelopoulos et al. 2020), Cross-CP (Vovk 2015), aggregated CP (Carlsson, Eklund, and Norinder 2014), APS (Romano, Sesia, and Candes 2020), and the jackknife (Miller 1974; Efron 1979). We compared ACP with the former three. Unlike full CP, all the above methods have weaker validity guarantees or they tend to attain less efficiency.

Cherubini, Chatzikokolakis, and Jaggi (2021) introduced exact optimizations for full CP for classification. Their method saves an order of magnitude in time complexity for many models, but is alas only applicable to models supporting incremental and decremental learning (e.g., k-NN). Crucially, it is unlikely extendable to neural networks

The closest in spirit to our approach is the Discriminative Jackknife (DJ) by Alaa and Van Der Schaar (2020), which uses IF to approximate jackknife+ confidence intervals for regression (Barber et al. 2021). We note several differences between ACP and DJ, besides their different goals (classification vs regression). While DJ approximates LOO w.r.t. the parameters, we introduce a direct approach to approximate the nonconformity scores in Equation 4. Whereas DJ only does decremental learning, which allows them to exploit Hessian-Vector-Products (Pearlmutter 1994), approximating full CP requires us to do both incremental and decremental learning (Algorithm 2). We also prove that our approximation error decreases w.r.t. the size of the training set.

7 Conclusion and Future Work

Full CP is a statistically sound method for providing performance guarantees on the outcomes of ML models. For classification tasks, CP generates prediction sets which contain the true label with a user-specified probability. Unfortunately, full CP is impractical to run for more than a few hundred training points. In this work, we develop ACP, a computationally efficient method which approximates full CP via influence functions; this strategy avoids the numerous recalculations that full CP requires. We prove that ACP is consistent: it approaches full CP as the training set grows. Our experiments support the use of ACP in practice.

There are many directions to improve ACP. For example, we assumed that the ERM solution is unique. While our approximation works well in practice, it would be a fruitful endeavour to relax this assumption; initial work towards this was done for IF by Koh and Liang (2017). Another direction is to build nonconformity scores on the studentized scheme, a middle way between deleted and ordinary (Vovk et al. 2017). Future work can also investigate when Condition 1 holds.

Finally, while we scale full CP to relatively large models, computing and inverting the Hessian becomes very expensive as the number of parameters increases. Recent tools to approximate the Hessian, like the Kronecker-factored Approximate Curvature (K-FAC) method (Martens and Grosse 2015; Ba, Grosse, and Martens 2017; Tanaka et al. 2020), might help further scale ACP to larger models like ResNets.

In conclusion, ACP helps scaling full CP to large datasets and ML models, for which running full CP would be impractical. Although split-based approaches like SCP and RAPS are less expensive to run, they do not attain the same efficiency as ACP. This makes the adoption of our method particularly appealing for critical real-world applications.
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