Prediction Intervals: 
Split Normal Mixture from Quality-Driven Deep Ensembles

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Abstract

Prediction intervals are a machine- and human-interpretable way to represent predictive uncertainty in a regression analysis. In this paper, we present a method for generating prediction intervals along with point estimates from an ensemble of neural networks. We propose a multi-objective loss function fusing quality measures related to prediction intervals and point estimates, and a penalty function, which enforces semantic integrity of the results and stabilizes the training process of the neural networks. The ensembled prediction intervals are aggregated as a split normal mixture accounting for possible multimodality and asymmetricity of the posterior predictive distribution, and resulting in prediction intervals that capture aleatoric and epistemic uncertainty. Our results show that both our quality-driven loss function and our aggregation method contribute to well-calibrated prediction intervals and point estimates.

1 INTRODUCTION

Quantifying predictive uncertainty of machine learning models is crucial in applications, e.g., mission-critical systems, where it is essential to know when the model is not able to provide accurate predictions. In decision support systems, providing the human with an additional information about the uncertainty of a prediction may decrease the response time and increase the accuracy of an action. It can also positively contribute in building trust and understanding towards the machine learning system and its correct facilitation. In this work, we focus on the quantification of predictive uncertainty for the regression task, specifically in the form of prediction intervals which have probabilistic interpretation and which are interpretable for both humans and machines.

Prediction interval (PI) is an estimate representing predictive uncertainty in the form of two values between which a future observation will fall with a certain probability. Well-calibrated/high-quality prediction intervals are as narrow as possible while attaining the desired coverage probability.

While neural networks (NNs) are powerful function approximators, they are poor at representing predictive uncertainty. Previously, methods adopting a Bayesian approach in the context of neural networks (e.g., Graves, 2011; Blundell et al., 2015; Hernandez-Lobato and Adams, 2015; Krueger et al., 2017; Louizos and Welling, 2017; Pawlowski et al., 2017; Wu et al., 2019; Izmailov et al., 2019) represented the state-of-the-art for providing...
predictive uncertainty estimates. See Yao et al. (2019) for an overview. Recently, a number of non-Bayesian yet probabilistic approaches provide competitive predictive uncertainty estimates (Lakshminarayanan et al., 2017; Pearce et al., 2018; Tagasovska and Lopez-Paz, 2019). In this paper, we focus on and extend the latter branch of research.

Our work builds on the findings of Pearce et al. (2018) which is inspired by works of Khosravi et al. (2011) and Lakshminarayanan et al. (2017). Pearce et al. (2018) presented a method based on aggregating ensembled PIs from a set of NNs optimized with respect to a so-called quality-driven loss function (see Section 2.4).

Although the aforementioned work has achieved promising results of well-calibrated PIs, it has several limitations that need to be addressed. In our continuation, we primarily address three main limitations of the state-of-the-art method: (1) The inability to generate point estimates, (2) the theoretically weakly justified method for aggregation of ensembled PIs, and (3) the fragile training process. With this in mind, the main contributions of this work can be summarized as follows:

- We retrofit the quality-driven loss function:
  - With a point estimate loss (particularly MSE or mean squared error) to be able to draw point estimates and PIs from the same generative distribution.
  - With a penalty function adding a constraint and thus enforcing the integrity of the results, i.e. avoiding crossing of PI boundaries or point estimates out of PI bounds, and consequently stabilizing/strengthening the training process.

- We propose a new aggregation method for ensembles of PIs with point estimates. It is fitting a split normal mixture (Wallis, 2014) providing tighter and theoretically well-founded aggregates of PIs.

- We propose an analytical approach to parameter initialization for the parameter fitting process of a split normal probability density function, thus increasing the success of fitting and accelerating it compared to random initialization.

In addition to the above, we provide important insights and guidelines for hyper-parameter search contributing to reproducibility and reliable model fitting, and we suggest directions for future research.

2 BACKGROUND

In this section, we formally introduce the predictive uncertainty and prediction intervals, we consider methods for quantifying predictive uncertainty in NNs, especially in the form of prediction intervals, and we provide insights into the work of Pearce et al. (2018) which we build on.

2.1 Predictive Uncertainty

The sources of predictive uncertainty can be categorized into aleatoric and epistemic uncertainty (Kiureghian and Ditlevsen, 2009).

The aleatoric or aleatory or data uncertainty is also known as the irreducible uncertainty, i.e. it cannot be reduced either through model or data. It arises from an inherent and irreducible data noise. The aleatoric uncertainty can be captured by learning the conditional distribution between the target and the input variables. The aleatoric uncertainty can further be characterized as homoskedastic (homoscedastic) if the irreducible noise is constant across random variables or as heteroskedastic (heteroscedastic) if contrary.

In contrast to aleatoric uncertainty, the epistemic uncertainty is reducible, and it can be further decomposed into model uncertainty and distributional uncertainty. Model uncertainty can be caused by model bias or parameter uncertainty due to insufficient data. Distributional uncertainty can be caused by the mismatch between training and test set (Malinin and Gales, 2018), and it is often described as a part of the model uncertainty.

Generally, the aim is to reduce epistemic uncertainties. However, constraints emerging from model or data usually do not allow it.

Please note that predictive uncertainty is a broad concept, and the literature is inconsistent in the terminology. We follow the taxonomy found in Kiureghian and Ditlevsen (2009; Pearce et al., 2018; Malinin and Gales, 2018).

2.2 Prediction Intervals

Given an input $x^{(i)}$, a prediction interval $[\hat{y}_L^{(i)}, \hat{y}_U^{(i)}]$ of a sample $i$ captures the future observation (target variable) $y^{(i)}$ with the probability equal or greater than $\gamma \in [0, 1]$ (eq. 1). The value of $\gamma$ is commonly set to 0.95 or 0.99. Common in the literature is an alternative notation with $\alpha$.

$$\Pr(\hat{y}_L^{(i)} \leq y^{(i)} \leq \hat{y}_U^{(i)}) \geq \gamma = (1 - \alpha) \quad (1)$$

Given $n$ samples, the quality of the generated prediction intervals is assessed by measuring the prediction interval
coverage probability (PICP)

\[ \text{PICP} = \frac{c}{n} \]

where

\[ c = \sum_{i=1}^{n} k_i \]  

for

\[ k_i = \begin{cases} 1 & \text{if } \hat{y}_{L}^{(i)} \leq y^{(i)} \leq \hat{y}_{U}^{(i)}, \\ 0 & \text{otherwise,} \end{cases} \]  

and by measuring the mean prediction interval width (MPIW)

\[ \text{MPIW} = \frac{1}{n} \sum_{i=1}^{n} (\hat{y}_{U}^{(i)} - \hat{y}_{L}^{(i)}) \]

or its normalized version NMPIW

\[ \text{NMPIW} = \frac{\text{MPIW}}{r} \]

where \( r = \max(y) - \min(y) \). Please note that MPIW assumes \( \hat{y}_{U}^{(i)} \geq \hat{y}_{L}^{(i)}, \) i.e. no crossing of PI bounds. Ignoring this fact can lead to misleading results.

It is desired to achieve \( \text{PICP} \geq \gamma \) while having MPIW as small as possible.

2.3 Related Work

Following is a concise exploration of methods based on NNs capable of quantifying predictive uncertainty in a regression analysis. Methods adopting the Bayesian approach (such as BNN or Bayesian neural networks) are a prominent group (Graves, 2011; Blundell et al., 2015; Hernandez-Lobato and Adams, 2015; Krueger et al., 2017; Louizos and Welling, 2017; Pawlowski et al., 2017; Wu et al., 2019; Izmailov et al., 2019; etc.) and represented for long the state-of-the-art to estimate predictive uncertainty. We explore alternative non-Bayesian yet probabilistic methods representing competitive and possibly a state-of-the-art alternative.

Interpreting Monte Carlo dropout (MC-dropout) at test time as approximate Bayesian inference (Gal and Ghahramani, 2016) has been a widely used method to quantify predictive uncertainty, mainly due to its scalability and simplicity. Interpretation of MC-dropout as an ensemble model combination motivated consecutive research on ensemble-based methods which have been shown to be superior in generating predictive uncertainty to MC-dropout or even Bayesian methods (Lakshminarayanan et al., 2017). Recently, it has been shown that deep ensembles with random initialization may explore different modes in function space, and therefore perform well in exploring model uncertainty. That is in contrast to subspace sampling methods (e.g. MC-dropout, weight averaging) which may generate a set of diverse functions but still in the vicinity of the starting point and thus generating insufficiently diverse predictions (Fort et al., 2019). Also recently, the robustness of uncertainty quantification methods under dataset shift was investigated, and although all compared methods have been deceptive to increasing dataset shifts, deep ensembles have shown the greatest robustness (Ovadia et al., 2019).

More recently, an alternative line of work employs a loss function to optimize for generating specifically prediction intervals. A method based on quantile regression is utilizing the so-called pinball loss function (Koenker, 2005) to optimize a single NN for the desired PIs and so-called Orthonormal Certificates to account for epistemic uncertainty (Tagasovska and Lopez-Paz, 2019). A method called LUBE (Khosravi et al., 2011) constructs prediction intervals using quality-driven loss function utilizing quality metrics PICP and NMPIW. LUBE applies simulated annealing for training a NN with respect to the quality-driven loss function. Inspired by Khosravi et al. (2011) and Lakshminarayanan et al. (2017), a slightly modified quality-driven loss function optimized for gradient descent was proposed (Pearce et al., 2018) and extended with ensembles to account for epistemic uncertainty. Although presenting favorable results, both methods deliver only prediction intervals without point estimates.

2.4 Quality-Driven Ensembles by [Pearce et al., 2018]

Our work builds on [Pearce et al., 2018] henceforth referred to as the original quality-driven ensembles or shortly original QDE (in results annotated as SEM-QD). First, the quality-driven loss function is described (annotated as QD). Second, the aggregation method is described (annotated as SEM).

2.4.1 Loss Function

Parameters of each NN in an ensemble of size \( m \) are optimized with respect to the loss function \( \mathcal{L}_{QD} \) (eq. [5]).

The loss function operates on mini-batches of size \( n \).

\[ \mathcal{L}_{QD} = \mathcal{L}_{MPIW} + \lambda \frac{n}{n} \frac{1}{n} \mathcal{L}_{PICP} \]  

\[ \mathcal{L}_{MPIW} \] defined in eq. [6] is optimizing the width of the PIs capturing an observation. Due to variables \( c \) and \( k_i \) from eq. [2] and [3], \( \mathcal{L}_{MPIW} \) only includes those samples for which the observation is inside the PI.

\[ \mathcal{L}_{MPIW} = \frac{1}{c} \sum_{i=1}^{n} (\hat{y}_{U}^{(i)} - \hat{y}_{L}^{(i)}) \cdot k_i \]
\( L_{PICP} \) defined in eq. (7) is optimizing the coverage probability of the PIs penalizing only if the PICP is below the desired \( \gamma \).

\[
L_{PICP} = \max(0, (1 - \alpha) - PICP)^2 \quad (7)
\]

Expanded, we get

\[
L_{QD} = \frac{1}{c} \sum_{i=1}^{n} (\tilde{y}_L^{(i)} - \hat{y}_L^{(i)}) \cdot k_i + \lambda \frac{n}{\alpha(1 - \alpha)} \max(0, (1 - \alpha) - PICP)^2.
\]

The hyper-parameter (Lagrangian) \( \lambda \) controls the importance of \( L_{PICP} \) with respect to \( L_{MPIW} \). The intuition behind the fraction \( \frac{n}{\alpha(1 - \alpha)} \) is that it should reflect the confidence of \( L_{PICP} \) with respect to \( n \) and \( \alpha \).

### 2.4.2 Aggregation Method

Given an ensemble of \( m \) NN models fitted with respect to \( L_{QD} \) [5], we acquire prediction intervals \([\hat{y}_L^{(i)}, \hat{y}_U^{(i)}]\) for sample \( i \in \{1 \ldots n\} \) and model \( j \in \{1 \ldots m\} \). The following calculations are taken to acquire the final predictions intervals \([\tilde{y}_L^{(i)}, \tilde{y}_U^{(i)}]\) that should account also for the epistemic uncertainty.

\[
\mu_L^{(i)} = \frac{1}{m} \sum_{j=1}^{m} \hat{y}_L^{(ij)},
\]

\[
\mu_U^{(i)} = \frac{1}{m} \sum_{j=1}^{m} \hat{y}_U^{(ij)},
\]

\[
\sigma_L^{(i)^2} = \frac{1}{m-1} \sum_{j=1}^{m} (\hat{y}_L^{(ij)} - \mu_L^{(i)})^2,
\]

\[
\sigma_U^{(i)^2} = \frac{1}{m-1} \sum_{j=1}^{m} (\hat{y}_U^{(ij)} - \mu_U^{(i)})^2.
\]

These quantities are then combined to generate the final PIs. In the paper [Pearce et al. 2018], the aggregation is done as follows:

\[
\tilde{y}_L^{(i)} = \mu_L^{(i)} - 1.96 \cdot \sigma_L^{(i)}, \quad (8)
\]

\[
\tilde{y}_U^{(i)} = \mu_U^{(i)} + 1.96 \cdot \sigma_U^{(i)}. \quad (9)
\]

The actual implementation differs in using a standard error of the mean (SEM) \( \sigma_L^{(i)} \) [10] and \( \sigma_U^{(i)} \) [11] instead of in the paper presented standard deviation \( \sigma_L^{(i)} \) [8] and \( \sigma_U^{(i)} \) [9] respectively; notice the introduction of the scalar \( 1/\sqrt{m} \) in both equations.

\[
\tilde{y}_L^{(i)} = \mu_L^{(i)} - 1.96 \cdot \sigma_L^{(i)} \cdot \frac{1}{\sqrt{m}} \quad (10)
\]

Aggregating PIs using the latter equations (10) and (11) results in narrower PIs. Both aggregation methods lack any theoretical justification: The lower and upper PI boundaries are aggregated independently, i.e. without mutual consideration, and we consider this as a flaw of the method. In fact, under a simple assumption of normal posterior distribution that is correctly captured by \( m \) NN models but with shifted PIs with correct \( \gamma \) coverage probability (PICP), it can be shown that both original aggregation methods in the above equations yield PIs resulting in PICP greater than \( \gamma \), and thus greater MPIW. In the results, the latter aggregation function (eq. [10] and [11]) was evaluated (annotated as SEM).

### 3 METHOD

In this section, we describe our method annotated as SNM-QD+ that provides a point prediction along with prediction interval as output. First, an ensemble of neural network models is trained with an extended quality-driven loss function annotated as QD+ (eq. [12]). Second, the results from each model in the ensemble are aggregated to provide a final result (incorporating the epistemic uncertainty) by a fitting split normal density functions [Wallis 2014] from each NN’s point estimate and PI, and aggregating them into a split normal mixture (annotated as SNM) from which the final PI of coverage probability \( \gamma \) is calculated.

#### 3.1 Quality-Driven Loss Function

By having a single model for prediction intervals and point estimates, we achieve a coherency of the results and so avoid the case of two disjoint models learning different function approximations for prediction intervals and point estimates. In other words, providing prediction intervals and point estimates as a result of two disjoint models may not capture the predictive uncertainty of the point estimate model.

We therefore propose a new loss function

\[
L_{QD+} = (1 - \lambda_1)(1 - \lambda_2) \cdot L_{MPIW} + \lambda_1(1 - \lambda_2) \cdot L_{PICP} + \lambda_2 \cdot L_{MSE} + \xi \cdot L_P
\]

where point estimates \( \hat{y}^{(i)} \) are optimized by

\[
L_{MSE} = \frac{1}{n} \sum_{i=1}^{n} (\hat{y}_i^{(i)} - y^{(i)})^2
\]
We retrofit a slightly simplified version of the loss function $\lambda$ fitted from an ensemble (dashed) to this end, we assume that the posterior predictive distribution from a single model is a split normal distribution (Wallis 2014). The split normal distribution or two-piece normal distribution is a result of joining halves of normal distributions with the same mode but different variances. The split normal probability density function (PDF) is defined as

$$f_{SN}(x; \mu, \sigma_1, \sigma_2) = \begin{cases} A \exp \left(-\frac{(x-\mu)^2}{2\sigma_1^2}\right) & \text{if } x < \mu, \\ A \exp \left(-\frac{(x-\mu)^2}{2\sigma_2^2}\right) & \text{otherwise}, \end{cases}$$

where $A = \sqrt{2/\pi} (\sigma_1 + \sigma_2)^{-1}$.

Given the lower and upper PI bounds $[\hat{y}_L^{(ij)}, \hat{y}_U^{(ij)}]$ and the point estimates $\hat{y}^{(ij)}$ from an ensemble, we can fit parameters of a split normal distribution for each ensemble result by optimizing the loss

$$L_{SN} = [F_{SN}(\hat{y}_L^{(ij)}; \cdot) - \alpha/2]^2 + [F_{SN}(\hat{y}_U^{(ij)}; \cdot) - (1 - \alpha/2)]^2$$

where the split normal cumulative density function (CDF) is defined as

$$F_{SN}(x; \mu, \sigma_1, \sigma_2) = \begin{cases} \frac{\sigma_1 + \exp \left(-\frac{1}{2\sigma_1^2}(x-\mu)^2\right)}{\sigma_1 + \sigma_2} & \text{if } x < \mu, \\ \frac{\exp \left(-\frac{1}{2\sigma_2^2}(x-\mu)^2\right)}{\sigma_1 + \sigma_2} & \text{otherwise}. \end{cases}$$

Through a gradient descent (GD) optimization of eq. (15) we estimate the parameters $\sigma_1^{(ij)}$ and $\sigma_2^{(ij)}$ with respect to $[\hat{y}_L^{(ij)}, \hat{y}_U^{(ij)}]$ (variable $x$), $\hat{y}^{(ij)}$ (variable $\mu$) and $\alpha$.

When fitting the parameters $\sigma_1^{(ij)}$ and $\sigma_2^{(ij)}$ with eq. (15) as the objective, initialization of $(\sigma_1^{(ij)}, \sigma_2^{(ij)})$ plays an important role for finding the optimal solution. A random initialization does not always yield the optimal solution. Therefore, following the hypothesis that the parameters of a split normal distribution are close to those of normal distribution, we initialize the $\sigma_1^{(ij)}$ and $\sigma_2^{(ij)}$ as follows:

$$\sigma_1^{(ij)} = \frac{\hat{y}_L^{(ij)} - \hat{y}^{(ij)}}{\sqrt{2} \text{erf}^{-1}(2p_L - 1)},$$

$$\sigma_2^{(ij)} = \frac{\hat{y}_U^{(ij)} - \hat{y}^{(ij)}}{\sqrt{2} \text{erf}^{-1}(2p_U - 1)},$$

where $p_L = \alpha/2$ and $p_U = 1 - \alpha/2$. The above is derived from the inverse CDF (quantile function) of a normal distribution:

$$F_N^{-1}(p) = \mu + \sigma \sqrt{2} \text{erf}^{-1}(2p - 1)$$

with $p \in (0, 1)$.

In our experiments, this has been proven as superior compared to random initialization.

Figure 2: An example illustrating split normal PDFs (dashed) fitted from an ensemble ($n = 5$) of PI and point estimates, and subsequently aggregated into a split normal mixture (black) from which the final PI is calculated.
As a result of the above steps, we acquire an ensemble of $m$ split normal PDFs \( \{ f_{SN}(x_i) \}_{i=1}^{n} \) (for a single sample $i$) from which we create a mixture PDF

\[
f^{(i)}(x) = \frac{1}{m} \sum_{j=1}^{m} f_{SN}(x; \theta^{(ij)})
\]

See Figure 2 for illustration. The final PIs \( \tilde{y}_{L}^{(i)}, \tilde{y}_{U}^{(i)} \) are calculated from the mixture distribution $f^{(i)}(x)$ by numerically solving

\[
y_{L}^{(i)} = F^{-1}(i) (\alpha/2) \quad \text{and} \quad y_{U}^{(i)} = F^{-1}(i) (1 - \alpha/2)
\]

where $F^{-1}(i)(p)$ is the inverse CDF of the mixture $f^{(i)}(x)$.

The final point estimates are calculated as an equally weighted combination (mean) of point estimates from an ensemble.

\[
\hat{y}^{(i)} = \frac{1}{m} \sum_{j=1}^{m} y^{(ij)}
\]

4 EXPERIMENTS

Our proposed loss function QD+ (Section 3.1) and aggregation method SNM (Section 3.2) are compared with the original QDE (Pearce et al. 2018), specifically with the loss function QD (Section 2.4.1) and aggregation method SEM (Section 2.4.2). We also compare against an ensemble of NNs optimizing the parametrization of a normal distribution (Lakshminarayanan et al. 2017), referred as a mean-variance estimator (MVE). Additionally, we compare point estimates generated by our method SNM-QD+ with an ensemble of NNs optimizing MSE (eq. 13) for point estimates solely.

We follow a similar experimental setup initially set by Hernandez-Lobato and Adams (2015) and also followed in related works (Gal and Ghahramani 2016, Lakshminarayanan et al. 2017, Pearson et al. 2018, Tagasovska and Lopez-Paz 2019).

4.1 Datasets

Ten open-access benchmark datasets from the UCI dataset repository are used (Dua and Graff 2017). For each dataset we have created 20 shuffled versions across which we have evaluated the methods, i.e. 20 trials. Exceptions are the Year dataset with 1 trial (without shuffling) and the Protein dataset with 5 trials. The datasets are standardized (to zero mean and unit variance) based on the training set. The input of evaluation measures is standardized based on the full dataset for comparability across different trials.

4.2 Hyper-Parameters

We optimize for 95% prediction intervals, i.e. $\alpha = 0.05$. To not add any competitive advantage to our method, we keep the NN sizes identical across different models. Note that this may be a disadvantage for our model, given the complexity of the loss function QD+. This potentially sub-optimal setting should be kept in mind as our comparison’s objective is feasibility when compared to models specialized either to generate point estimates or PIs (and not both at the same time).

The ensemble size is $m = 5$. The size of mini-batches is $n = 100$ ($n = 1000$ for the Year dataset). All NNs have 2 hidden layers with 50 units (100 units for Protein and Year datasets) and ReLU activation functions.

For the hyper-parameter search (HPS), we do not follow the legacy of Hernandez-Lobato and Adams (2015). The HPS is performed only on a single concrete shuffled version of a dataset with excluded 10% test set, i.e. training set remains. Given the training set, the hyper-parameters (HPs) are validated on 5 shuffled 90% and 10% splits (i.e. 81% and 9% of the complete dataset) for training and validation, respectively. This deviates opposed to the originally suggested single 80% and 20% split (i.e. 72% and 18% of the complete dataset) without cross-validation. The change of the hyper-parameter search setup was motivated by the difficulty to find good HPs using the original setup, especially in smaller datasets (such as Boston, Concrete, Energy, Wine and Yacht).

A random HPS was performed on the following hyper-parameters (depending on the model): learning rate, decay rate, $\lambda_1$, $\lambda_2$, epochs. $\xi$ is set to 10 for all experiments. The remaining settings are consistent with those of Pearce et al. (2018). The applied heuristic for selecting the best hyper-parameters was as follows: If mean PICP is equal to $\gamma \pm 0.01$, then HPs with the lowest MPIW and MSE were selected. Alternatively, also heuristic considering mean and standard deviation of PICP proved good results. If neither of the criteria were fulfilled, the best achieved PICP was selected. Also, the training process on the validation set was analyzed for steady convergence and overfitting. Note that the loss cannot be used to select HPs because $\lambda_1$, $\lambda_2$ and $\xi$ scale parts of the loss function; hence the loss is associated with particular values of $\lambda_1$, $\lambda_2$ and $\xi$.

Due to a smaller number of HPS trials (max. 300), we did not always find well-performing HPs. Therefore, some experiments (QD+ for the datasets Energy, Kin8nm, Naval, Protein and Year) were manually fine-tuned. Adjusting hyper-parameters ($\lambda_1$ and $\lambda_2$, learning rate, decay rate...
Table 1: Ultimately, PIs are compared between SNM-QD+, SEM-QD and MVE, and point estimates are compared between SNM-QD+, MSE and MVE in Table 2. Additionally, aggregation methods are compared in SNM-QD+ vs. SEM-QD+. All methods are ensembles of 2-layer NNs. The values stand for mean ± standard error of the mean. The best results (with standard error of the mean taken in consideration) are shown in bold.

| Dataset  | SNM-QD+ | SEM-QD+ | SEM-QD | MVE  | SNM-QD+ | SEM-QD+ | SEM-QD | MVE  |
|----------|---------|---------|--------|------|---------|---------|--------|------|
| Boston   | 0.95 ± 0.01 | 0.97 ± 0.01 | 0.95 ± 0.01 | 0.88 ± 0.01 | 1.58 ± 0.06 | 1.82 ± 0.07 | 1.52 ± 0.06 | 0.89 ± 0.01 |
| Concrete | 0.94 ± 0.01 | 0.96 ± 0.01 | 0.97 ± 0.00 | 0.96 ± 0.00 | 0.99 ± 0.04 | 1.14 ± 0.05 | 1.36 ± 0.02 | 1.14 ± 0.02 |
| Energy   | 0.99 ± 0.00 | 0.99 ± 0.00 | 0.99 ± 0.01 | 0.98 ± 0.01 | 0.29 ± 0.01 | 0.31 ± 0.02 | 0.48 ± 0.03 | 0.16 ± 0.00 |
| Kin8nm   | 0.97 ± 0.00 | 0.98 ± 0.00 | 0.99 ± 0.00 | 0.97 ± 0.00 | 1.07 ± 0.01 | 1.21 ± 0.01 | 1.29 ± 0.01 | 1.04 ± 0.01 |
| Naval    | 1.00 ± 0.00 | 1.00 ± 0.00 | 0.99 ± 0.00 | 1.00 ± 0.00 | 0.99 ± 0.00 | 0.10 ± 0.00 | 0.36 ± 0.02 | 0.05 ± 0.00 |
| Power    | 0.95 ± 0.00 | 0.96 ± 0.00 | 0.96 ± 0.00 | 0.96 ± 0.00 | 0.60 ± 0.00 | 0.85 ± 0.00 | 0.87 ± 0.00 | 0.87 ± 0.01 |
| Protein  | 0.95 ± 0.00 | 0.97 ± 0.00 | 0.95 ± 0.00 | 0.98 ± 0.00 | 2.12 ± 0.01 | 2.31 ± 0.01 | 2.24 ± 0.01 | 2.82 ± 0.10 |
| Wine     | 0.94 ± 0.01 | 0.95 ± 0.01 | 0.92 ± 0.01 | 0.94 ± 0.00 | 2.62 ± 0.06 | 2.92 ± 0.07 | 2.06 ± 0.03 | 2.94 ± 0.02 |
| Yacht    | 0.94 ± 0.01 | 0.96 ± 0.01 | 1.00 ± 0.00 | 0.99 ± 0.00 | 0.12 ± 0.00 | 0.13 ± 0.00 | 0.25 ± 0.02 | 0.40 ± 0.04 |
| Year     | 0.94 ± NA   | 0.96 ± NA   | 0.95 ± NA   | 0.96 ± NA   | 2.34 ± NA   | 2.54 ± NA   | 2.29 ± NA   | 2.75 ± NA   |

Table 2: Extension of Table 1 evaluating point estimates of SNM-QD+, MSE and MVE.

| Dataset  | SNM-QD+    | SEM-QD+    | MVE  |
|----------|------------|------------|------|
| Boston   | 0.112 ± 0.013 | 0.115 ± 0.013 | 0.127 ± 0.019 |
| Concrete | 0.056 ± 0.003 | 0.053 ± 0.003 | 0.077 ± 0.004 |
| Energy   | 0.001 ± 0.000 | 0.001 ± 0.000 | 0.001 ± 0.000 |
| Kin8nm   | 0.060 ± 0.001 | 0.059 ± 0.001 | 0.062 ± 0.001 |
| Naval    | 4.865 ± 0.000 | 1.3e-4 ± 0.000 | 1.2e-4 ± 0.000 |
| Power    | 0.042 ± 0.001 | 0.050 ± 0.001 | 0.048 ± 0.001 |
| Protein  | 0.310 ± 0.002 | 0.361 ± 0.004 | 0.445 ± 0.023 |
| Wine     | 0.597 ± 0.015 | 0.616 ± 0.018 | 0.621 ± 0.015 |
| Yacht    | 0.602 ± 0.000 | 0.001 ± 0.000 | 0.003 ± 0.000 |
| Year     | 0.637 ± NA   | 0.636 ± NA   | 0.639 ± NA   |

and number of epochs) has proven to be intuitive by analyzing PICP, MPIW and MSE on a validation set. We resorted to manual fine-tuning only if a model was considerably under-performing compared to the original QDE or compared to MVE. There was no attempt to optimize to the model’s full potential.

4.3 Evaluation

The input of evaluation measures is standardized based on the full dataset for comparability across different trials. The following measures are used to assess and compare models (QD+, QD and MVE) and aggregation methods (SNM and SEM): PICP, MPIW and mean squared error (MSE).

4.4 Results

Tables 1 and 2 show the results of comparing the two models QD+ and QD, combined with the aggregation methods SNM and SEM. We have also included a model that generates only point estimates (denoted MSE), and MVE generating both PIs and point estimates. Consider SNM-QD+ vs. SEM-QD for a comparison between our method and the original QDE method [Pearce et al. 2018], respectively. SNM-QD+ vs. SEM-QD+ gives a comparison between our split normal aggregation method and the original aggregation method, respectively. SNM-QD+ vs. MVE provides a comparison of methods generating both PIs and point estimates. Finally, Table 2 shows the evaluation of the point estimates between SNM-QD+, MSE and MVE. The best results (with standard error of the mean taken in consideration) are presented using bold font. Supplement [A] provides the evaluation of non-aggregated PIs.

Overall, the results demonstrate that a complex multi-objective quality-driven loss function (QD+) can deliver well-calibrated PIs when the split normal mixture (SNM) is employed as the aggregation method. The proposed method SNM-QD+ performs best with respect to PIs and competitively with respect to point estimates. We argue that the quality of the point estimates can be improved by increasing the capacity of NNs or by hyper-parameter fine-tuning.

The results clearly show that the split normal aggregation method (SNM) yields well-calibrated PIs, while the original aggregation method (SEM) generally tends to generate wider PIs. Contrary to SEM, SNM aggregation method does not treat PI boundaries independently. It fits flexible enough (asymmetric) distributions within a versatile mixture distribution that is, as the results show, a sufficient approximation of the posterior predictive distribution. Note that the SNM aggregation method is not necessarily tied to the QD+ loss, but can be applied in other situations as well. A potential drawback of SNM is the apparent computational overhead of fitting the split normal mixture when the number of samples is large. However, this learning task is parallelizable and distributable; hence the wall-clock time spent on the task is negligible.

The training process of QD+ is, contrary to the original QD, quite robust. To give a particular example with the
Protein dataset, training 2-layer NNs with the original QDE required approximately one retry per run due to interval crossing or high loss value (stuck in local minimum). No retries were required when using QD+. Details are given in Supplement B. Furthermore, the original QD is sensitive to parameter initialization. Overall, the penalty function \( L_P \) (eq. 14) in QD+ results in significant improvements of the stability of the training process and strengthens the integrity of the output (mitigates the undesired interval crossing and point estimates out of the PI bounds). Supplement C provides a sensitivity analysis together with an ablation study clearly demonstrating the importance of the penalty function.

Since we compare our results with Pearce et al. (2018) and Lakshminarayanan et al. (2017), our work is also comparable with Bayesian approach by Hernandez-Lobato and Adams (2015), and partly also with Tagasovska and Lopez-Paz (2019).

4.5 Implementation

Our implementation is shared\footnote{https://github.com/tarik/pi-snm-qde} and results fully reproducible. To embrace reproducibility, we use the Python package Sacred (Greff et al., 2017). PyTorch (Paszke et al., 2019) is used as the main NN framework and the fitting of a split normal mixture is implemented using JAX (Bradbury et al., 2018).

5 CONCLUSIONS

The main finding of this paper is that models delivering point estimates together with prediction intervals are competitive to models providing only one or the other. Motivated by the results of the quality-driven deep ensembles (Pearce et al., 2018) as an alternative to Bayesian methods, we endeavored to address the three key limitations: (1) The inability to generate point estimates; (2) the weakly justified method for aggregation of prediction interval ensembles; (3) the fragile training process.

We propose a new quality-driven loss function generating both prediction intervals and point estimates, and we dramatically increase the robustness of the training process by integrating a penalty function.

A unique and well-founded method fitting a split normal mixture as the aggregate of ensembled neural network output generates well-calibrated prediction intervals accounting for aleatoric and epistemic uncertainty. Moreover, an analytical approach for initializing parameters of a split normal probability density function is proposed that leads to acceleration and dramatically increased success of the fitting process.

With this work, we extend the practitioners toolset for quantifying predictive uncertainty in the regression task.

6 FUTURE WORK

We end the paper with the following suggestions for future research or possible improvements:

- We have shown that a single model architecture can be used to generate both point estimates and prediction intervals. We therefore envision a two-step learning process, where one first optimizes the model to generate point estimates and consequently learns to generate prediction intervals. To avoid catastrophic forgetting, it seems appropriate to treat this as a transfer learning problem. The key benefit of this operation is that it will allow us to reuse (pre-trained) NN models that currently only generate point estimates.

- It seems reasonable to use the NMPIW measure (eq. 4) in place of MPIW in the definition of the loss function \( L_{MPIW} \) (eq. 6). While it is already relatively intuitive to find HPs manually, and the HPS can be automated efficiently, we hypothesise that using NMPIW may lead to better-scaled hyper-parameters and possibly also to smaller variations across different datasets. The current work uses MPIW to ease the comparison between QD and QD+.

- The current approach learns to output a reasonable PI that is later used to fit a split normal PDF. This could alternatively be simplified so that the output of the neural network was the parameterization of the split normal PDF (or a mixture of split normal PDFs) directly. This approach is relevant for use-cases where a full predictive distribution is needed and would require a training objective defined for such situations instead of QD+. Again, the implemented approach was chosen to extend the non-Bayesian branch of research and ease comparison with the work of Pearce et al. (2018).

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SUPPLEMENTARY MATERIAL

A Additional Results

Related to Table 1, Table A shows the evaluation of non-aggregated PIs (not accounting for epistemic uncertainty) for the QD+ and QD models.

Table A: Evaluation of non-aggregated PIs (2-layer NNs).

| Dataset | PICP | QD | MPIW |
|---------|------|----|------|
| Boston  | 0.85 ± 0.01 | 0.88 ± 0.01 | 1.22 ± 0.04 | 1.12 ± 0.02 |
| Concrete | 0.86 ± 0.01 | 0.89 ± 0.00 | 0.79 ± 0.03 | 1.03 ± 0.01 |
| Energy  | 0.90 ± 0.01 | 0.93 ± 0.00 | 0.21 ± 0.01 | 0.27 ± 0.01 |
| K88nm   | 0.91 ± 0.00 | 0.92 ± 0.00 | 0.95 ± 0.01 | 1.01 ± 0.01 |
| Naval   | 0.98 ± 0.00 | 0.94 ± 0.00 | 0.08 ± 0.00 | 0.25 ± 0.00 |
| Power   | 0.94 ± 0.00 | 0.93 ± 0.00 | 0.79 ± 0.00 | 0.75 ± 0.00 |
| Protein | 0.93 ± 0.00 | 0.93 ± 0.00 | 2.03 ± 0.01 | 2.11 ± 0.01 |
| Wine    | 0.91 ± 0.00 | 0.91 ± 0.00 | 2.31 ± 0.03 | 1.81 ± 0.01 |
| Yacht   | 0.76 ± 0.01 | 0.89 ± 0.01 | 0.07 ± 0.00 | 0.15 ± 0.01 |
| Year    | 0.93 ± 0.00 | 0.93 ± 0.00 | 2.27 ± 0.00 | 2.07 ± 0.02 |

In Table B, we show the reproduced results of the original QDE method following the exact experimental setting as implemented in [Pearce et al. 2018]. These results are included to offer a comparison between the 1-layer and 2-layer models (Table B and Table 1, respectively). It also serves as a verification of our re-implementation of the method by [Pearce et al. 2018].

Table B: Original QDE with 1-layer NNs as a reference.

| Dataset | PICP | QD | MPIW |
|---------|------|----|------|
| Boston  | 0.90 ± 0.01 | 0.80 ± 0.01 | 1.01 ± 0.01 | 0.81 ± 0.01 |
| Concrete | 0.92 ± 0.01 | 0.84 ± 0.00 | 1.01 ± 0.01 | 0.83 ± 0.00 |
| Energy  | 0.97 ± 0.00 | 0.91 ± 0.00 | 0.45 ± 0.01 | 0.38 ± 0.01 |
| K88nm   | 0.96 ± 0.00 | 0.89 ± 0.00 | 1.24 ± 0.00 | 1.02 ± 0.00 |
| Naval   | 0.99 ± 0.00 | 0.96 ± 0.00 | 0.25 ± 0.02 | 0.17 ± 0.00 |
| Power   | 0.95 ± 0.00 | 0.94 ± 0.00 | 0.85 ± 0.00 | 0.81 ± 0.00 |
| Protein | 0.95 ± 0.00 | 0.94 ± 0.00 | 2.26 ± 0.00 | 2.17 ± 0.00 |
| Yacht   | 0.93 ± 0.01 | 0.91 ± 0.00 | 2.31 ± 0.02 | 2.04 ± 0.01 |
| Year    | 0.95 ± NA | 0.93 ± 0.00 | 2.41 ± NA | 2.22 ± 0.01 |

Table C considers a simple baseline with point estimates constructed as the mean of PIs from SEM-QD (annotated SEM-QD*). This simple baseline is underperforming compared to the results in Table 2.

Table C: Extension of Table 2 provides a simple baseline with point estimates constructed as the mean of PIs from SEM-QD (annotated SEM-QD*). Also, the sizes |D| and input dimensions d of the datasets are presented.

| Dataset | Failures/Retries | Target Model Count |
|---------|-----------------|--------------------|
| Boston  | 0               | 5 · 20             |
| Concrete | 0   | 5 · 20             |
| Energy  | 0               | 5 · 20             |
| K88nm   | 0               | 5 · 20             |
| Naval   | 0               | 5 · 20             |
| Power   | 0               | 5 · 20             |
| Protein | 0               | 5 · 20             |
| Wine    | 0               | 5 · 20             |
| Yacht   | 0               | 5 · 20             |
| Year    | 0               | 5 · 1              |

In Table 3, we show the number of failed/repeated training attempts of QD+ and QD. The targeted model count equals to an ensemble size m times number of trials t.

| Dataset | Failures/Retries | Target Model Count |
|---------|-----------------|--------------------|
| Boston  | 0               | 5 · 20             |
| Concrete | 0   | 5 · 20             |
| Energy  | 0               | 5 · 20             |
| K88nm   | 0               | 5 · 20             |
| Naval   | 0               | 5 · 20             |
| Power   | 0               | 5 · 20             |
| Protein | 0               | 5 · 20             |
| Wine    | 0               | 5 · 20             |
| Yacht   | 0               | 5 · 20             |
| Year    | 0               | 5 · 1              |

C Sensitivity Analysis

Figure A provides a sensitivity analysis with respect to hyper-parameters λ1, λ2 and ξ. It visualizes the measures PICP, NMPIW and MSE in relation to changing hyper-parameters of the QD+ model. The analysis was performed on the Yacht dataset. The sensitivity analysis is also an ablation study since the border values, i.e. λ1, λ2 ∈ {0, 1} and ξ = 0, disable certain parts of the LQD+ loss function (eq. 12).

Figure A would guide us to choose λ1 around 0.975 and λ2 around 0.05. Leaving out the penalty function from the loss function (i.e. ξ = 0) leads to a non-converging training process for λ2 below approx. 0.7 and more noisy measures in the remaining hyper-parameter space (Figure A). It demonstrates the critical importance of the penalty function L_P (eq. 14), thus, supporting the claims about the robustness together with the evidence in Table 2.

B Robustness

The robustness of training process of QD+ vs. the fragility of training process of QD is shown in Table D. The contribution of the penalty function L_P to the robustness is shown in Figure A of the following section.
Figure A: Sensitivity analysis of QD+ on Yacht dataset with respect to hyper-parameters $\lambda_1$ and $\lambda_2$ with (a) and without (b) an active penalty function $\mathcal{L}_P$. Gray color stands for a failed training process, i.e. high loss or outputs violating the semantic integrity (PI crossing or point estimates outside the PI bounds).