# POST-HOC UNCERTAINTY QUANTIFICATION FOR QT INTERVAL MEASUREMENTS WITH ENSEMBLES OF ELECTROCARDIOGRAPHIC LEADS AND DEEP MODELS

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#### ABSTRACT

Standard electrocardiography (ECG) allows to record the electrical activity of the heart from different angles called leads. The QT interval, which corresponds to the time elapsed between the onset of ventricular contraction and the end of ventricular relaxation, is an ECG biomarker of drug cardiotoxicity. Deep neural networks (DNNs) have achieved state-of-the-art performance in QT interval measurement but are missing uncertainty quantification, which is necessary for safer decision making. Uncertainty is usually encoded in DNNs through probability distributions over model weights. In this paper, we combine this approach with notions of multisensory integration whereby neural systems account for uncertainty by optimally integrating all available sensory inputs. We thus approximate the posterior predictive distribution of the QT interval given a multi-lead ECG as a weighted average across leads (lead integration) and models (deep ensembling) and derive  $100(1 - \alpha)$ % Bayesian prediction intervals (PIs). We apply this method to QT-based cardiac drug safety monitoring and compare it to an adapted version of conformal prediction. The Bayesian and conformal approaches yield comparable empirical coverage (77%-82% for mean PI widths of ~28 milliseconds,  $\alpha = 0.1$ ). The former is more straightforward and shows better error-based calibration. Data and code implementation are available at https://github.com/mouslyddiaw/qt-uncertainty.

# **1** INTRODUCTION

Electrocardiography (ECG) typically requires 10 on-skin electrodes to record the cardiac electrical activity from 12 different angles or leads: I, II, III, aVR, aVL, aVF (limb leads) and V1-V6 (chest leads). The QT interval, measured from the start of the QRS complex to the end of the T wave (cf. ECG annotations on Lead I, Figure 1), represents the duration of ventricular contraction and relaxation. Its prolongation is a surrogate biomarker for the risk of torsades de pointes (TdP), a life-threatening arrhythmia. ECG monitoring is therefore essential to the prevention of TdP induced by otherwise useful medications.

The American Heart Association provided guidelines for such ECG monitoring (Drew et al., 2004; 2010; Tisdale et al., 2020) but their use is yet to be widespread (Putnikovic et al., 2022). Given the difficulty and unreliability of manual QT measurements (Malik, 2004; Lester et al., 2019), robust algorithms could allow accurate and real-time monitoring, specially amongst non-cardiologists—for instance, drug-induced QT prolongation is prevalent in psychiatry (Ali et al., 2020). Deep Learning (DL) has achieved state-of-the-art ECG interval measurement performance (Giudicessi et al., 2021; Hicks et al., 2021; Diaw et al., 2022). However, uncertainty quantification (UQ) of DL-based ECG interval measurements, important for safe decision making, remains under-explored.

In DL, predictive uncertainty has mostly been quantified through the posterior over model weights using Bayesian neural networks (BNNs) or approximates (Gal & Ghahramani, 2016). Deep ensembling (Lakshminarayanan et al., 2017), which also fits within approximate Bayesian inference (Wilson & Izmailov, 2020), is fast becoming the gold standard for UQ in DNNs as it has outperformed



Figure 1: Construction of QT prediction intervals for a standard 12-lead ECG using a deep ensemble

approximate BNNs (Ovadia et al., 2019; Ashukha et al., 2020). Less common UQ approaches in DL include test-time data augmentation, mostly used in medical image processing (Ayhan & Berens, 2018; Wang et al., 2019). Conformal prediction (Vovk et al., 2005) has been of great interest for the machine learning (ML) community at large (Angelopoulos et al., 2020; Ndiaye, 2022) as it is distribution-free and model-agnostic.

In cognitive science (e.g. study of perception), uncertainty is usually encoded at input level of neural systems rather than in synaptic weights (equivalent of DNN weights). Pearce (2020) suggests that incorporating this paradigm in current UQ methods for DL could push the field forward. This is relevant to QT interval measurement that starts, in common practice, with an efficient visualization of all leads of a given ECG recording, similar to how the visual system optimally perceives an object by combining all available sensory signals (Jacobs, 1999). Here, uncertainty depends more on the object being observed (external or aleatoric uncertainty) than on the observer (internal or epistemic). In this paper, we consider approximate Bayesian and conformal approaches to generate QT prediction intervals (PIs). Our contributions are: (i) We draw insights from how the visual system accounts for uncertainty to propose a probabilistic model of QT measurement on multi-lead ECG recordings (lead integration). We combine lead integration with deep ensembling to approximate the posterior predictive distribution from which a PI is derived (cf. Figure 1). (ii) We build adaptative conformal PIs by leveraging the ECG features learned by a deep ensemble (iii) We demonstrate the Bayesian and conformal approaches on real-world clinical data. The Bayesian method presents interesting features for clinical decision support as it yields high-quality PIs, like the conformal predictor, and is easier to implement.

# 2 UNCERTAINTY QUANTIFICATION

#### 2.1 APPROXIMATE BAYESIAN INFERENCE

Standard ECG recordings have multiple leads, each sensing the cardiac electrical activity from a different spatial viewpoint. Denote  $D = \{(X_t^i, y_i)\}_{i=\{1,...,n\}}$  a dataset of n ECG samples where  $X_t = \{x_t^l\}_{l=\{1,...,L\}}$  represents a L-lead ensemble of ECG beats (typically, L = 12) and y the corresponding QT interval. Denote  $f_{\theta} : x_t \to y$  a QT estimator parameterized by  $\theta$ , optimized on D and capable of analyzing all L types of lead. We propose to approximate the posterior predictive distribution  $Y = p_{\theta}(y|X_t, D)$  by generating multiple QT estimates  $f_{\theta}(x_t^l)$ . We subsequently derive a PI of level  $1 - \alpha$ ,  $\alpha \in [0, 1]$ , following the equal-tailed method,  $PI = [y^{(\alpha/2)}, y^{(1-\alpha/2)}]$  where  $y^{(\alpha/2)}$  and  $y^{(1-\alpha/2)}$  are the  $\alpha/2$  and  $1 - \alpha/2$  quantiles of the predictive distribution Y.

Lead integration. In its simplistic Bayesian model, the human visual system encodes uncertainty by linearly combining all available cues or sensory signals, each weighed in proportion to their reliability (cf. Appendix A.1). Human ECG annotators usually solve the task of finding the QT interval y by superimposing all lead components of  $X_t$  to better define the beginning of the QRS complex and end of the T wave. This resembles an optimal cue integration system as the annotator aggregates all information provided by the different leads, which can have various degrees of reliability. For instance, some leads might have lower amplitude T waves or be noisier. We propose to encode this notion of lead reliability in automated measurement using a linear pooling model (Stone, 1961),

$$Y = p_{\theta}(y|X_t, D) = p_{\theta}(y|x_t^1, \dots, x_t^L, D) = \sum_{l=1}^L w_l p_{\theta}(y|x_t^l, D)$$
(1)

where the weight  $w_l$  denotes the reliability of lead  $x_t^l$  relative to the other leads, all weights summing to 1. This formulation requires prior knowledge on lead reliability.

**Implicit combination with deep ensembling (UQ-EL)**. Denote  $\{f_{\theta_1}, \ldots, f_{\theta_k}\}$  an ensemble of k single-lead DL models parameterized by  $\theta_1, \ldots, \theta_k$  respectively. We can define the mean QT  $1 \sum_{k=1}^{k} \frac{1}{k}$ 

estimator  $f_{\theta} = \frac{1}{k} \sum_{j=1}^{k} f_{\theta_j}$ . Assuming that, a priori, all leads have fairly equal reliability ( $w_l \approx 1/L$ ),

we then approximate Y (Equation 1) with the L estimates  $f_{\theta}(x_t^l)$ .

**Explicit combination with deep ensembling (UQ-ELM**<sup>1</sup>). The posterior distribution of the deep ensemble is approximatively a uniformly weighted mixture model (Lakshminarayanan et al., 2017),

$$p_{\theta}(y|x_t, D) \approx \frac{1}{k} \sum_{j=1}^{k} p(y|x_t, \theta_j)$$
(2)

With  $w_l \approx 1/L$ , we can rewrite the predictive distribution  $p_{\theta}(y|X_t, D)$  using Equations 1 and 2 as

$$Y = p_{\theta}(y|X_t, D) \approx \frac{1}{L} \sum_{l=1}^{L} p_{\theta}(y|x_t^l, D) \approx \frac{1}{k \times L} \sum_{l=1}^{L} \sum_{j=1}^{k} p(y|x_t^l, \theta_j)$$
(3)

Here, we approximate Y using all the  $k \times L$  estimates  $\hat{y}_{l,j} = f_{\theta_j}(x_t^l)$  instead of the L model averages used in UQ-EL.

# 2.2 LOCALLY ADAPTATIVE SPLIT CONFORMAL PREDICTION (LASCP)

Given n past observations and a prespecified miscoverage rate  $\alpha$ , CP consists in fitting a model  $f_{\theta}$  on the n samples and building for a new observation  $Y_{n+1}$  a predictive interval  $PI_{\alpha}$  such that  $P(Y_{n+1} \in PI_{\alpha}) \geq 1 - \alpha$ . In split CP (SCP) (Papadopoulos et al., 2002), the n samples are split in 2 sets for model fitting (performed on, say,  $D_{train}$ ) and nonconformity calibration (performed on  $I_2$  of size  $n_2$ ). Nonconformity scores are computed as absolute residuals  $\epsilon = |y - \hat{y}|$  and the  $[(n_2 + 1)(1 - \alpha)]/n_2$  quantile of the scores  $(\hat{q}_{n_2})$  calibrated on  $I_2$  is used to define constant-length PIs  $[\hat{y} - \hat{q}_{n_2}, \hat{y} + \hat{q}_{n_2}]$ . LASCP (Papadopoulos et al., 2008; 2011; Lei et al., 2018) creates adaptative intervals by first fitting a model  $r : x \to \epsilon$  (residual fitting) on another calibration dataset  $I_1$ . In the traditional LASCP algorithm, f and r have the same explicit input features. In this paper, we use the ECG features learned by a deep ensemble to train a shallow ML model for residual fitting. The proposed framework for LASCP calibration on L-lead ECG data using a deep ensemble is detailed in Appendix A.2.

# **3** EXPERIMENTS AND RESULTS

**Clinical data**<sup>2</sup>. Study 1 (**S1**) stems from a prospective randomized placebo-controlled clinical trial including 22 healthy subjects (Johannesen et al., 2014). Each subject was followed during 5 periods during which they received a placebo (*P*) or one of the following drugs: Dofetilide (*D*), Quinidine (*Q*), Ranolazine (*R*), and Verapamil (*V*). For each period, 12-lead ECGs were recorded for 24 hours and 3 10-second ECGs were extracted at 16 timepoints (1 point pre-dose and 15 points post-dose) leading to a total of 5219 10s ECG recordings. Similarly, Study 2 (**S2**) aimed at analyzing the ECG effects of *D*, Lidocaine + *D*, Mexiletine + *D* and Moxifloxacin + Diltiazem (Johannesen et al., 2016). ECGs were extracted at 14 timepoints, leading to a total of 4211 10s ECG recordings. Semi-automated QT annotations made on representative median beats are available for each 12-lead ECG in S1 and S2.

**Deep ensembling with 5-fold cross-validation**. Resampling techniques can be used to generate multiple distinct models from a single training set. We constitute a training set using ECGs recorded before/after administration of V (non QT-prolonging drug) and Q (QT-prolonging) so as to learn a wide range of ECG morphologies. We refer to this set as **S1a** and the remaining data in S1 as **S1b**. We split in 5 groups the 22 subjects in S1a for patient-stratified cross-validation. We generate 5 versions of the single-lead residual neural network (ResNet) proposed by Diaw et al. (2022). Data

<sup>&</sup>lt;sup>1</sup>Stands for UQ with an Ensemble of Leads and Models

<sup>&</sup>lt;sup>2</sup>Available at https://physionet.org/content/ecgrdvq/1.0.0/ and https://physionet.org/content/ecgdmmld/1.0.0/

preparation and DNN optimization are conducted as in the original paper (Diaw et al., 2022). For a single-lead ECG input (1.2 second average beat sampled at 500 Hz), the ResNet yields d = 32 global features by average pooling the last feature maps before the regression head.

**LASCP calibration**. We split S1b in two distinct sets, S1b =  $I_1 \cup I_2$  (cf. Appendix A.3 for a summary of the subsets). We train a gradient boosting regressor on  $I_1$  for residual fitting. Model parameters and residual RMSE scores are detailed in Appendix A.4.

**Evaluation metrics**. We compute the prediction interval coverage probability (CP), defined as the proportion of target QT intervals that fall within the PI, the mean prediction interval width (MW), and the mean absolute deviation (MAD), which measures how far the target QT intervals not covered by the PIs are from the closest PI bounds (lower or upper).



Data	Method	СР	MW (ms)	MAD (ms)
S1b (N = 3163)	UQ-EL	82%	<b>29.49</b>	<b>7.20</b>
	UQ-ELM	<b>95%</b>	43.91	9.81
S2 (N = 4211)	UQ-EL	77%	<b>28.27</b>	3.95
	UQ-ELM	<b>90%</b>	40.47	<b>3.26</b>
	LASCP	82%	28.67	3.50

Table 1: Evaluation of the 90% PIs ( $\alpha = 0.1$ ) on N 12-lead ECG ensembles



Figure 3: Error-based calibrationFigure 4: 24-hour QT monitoring following drug intake (90% PIs)

**Results**. We evaluated the Bayesian methods (UQ-EL and UQ-ELM) on S1b and S2 and the LASCP method on S2. As shown in Figure 2 where coverage is plotted as a function of  $\alpha$  for each study and method, the 3 methods are overall well-calibrated, specially UQ-ELM, which always achieves above nominal coverage, i.e.  $\geq 100(1 - \alpha)\%$ . Table 1 details the performance of the 3 methods for  $\alpha = 0.1$  based on the aforementioned evaluation metrics. The results suggest that UQ-EL and LASCP yield PIs of comparable quality and though their coverage is lower than that of UQ-ELM, the out-of-bound target QT intervals do not fall far from the PIs as reflected by the low MADs.

Figure 3 shows the error-based calibration plots of the 3 methods obtained by first dividing the set of absolute differences between ground-truth and mean prediction (errors) in S2 into bins of 100 samples and then computing the MW and mean absolute error (MAE) within each bin. PI width seems to increase more consistently with model error with the Bayesian methods than with LASCP, which makes them useful for human review of less accurate DL-based QT measurements flagged based on PI width. Figure 4 illustrates patient-specific QT-based drug safety monitoring based on PIs generated with UQ-EL and LASCP. Both methods yield 90% PIs that contain, most of the time, the actual QT interval and provide reliable information on drug-induced QT prolongation (or lack thereof). We refer to Appendix A.5 for illustrations of the effects on the ECG of Dofetilide, known to significantly prolong the QT interval, and the impact on model performance.

# 4 CONCLUSION

TdP risk could be better managed with individualized, frequent and automated QT monitoring. In this paper, we improve the trustworthiness of DL-based QT estimators with Bayesian and conformal approaches to UQ. While we focus in providing high-quality PIs for our application, we could additionally leverage uncertainty to improve predictive performance, which is one of the main goals of UQ in DL, as done in Lakshminarayanan et al. (2017). We could also study the impact of other ensembling techniques on UQ-E(L)M and investigate methods for weighing lead reliability and model confidence so as to provide, when necessary, point estimates more accurate than averages. Zhu et al. (2014) conducted a similar study to better aggregate crowd-sourced ECG annotations.

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# A APPENDIX

#### A.1 PROBABILISTIC INFERENCE FOR MULTISENSORY INTEGRATION

The brain must reason probabilistically for optimal performance as uncertainty is intrinsic to most tasks. To this aim, the human visual system performs sensory cue (or multisensory) integration over space and time by first making estimates based on each available cue or sensory signal alone before linearly combining all estimates, weighing each cue in proportion to their reliability. This phenomenon, termed weak fusion (Clark & Yuille, 1990), has been modelled using Bayesian probability but the notion of cue reliability is fairly intuitive. Take the experiment conducted by Ernst & Banks (2002) on height estimation based on visual and tactile cues. We understand that these two input signals are not equally reliable. For instance, in complete darkness, any estimate based on vision is nothing but noise. Similar ideas are expressed in the field of probabilistic opinion pooling when individual opinions are aggregated to form a consensus (Stone, 1961).

One of the main results of perception modelling is that when the uncertainty associated with each cue available to the visual system is approximated by a Gaussian likelihood function, the average estimate made by the ideal observer is a weighted average of the average estimates that would be derived from each cue alone. More formally, denote s the parameter being estimated and  $\{s_1, \ldots, s_n\}$  a set of n cues. Assuming that the cues are conditionally independent, we can write from Bayes' rule that

$$p(s|s_1,\ldots,s_n) \propto \left(\prod_{i=1}^n p(s_i|s)\right) p(s) \tag{4}$$

Then assuming that the prior p(s) is uniform, i.e. all values of s are equally probable before observation, we write

$$p(s|s_1,\ldots,s_n) \propto \prod_{i=1}^n p(s_i|s)$$
(5)

Under Gaussian assumption, reasonable in light of the central limit theorem, we can define the average (or maximum likelihood) estimate  $\hat{s}$ , often referred to as optimal cue integration,

$$\hat{s} = \sum_{i=1}^{n} w_i \hat{s}_i \tag{6}$$

where  $w_i = r_i / \sum_{i=1}^n r_i$  denotes the weight of cue  $s_i$ ,  $r_i = 1/\sigma_i^2$  its reliability and  $\sigma_i^2$  its variance.

Equation 6 is at the basis of the experiments conducted in vision research to understand how the human visual system integrates multisensory information (Jacobs, 1999; Ernst & Banks, 2002; Knill & Saunders, 2003; Fetsch et al., 2012).

### A.2 LASCP ALGORITHM

We keep the same notations as in Section 2. The proposed algorithm for LASCP calibration on *L*-lead ECG data using a deep ensemble is as follows:

- 1. For each *L*-lead ensemble in the first calibration set  $I_1 = \{(X_t^i, y_i)\}_{i=\{1,...,n_1\}}$  of size  $n_1$ , compute the absolute residuals  $\epsilon_i = |y_i \bar{y}_i|$  with  $\bar{y}_i = \frac{1}{k \times L} \sum_{l=1}^L \sum_{j=1}^k \hat{y}_{l,j}$
- 2. Train a model  $r : (\mathbb{R}^d)^{k \times L} \to \mathbb{R}$  on  $\{(G_i, \epsilon_i)\}_{i \in \{1, \dots, n_1\}}$  with  $G_i$  the set of d-dimensional single-lead ECG features extracted by the ensemble of k models.
- 3. For every sample in the second calibration set  $I_2 = \{(X_t^i, y_i)\}_{i=\{1,...,n_2\}}$ , compute the nonconformity score  $s_i = \frac{|y_i \bar{y}_i|}{r(G_i)}$ . Then, compute the aforementioned quantile  $\hat{q}_{n_2}$  of  $S = \{s_i\}_{i=\{1,...,n_2\}}$ . For each new ECG sample, derive  $PI = [\bar{y} \hat{q}_{n_2}r(G), \bar{y} + \hat{q}_{n_2}r(G)]$ .

## A.3 DATA SPLITTING

Table 2 details the number of subjects and 12-lead ECG recordings in the subsets sampled from S1 and S2.

Table 2: Number of subjects and 12-lead ECG recordings in the datasets

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# A.4 CONFORMAL RESIDUAL FITTING

We trained a gradient boosting regressor using Python's Scikit-learn library (v1.1.2). The hyperparameters were optimized by cross-validated grid-search, resulting in 500 boosting stages to perform ( $n\_estimators = 500$ ) and a learning rate of 0.01. Table 3 details the resulting RMSE scores.

Table 3: Residual fitting evaluation

Split	RMSE (ms)
$I_2$	6.366

#### A.5 ECG ILLUSTRATIONS

Figure 5 illustrates prediction intervals on 2 ECGs from the same subject before the administration of the QT-prolonging Dofetilide, i.e. at baseline, and 12 hours afterwards. The PI is tighter at baseline where the T wave on all ECG beats are more or less prominent and end around the same time. 12 hours later, the drug affects the ECG by lengthening the T wave and lowering its amplitude, each lead in its own way, which increases uncertainty.



Figure 5: Examples of PIs on ECGs recorded before and 12 hours after administration of Dofetilide (drug known to prolong the QT interval) by a subject whose QT profile is illustrated in Figure 4 (id:2012). For illustration purposes, we assume that the uncertainty in the estimation of the end of the T wave is dominant compared to that of the QRS onset.