# Deterministic Calibration Function Development

Georgi Tancev

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## Case Example: Introduction

- ▶ Sensor generates voltage due to reaction(s) with chemical compounds.
- ▶ Voltage is proportional to amount of reactants.
- ▶ Mapping sensor signal to gas reference concentration  $(f: S \rightarrow R)$ , but accounting for cross-sensitivities and other phenomena.

## Case Example: Conditions

- ▶ The aim is to gather prior information from published data.
- Experimental data from DeVito (2009): Measurement campaign took place at a main street in the center of an Italian city characterized by heavy car traffic.
- Raw sensor data of CO<sub>2</sub>, CO, NO<sub>x</sub>, NO<sub>2</sub>, O<sub>3</sub>, temperature, and absolute/relative humidity.
- ▶ Reference data of NO<sub>x</sub>, NO<sub>2</sub>, CO, C<sub>6</sub>H<sub>6</sub>.
- Each sensor/reference is stationary without replicates and a sampling rate of 1 h<sup>-1</sup> for a period of one year.
- Is it possible to map raw sensor data to reference, and if yes, how difficult is it?

## Case Example: Constraints

- Model inference should be feasible with respect to time and space complexity in case model and/or calibration are run locally.
- ▶ Backpropagation, which is used for neural network inference, has a time complexity of  $\mathcal{O}(n \cdot m \cdot h^k \cdot o \cdot i)$  with n training samples, m features, k hidden layers with h latent variables each, o outputs, and i iterations.
- ▶ Gradient descent has time complexity of  $\mathcal{O}(n \cdot m \cdot o \cdot i)$ , since a neural network without latent variables and with linear activation is equal to a multi-linear regression.
- Stochastic gradient descent has even lower time complexity due to smaller batch size n.
- ▶ In addition, space complexity has to be considered as some algorithms are memory-based and need to store all data.
- ▶ Ideally, model  $M_{t-1}$  from previous calibration time point t-1 is retrained by performing several iterations of (stochastic) gradient descent starting from parameters  $W_{t-1}$ . This ensures that the new model  $M_t$  with parameters  $W_t$  does not change too much from its parameter distribution p(W).

## Case Example: Workflow

- Selecting Scikit-Learn (and TensorFlow for comparison) as tool for model development.
- Starting with a more complex model and trying to predict reference concentrations from sensor voltages.
- 3. Tuning hyperparameters to obtain a good benchmark.
- 4. Pruning model by keeping loss approximately constant.
- 5. Inspecting model to understand its behavior.

## Methods: Preprocessing

- Drift is unknown, assuming first six months as free from drift and discarding the rest.
- Standard scaling to zero mean and unit variance.
- Dropping missing values instead of imputing.
- Removing outliers with isolation forest.
- Shuffling.
- Final dataset consists of N = 2'700 samples over six months.
- ► Splitting data in  $\frac{7}{10}N$  training and  $\frac{3}{10}N$  test set.

### Methods: Model Development

- Choosing a neural network (multi-layer perceptron) with rectifier activation function as model due to its capability to learn non-linearities.
- Performing a randomized grid search to optimize bias/variance by minimizing mean squared error via 5-fold cross-validation with respect to
  - L<sub>2</sub>-regularization parameter  $\alpha \in \{10^{-2+0.08j} \mid j \in \mathbb{N}, 0 \le j \le 50\}$ ,
  - ▶ number of hidden layers  $k \in \{1, 2, 3\}$  ,
  - ▶ and number of latent variables  $h \in \{5, 10, 15\}$ .
- This leads to a deterministic machine (maximum likelihood estimate), although
  - data is noisy (aleotoric uncertainty)
  - ▶ and sensors come from a population (epistemic uncertainty).
- Pruning model by removing layers and latent variables unless loss changes significantly from benchmark.

- Final model consists of 1 layer with 10 latent variables and regularization parameter  $\alpha=1.6$
- Amount of data seems reasonable.

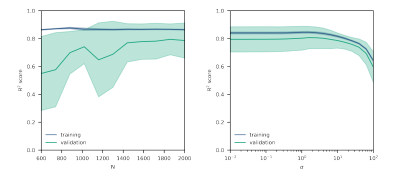


Figure: Training and validation curves for final model.

- ▶  $R^2$  score is 0.83,  $\beta$  is 0.81, and mean squared error is 0.10 averaged over all references on test data, which is good compared to literature.
- ► C<sub>6</sub>H<sub>6</sub> has no reference but shows best prediction performance.

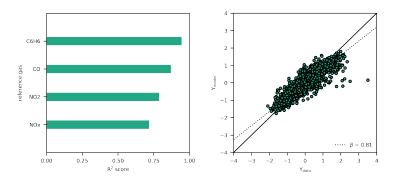


Figure: Model performance with respect to individual references.

- Only two sensors are important and rest is apparently redundant.
- The instruments of low importance could suffer from correlations since permutations can be compensated by correlated features.

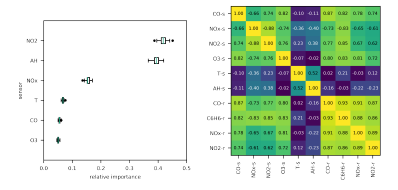


Figure: Importance of individual sensors and Spearman rank correlation of signals.

- One sensor detects and measures several gases in cases of highly correlated signals and/or references.
- Performing laboratory experiments with varying conditions helps decorrelating signals and removing sampling bias.

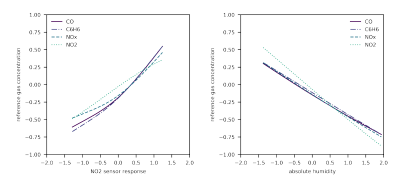


Figure: Partial dependence plots of references on sensors.

- Non-linearities are not very pronounced in this particular case.
- Non-linearities can also be modeled using parametric linear models by performing some feature engineering, e.g. basis expansion, or non-parametric random forests with deep trees.

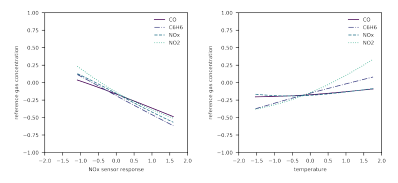


Figure: Partial dependence plots of references on sensors.

Underlying pollutant generation processes are not independent, and hence, chemical concentrations are conditional dependent.

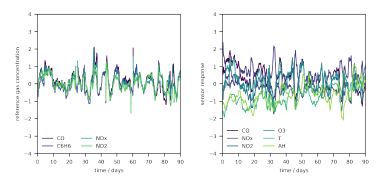


Figure: Time series of references and sensor responses.

- Comparison between Scikit-Learn and TensorFlow, since the latter is specialized on graphical models and neural networks.
- ▶ Same architecture with early stopping instead of regularization.
- ▶  $R^2$  score is 0.86,  $\beta$  is 0.86, and mean squared error is 0.09 averaged over all references on test data, which is consistent with previous result.

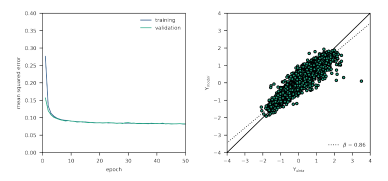


Figure: Model summary with loss and metrics.

- ▶ Comparison of different algorithms  $A = \{\text{neural network, random forest, linear regression}\}$  is mandatory.
- Performing basis expansion and hyperparameter optimization to control for bias and variance.
- Performance differences are not that significant compared to differences with respect to time and space complexity.

Table: Model performance on test set data of different algorithms.

metric	neural network	random forest	linear regression
mean squared error	0.09	0.09	0.10
R <sup>2</sup>	0.86	0.86	0.85
β	0.86	0.82	0.84

### Conclusion

- Inspecting model is a key aspect in machine learning.
- Examining data distribution and quality is of high importance.
- Uniform sampling under laboratory conditions with some design of experiments as complementary data acquisition method is unavoidable for a good model.
- Choice of model class is less likely to be relevant with respect to performance, which would be consistent with the ambiguity in the literature.

Probabilistic Calibration Function Development

## Probabilistic Modeling with TensorFlow Probability

- ▶ Modeling of aleotoric uncertainty (noise) by Monte-Carlo experiments.
- ▶ To obtain a prediction  $y_i$  for measurement  $x_i$ , k samples from posterior distribution  $p(y|x_i)$  are drawn and expectation  $y_i = \mu_{y|x_i}$  is calculated.
- Each prediction  $y_i$  has information about uncertainty due the standard deviation  $\sigma_{y|x_i}$  from the k draws.

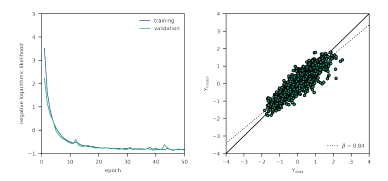


Figure: Model summary with loss and metrics.

## Probabilistic Modeling with TensorFlow Probability

- Case example with synthetic time series data from three sensors and three references.
- ▶ Prediction is an aggregated result of  $\mu_{y|x_i} \pm 1.96 \cdot \sigma_{y|x_i}$  (true value lies in this interval with a probability of 95%).

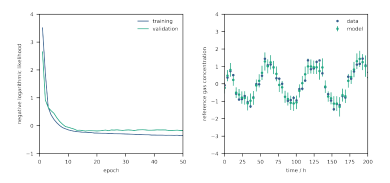


Figure: Training curve with sample prediction.