Predicting Superconducting Critical Temperatures with Uncertainty Quantification using Supervised Machine Learning

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Abstract will be added here.

I. INTRODUCTION

Superconductors are materials that lose all electrical resistance at low temperatures. These materials have a critical temperature (T_C) at which they lose their resistance. Most have very low critical temperatures, but "unconventional superconductors" can have critical temperatures as high as room temperature under non-atmospheric conditions.

Electrons in superconductors form Cooper Pairs below their critical temperature. These pairs of electrons are held together with phonouns, which are atomic-level collective excitations. Phonouns are similar to photons in that they also have particle-like properties [1].

Unconventional superconductors are still not well understood and remain an open question in Physics. Understanding them could lead to the discovery of superconducting materials stable at room temperature under atmospheric conditions. Such a material would have large implications, such as super efficient electricity transfer and vast efficiency improvements for applications like particle accelerators and power lines.

II. METHODOLOGY

II.1. Database

We chose to use one of the most popular experimental datasets, the supercon database from Japan's National Institute for Materials Science. This datasets contains 16,414 superconductor chemical compositions and their experimentally measured critical temperatures. Unfortunately, the database is not currently available on their website for unspecified reasons, so we obtained the dataset from a github repository that used this data [2].

In this dataset, there are 10,154 samples with a critical temperature below 10K and 6,210 samples above 10K. There are 159 samples with temperatures above 100K and 0 samples above 260K. A histogram represents this information graphically in Figure 1. Since most of this data is at low temperatures, we can expect less accuracy in our models for higher critical temperature superconductors.

Most superconductor databases do not include enough information to train an effective machine learning model, but such data can be extracted from the data they do provide. We use matminer to produce our features from

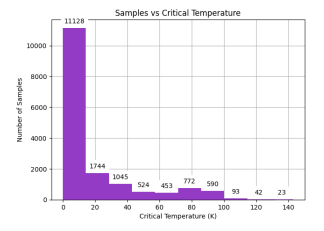


FIG. 1. A histogram of critical temperatures in the dataset. Most superconductors are found at low temperatures and this dataset is representative of that.

the provided material data. Matminer is a python library that generates data from various measured properties of a material [3]. Matminer collects existing calculations into a machine learning friendly python package.

Our database only provides the superconductor composition data. Matminer's featurizers can generate 53 features from the composition of a material. If we had band structure or other data, we could produce more information that we could use in our model.

II.2. Machine Learning

Previous papers have used random forest models to predict critical temperature [citation needed], but this paper will examine eight models before settling on two for further investigation. All models are implemented with Scikit-Learn, with the notable exception of a mlens superlearner [4, 5]. We will also use MAPIE models for uncertainty, discussed in Section II.5.

We decided to test a variety of machine learning models to ensure we would find an optimal model. These models are listed below:

- Linear Regression
- Support Vector Regression (SVR)
 - SVR makes predictions with decision boundaries, which are lines parallel to the regression line or curve. The model aims to maximize the amount of data within the decision boundaries

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and has hyperparameters to modify sensitivity to prevent overfitting. 1

• Elastic Net

Elastic Net uses L1 and L2 penalties to stabilize a linear model.

• Bayesian Ridge

 Bayesian Ridge uses probability distributors instead of point estimates for a linear model.

• Decision Tree

Decision trees are very interpretable - they break predictions into nodes of the tree, eventually leading to a prediction value. These trees can be represented graphically and show how they produce results, unlike most machine learning models.

• KNeighbors (KNN)

- KNN models are a little different, they store all the data and predict values based on a similarity measure. The model looks at a specified number of similar neighbors to produce a prediction.

• Random Forest Regression (RFR)

RFR is an ensemble method that uses numerous decision trees and subsamples the data with replacement. This means that the model replace data after using it in a subset.

• Extra Trees

- Extra Trees is like RFR, but it does not replace the data after use in a subset.

• Superlearner

 Our superlearner is a mlens model that can combine multiple high-scoring Scikit-Learn model predictions and sometimes improve the performance from the individual models.

II.3. Code Structure

The source code used for this paper is available publicly on github at https://github.com/sylphrena0/classe2022. This repository also includes the source files for this latex paper, data files, images, and documentation files.

Our research uses numpy and pandas throughout our code to handle arrays and tabular data [6, 7]. We also use matplotlib and seaborns to generate our graphs [8, 9].

The code is split into multiple python files so processes could be completed in stages and to maintain readability in the code. Most of our testing and final training was completed in juypter notebooks, but some computations were highly computationally expensive and needed to be run remotely. For these jobs, we created simple python files and made bash scripts to run them on Cornell's CLASSE compute farm. We also made several bash aliases and functions to simplify the compute farm workflow, which are also available on the github repository.

Since we used multiple files, we chose to create shared dependencies files where we defined functions to import data, train models, and generate our graphs. These files are then imported in all the relevant scripts to reduce redundancy. More detailed explainations of the purpose of each file is available in the github readme file and documentation within the files.

II.4. Code Evaluation

First, the featurizer script imports the dataset, extracts features from the material compositions, and exports the csv data. This script is one of the most computationally expensive and takes several hours to run on the CLASSE compute farm with 64 dedicated cores.

After the features are exported, our analysis jupyter notebook imports the data with the shared import function and exports histograms and a correlation matrix.

Next, the training_single jupy ter notebook or script can train individual models with the shared evaluation functions. This is used to get a landscape of initial performance before optimization. After training, the function plots the actual T_{C} versus the model prediction, using a heatmap to visualize the difference from the ideal prediction.

The optimizer script then uses a grid of manually defined hyperparameters to optimize models based on R2 score. This allowed significant improvements to baseline models. After optimization, the optimized models can be plotted in our single training notebook. After confirmation that the model is better than the baseline, the models can then be plotted together in a single graph using our bulk training notebook.

We evaluated our models using several metrics - R2 scores (R2) for regression evaluation, Mean Squared Error (MSE) and Mean Absolute Error (MAE) for error evaluation, and MAPIE Effective Mean Widths (MWS) for uncertainty evaluation.

II.5. Uncertainty

Our evaluation functions can produce uncertainty calculations using forestci, mapie, or lolopy [10–12].

Forestci is python implementation of an algorithm from [13] that predicts confidence intervals for random forest models. It is the fastest of the uncertainty methods listed.

Overfitting occurs when a model is trained to be too specific to a particular dataset and is not generalizable.

The Model Agnostic Prediction Interval Estimator (MAPIE) python library is more recent implementation of jackknife based on [14]. MAPIE uses various resampling methods. Most methods require the use of MAPIE's own MapieRegressor, which accepts an Scikit-Learn regressor and keeps track of uncertainty as the model is trained. MAPIE also has a prefit method, but it is difficult to extract uncertainty bars for individual points from this data - it splits a celebration set off the test set to generate uncertainty, so it can't be easily added to our plot of test set predictions. Thus, we will only compare the normal MAPIE methods with the other libraries. MAPIE trains much slower than other models, particularly on our superlearner, but it is still considerably faster than our final uncertainty model, lolopy.

Lolo is a Scala random forest machine learning library and is not a native python implementation. Lolopy is a python wrapper for lolo, but this implementation is very slow for large datasets.

III. RESULTS

III.1. Model Optimization

Each model's hyperparameters² were optimized with various optimization methods. Optimization can be computationally expensive, so we chose to optimize on a randomly selected subset of 2,000 materials, using a numpy random state for reproducibility. To start, we used Scikit-Learn's GridSearchCV, which tests combinations from a grid of hyperparameters and returns the best performing model based on a specified metric.

We also implemented Bayesian optimization using Gaussian Processes methods from the Scikit-Optimize library [15]. Bayesian optimization attempts to optimize models intelligently, instead of randomly testing specified hyperparameters. We provide a range of hyperparameters values for Bayesian optimization to test, and the algorithm uses acquisition functions to decide which specific values to use within the specified range. This is different from GridSearchCV, which is simpler but can take much longer to find optimial hyperparameters. We only implemented Bayesian optimization on our top models.

The performance of each optimizer on selected high performance models is shown in Table I. Note that the optimal method for each model (shown in bold), was found with a different method each time. The Bayesian optimization was much less computationally expensive than GridSearchCV, however, and is the recommended method for large datasets.

| Model | Optimizer | B2 | MSE | MAE | MWS |
|------------------|----------------------------|-------|---------|-------|--------|
| Widdei | Base Model | | | 5.791 | 53.135 |
| Random Forest | | | | 0 | |
| | GridSearchCv | | - | 5.816 | 52.416 |
| | Bayesian – PI | 0.815 | 134.509 | 5.72 | 50.985 |
| | Bayesian – EI | 0.814 | 135.267 | 5.763 | 51.533 |
| ₩ | Bayesian – gp_hedge | 0.815 | 134.785 | 5.747 | 51.021 |
| Extra Trees | Base Model | 0.818 | 132.395 | 5.214 | 48.153 |
| | GridSearchCv | 0.818 | 132.374 | 5.215 | 48.231 |
| | Bayesian – PI | 0.818 | 132.104 | 5.225 | 48.154 |
| 鱼色 | Bayesian – EI | 0.816 | 133.5 | 5.257 | 48.757 |
| | ${\bf Bayesian-gp_hedge}$ | 0.819 | 131.783 | 5.202 | 47.732 |
| KNN | Base Model | 0.646 | 257.108 | 8.563 | 75.612 |
| | $\mathbf{GridSearchCv}$ | 0.703 | 216.186 | 7.515 | 69.758 |
| | Bayesian – PI | 0.652 | 253.137 | 8.201 | 74.177 |
| Ξ | Bayesian – EI | 0.652 | 253.137 | 8.201 | 74.177 |
| | Bayesian – gp_hedge | 0.566 | 315.238 | 8.159 | 84.546 |

TABLE I. Comparison of optimization methods by model.

III.2. Base Models

Our base models provided good results and the top two models, Random Forest and Extra Trees, only improved their scores marginally with optimization. Our numerical results of the optimized and optimized models are shown in Table II.

| | Model | R2 | MSE | MAE | MWS |
|-------------|--------------------------|-------|---------|--------|---------|
| Unoptimized | Extra Trees Regression | 0.818 | 132.395 | 5.214 | 48.153 |
| | Random Forest Regression | 0.807 | 140.482 | 5.791 | 53.135 |
| | KNeighbors Regression | 0.646 | 257.108 | 8.563 | 75.612 |
| | Decision Tree Regression | 0.644 | 258.945 | 6.886 | 76.715 |
| | Bayesian Regression | 0.392 | 441.925 | 14.52 | 90.7 |
| | Linear Regression | 0.392 | 442.163 | 14.506 | 90.705 |
| | Elastic Net Regression | 0.328 | 488.242 | 15.603 | 97.938 |
| | Support Vector Machines | 0.084 | 666.16 | 15.511 | 134.628 |
| Optimized | Extra Trees Regression | 0.819 | 131.624 | 5.205 | 48.088 |
| | Random Forest Regression | 0.816 | 133.87 | 5.714 | 50.759 |
| | KNeighbors Regression | 0.703 | 216.186 | 7.515 | 69.758 |
| | Decision Tree Regression | 0.664 | 244.095 | 7.152 | 74.834 |
| | Elastic Net Regression | 0.392 | 442.133 | 14.487 | 90.713 |
| | Bayesian Regression | 0.392 | 441.925 | 14.52 | 90.7 |
| | Linear Regression | 0.392 | 442.163 | 14.506 | 90.705 |
| | Support Vector Machines | 0.325 | 490.661 | 14.186 | 106.946 |

TABLE II. CV scores of the models using all features and data, comparing optimized and base hyperparameters.

III.3. Cleaned Models

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III.4. Error Comparison

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III.5. Feature Importance

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 $^{^2}$ Hyperparameters are machine learning parameters that change how a model is trained.

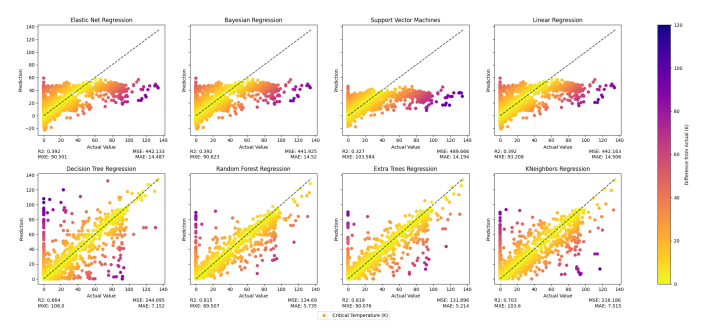


FIG. 2. Graph of optimized model predictions versus actual critical temperatures for each base model.

IV. CONCLUSION

IV.1. Future Work

ACKNOWLEDGMENTS

I would like to thank my mentor, Suchismita Sarker, for her guidance and support with this project. This work is supported by the U.S. National Science Foundation under award number NSF PHY-2150125, REU Site: Accelerator Physics and Synchrotron Radiation Science.

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