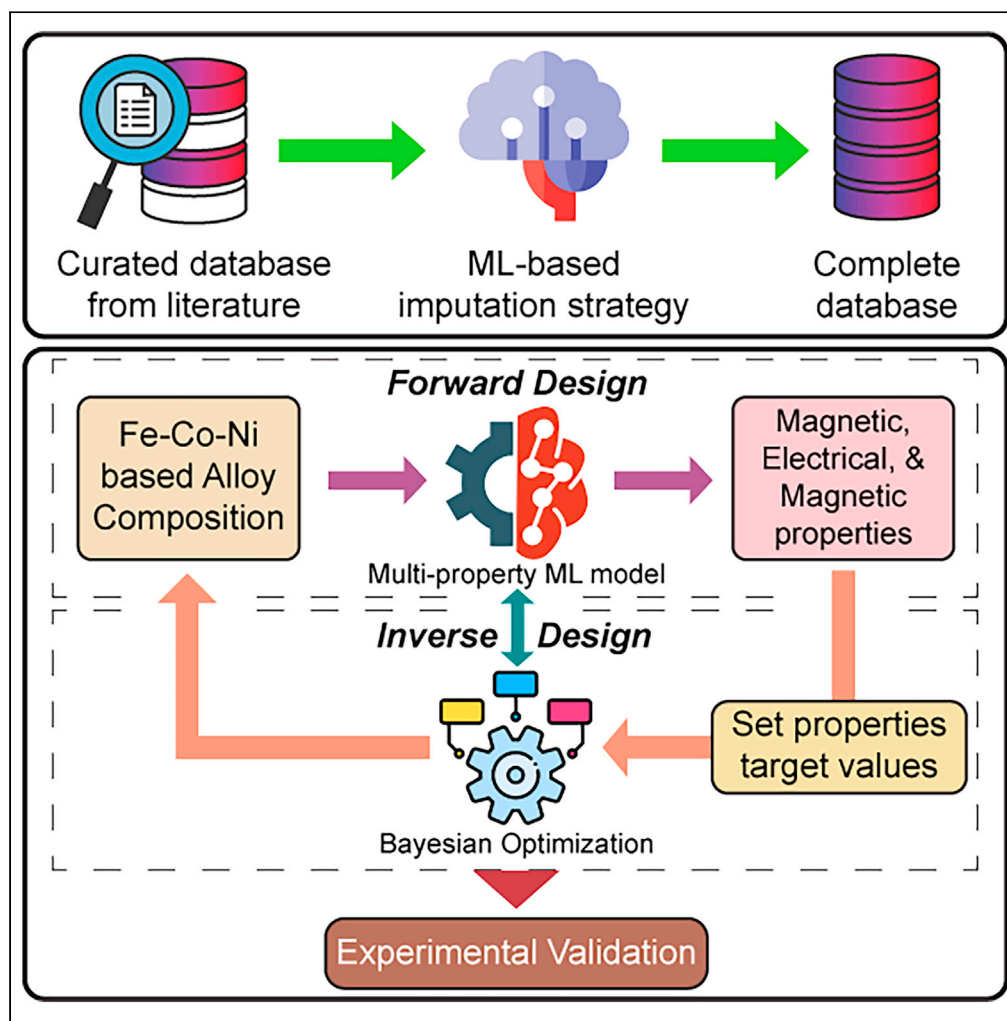


Article

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Highlights

A machine learning framework for multi-property optimized Fe-Co-Ni alloy design

Predictive models and multi-objective Bayesian optimization predict promising alloys

Experimental validation confirms superior multi-property performance of the alloys

$\text{Fe}_{66.8}\text{Co}_{28}\text{Ni}_{5.2}$ and $\text{Fe}_{61.9}\text{Co}_{22.8}\text{Ni}_{15.3}$ are the identified alloy compositions

Padhy et al., iScience 27,
109723
May 17, 2024 © 2024 The
Authors. Published by Elsevier
Inc.
[https://doi.org/10.1016/
j.isci.2024.109723](https://doi.org/10.1016/j.isci.2024.109723)

Article

Experimentally validated inverse design of multi-property Fe-Co-Ni alloys

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SUMMARY

This study presents a machine learning (ML) framework aimed at accelerating the discovery of multi-property optimized Fe-Ni-Co alloys, addressing the time-consuming, expensive, and inefficient nature of traditional methods of material discovery, development, and deployment. We compiled a detailed heterogeneous database of the magnetic, electrical, and mechanical properties of Fe-Co-Ni alloys, employing a novel ML-based imputation strategy to address gaps in property data. Leveraging this comprehensive database, we developed predictive ML models using tree-based and neural network approaches for optimizing multiple properties simultaneously. An inverse design strategy, utilizing multi-objective Bayesian optimization (MOBO), enabled the identification of promising alloy compositions. This approach was experimentally validated using high-throughput methodology, highlighting alloys such as Fe_{66.8}Co₂₈Ni_{5.2} and Fe_{61.9}Co_{22.8}Ni_{15.3}, which demonstrated superior properties. The predicted properties data closely matched experimental data within 14% accuracy. Our approach can be extended to a broad range of materials systems to predict novel materials with an optimized set of properties.

INTRODUCTION

Material innovation has often been the key to advances in technology. Multi-component alloys, polymers, composites, semiconductors, superconductors, and biocompatible materials have promoted a revolution in construction, transportation, energy, bioengineering, microelectronics, and space technologies.^{1,2} Research and development in the field of materials science can be divided into four major paradigms: empirical trial and error methods, theoretical models and laws, computational simulations, and data-driven science.^{3–6} The first three paradigms are experiment, theory, and computation and constitute the traditional methodology of discovering and developing materials. From research to applications, traditional methods typically take decades, require high expenditures, are highly inefficient, and usually do not result in the best composition for a given application.^{5,7–9} On the other hand, data-driven science using machine learning (ML) of mined material data integrated with high-throughput experiments and computing can be used to design and discover new materials, and a vast space of materials and processing conditions can be studied with minimum time and cost.^{6,7,9–21}

One such important class of materials are alloys used in essential components of electro-magnetic energy conversion devices, for e.g., electrical motors, generators, transformers, inductors, actuators, magnetic-MEMS, and magnetic recording media.^{2,22–25} Due to increased global electricity consumption, e.g., an increase of 1.7% from 2018 to 2019²⁶ and a projected growth rate of 3–4% per annum from 2019 to 2050,²⁷ there is an urgent need for developing energy-efficient electrical technologies. Next-generation superior alloys hold the key to developing high efficiency, low cost, and compact devices that are also suitable to operate in extreme service conditions, for e.g., high frequency rotating electrical machines.^{23,28–31} Additionally, an improvement of only 1% in the efficiency of electric motors will reduce global power consumption by 94.5 TW-hours and carbon dioxide emissions by 60 million metric tons.^{32,33}

However, there is no single commercial magnetic alloy that is commercially used with optimized multiple properties for current technological applications. Recently, a novel alloy (Fe_{32.6}Ni_{27.7}Co_{27.7}Ta₅Al₇) was reported with a high tensile strength of 1336 MPa and a low coercivity of 78 A/m.³⁴ This combination of properties was achieved by developing precipitates, enriched in nickel, tantalum, and aluminum with a controlled size and by tuning the particle/matrix interfacial coherency.^{34,35} The precipitates developed in these alloys impeded dislocation motion, thereby improving strength, and did not significantly increase the pinning effect of the precipitates on the domain wall movement, maintaining low coercivity. This phenomenon was achievable due to the use of various constituent elements in the alloy system, which

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<https://doi.org/10.1016/j.isci.2024.109723>



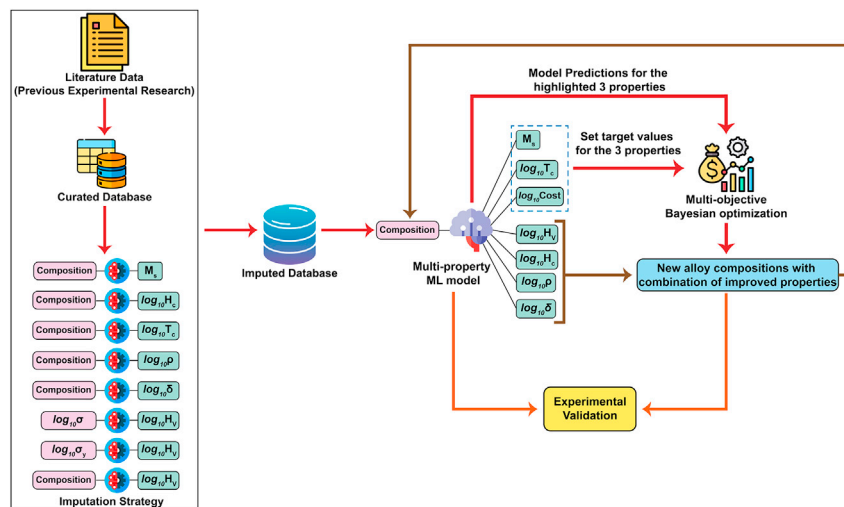


Figure 1. Overview of the work

Database Curation, Machine Learning (ML) based imputation strategy used to complete the database, multi-property ML regression model [Forward Design of alloys], Multi-objective Bayesian optimization (MOBO) [Inverse Design of alloys], and the Experimental Validation of forward design and predicted new alloy composition from inverse design.

facilitates a balance of multiple properties.³⁵ However, the alloy exhibited lower saturation magnetization than those of the best soft magnetic materials because the precipitates formed were paramagnetic in nature.³⁵ Therefore, the right choice of the principal constituent elements and their content in a multi-component alloy system is essential in developing next-generation magnetic alloys.

In this work, Fe-Co-Ni-based alloy compositions were chosen as the three elements are ferromagnetic in nature and certain alloy compositions can potentially provide a better combination of properties. Binary alloys of these ferromagnetic elements exhibit certain excellent properties. For e.g., Ni₇₈Fe₂₂ (78-Permalloy) and Ni₇₈Fe₁₇Mo₅ (Supermalloy) exhibit high permeability (μ) and low coercivity (H_c). Co exhibits the highest Curie temperature (T_c) of 1394 K, while Fe₆₅Co₃₅ exhibits the highest saturation magnetization (M_s) of 2.47 T followed by Fe₄₉Co₄₉V₂. On the other hand, Fe-Si alloys with 2–4.5 wt % Si exhibit low H_c , low magnetocrystalline anisotropy and high electrical resistivity (ρ).^{2,22,23,36} The desired set of technical properties required for next-generation alloys for rotating electric machines include magnetic, electrical, and mechanical properties, e.g., high M_s , low H_c , high T_c , high μ , high ρ , high tensile (σ) and yield (σ_y) strength, high ductility (δ), and high hardness (H_v). Thus, multi-property optimization and new alloy compositions are required. Machine learning (ML) can reveal novel alloy compositions with optimized properties.

ML has been applied to alloy design mainly focusing on a single property,^{13,14,37–40} and a few studies have been performed to optimize multiple properties.^{8,41–47} In the present study, we collect multiple property data on Fe-Co-Ni alloys from the literature to create a curated database. We combined ML with Bayesian optimization (BO) to identify Fe-Co-Ni alloys with the desired multiple property set. The curated database contained gaps in the property data that were filled using an ML-based imputation technique. This imputed database was used to develop a multi-property ML model (forward design). Multi-objective BO (inverse design) was implemented to identify alloys with superior property sets. Both the forward and inverse design results were validated using experimental data obtained using a high-throughput methodology. The compositions, Fe_{65.84}Co_{28.66}Ni_{5.5} and Fe_{61.5}Co_{23.14}Ni_{15.35}, revealed by BO, demonstrate high values of M_s , T_c , and H_v , moderately low values of H_c and moderately high values of ρ , a superior property set compared to conventional Fe-Co-Ni magnetic alloys. The strategy is shown in Figure 1. The proposed approach is applicable to a wide variety of multi-property optimization materials design problems.

METHODOLOGY

The Python programming language was used for exploratory data analysis, model building, and optimization. The Pandas^{48–50} library was used to handle data. Matplotlib,⁵¹ seaborn,⁵² and plotly⁵³ libraries were used for plotting. Keras⁵⁴ from TensorFlow⁵⁵ library was used to implement Deep Neural Network (DNN) models, XGBoost⁵⁶ library was used to implement XGBoost Regression (XGBR), and Scikit-Learn⁵⁷ library was used for the rest of the models. For BO, gp_minimize from the Scikit-Optimize⁵⁸ library was implemented.

The Fe-Co-Ni-based alloy database along with the codes for database analysis, the eight imputation steps, multi-property models, multi-objective Bayesian optimization (MOBO), and plotting of experimental validation results is publicly available.⁵⁹

Database Curation and analysis

A database of Fe-Co-Ni-based alloys (with the optional addition of other elements up to a combined at % less than 10) and their corresponding magnetic (M_s , H_c , T_c), electrical (ρ), and mechanical (σ , σ_y , δ , Vickers Hardness (H_v)) properties were collected from the literature. From thousands of publications and hundreds of handbooks describing Fe-Co-Ni-based alloy composition(s) and their properties, data was

extracted from selected sources based on the criterion that two or more properties for the alloy composition were experimentally determined. For research publications, Clarivate Web of Science was used to search for articles with various search strings for the year range from 1900 to 2021. Further, relevant publications were selected for each search string by first screening various abstracts and then screening the publication for information related to Fe-Co-Ni-based alloys. Sources used for database building include:

- (1) Books – *Ferromagnetism*,⁶⁰ *Magnetism and Magnetic Materials*,⁶¹ *Bibliography of Magnetic Materials and Tabulation of Magnetic Transition Temperature*⁶²;
- (2) Handbooks – *Powder Metallurgy Data Part 1: Metals and Magnets*,⁶³ *ASM Handbook Volume 2*,⁶⁴ *Handbook of Materials Selection*^{65,66};
- (3) Research/Review articles^{67–80,81–112}.

A total of 1208 data entries were curated from the above-mentioned references from text, tables, and graphical images. The information from graphical images were extracted using graph digitizing software such as Graph Grabber (v2.0.2, Quintessa Ltd., UK), Engauge Digitizer,¹¹³ and Digitizer Tool of OriginPro 2020b (OriginLab Corporation, USA). The content of elements in the alloy composition was converted to atomic percentage (at. %). If the reported composition was in weight percentage (wt. %), it was converted to at.% before adding to the database. The properties were converted to SI units, i.e., M_s in Tesla (T), H_c in Ampere/meter (A/m), T_c in Kelvin (K), ρ in Micro ohm centimeter ($\mu\Omega \cdot \text{cm}$), σ and σ_y in Megapascals (MPa), δ in %, and H_V in HV. Using the elemental content information and the individual element price,¹¹⁴ overall raw material cost (US\$/kg) for each alloy was calculated and added to the database. For database analysis, the distribution of the elements and properties was plotted. The Pearson correlation coefficients for each property versus the alloy composition as well as between the properties were calculated and plotted. Finally, the percentage of the property data filled in the database was calculated for each property.

Imputation strategy

An imputation strategy was developed to fill the empty cells in the properties columns so that the imputed database can be used to build a multi-property model mapping the composition to the properties. 12 different regression models were built, mapping the composition to each of the responses, by training on 80% of the available property data. These are Linear Regression (Linear), Ridge Regression (Ridge), Lasso Regression (Lasso), K-Nearest Neighbors Regression (KNNR), Support Vector Regression (SVR), Decision Tree Regression (DTR), Gradient Boosting Regression (GBR), Random Forest Regression (RFR), Extra Trees Regression (ETR), XGBoost Regression (XGBR), Deep Neural Network Regression with same number of units in all the hidden layers (DNNR_rectangle), and Deep Neural Network Regression with different number of units in the hidden layers (DNNR_random). The details of the hyperparameters optimization and model building are discussed in [Section S2](#) of supplementary information. After all the models were built, the best model with the lowest mean-absolute error (MAE) for the testing data was selected for the imputation of that respective property column.¹¹⁵ Those models also exhibited low root mean-squared error (RMSE) and high coefficient of determination (R^2).

For all the properties except $\log_{10}H_V$, imputation was performed directly using the best model mapping from composition to the property as explained above. For $\log_{10}H_V$, imputation was done in 3 steps.

- (1) 12 regression models were built mapping from $\log_{10}\sigma$ to $\log_{10}H_V$. The model with the lowest MAE was chosen to fill up the empty cells in the $\log_{10}H_V$ column for which σ values were reported in the literature but not the H_V values.
- (2) 12 regression models were built mapping from $\log_{10}\sigma_y$ to $\log_{10}H_V$, the model with the lowest MAE was chosen to impute the $\log_{10}H_V$ values for which σ_y were reported but not H_V .
- (3) Using both the experimental and imputed $\log_{10}H_V$ values, all the 12 regression models were built mapping from the composition to $\log_{10}H_V$, the model with the lowest MAE was chosen for the imputation of the $\log_{10}H_V$ column.

Multi-property model for the forward design of the multi-property alloy

The imputed database was next used to build multi-property predictive models mapping the composition to M_s , $\log_{10}H_c$, $\log_{10}T_c$, $\log_{10}\rho$, $\log_{10}\delta$, $\log_{10}H_V$, and $\log_{10}\text{Cost}$. Six ML models, i.e., RFR, ETR, neural network regression with 3 hidden layers (NNR_3layers), neural network regression with 4 hidden layers (NNR_4layers), DNNR_rectangle, and DNNR_random were deployed. The hyperparameters for each of the models are discussed in [Section S3](#) of supplementary information. Further, using the same hyperparameters, another set of six multi-property predictive models was built mapping the composition and Wen alloy features³⁸ to the 7 properties. Further, the MAE of all the different models were compared with each other to select one tree-based model and one neural network model with the lowest MAE as multi-property predictive models. These models were experimentally validated using 40 Fe-Co-Ni compositions which were synthesized and characterized in a high-throughput manner.

Multi-objective Bayesian optimization for inverse design of the multi-property alloy

To design alloy compositions possessing an optimum balance of properties using the multi-property model, the implementation of a robust and computationally low-cost optimization algorithm is required. In this work, Bayesian optimization (BO) is used for fast, low cost and effective global optimization of those objective functions which are expensive and time-consuming to evaluate, i.e., expensive black box functions.^{116–119}

A Gaussian Process (GP) based BO was used to obtain the multi-property regression (MPR) model predictions closer to the target values. The global minima/maxima composition was determined from the defined search space of elements. BO uses prior and evidence knowledge to define a posterior distribution over function space. The function space is maximized/minimized at each step such that the expected maximum/minimum of the given model moves closer to the global maximum/minimum.¹²⁰ In this work, *gp_minimize* from *Scikit-Optimize* was used for BO which minimizes the defined objective function. The acquisition function set in *gp_minimize* was “EI,” which denotes a negative expected improvement. The expected improvement determines the global maximum of the objective function, which is explained in [Section S5](#) of supplementary information. Thus, a negative expected improvement finds the global minima of the objective function.

The objective function defined for the BO to design alloy composition containing only Fe, Co, and Ni is described by the equations later in discussion,

$$x_n = \frac{x}{\sum x} \times 100 \quad (\text{Equation 1})$$

$$x_{\text{transformed}} = \text{StandardScaler}(x_n) \quad (\text{Equation 2})$$

$$\text{pred} = \text{MPR}(x_{\text{transformed}}) \quad (\text{Equation 3})$$

$$\text{objective} = \sum w_i \times \text{abs}(\text{target}_i - \text{pred}_i), \text{ if } \text{target}_i \neq \text{Nan} \quad (\text{Equation 4})$$

where x in [Equation 1](#) is the compositional array containing at.% values of Fe, Co, Ni, and other elements determined by the BO algorithm from the defined ranges: Fe, Co, Ni $\in [0, 100]$ and other elements $\in [0, 0.001]$, x_n in [Equation 1](#) is the normalized compositional array to always maintain the sum of at.% of elements at 100%, *StandardScaler* in [Equation 2](#) is the standard scaling function used on the training set for building the MPR model, *pred* in [Equation 3](#) is the array of predicted property values by the MPR model, w_i in [Equation 4](#) are the weights for optimizing each property which were defined manually, *Nan* in [Equation 4](#) stands for Not a number, and target_i and pred_i in [Equation 4](#) are the set target value and MPR model predicted value of each property. The idea of using weights is to emphasize the optimization of the desired property set close to the desired target value compared to other property set. This would be useful for discovering the optimum alloy composition for a particular application.

Experimental validation of forward and inverse design

40 Fe-Co-Ni alloy compositions were prepared using high-throughput experimental scheme using high-throughput spark plasma sintering (SPS) of ball-milled alloy powders which is explained elsewhere.^{112,121} The alloy powders were prepared via ball milling (Fritsch Pulverisette-7) of Fe, Co, and Ni powders (purity $\geq 99.95\%$, Tosoh SMD Inc.) using tungsten carbide vials and balls. 10 mm diameter balls were used and the ball-to-powder ratio was maintained at 12:1. The bulk alloy samples were prepared via SPS (Fuji Electronic Industrial, SPS-211LX) in compositionally graded cylindrical samples by stacking ball-milled powders and sintering them for 15 min at 950°C, 40 MPa, at vacuum level less than 8 Pa. The composition was measured using an energy-dispersive X-ray spectrometer attached to the JEOL JSM-7800F-SEM. The magnetic properties were measured using a physical property measurement system (EverCool-II, QD). The Curie temperature was measured in an accelerated manner using a modified thermogravimetric analysis (TGA) by placing a magnet near the TGA pan.^{109,112,121,122} The Vickers hardness and electrical resistivity were measured using a Vickers hardness tester (Future-Tech) and a four-point probe (4PP) tester (Keithlink), respectively. The Cost for these compositions was calculated, as described in [Section database curation and analysis](#).

For the experimental validation of the forward design methodology, the 40 Fe-Co-Ni compositions were used in the selected tree-based model and neural network model to predict the property values. These predicted values were compared with the experimental values of the properties and the MAE for all the properties for both models were calculated.

For the experimental validation of the inverse design methodology, the predicted alloy compositions by BO for the set target values of the properties were plotted and compared using interactive ternary contour plots of the properties of 40 rapidly synthesized Fe-Co-Ni compositions. For the predicted compositions, the properties were predicted using the compositions in the model which was used in BO.

RESULTS AND DISCUSSION

Database curation and analysis

15 elements were present in the database, the distribution is shown in [Figure 2A–2N](#). The elements are Fe, Co, Ni, V, Mo, Cr, Cu, Mn, C, W, Ta, Nb, Al, Ti, and Si. According to the distribution of Fe shown in [Figure 2A](#), there are more than 250 alloys containing Fe between 46.33 at.% and 52.67 at.%. From the distribution of Co shown in [Figure 2B](#), a similar number can be inferred. The compositions containing Ni are broadly distributed as shown in [Figure 2C](#). From the distribution of V and Cu in [Figures 2D and 2G](#) the alloy composition containing V or Cu is usually ~ 2 at.%. The 15 elements were considered as features for the multi-property ML model, as described in [Table 1](#).

The distribution of properties is discussed in [Section S1](#) of supplementary information. The values of all the properties except for M_s were converted to \log_{10} scale to reduce the skewness in the data and to make different property values comparable to each other. The distribution of the modified properties and cost is shown in [Figures 2O–2W](#). The skewness of the distribution of the \log_{10} scale converted properties decreased compared to the distribution in [Figure S1](#), except for T_c and Cost.

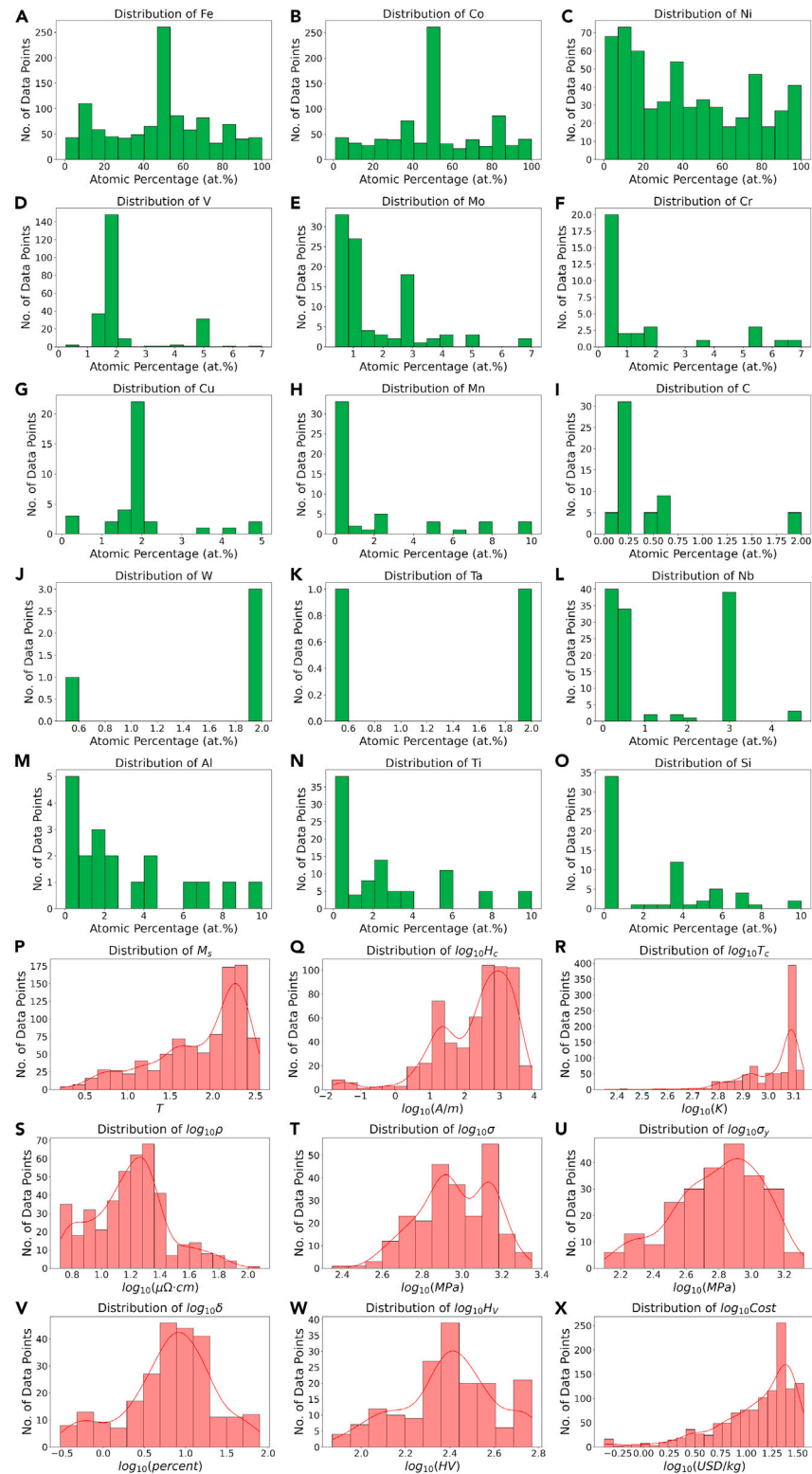


Figure 2. Distribution analysis of the database

Distribution of (A–O) 15 elements atomic percentage (at.%), (O–Q) magnetic properties (M_s , $\log_{10}H_c$, $\log_{10}T_c$), (R) electrical resistivity ($\log_{10}\rho$), (S–V) mechanical properties ($\log_{10}\sigma$, $\log_{10}\sigma_y$, $\log_{10}\delta$, $\log_{10}HV$), and (W) $\log_{10}Cost$ in the database. In (P–X) the solid continuous lines are kernel density estimations which show the probability density function of the data entries for each property.

Table 1. Table of features and responses used for multi-property machine learning (ML) model

Features	Fe, Co, Ni, V, Mo, Cr, Cu, Mn, C, W, Ta, Nb, Al, Ti, Si
Responses	M_s , $\log_{10}(H_c)$, $\log_{10}(T_c)$, $\log_{10}(\rho)$, $\log_{10}(\delta)$, $\log_{10}(H_V)$, and $\log_{10}(\text{Cost})$

To understand the relationship between the properties and alloy composition, the Pearson correlation coefficient was calculated for each property against all the 15 elements, as shown in Figure 3A. The Pearson correlation coefficient was also calculated between the properties to understand the inter-property relationships as shown in Figure 3B. From Figure 3A, it can be observed that the Fe content has a positive correlation with M_s while the Ni content has a negative correlation, implying that the addition of Fe will increase M_s whereas the addition of Ni will decrease M_s .¹²³ For $\log_{10}H_c$, the Co content has a moderately positive correlation whereas the Ni content has a moderately negative correlation, which implies that the addition of Co increases $\log_{10}H_c$ while the addition of Ni decreases $\log_{10}H_c$. To support this inference, there have been studies in the past reporting that permalloy and moly permalloy exhibit lower coercivity values compared to Fe-Co alloys.^{23,106,109,112}

The Co content has a positive correlation with $\log_{10}T_c$ while the Ni content has a negative correlation, implying that the addition of Co increases $\log_{10}T_c$, on the other hand, the addition of Ni decreases $\log_{10}T_c$. This is consistent with the fact that Co has the highest T_c of 1394 K,^{60,124} Ni has a much lower T_c of 631 K.¹²⁵ This change in T_c values can also be rationalized by the mean field model and the Bethe-Slater curve.^{18,126} $\log_{10}H_c$ has a positive linear correlation with $\log_{10}T_c$, $\log_{10}\sigma$, and $\log_{10}\sigma_y$ (Figure 3B). The positive linear correlation between $\log_{10}H_c$ and $\log_{10}T_c$ can be attributed to an increase in Co content (Figure 3A).

Moreover, interestingly, Fe has a moderately negative correlation with $\log_{10}\delta$ implying, with all other factors unchanged, δ decreases with an increase in Fe content. Further, V and C have a moderately positive correlation with $\log_{10}H_V$, which implies that the addition of V and C increases H_V . From Figure 3B, it is evident that $\log_{10}\sigma$, $\log_{10}\sigma_y$, and $\log_{10}H_V$ have a strong positive linear correlation with each other. Therefore, $\log_{10}H_V$ was utilized as the main mechanical property prediction and experimental validation in this study.

Thus, seven properties – M_s , $\log_{10}H_c$, $\log_{10}T_c$, $\log_{10}\rho$, $\log_{10}\delta$, $\log_{10}H_V$, and $\log_{10}\text{Cost}$ – were chosen as the responses for the multi-property ML model, as listed in Table 1. Before building the multi-property model, the experimental data available in the database for the responses was analyzed, as shown in Figure 3C. The Cost was calculated from the alloy composition; hence, the $\log_{10}\text{Cost}$ column is 100% filled in the database. M_s and $\log_{10}T_c$ have the most experimental data, i.e., 75% and 72% of the data entries, respectively. In the case of T_c , the data for the binary alloy compositions was obtained from the phase diagram, the data for commercial alloy compositions such as Hiperco was obtained from the reported sources.

$\log_{10}H_V$ has the least experimental data, which is 14.5% of the total data entries, which shows that only a limited number of mechanical property studies have been performed on Fe-Co-Ni-based magnetic alloys. Figure 3C shows that only a small number of rows in the database are complete, i.e., 0.66% of the data entries, which corresponds to just 8 rows. Therefore, an ML-based imputation strategy was developed to fill the empty cells of the responses column to build a multi-property model, which is discussed in Section [imputation strategy](#).

Imputation strategy

Figure 4 shows the heatmap of MAE, RMSE, and R^2 values of the all the regression models built to find the best model for the imputation of the property. The best regression models used for stepwise imputation are shown in Table 2 along with their MAE, RMSE, and R^2 values.

Forward design of the multi-property alloy

Multi-property models

Figure 5A–5C shows the heatmap of the performance metrics (MAE, RMSE, and R^2) of the six models with composition as the input features to the six models. Among them, the ETR model was found to be the best predictive model with the lowest MAE and high R^2 for all the properties. The ETR model can predict a novel composition within the limits of each property value in the database which achieves an optimum balance of properties.^{43,127,128} The performance of the ETR model in predicting M_s , $\log_{10}T_c$, and $\log_{10}\text{Cost}$ is better and more accurate, the model exhibits $R^2 > 0.95$ and MAE < 0.05 . This can be attributed to the strong compositional dependence of M_s and T_c . As Cost is calculated using composition and individual element cost, it explains the good performance of the ETR model. The performance of the ETR model for predicting $\log_{10}\delta$ is moderately good and accurate with a good R^2 , in the range of 0.85–0.9, and low MAE. For $\log_{10}H_c$, the performance of the ETR model is less accurate with good R^2 , in the range of 0.85–0.9, and a high MAE. For $\log_{10}\rho$ and $\log_{10}H_V$, the performance of the ETR model is moderately accurate and precise with decent R^2 , in the range of 0.75–0.8, and low MAE. H_c , ρ , and H_V are more process-dependent properties compared to composition, which explains this moderate performance of the ETR model.

Among the NN architectures, NNR_4layers performed the best with lowest MAE for all the properties and highest R^2 for M_s , $\log_{10}\delta$, and $\log_{10}\text{Cost}$. However, the model exhibited the second lowest R^2 values for $\log_{10}H_c$, $\log_{10}T_c$, and $\log_{10}\rho$ and the second highest R^2 value for $\log_{10}H_V$. Wen et al. designed a high entropy alloys with hardness higher than the training database by using 2 iterations of ML combined with experiments in which they implemented the ML model with only compositional input features in the first loop and the ML model with compositional and physical features in the second loop.³⁸ They reported that designing alloys with both compositions and physical features was more effective than designing alloys with only compositional features. Hence, WenAlloys featurizer class in Matminer¹²⁹ package was used to evaluate the physical features for all the data in the imputed database and built all six models with the same hyperparameters to

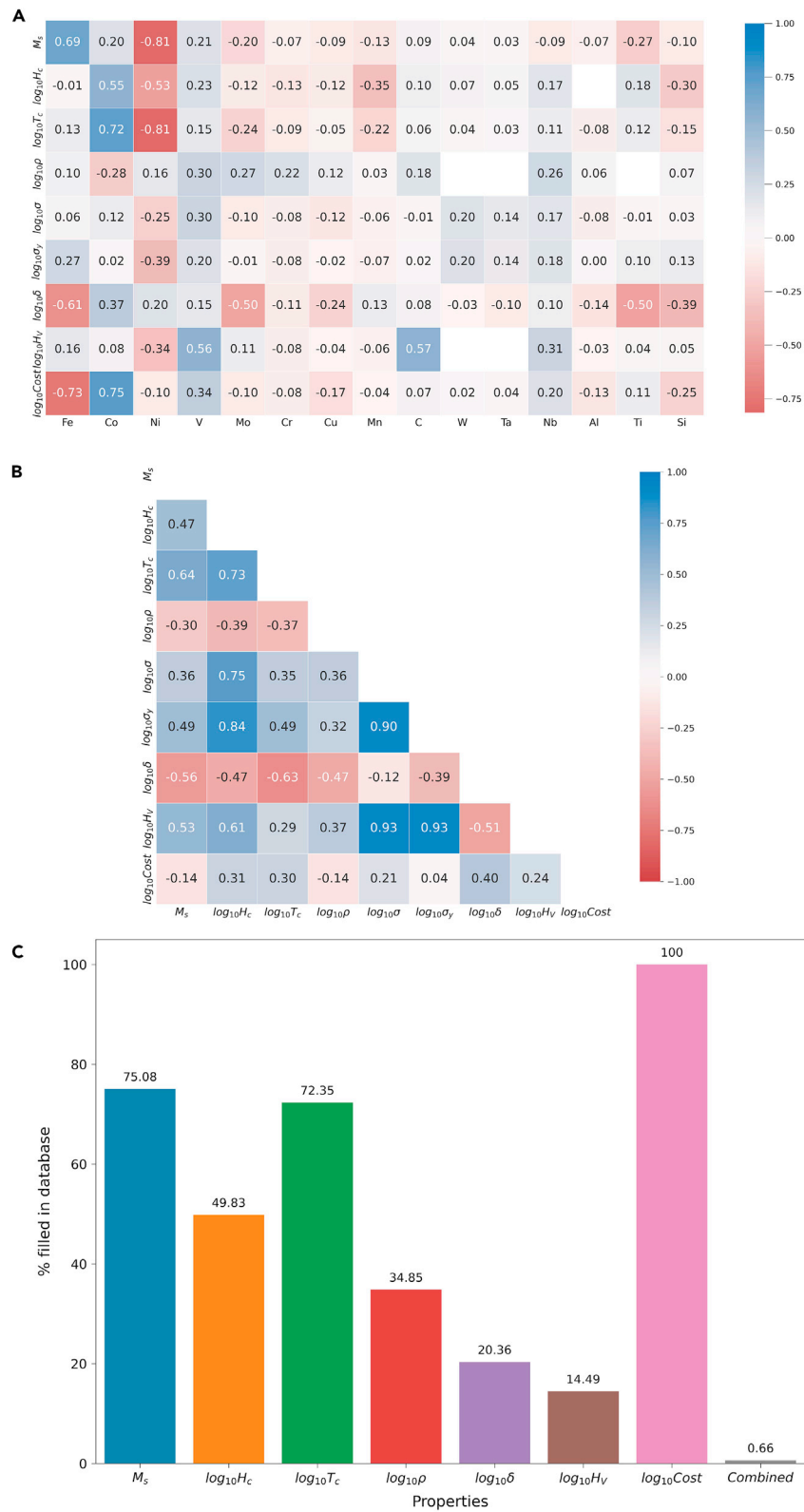


Figure 3. Exploratory data analysis

Pearson correlation coefficients heatmap of (A) properties vs. elements and (B) properties vs. properties. (C) Percentage of properties data available for the alloy compositions in the database.

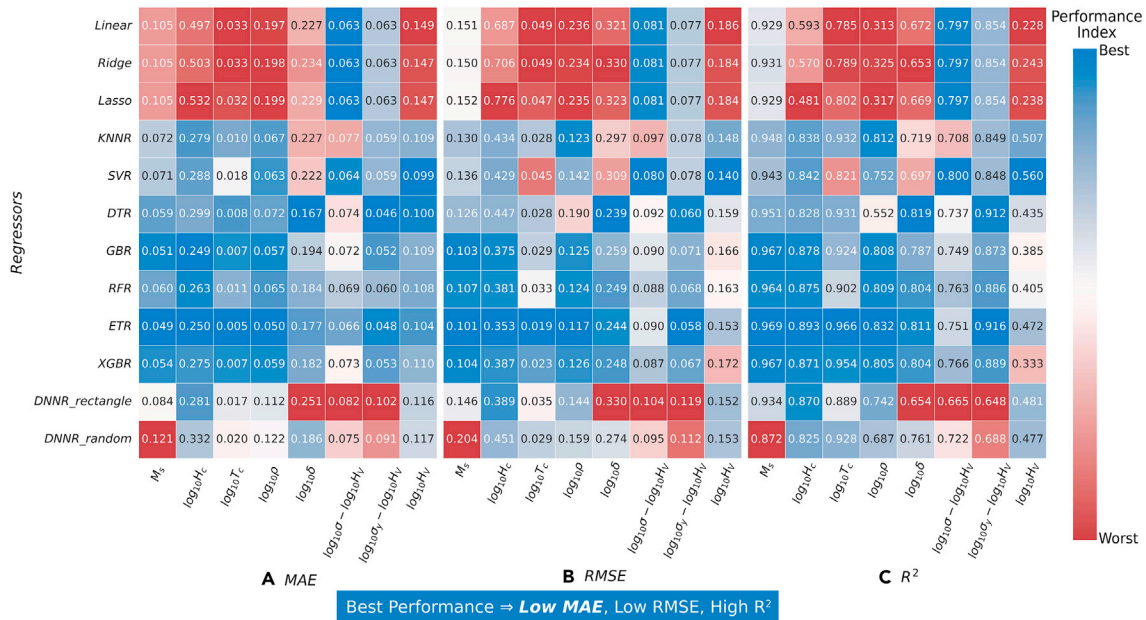


Figure 4. Heatmap of performance metrics

(A) mean absolute error (MAE), (B) root mean squared error (RMSE), and (C) coefficient of determination (R^2) of various ML regressor models for the imputation of each property. The lowest MAE model is chosen for the imputation of each property which is indicated with the darkest shade of blue in (A) for each column.

compare their performance with the models with only compositional features as input. Figures 5D–5F shows the heatmap of the performance metrics (MAE, RMSE, and R^2) of the six models with composition and physical descriptors as the input features for the six models.

It can be observed that the performance of tree-based models does not change with the addition of physical features as input, as shown in the bar plots in Figures 5G–5I. However, the performance of the NN models is improved after the addition of physical features as input, especially for NNR_4layers. The comparison of the performance of NNR_4layers and NNR_4layers_Wen (NNR_4layers with composition and Wen alloy features as input) is shown in the bar plots in Figures 5G–5I. The predicted vs. experimental values plots along with their performance metrics of each property for these 4 models are presented in Section S3 of supplementary information. The NNR_4layers_Wen model performed better in predicting properties compared to other NN architecture models but not better than the ETR model. Thus, the ETR and NNR_4layers_Wen models were selected for experimental validation.

Experimental validation

The predictions of the multi-property ETR and NNR_4layers_Wen models were compared with the experimental values of the properties as shown in Figure 6. In Figure 6A, for $0.9 T < M_s < 1.4 T$, both the models predicted values within $\pm 10\%$ range of the experimental values. For $M_s > 1.4 T$, both the models overpredicted compared to the experimental values and approximately 55% of the predictions by ETR and 72.5% of the predictions by NNR_4layers_Wen have more than 10% error from experimental values. The overprediction by the models could be due to the fact that the high M_s alloys in the database are mainly binary FeCo alloys or FeCo alloys with the addition of V, while the predictions performed were for ternary Fe-Co-Ni alloys. In Figure 6B, it can be observed that the models underpredicted the $\log_{10}H_c$ values which can be

Table 2. Best ML regressor model used for the imputation of each property

Step No.	Mapping	Best ML Regressor for imputation (MAE, RMSE, R^2)
1	Composition to M_s	Extra Trees (0.049, 0.101, 0.97)
2	Composition to $\log_{10}H_c$	Extra Trees (0.249, 0.353, 0.89)
3	Composition to $\log_{10}T_c$	Extra Trees (0.006, 0.02, 0.97)
4	Composition to $\log_{10}\rho$	Extra Trees (0.05, 0.117, 0.83)
5	Composition to $\log_{10}\delta$	Decision Tree (0.167, 0.239, 0.82)
6	$\log_{10}\sigma$ to $\log_{10}H_v$	Linear (0.063, 0.081, 0.8)
7	$\log_{10}\sigma_y$ to $\log_{10}H_v$	Decision Tree (0.06, 0.047, 0.91)
8	Composition to $\log_{10}H_v$	Support Vector (0.14, 0.099, 0.56)

Table 3. Selected properties of commonly used commercial magnetic alloys

Alloy	M_s (T)	H_c (A/m)	T_c (K)	ρ ($\mu\Omega \cdot \text{cm}$)	H_V (HV)	Calculated Cost (\$/kg)	Reference
Fe _{94.21} Si _{5.79}	2.03	6.4	1018	45–48	170–195	0.46	Moses ^{130,131}
Fe _{87.86} Si _{12.14}	1.8	–	973	82	395	0.51	Kasai and Abe et al. ^{132,133}
Fe ₄₉ Co ₄₉ V ₂	2.4	16–398	1203	27	180–220	23.35	Gutfleisch Fiorillo and Tan et al. ^{2,134,135}
Fe	2.16	4–80	1044	10	150	0.42	Gutfleisch et al. ^{2,64}
Fe ₅₀ Ni ₅₀	1.4–1.6	4–20	753	40–50	120–180	7.33	Gutfleisch Fiorillo, Tan et al ^{2,134–136}
Ni ₇₈ Fe ₁₇ Mo ₅	0.65–0.82	0.25–0.64	673	60	160	13.86	Gutfleisch et al. and ASTM International ^{2,137}
Fe ₅₃ Ni ₃₀ Co ₁₇	1.2	63.66–71.62	703	43	160–230	10.24	Special Metals The NILO® and NILOMAG® Nickel-Iron Alloys ¹³⁸

due to the high bias in the curated database a containing higher number of reported H_c values for sheets and ingot samples prepared by the casting method and the experimental procedure used for validation by powder metallurgy techniques. Similarly, the models underpredicted $\log_{10} \rho$ values too as shown in Figure 6D. Moreover, H_c and ρ depend on the processing conditions of the alloys which influence the structure of the alloys, which is not considered in both the models.

For $\log_{10} T_c$ and $\log_{10} H_V$, it can be observed in Figures 6C and 6E, the ETR model predicted all the values within $\pm 10\%$ range of the experimental values. However, the NNR_4layers_Wen model predicted 100% of $\log_{10} T_c$ and 97.5% of $\log_{10} H_V$ values. The predictions for $\log_{10} T_c$ by both the models matched well the experimental values. For $\log_{10} H_V$, both the models overpredicted compared to experimental values which can be attributed to the bias in the database containing a higher number of reported H_V values for sintered binary alloys with the addition of other elements and few reported H_V values for ternary alloys in the form of sheets. Finally, for $\log_{10} \text{Cost}$, both the models predicted 97.5% values within the $\pm 10\%$ range of the experimental values. As shown in Figure 6F, for $\log_{10} \text{Cost} < 0.95$, the ETR model underpredicted and the NNR_4layers_Wen model overpredicted. As expected, for $\log_{10} \text{Cost} > 0.95$, both the models made accurate predictions. The experimental values and the predicted values for all the 40 compositions and calculated MAE of each property for both the models are tabulated in Section S4 of the supplementary information.

This shows that for a given Fe-Co-Ni alloy composition, both ETR and NNR_4layers_Wen model would predict the $\log_{10} T_c$ and $\log_{10} \text{Cost}$ with high accuracy and the $\log_{10} H_V$ and M_s with moderate accuracy. Overall, the performance of both models deployed on the experimental data are comparable to each other with NNR_4layers_Wen performing better than the ETR model. However, on the main database, the performance of the ETR model was better than that of NNR_4layers_Wen model. Thus, both the models were considered for designing multi-property alloys using Bayesian optimization.

Inverse design of the multi-property alloy

Multi-objective Bayesian optimization

For designing alloy compositions using the ETR model, the objective function defined in Equations 1, 2, 3, and 4 is used in BO. For using the NNR_4layers_Wen model, the objective function is modified to include the Wen alloy features between Equations 4 and 6 and is described by the equations later in discussion,

$$\text{add}_{\text{feat}} = \text{WenAlloys.featurize}(\text{Composition}(x_n)) \quad (\text{Equation 5})$$

$$x_{\text{all}} = \text{concatenate}(x_n, \text{add}_{\text{feat}}) \quad (\text{Equation 6})$$

$$x_{\text{transformed}} = \text{StandardScaler}(x_{\text{all}}) \quad (\text{Equation 7})$$

where add_{feat} in Equation 5 is the Wen alloy features array calculated using WenAlloys featurizer in matminer, x_n in Equations 5 and 6 is the normalized compositional array, StandardScaler in Equation 7 is the standard scaling function used on the train set for building multi-property NNR_4layers_Wen model.

For the acquisition function “EI” (as mentioned in Section multi-objective bayesian optimization for inverse design of the multi-property alloy), the ξ parameter value in gp_minimize was assigned the default value of 0.01. The ξ parameter controls how much the acquisition value favors exploration and exploitation. For the mentioned set value, the acquisition function will favor more exploitation and less exploration. This means the algorithm will search for a promising region in the search and probe to improve a promising solution. For the case of searching ternary Fe-Co-Ni alloy composition, this is beneficial as the search space is not as vast as a high entropy alloy or complex multi-component alloy.

The properties used in the objective function of MOBO (described in Equation 7) to design Fe-Co-Ni alloy compositions are M_s , T_c , and Cost because M_s and T_c are composition-dependent properties. Cost is calculated using the composition and individual element price which

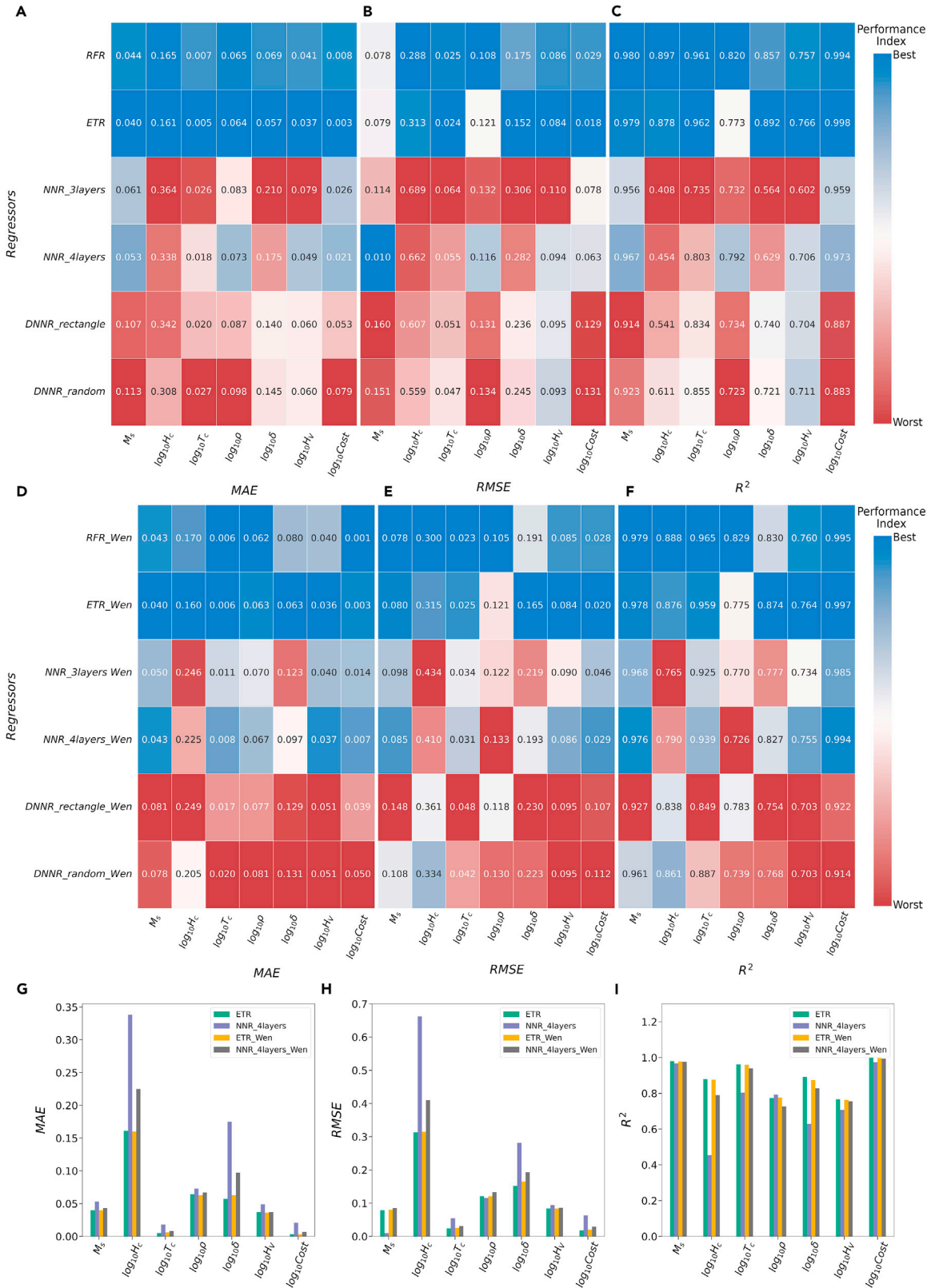


Figure 5. Performance of multi-property regression models

(A) mean absolute error (MAE), (B) root mean squared error (RMSE), and (C) coefficient of determination (R^2) of various multi-property regressor (MPR) models with composition as model features.

(D) MAE, (E) RMSE, and (F) R^2 of the MPR models with composition and Wen alloy features as model features.

(G) MAE, (H) RMSE, (I) R^2 of extra trees regressor and neural network regression consisting of 4 hidden layers with and without Wen alloy features as model features.

makes it a composition-dependent property too. Moreover, in Section [forward design of the multi-property alloy](#), it was observed that both the models are quite accurate in predicting these three properties compared to the predictions of other properties. The weights for optimizing these three properties are each set to 1 so that equal emphasis is given while designing the alloy composition. Thus, [Equation 7](#) can be rewritten as the equation later in discussion.

$$\text{objective} = \text{abs}(\text{target}_{M_s} - \text{pred}_{M_s}) + \text{abs}(\text{target}_{T_c} - \text{pred}_{T_c}) + \text{abs}(\text{target}_{\text{Cost}} - \text{pred}_{\text{Cost}}) \quad (\text{Equation 11})$$

To set the target values of M_s , T_c , and Cost for the inverse design of alloys, a survey of the target properties of the most commonly used alloys for various rotating electrical machines applications was conducted and is tabulated in [Table 3](#). Based on the values of M_s , T_c , and Cost in the table, target values of M_s , T_c , and Cost were determined to be set in the range of 1.5–2.4 T, 970–1200 K, and 10\$–19\$/kg for discovering Fe-Co-Ni alloy compositions with a moderate to high M_s , high T_c , and low Cost (Cost values based on the individual elemental cost of Fe, Co, and Ni).

The set target values of M_s , T_c , and Cost for the inverse design are as follows:

- (1) BO Set 1: M_s – 1.5 T, T_c – 970 K, and Cost – 15 \$/kg.
- (2) BO Set 2: M_s – 1.75 T, T_c – 1000 K, and Cost – 11.5 \$/kg.
- (3) BO Set 3: M_s – 1.75 T, T_c – 1143 K, and Cost – 19 \$/kg.
- (4) BO Set 4: M_s – 2 T, T_c – 1070 K, and Cost – 10.5 \$/kg.
- (5) BO Set 5: M_s – 2.2 T, T_c – 1173 K, and Cost – 10 \$/kg.
- (6) BO Set 6: M_s – 2.4 T, T_c – 1173 K, and Cost – 10 \$/kg.

For each of the six sets of target values, our Bayesian optimization strategy was used to predict alloy compositions by performing 200 iterations of the acquisition function and among them 5 alloy compositions with the lowest acquisition function values were selected for experimental validation. For instance, the evolution of acquisition function value for ETR model is shown in [Section S5](#) of the supplementary information.

Experimental validation

The predicted Fe-Co-Ni alloy compositions from Bayesian optimization and their respective predicted properties from ETR and NNR_4layers_Wen models are shown in [Figure 7](#). These predicted alloy compositions were compared to the 40 rapidly synthesized Fe-Co-Ni compositions and their properties as shown in the ternary contour plots of M_s , T_c , H_c , ρ , H_V and Cost.

Comparing the composition predictions of target sets BO set 1 and BO set 2 for both models, it can be observed that Ni content decreased in the predicted composition with an increase in the set values of M_s and T_c . The Fe content increased with a decrease in the set value of the Cost, as shown in [Figures 7A–7C](#). In the composition predictions for BO set 2 for both models, the Fe content was significantly high compared to the BO set 1 as the set values of M_s and Cost were higher and lower, respectively, as shown in [Figures 7A and 7C](#).

For the BO set 1, the composition predictions for both models were similar to each other, and all the 5 predicted compositions were quite close to each other for both models, as shown in [Figure 7](#). The predicted compositions for BO set 1 for the ETR model exhibited M_s values in the range of 1.54–1.58 T, T_c in the range of 900–912 K, and Cost in the range of 10.6\$–11.3\$/kg according to the experimental contour plots. Similarly, the predicted compositions for BO set 1 for the ETR model exhibited M_s in the range of 1.54–1.58 T, T_c in the range of 900–912 K, and Cost in the range of 11.4\$–11.7\$/kg, according to the experimental contour plots. The predicted compositions for BO set 1 for the NNR_4layers_Wen model consisted of higher Ni content compared to that for ETR model. The predicted compositions for BO set 2 for both the models exhibited M_s values in the range of 1.46–1.5 T, T_c in the range of 1000–1010 K, and Cost in the range of 15\$–15.5\$/kg, according to the experimental contour plots.

Comparing the composition predictions of target sets for BO set 2 and BO set 3 for both models, the Co content increased significantly in the predicted composition with an increase in set values of T_c and Cost, as shown in [Figures 7B and 7C](#). Moreover, it can be observed that, for both models, the predicted compositions for BO set 3 consisted of a higher content of Co and a lower content of Ni when compared to that of BO set 1 due to the increase in set value of T_c and M_s , as shown in [Figures 7A and 7B](#). The predicted compositions for BO set 3 for the ETR model exhibited M_s values in the range of 1.58–1.6 T, T_c in the range of 1090–1110 K, and Cost in the range of 18.5\$–19\$/kg, according to the experimental contour plots. Similarly, the predicted compositions for BO set 3 for the NNR_4layers_Wen model exhibited M_s in the range of 1.73–1.76 T, T_c in the range of 1100–1160 K, and Cost in the range of 18.5\$–20\$/kg, according to the experimental contour plots. The predicted compositions for BO set 3 for the NNR_4layers_Wen model consisted of higher Fe and Co content and lower Ni content compared to that for ETR model. The predicted composition with the lowest objective value for BO set 3 for the NNR_4layers_Wen model exhibited M_s in the range of 1.74 T, T_c of 1148 K, and Cost of 19.5 \$/kg according to the experimental contour plots.

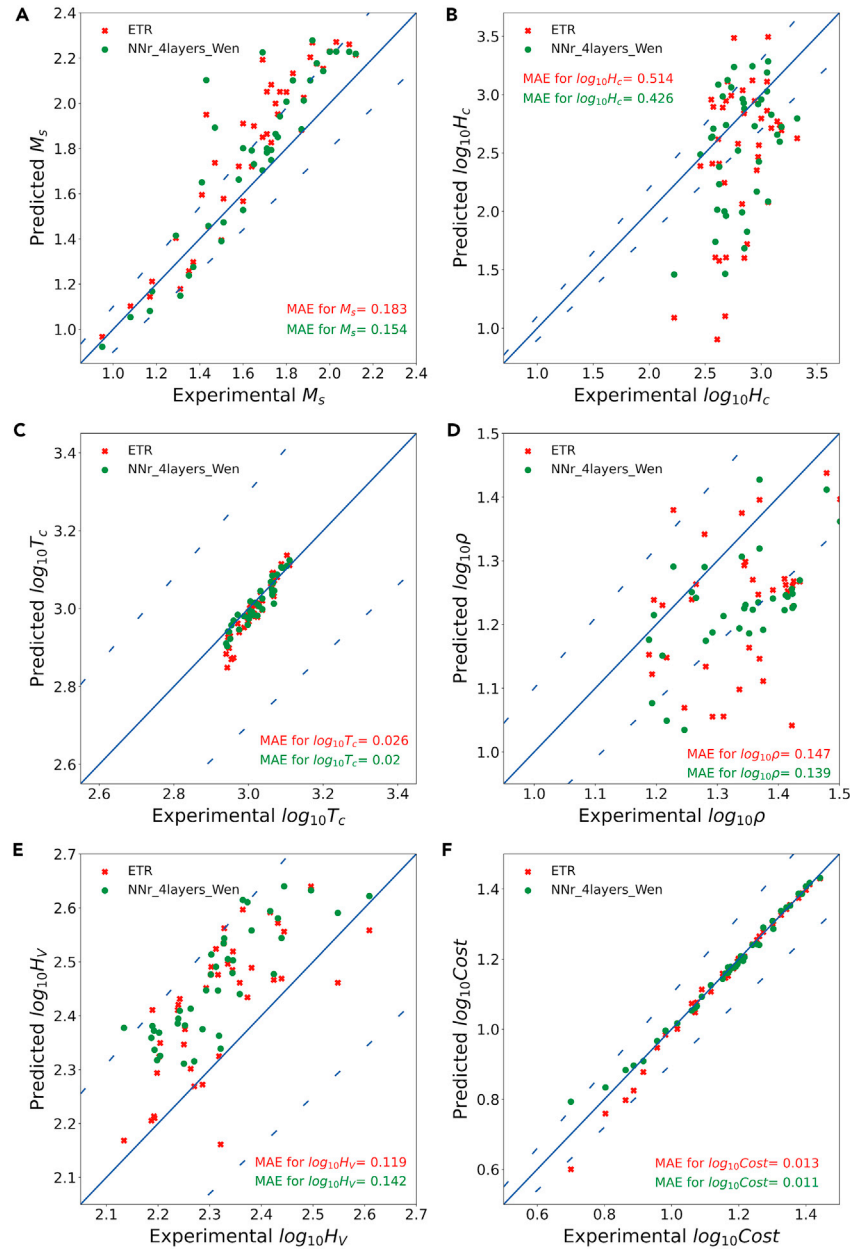


Figure 6. Experimental validation of selected multi-property models

Plot of experimental values vs. predicted values of (A) M_s , (B) $\log_{10}H_c$, (C) $\log_{10}T_c$, (D) $\log_{10}\rho$, (E) $\log_{10}H_v$, and (F) $\log_{10}Cost$ with the $\pm 10\%$ error lines (dashed) for extra trees regressor with composition as input features (green) and neural network regressor with composition and Wen alloy features as input features (red).

The predicted compositions for BO sets 4, 5, and 6 for ETR model consisted of higher Fe and lower Ni content compared to that for BO sets 1, 2, and 3 as the set target value of M_s is higher and Cost is lower as shown in Figure 7. For the NNR_4layers_Wen model, the same trend was exhibited by BO sets 5 and 6 but the predicted compositions for BO set 4 consisted of higher Fe and Ni content compared to that for BO set 3, as shown in Figure 7. Further, it can be observed that in the predicted compositions for BO sets 4, 5, and 6 for both models, Fe content increased, and Ni content decreased with an increase in the set target values of M_s ; Co content was higher for an increase in the set target values of T_c . These trends correlate well with the trends expected with domain expertise and the subset of these properties with Fe, Co, and Ni observed in the database, as explained in Section database curation and analysis. Also, it can be observed that the predicted compositions for BO sets 4, 5, and 6 for the NNR_4layers_Wen model were spread out over a wide range of compositions compared to that for the ETR model. This can be attributed to the multi-numbered and multi-level decision-based nature of the ETR model which would make the BO approach for

Table 4. Predicted compositions by MOBO for the six target sets and their respective experimental properties for BO sets 1 to 5 and predicted properties for BO set 6 using ETR model

BO Set No.	Alloy Composition	M_s (T)	H_c (A/m)	T_c (K)	ρ ($\mu\Omega\cdot\text{cm}$)	H_V (HV)	Cost (\$/kg)
1	Fe _{27.3} Co ₂₅ Ni _{47.7}	1.47	385	1002	22.8	198	15.28
2	Fe _{44.7} Co _{15.4} Ni _{39.9}	1.58	570	890	40.3	210	11.1
3	Fe _{23.5} Co _{40.8} Ni _{35.7}	1.6	290	1103.7	31.5	188	18.75
4	Fe ₅₀ Co _{20.8} Ni _{29.2}	1.77	770	1002	30	248	11.48
5	Fe _{61.9} Co _{22.8} Ni _{15.3}	2.01	1350	1056.8	24.5	365	10.2
6	Fe _{66.8} Co ₂₈ Ni _{5.2}	2.38	362.1	1190.2	17.3	254.2	10.5

the compositions closer to each other. However, the complex non-linear inter-connected nature of the NNR_4layers_Wen model would give BO freedom to approach different compositions, some of which may be the local optimum points.

In Figures 7A–7C, it can be observed that the predicted compositions for BO set 4 exhibited lower M_s values according to the experimental contour plot compared to the set target value. However, for BO set 5, the predicted compositions exhibited similar M_s values according to the experimental contour plot compared to the set target value. The predicted compositions for BO set 4 for the ETR model exhibited M_s in the range of 1.73–1.78 T, T_c in the range of 990–1010 K, and Cost in the range of 10.8\$–11.8\$/kg, similar to the experimental contour plots. The predicted compositions for the BO set 5 for the ETR model exhibited M_s in the range of 1.98–2 T, T_c in the range of 1040–1065 K, and Cost in the range of 9.8\$–10.4\$/kg, according to the experimental contour plots. The predicted composition with the lowest objective value for the BO set 4 for the NNR_4layers_Wen model exhibited a M_s of 1.79T, T_c of 1020 K, and Cost of 12.7 \$/kg, similar to the experimental contour plots. The predicted composition with the lowest objective value for BO set 5 for the NNR_4layers_Wen model was almost overlapping with the existing experimental composition of Fe_{61.3}Co_{19.3}Ni_{19.4}, which exhibited a M_s of 1.94 T, T_c of 1000 K, and Cost of 9.6 \$/kg.

For the predicted compositions for all the BO sets for ETR model, the color of the predicted H_c values was blue according to the experimental contour plot color bar, as shown in Figure 7D. The range of predicted H_c values for the predicted compositions using the ETR model was as follows: BO set 1 in the range of 3–4.2 A/m, BO set 2 in the range of 38–45 A/m, BO set 3 in the range of 353–369 A/m, BO set 4 in the range of 311–333.5 A/m, BO set 5 in the range of 549–569 A/m, and BO set 6 in the range of 362–434 A/m. Considering the experimental contour lines, the predictions for BO set 3 using the ETR model were closest to experimental values.

The range of predicted H_c values for the predicted compositions using the NNR_4layers_Wen model was as follows: BO set 1 in the range of 109–122 A/m, BO set 2 in the range of 55–60 A/m, BO set 3 in the range of 805–964 A/m, BO set 4 in the range of 97–379 A/m, BO set 5 in the range of 506–1306 A/m, and BO set 6 in the range of 894–1146 A/m. Considering the experimental contour lines, the predictions for H_c for the predicted compositions using the NNR_4layers_Wen model were not close to the experimental values.

For ρ , the predicted values for the predicted compositions for BO sets 1–5 using the ETR model were in the range of 18–24 $\mu\Omega\cdot\text{cm}$, which implies that it is challenging to discover Fe-Co-Ni compositions having ρ greater than 30 $\mu\Omega\cdot\text{cm}$ using the ETR model. The prediction of ρ for all BO sets compositions were also lower than the estimated actual values from the contour plot in Figure 7E. For the BO set 6, the predicted ρ for the predicted compositions was in the range 16–17.3 $\mu\Omega\cdot\text{cm}$, which can be close to the actual value.

Similarly, the predicted ρ values for predicted compositions for BO sets 1–5 using the NNR_4layers_Wen model were in the range of 14–23 $\mu\Omega\cdot\text{cm}$, which implies that it is challenging to discover Fe-Co-Ni compositions having ρ greater than 30 $\mu\Omega\cdot\text{cm}$ using the NNR_4layers_Wen model. The ρ values in the database would be in this range and most of the samples are prepared starting from the casting method which is a different method of synthesis compared to the experimental validation method.

For H_V , it can be observed from Figure 7F that the predicted values for the predicted compositions of all the BO sets using the ETR model were closer to the experimental values than those using the NNR_4layers_Wen model by comparing the color trend of the contour lines.

To analyze further the performance of the inverse design strategy using both MPR models, the predicted composition with the lowest objective value for the BO sets 1–5 for both models was considered. The predicted properties were compared to the experimental values obtained from the contour plots to calculate the deviation percentages. These results are tabulated in Section S5 of the supplementary information. The composition predicted for BO set 1 for the ETR model has a 2% lower experimental M_s value and for BO sets 2–5 have 8.5–11.5% lower experimental M_s values compared to the predicted M_s values. Similarly, the composition predicted for BO set 1 for the NNR_4layers_Wen model has a 2% lower experimental M_s value, for BO set 3, it has a 1.1% lower experimental M_s value and for BO sets 2, 4, and 5, it has 11% lower experimental values compared to predicted M_s values.

The composition predicted for BO sets 1–3 for the ETR model have 4–7% higher experimental T_c values, for BO set 4 has 14.8% higher experimental T_c value, and for BO set 5 has just 0.9% higher experimental T_c value compared to the predicted T_c values. The composition predicted for BO sets 1–5 for the NNR_4layers_Wen model have 5–9% higher experimental T_c values compared to the predicted T_c values.

For Cost, the composition predicted for the BO sets 1–5 for both models has predicted values quite similar to experimental values. This is because Cost is calculated using the weighted average of elemental costs.

The average deviation of the predicted M_s , T_c , and Cost values of the predicted compositions were 8.2% higher, 6.4% lower, and 1.1% lower compared to the experimental values, respectively, for the ETR model. The average deviation of predicted M_s , T_c , and Cost values

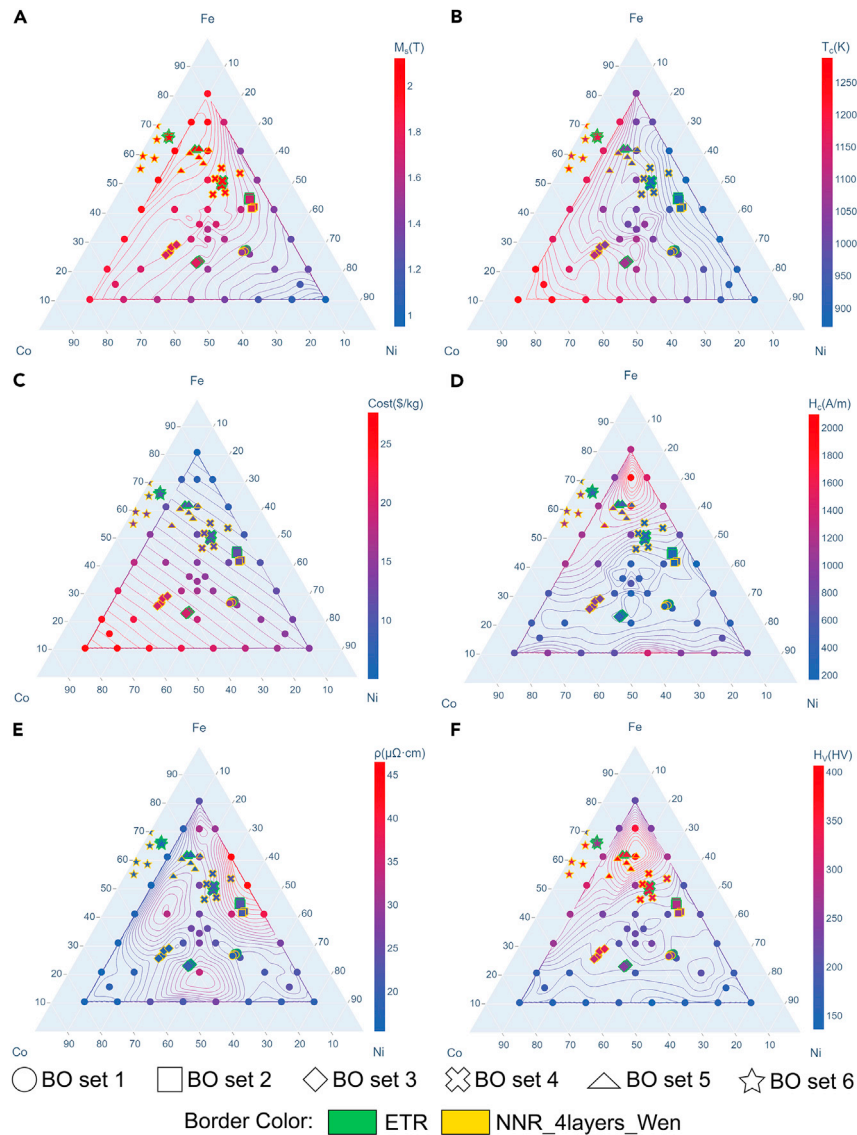


Figure 7. Experimental validation of inverse design strategy

Ternary contour plot of (A) M_s , (B) $\log_{10} T_c$, (C) $\log_{10} Cost$, (D) $\log_{10} H_c$, (E) $\log_{10} \rho$, and (F) $\log_{10} H_V$ of 40 experimentally synthesized Fe-Co-Ni alloy compositions with predicted compositions from inverse design strategy with their respective predicted property values from ETR (points with green border) and NNR_4layers_Wen (points with yellow border) for 6 target sets.

of the predicted compositions were 7.2% higher, 6.6% lower, and 0.4% lower compared to the experimental values, respectively, for the ETR model.

The deviations of the predicted H_c values of the predicted compositions for the BO sets 1–5 for both models were quite high compared to the experimental H_c values. This could be attributed to the well-known dependence of H_c on processing parameters, microstructure, and heat treatment parameters. Similarly, the average deviations of predicted ρ values of the predicted compositions for the BO sets 1–5 for both models were 45–46% higher compared to the experimental ρ values. Finally, the predicted H_V values of the predicted compositions for the BO sets 1 and 5 for the ETR model were very close to the experimental H_V values, with a deviation of 1.8% and 5.2%, respectively. The predicted H_V values of the predicted compositions for the BO sets 2, 3, and 4 for the ETR model were 20–29% higher compared to the experimental H_V values. The average deviation of the predicted H_V values of the predicted compositions for the ETR model was 14.2% higher compared to the experimental H_V values. On the other hand, the predicted H_V values of the predicted compositions for the NNR_4layers_Wen model exhibited higher deviation percentages compared to that for the ETR model except for BO set 5. Overall, the average deviation of predicted H_V values of the predicted compositions for the NNR_4layers_Wen model was 22.9% higher compared to the experimental H_V values.

It can be inferred that the BO composition predictions and their predicted property values using the ETR model performed slightly better compared to that from the NNR_4layers_Wen model with a similar prediction performance for M_s , T_c , and Cost and better prediction performance for H_V . The predicted composition with the lowest objective value of MOBO for each target set using the ETR model and their respective approximate experimental properties for BO sets 1 to 5 and predicted properties for BO set 6 are tabulated in Table 4. Comparing the predicted property values of the BO predicted compositions for the ETR model, $\text{Fe}_{66.8}\text{Co}_{28}\text{Ni}_{5.2}$ (predicted M_s – 2.38 T, T_c – 1190.2 K, H_c – 362.1 A/m, ρ – 17.3 $\mu\Omega$ cm, H_V – 254.2 HV and Cost – 10.5\$/kg) and $\text{Fe}_{61.9}\text{Co}_{22.8}\text{Ni}_{15.3}$ (predicted M_s – 2.22 T, T_c – 1047.5 K, H_c – 549.5 A/m, ρ – 18.7 $\mu\Omega$ cm, H_V – 347.1 HV and Cost – 10\$/kg) demonstrate superior multi-property performance compared to previous reports, and performance comparable to recently designed Fe-Co-Ni alloys using accelerated experimental methodology.^{123,135}

Conclusion and outlook

We curated a database of bulk Fe-Co-Ni-based magnetic alloys containing magnetic, mechanical, and electrical properties. The database is heterogeneous as alloys synthesized using different methods are included. The gaps in the database were filled using a unique ML-based multi-step imputation strategy. A comparative study of the performance of 12 different ML regression models, from linear regression models to various tree-based models to DNNs, mapping from composition to each property and mapping between mechanical properties was performed. The best model with the lowest MAE was chosen to fill the empty spaces for that property column. This imputed database was used to develop a multi-property model (forward design) mapping from composition to all the properties by using 6 different models. Apart from using only compositional input features, physical descriptors (Wen alloy features) were also used as input features to develop multi-property models using these 6 models. The best model with the lowest MAE, i.e., the ETR model, and the best neural network model with lowest MAE, i.e., NNR_4layers_Wen was chosen.

Further, a multi-objective BO (inverse design) was developed to predict alloy compositions by providing a target set of properties. Both the forward and inverse design methodology were experimentally validated by using 40 Fe-Co-Ni alloy compositions synthesized and characterized using high-throughput techniques. Overall, the ETR model performed better compared to the NNR_4layers_Wen model in both forward and inverse design strategies. Finally, two alloy compositions $\text{Fe}_{65.84}\text{Co}_{28.66}\text{Ni}_{5.5}$ and $\text{Fe}_{61.5}\text{Co}_{23.14}\text{Ni}_{15.35}$ were obtained from inverse design using ETR model with the optimum combination of $M_s > 2$ T, $T_c > 1050$ K, $H_V > 250$ HV, $H_c < 1400$ A/m, $\rho > 17$ $\mu\Omega$ cm, and Cost ≈ 10 \$/kg. The predicted properties value of the predicted compositions by MOBO were within 14% of the experimental values for M_s , T_c , Cost, and H_V . Our proposed approach can be used to design any class of materials using a heterogeneous database.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
 - Lead contact
 - Materials availability
 - Data and code availability
- EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS
- METHOD DETAILS
- QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.isci.2024.109723>.

ACKNOWLEDGMENTS

This work is supported by the AME Programmatic Fund by the Agency for Science, Technology and Research, Singapore under Grant No. A1898b0043 and Production Area of Advance (AoA) at Chalmers University of Technology. We would like to acknowledge the Facility for Analysis, Characterization, Testing and Simulation, Nanyang Technological University, Singapore, for use of their EDS facility.

AUTHOR CONTRIBUTIONS

Conceptualization - S.P.P., V.C., Y.F.L., K.H., and R.V.R.; methodology – S.P.P., V.C., Y.F.L., and K.H.; data curation – S.P.P.; formal analysis – S.P.P., V.C., and K.H.; investigation – S.P.P. and V.C.; software – S.P.P., Y.F.L., R.Z., and M.T.; visualization – S.P.P. and K.H.; writing (original draft) – S.P.P.; writing (review and editing) – S.P.P., V.C., Y.F.L., K.H., and R.V.R.; supervision – V.C., Y.F.L., K.H., and R.V.R.; funding acquisition – R.V.R.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: January 4, 2024

Revised: March 11, 2024

Accepted: April 8, 2024

Published: April 10, 2024

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited data		
GitHub repository with literature curated and experimental data and code (https://github.com/Shakti-95/Data-and-Codes-for-Experimentally-Validated-Inverse-design-of-Multi-Property-Fe-Co-Ni-alloys)	This paper; Zenodo	https://doi.org/10.5281/zenodo.10686272
Software and algorithms		
Python 3.9	Python Software Foundation	https://www.python.org/
Jupyter notebook	Jupyter	https://jupyter.org/
WenAlloys featurizer (Matminer)	Wen et al., ³⁸ 2019	https://doi.org/10.1016/j.actamat.2019.03.010

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources should be directed to the lead contact, Prof. Raju V. Ramanujan (ramanujan@ntu.edu.sg).

Materials availability

This study did not generate new materials.

Data and code availability

- The curated data from literature, imputed data, and experimental data have been deposited in the GitHub repository (<https://github.com/Shakti-95/Data-and-Codes-for-Experimentally-Validated-Inverse-design-of-Multi-Property-Fe-Co-Ni-alloys>) and are publicly available.
- All original codes have been deposited in the GitHub repository (<https://github.com/Shakti-95/Data-and-Codes-for-Experimentally-Validated-Inverse-design-of-Multi-Property-Fe-Co-Ni-alloys>) and are publicly available as of the date of publication. DOIs are listed in the [key resources table](#).
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

This work did not need any unique experimental model.

METHOD DETAILS

The complete Methodology section in the main text should be under this section heading.

QUANTIFICATION AND STATISTICAL ANALYSIS

All the analysis and plots were performed in Python. The figures were all prepared using licensed version of Adobe Illustrator.