# <sup>1</sup> Evidence-based data mining method to reveal similarities between <sup>2</sup> materials based on physical mechanisms

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Measuring the similarity between materials is essential for estimating their properties and revealing the asso-17 ciated physical mechanisms. However, current methods for measuring the similarity between materials rely 18 on theoretically derived descriptors and parameters fitted from experimental or computational data, which 19 are often insufficient and biased. Further, outliers and data generated by multiple mechanisms are usually 20 included in the dataset, making the data-driven approach challenging and mathematically complicated. To 21 overcome such issues, we apply the Dempster-Shafer theory to develop an evidential regression-based similar-22 ity measurement (eRSM) method, which can rationally transform data into evidence. It then combines such 23 evidence to conclude the similarities between materials, considering their physical properties. To evaluate the 24 eRSM, we used two materials datasets, including 3d transition metal-4f rare-earth binary and quaternary 25 high-entropy alloys with target properties, Curie temperature and magnetization. Based on the informa-26 tion obtained on the similarities between the materials, a clustering technique is applied to learn the cluster 27 structures of the materials that facilitate the interpretation of the mechanism. The unsupervised learning 28 experiments demonstrate that the obtained similarities are applicable to detect anomalies and appropriately 29 identify groups of materials whose properties correlate differently with their compositions. Furthermore, 30 significant improvements in the accuracies of the predictions for the Curie temperature and magnetization 31 of the quaternary alloys are obtained by introducing the similarities, with the reduction in mean absolute 32 errors (MAE) of 36% and 18%, respectively. The results show that the eRSM can adequately measure the 33 similarities and dissimilarities between materials in these datasets with respect to mechanisms of the target 34 properties. 35

### INTRODUCTION 36

The concept of machine learning has great potential 37 for application in several areas of materials science, espe-38 cially for discovering new materials. In materials science, 39 a number of the problems addressed by data-driven ap-40 proaches require the effective utilization of existing ma-41 terial data for predicting the properties of new mate-42 rials and understanding the underlying physicochemical 43  $_{44} \text{ mechanisms}^1$ .

From an engineering point of view, developing a data-45 46 driven model that quickly and accurately predicts the physical properties of possible materials from accumu-47 48 49 50 52 If there are sufficient independent supervised data from 72 rial data can accelerate the process of describing, inter-53 the distribution of the target material data, a model with 73 preting, and understanding the physicochemical mech-54 high prediction accuracy can be built using state-of-the-74 anisms underlying the observed physical phenomena of

55 art data-driven techniques. However, because materi-56 als research and development aim to develop materials 57 that are superior to existing ones, the distribution of the 58 target prediction data may be completely different from <sup>59</sup> the distribution of the original training data. Therefore, 60 there are concerns about whether data-driven models can 61 accurately predict the physical properties of new materi-62 als.

On the contrary, considering the history of materi-64 als science, researchers have discovered various materi-65 als through a loop of hypothesis and verification based 66 on their knowledge, experience, and serendipity. Partic-67 ularly, hypothesizing relies heavily on describing, interlated data can reduce the time required for material de- or preting, and understanding the underlying physicochemvelopment. By applying a data-driven model to screen on ical mechanisms of the observed physical phenomena of materials *in-silico*, we narrow down the candidates that 70 materials. Scientifically, applying a data-driven approach require expensive calculations and experiments to verify. 71 to extracting knowledge from existing complicated mate-

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1 materials. This reduces the time required for material 59 nisms is necessary to assess the similarity between the <sup>4</sup> and understandable to humans must be developed.

One of the most intuitive and interpretable data-driven 5 6 approaches for humans is analogy-based inductive rea-7 soning, which infers the properties of a new instance uss ing the information of the observed instances that are  $\circ$  most similar to it<sup>2-5</sup>. By applying analogy-based mod-10 els, we can easily explain the reasoning process behind the predictions and reveal the physicochemical mecha-11 nisms rationalizing the observations<sup>6,7</sup>. Materials scien-12 tists have resolved different problems in materials science 71 apply data-driven approaches to materials science. 13 by systematizing information about analogies in compo-14 sition or structure between materials that exhibit similar 15 physicochemical properties<sup>8–11</sup>. 16

Especially, in a discipline based on fundamental prin-17 18 ciples, such as condensed matter physics, it is essential to elucidate the physical mechanisms and which materi-19 als are manifested through each of these physical mecha-20 nisms. However, despite several new materials and supe-21 rior properties having been discovered, it is still difficult 22 to appropriately quantify the similarities between mate-23 rials to elucidate the underlying physicochemical mech-24 anisms of these properties. Furthermore, this difficulty 25 arises from the fact that the mechanisms of materials' 26 properties are typically interpreted in terms of physico-27 chemical concepts based on relative criteria. 28

The phenomenon of superconductivity in materials, 29 which originates from the instability of metals, is a well-30 known example of the above difficulty. One of the most 31 successful theories that describe the microscopic mech-32 anisms is the Bardeen-Cooper-Schrieffer (BCS) theory 33 for superconductivity<sup>12</sup>, the origin of which is electron-34 phonon interactions. However, there also exist other mechanisms. For example, one of the most plausive 36 origins of superconductivity in the high- $T_C$  cuprates is 37 electron-electron interactions. Nevertheless, it is not easy 38 to achieve a consensus of classifying the superconducting 39 mechanism of materials among researchers as the ori-40 gins. Although the emergence of superconductivity is 41 basically due to the instability in the metallic phase, it 42 is not easy to achieve the consensus because both the 43 mentioned and other mechanisms can contribute cooper-44 <sup>45</sup> atively in increasing the  $T_C$  value, for example. Although it is challenging to classify individual materials when con-46 sidering phenomena that cause such a situation, it is expected that the underlying physical mechanisms can be 48 discovered if we can inductively quantify the similarities 49 50 between the materials of interest and group similar materials using all observation data. 51

Incidentally, inductive reasoning with inefficient sim-52 110 ilarity assessment can lead to misidentification of 53 outliers<sup>13</sup> and difficulty in explaining the underlying 111 physicochemical mechanisms of datasets using single 112 55 56 models. Therefore, regarding predefined material de- 113 57 scriptors, an exhaustive examination of all possible hy-58 potheses about the unknown physicochemical mecha- 114

<sup>2</sup> development. Hence, to be effectively applied to materials. Furthermore, similarity measures are usually 3 als science, data-driven approaches that are interpretable 61 context-dependent. Because the context changes, the 62 similarity measure must be modified to adequately cap- $_{63}$  ture the phenomena under study<sup>14,15</sup>. Thus, a quanti-64 tative measure of similarity needs to consider the uncertainty arising from the context or the measurement itself, 65 especially in situations where material data are often in-66 67 sufficient and heavily biased. Moreover, similarities from 68 different contexts may not be directly comparable in the 69 integration to draw conclusions about the similarity between materials. These reasons make it challenging to 70

> To overcome such issues and efficiently extract knowl-72 73 edge from the data, we propose a new approach that 74 shifts from measuring the similarity between materials 75 to quantitatively measure the confidence in their similarities. We adopt the Dempster–Shafer theory  $^{16-18}$ , re-76 ferred to as the evidence theory, to develop an eviden-77 78 tial regression-based similarity measurement (eRSM) for detecting subgroups of materials such that leaned mod-79 els from the subgroups show high correlations between 80 s1 descriptors and the target property of the constituent 82 materials. Further analysis of models describing the subgroups provide valuable information to extract, interpret, 83 84 and understand physical mechanisms. The Dempster-Shafer theory can be regarded as a generalization of the 85 86 Bayesian approach for solving the problem of incomplete 87 and insufficient information. Moreover, it is suitable for solving material data problems<sup>19,20</sup>. The measure of sim-<sup>89</sup> ilarity here refers to whether the observed physical prop-90 erties of the materials under study are described using the same hidden mechanism that has not yet been re-91 92 vealed. In other words, we consider any pair of materials (in the dataset) as similar if their physical properties can 93 be described by the same hidden mechanism; otherwise, 94 95 the pair of materials is considered dissimilar. We then 96 first generate numerous hypothetical mechanisms by ran-97 domly choosing subsets of data instances and construct-98 ing regression models for each subset. Each regression 99 model is considered a source of evidence of the similari-100 ties between materials. Thereafter, the Dempster–Shafer 101 theory<sup>16–18</sup>, which has a foundation for modeling and 102 combining the uncertainty of evidence, is applied to inte-<sup>103</sup> grate the collected pieces of evidence to draw conclusions about the similarities between materials. The eRSM con-105 sists of three main steps as follows:

- 1. Collect sources of evidence: Hypothetical mechanisms are collected from a dataset by applying regression analysis with single or mixture models and are used as sources of evidence to rationalize the similarity states of materials.
- 2. Model similarity evidence: An appropriate mass function is designed to model the obtained evidence within the framework of the evidence theory.
- 3. Combine pieces of evidence: Dempster's rule of

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FIG. 1. Illustrative figures of the three possible similarity states between two data instances (blue circles), including similar (a), dissimilar (b), and uncertain (c), considering a referential regression model  $f_r$  (black line). The gray region is the interval that determines whether a data instance can be considered to have been generated by regression model  $f_r$ .

combination is used to integrate the pieces of the evidence. 2

The steps of the eRSM are explained in detail in Sec-3 4 tion II. Regarding the framework of the evidence theory, 5 the essential contributions of the eRSM are collecting 6 sources of evidence about the similarities between mate-7 rials from datasets and designing suitable mass functions s to model the pieces of evidence rationally. The effective-• ness of obtained similarities using the eRSM for subdi-<sup>10</sup> viding alloys from datasets into homogenous subgroups 11 is supported by experiments on 1) a dataset of binary 12 alloys with their Curie temperature as a target property (Section IIIB); and 2) two dataset of quaternary alloys 13 with their magnetization (Section IIIC) and Curie tem-14 <sup>15</sup> perature (Section III D) as the target properties. Further 16 analysis of the detected subgroups to interpret the under-17 lying physical mechanisms is shown in Section IIIE

### METHODOLOGY 18 II.

We consider a dataset  $\mathcal{D}$  consisting of p data in-19 20 stances. We assume that a data instance with index i in  $\mathcal{D}$  is described by n predefined descriptors and 21  $_{22}$  is represented by an *n*-dimensional numerical vector, 23  $x_i = (x_i^1, x_i^2, \dots, x_i^n) \in \mathbb{R}^n$ . The target property of <sup>24</sup> the data instance  $x_i$  is  $y_i \in \mathbb{R}$ . Thereafter, the dataset 25  $\mathcal{D} = \{(x_1, y_1), (x_2, y_2) \dots (x_p, y_p)\}$  is represented using 26 a  $(p \times (n+1))$  matrix. In this study, we consider that <sup>27</sup>  $\mathcal{D}$  may contain pairs of data instances  $x_i$  and  $x_j$ , where **28**  $x_i \approx x_j$ ; however, the value of  $y_i$  is far from  $y_j$ .

### Collecting sources of similarity evidence Δ 29

30 31  $_{33}$  in  $\mathcal{D}$ . Considering each sample, we obtain two datasets:  $_{34}$  determined that it does not belong to  $f_r$ . By increasing  $\mathcal{I}_{24}$  the reference dataset,  $\mathcal{D}_{ref}$ , and the evaluation dataset,  $\mathcal{I}_{25}$  or decreasing the value of the parameter  $\alpha$ , the condition

35  $\mathcal{D}_{eval}$   $(\mathcal{D}_{ref} \cap \mathcal{D}_{eval} = \emptyset$  and  $\mathcal{D}_{ref} \cup \mathcal{D}_{eval} = \mathcal{D})$ . Con-36 sidering  $\mathcal{D}_{ref}$ , we can generate a single or multiple ref-37 erence functions  $f_r: \mathbb{R}^n \to \mathbb{R}$  using a Gaussian process  $_{38}$  (GP)<sup>21</sup> or a mixture of Gaussian processes (MGP)<sup>22</sup>, re-39 spectively. This study applies GP- or MGP-based models 40 instead of other nonlinear regression models such as ker-<sup>41</sup> nel ridge regression<sup>23</sup>, random forest regression<sup>24</sup>, or arti-<sup>42</sup> ficial neural networks<sup>25</sup> because GP or MGP can quantify 43 the uncertainty of its prediction without introducing any 44 other statistical validation. The sampling ratios of  $\mathcal{D}_{ref}$ from  $\mathcal{D}$  are fixed at 0.3 and 0.7 for the experiments with 45 46 GP and MGP, respectively. Each reference function  $f_r$ 47 is considered as a source to provide pieces of evidence 48 for the similarity between  $(x_i, y_i)$  and  $(x_i, y_i)$  in  $\mathcal{D}_{eval}$ . The function  $f_r$  is not used to provide any information 49 50 about the similarities between the data instances in  $\mathcal{D}_{ref}$ <sup>51</sup> or between a data instance in  $\mathcal{D}_{ref}$  and a data instance 52 in  $\mathcal{D}_{eval}$ . This is to exclude self-evaluation to ensure the 53 objectivity of the evidence. Regarding a reference func-54 tion  $f_r$ , we consider the state of the similarity between **55**  $(x_i, y_i)$  and  $(x_j, y_j)$  as:

- Similar: Both data instances can be considered to have been generated by the function  $f_r$  (Fig. 1 a).
- Dissimilar: Only one of the data instances can be considered to have been generated by the function  $f_r$  (Fig. 1 b).
- Uncertain: Neither of the data instances can be considered to have been generated by the function  $f_r$  (Fig. 1 c). The uncertain state indicates that  $f_r$ does not provide any information about the similarity between  $(\boldsymbol{x}_i, y_i)$  and  $(\boldsymbol{x}_j, y_j)$ .

To quantitatively evaluate whether  $(\boldsymbol{x}_i, y_i)$  can be con-66 <sup>67</sup> sidered to have been generated by the regression function 68  $f_r$ , we use the likelihood  $p(O_i|f_r)$ , the probability of event 69  $O_i$  that a data instance  $(\boldsymbol{x}_i, y_i)$  is observed, considering 70  $f_r$ . The likelihood  $p(O_i|f_r)$  is modeled using a normal 71 distribution with mean and standard deviation depend-<sup>72</sup> ing on the predicted target value  $\hat{y}_i = f_r(\boldsymbol{x}_i)$  and the cor-73 responding standard error  $\sigma_{\boldsymbol{x}_i}$  by  $f_r$ , respectively. This 74 is expressed as:

$$p(O_i|f_r) = \begin{cases} 1 & \text{if } \Delta_i \leq 3\bar{\sigma} \\ 2 \times \int_{\Delta_i - 3\bar{\sigma}}^{+\infty} \mathcal{N}\left(u|0, \alpha \, \sigma_{\boldsymbol{x}_i}\right) du & \text{otherwise} \end{cases},$$
(1)

75 where  $\Delta_i = |y_i - \hat{y}_i| = |y_i - f_r(\boldsymbol{x}_i)|$  is the deviation 76 from the true to the predicted target values of data in- $\tau$  stance *i* using  $f_r$ , and  $\bar{\sigma}$  is the average of the predictive 78 standard error of all the data instances in  $\mathcal{D}_{ref}$ .  $\alpha$  is 79 the hyperparameter used to adjust the condition that reso stricts the data instances belonging to the function  $f_r$ . We perform random subset sampling of the data in- a In other words, the interval that determines the probastances without replacement to collect a large amount of  $s_2$  bility that a data instance  $(x_i, y_i)$  belongs to  $f_r$  is  $\alpha \sigma_{x_i}$ . evidence of the similarity between pairs of data instances and if the data instance falls outside this interval, it is

 $_{3}$  smaller, respectively. Optimal values of  $\alpha$  can be chosen 4 using statistical criteria and appropriate validation meth-5 ods; however, we set  $\alpha = 2$  for all experiments in this • work to reduce model complexity. We consider  $p(O_i|f_r)$  $\tau$  as the probability that  $(\boldsymbol{x}_i, y_i)$  is generated by  $f_r$ , and s  $p(O_i|f_r) = 1 - p(O_i|f_r)$  is the probability that  $(\boldsymbol{x}_i, y_i)$  is • not generated by  $f_r$ . Supplementary Figure 1 illustrates 10 the process of modeling the probability  $p(O_i|f_r)$ .

Events where  $(\boldsymbol{x}_i, y_i)$  or  $(\boldsymbol{x}_i, y_i)$  is generated by the 11 12 function  $f_r$  are independent events. Therefore, consider-13 ing the function  $f_r$ , we can evaluate the joint probabilities 14 of observing:

• Both data instances: 15

$$p(O_i, O_j | f_r) = p(O_i | f_r) \times p(O_j | f_r);$$
(2)

• Only one of the data instances: 16

$$p(O_i, \overline{O}_j | f_r) + p(\overline{O}_i, O_j | f_r) = p(O_i | f_r) \times p(\overline{O}_j | f_r) + p(\overline{O}_i | f_r) \times p(O_j | f_r);$$
(3)

• Neither of the data instances: 17

$$p(\overline{O}_i, \overline{O}_j | f_r) = p(\overline{O}_i | f_r) \times p(\overline{O}_j | f_r) = 1 - p(O_i, O_j | f_r) - p(O_i, \overline{O}_j | f_r) - p(\overline{O}_i, O_j | f_r).$$
(4)

### Modeling evidence by mass functions 18 B.

Considering the Dempster–Shafer theory framework<sup>16</sup>, 19 <sup>20</sup> we begin by defining the frame of discernment  $\Omega$ . Let <sup>21</sup>  $\Omega = \{s, ds\}$  be the universal set representing the similar-<sup>22</sup> ity states of any two data instances  $(x_i, y_i)$  and  $(x_j, y_j)$ . s and ds denote the similarity and dissimilarity states 23 between the two data instances, respectively. 24

According to the Dempster–Shafer theory, the evidence 25 of the similarity states between these two data instances 26 <sup>27</sup> is represented by a mass function  $m^{i,j}$  (or a basic proba-<sup>28</sup> bility assignment)<sup>16</sup>. This assigns probability masses to <sup>29</sup> all the nonempty subsets of  $\Omega$  ( $\mathcal{X} = \{\{s\}, \{ds\}, \{s, ds\}\}$ ). 30 It is defined as follows:

$$m^{i,j}: \mathcal{X} \to [0,1] \text{ with } \sum_{E \in \mathcal{X}} m(E) = 1.$$
 (5)

<sup>31</sup> The masses assigned to  $\{s\}$  and  $\{ds\}$  reflect the degrees of 32 belief exactly committed to the evidence to support the 33 similarity and dissimilarity between  $(\boldsymbol{x}_i, y_i)$  and  $(\boldsymbol{x}_j, y_j)$ , <sup>34</sup> respectively. The weight assigned to  $\{s, ds\}$  expresses the <sup>35</sup> degree of belief that the evidence provides no information 36 about the similarity (or dissimilarity) between  $(x_i, y_i)$ and  $(\boldsymbol{x}_j, y_j)$ . 37

$$m_{f_r}^{i,j}(\{s\}) = \frac{p(O_i, O_j | f_r)}{\gamma_{i,j}}$$
(6)

$$n_{f_r}^{i,j}(\{ds\}) = \frac{p(O_i,\overline{O}_j|f_r) + p(\overline{O}_i,O_j|f_r)}{\gamma_{i,j}}$$
(7)

$$m_{f_r}^{i,j}(\{s,ds\}) = 1 - \frac{1}{\gamma_{i,j}} + \frac{p(\overline{O}_i,\overline{O}_j|f_r)}{\gamma_{i,j}},\qquad(8)$$

<sup>41</sup> where  $\gamma_{i,j} = (e^{\frac{\bar{\sigma}}{\Delta_y}} + 1) \times (\frac{\sigma_{x_i}}{\bar{\sigma}} + 1) \times (\frac{\sigma_{x_j}}{\bar{\sigma}} + 1)$  is a discount-<sup>42</sup> ing factor<sup>16,26</sup>, which describes the unreliability of evi-43 dence about the similarity between  $(\boldsymbol{x}_i, y_i)$  and  $(\boldsymbol{x}_j, y_j)$ 44 collected from a source of evidence  $f_r$ .  $\Delta_y$  is the varia-45 tion range of the target variable y in the dataset  $\mathcal{D}$ . The 46 smaller  $\bar{\sigma}$  is relative to  $\Delta_y$ , the more reliable the learned 47 regression function  $f_r$  is. Also, when  $\sigma_{\boldsymbol{x}_i}$  and  $\sigma_{\boldsymbol{x}_i}$  are 48 smaller than  $\bar{\sigma}$ ,  $f_r$  can provide reliable evidence for the 49 relationship between  $(\boldsymbol{x}_i, y_i)$  and  $(\boldsymbol{x}_i, y_i)$ . By contrast, 50 when  $\sigma_{\boldsymbol{x}_i}$  and  $\sigma_{\boldsymbol{x}_i}$  are large compared to  $\bar{\sigma}$ ,  $f_r$  cannot pro-51 vide reliable evidence for the relationship between  $(\boldsymbol{x}_i, y_i)$ 52 and  $(\boldsymbol{x}_j, y_j)$ . A detailed explanation of each component 53 in  $\gamma_{i,j}$  is provided in Supplementary Section I.

## 54 C. Dempster's rule in combining evidence

Assuming that we can collect q pieces of evidence from 56  $\mathcal{F}_r = \{f_r^1, \ldots, f_r^q\},$  a set of q reference functions is gen- $_{57}$  erated from  $\mathcal{D}$  to evaluate the similarity between a pair 58 of data instances with indices i and j. According to the 59 Dempster–Shafer theory framework, any two pieces of ev-60 idence collected from the reference functions  $f_r^l$  and  $f_r^k$ , 61 which are modeled by the corresponding mass functions 62  $m_{f_r}^{i,j}$  and  $m_{f_r}^{i,j}$ , respectively, can be combined using the 63 Dempster rule of combination to assign the joint mass 64  $m^{i,j}_{\{f^l_r,f^k_r\}}$  to each nonempty subset E of  $\Omega$  as follows:

$$m_{\{f_r^{i}, f_r^{k}\}}^{i,j}(E) = (m_{f_r^{i}}^{i,j} \oplus m_{f_r^{k}}^{i,j})(E) = \frac{\sum_{E_t \cap E_v = E} m_{f_r^{i}}^{i,j}(E_t) \times m_{f_r^{k}}^{i,j}(E_v)}{1 - \sum_{E_t \cap E_v = \emptyset} m_{f_r^{i}}^{i,j}(E_t) \times m_{f_r^{k}}^{i,j}(E_v)},$$
(9)

65 where  $E, E_t$ , and  $E_v$  are nonempty subsets of  $\Omega$ . Demp-66 ster's rule is commutative and associative.

67 Based on Dempster's rule, the obtained mass functions  $_{68}$  corresponding to the q pieces of evidence are combined 69 to assign the final mass  $m_{\mathcal{F}_r}^{i,j}$  as follows:

$$m_{\mathcal{F}_r}^{i,j}(E) = \left(m_{f_r^1}^{i,j} \oplus m_{f_r^2}^{i,j} \oplus \dots \oplus m_{f_r^q}^{i,j}\right)(E).$$
(10)

We perform similar analyses for all pairs of data in-70 Therefore, the mass function  $m_{f_r}^{i,j}$ , which models a  $\tau_1$  stances in  $\mathcal{D}$  to construct symmetric matrices M complete of evidence of the similarity between  $(\boldsymbol{x}_i, y_i)$  and  $\tau_2$  prising the similarities  $(M[i, j] = M[j, i] = m_{\mathcal{F}_r}^{i,j}(\{s\}))$  51

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<sup>1</sup> between them. Thereafter, the obtained matrix is ap-<sup>2</sup> plied for further unsupervised data mining analysis, such

<sup>3</sup> as clustering or data visualization.

### **EXPERIMENTS AND RESULTS** ₄ III.

In this section, we perform three experiments to 5 6 demonstrate the application of our similarity measure-7 ment in dealing with outliers and data generated by mul-<sup>8</sup> tiple mechanisms when designing materials descriptors. • We apply the eRSM to measure similarities between mag-10 netic of three datasets for detecting subgroups of materials: 1) The experimentally observed Curie tempera-<sup>12</sup> ture dataset  $(\mathcal{D}_{binary})$  of binary alloys for transitioning 13 rare earth metals, 2) Dataset of calculated magnetiza-14 tion of quaternary high-entropy alloys  $(\mathcal{D}_{quaternary}^{Mag})$ , and 15 3) Dataset of calculated Curie temperature of quaternary 16 high-entropy alloys  $(\mathcal{D}_{quaternary}^{T_C})$ . Note that the datasets 17  $\mathcal{D}_{quaternary}^{Mag}$  and  $\mathcal{D}_{quaternary}^{T_C}$  contain similar alloys and 18 differ only in the target properties.

### Datasets Α. 19

The details of the datasets investigated in this study 20 are as follows. 21

• Binary alloys dataset  $\mathcal{D}_{binary}^{27}$ : A material dataset 22 containing 100 transition-rare earth metal binary 23 alloys, comprising nickel (Ni), manganese (Mn), 24 cobalt (Co), or iron (Fe), and the corresponding 25 Curie temperatures  $(T_C)$ . This dataset was col-26 lected from the Atomwork database of the National 27 Institute of Materials Science<sup>28,29</sup>. Each binary al-28 loy in  $\mathcal{D}_{binary}$  is represented using seven descrip-29 tors: (1,2) the atomic number of transition metal 30  $(Z_T)$  and rare-earth  $(Z_R)$  constituents; (3) projec-31 tion of the spin magnetic moment onto the total 32 angular moment of the 4f elections  $(J_{4f}(1-g_j));$ 33 (4, 5) covalent radius  $(r_{covT})$  and first ionization 34  $(IP_T)$  of the transition metal; (6, 7) concentration 35 of the transition metal  $(C_T)$  and rare-earth metal 36  $(C_R)$ . The selection of these seven descriptors has 37 been discussed in detail in previous studies<sup>10,30</sup>. 38

• Quaternary high-entropy alloys 39  $\mathcal{D}_{quaternary}^{27}$ : 40 990 equiatomic quaternary high-entropy alloys, 75 41 42 43 44 45 46 47 48 49  $\mathcal{D}_{quaternary}$  is represented using 135 compositional st the model, the other Co-based alloys are often overesti-50



FIG. 2. (a) Observed and predicted Curie temperature of alloys in the dataset  $\mathcal{D}_{binary}$  using model generated for nickel (Ni), iron (Fe), and manganese (Mn)-based alloys. The blue and gray points indicate cobalt (Co)-based alloys and alloys of other transition metals (Ni, Fe, Mn), respectively. (b) Prediction error of Co-based alloys when excluding (top) or including (bottom) data of other Co-based alloys to the training dataset.

descriptors, including the means, standard deviations, and covariance of the atomic representations of their constituent elements<sup>13</sup> and four categorical features indicating the elements comprising the quaternary alloy. The feature selection process applied to this dataset has been discussed in detail in Supplementary Section III.

## 58 B. Assessment of the similarity between transition-rare earth 59 metal binary alloys based on mechanisms of Curie temperature

In the first experiment, we show the versatility of the 60 eRSM for detecting outliers and identifying a mixture of 61 mechanisms. We apply the eRSM to assess the similari-62 ties between 100 transition rare earth metal binary alloys 63 comprising nickel (Ni), manganese (Mn), cobalt (Co), or 64 iron (Fe) in the dataset  $\mathcal{D}_{binary}$  based on their Curie tem-65 peratures. We can construct a regression model using a 66 67 Gaussian process by considering the data instances in  $\mathcal{D}_{binary}$ . This shows a high prediction accuracy with an 68 69  $R^2$  score of 0.963 and an MAE of 40 (K) in ten-fold 70 cross-validation. However, such a nonparametric regres-71 sion model does not guarantee the reliability of the model 72 in the subsequent exploratory predictions. This is bedatasets 73 cause the number of observable alloys is relatively small A material dataset contains 74 compared to the number of possible alloys.

Figure 2 (a) shows the results of the exploratory predicwhich comprise 14 transition metals {Ag, Cd, Co, 76 tion of the Curie temperature of the Co-based binary al-Cr, Cu, Fe, Mn, Mo, Ni, Pd, Rh, Ru, Tc, Zn, 77 loys in  $\mathcal{D}_{binary}$  using a Gaussian process regression model and the corresponding calculated magnetizations 78 constructed from the data of binary alloys of Ni, Mn, and and Curie temperatures in the BCC phase. The 79 Fe. The regression model constructed from the data of dataset was collected from an original dataset so binary alloys of Ni, Mn, and Fe shows a high predicof 147,630 equiatomic quaternary high-entropy  $a_1$  tion accuracy in ten-fold cross-validation ( $R^2 = 0.946$ alloys calculated using Korringa-Kohn-Rostoker  $_{32}$  and MAE= 35 (K)). Although the Co-based alloys with coherent approximation method<sup>31</sup>. Each alloy in as high Curie temperature tend to be underestimated by



FIG. 3. (a) Heatmap illustrating the similarity matrix  $M_{binary}$  extracted for all the data instances in the  $\mathcal{D}_{binary}$ . (b) Confusion matrices measuring the regression-based similarities between alloys in four groups  $G_1$ - $G_4$  and the dissimilarities between the models generated for alloys in different groups.



FIG. 4. Dependence of  $T_C$  on the concentration of the transition metal  $(C_T)$  in alloys. Red, blue, green and yellow scatters indicate alloys containing cobalt (Co), iron (Fe), manganese (Mn), and nickel (Ni). Alloys in  $G_1$  are highlighted by triangles.

7 to clarify the mixture mechanism from this dataset.

By applying the eRSM on the dataset  $\mathcal{D}_{binary}$ , we ob-• tain a similarity matrix  $M_{binary}$  with moderately high similarity values among the data instances (Fig. 3 a). 10 Thus, approximately all the data instances can be re-11 gressed by a relatively smooth function. This is consis-12 tent with the high prediction accuracy of ten-fold cross-13 validation for all the alloys in the dataset. Considering 14 the exploratory data analysis, to avoid false intuition or <sup>16</sup> misunderstanding, the grouping of alloys in  $\mathcal{D}_{binary}$  is 17 done such that the similarities between the alloys in each <sup>18</sup> group are high. Moreover, one alloy can belong to more <sup>19</sup> than one group simultaneously, or it can be in none of the groups. We apply a graph-based clustering method<sup>32</sup> to 20 the extracted similarity matrix to detect overlapping sub-21 groups of materials. As a result, we observe four groups 22 of alloys, denoted as  $G_1$ ,  $G_2$ ,  $G_3$ , and  $G_4$ , which show 23 <sup>24</sup> high intragroup similarities, exceeding 0.7 (Fig. 3 a). <sup>25</sup> Nevertheless, the similarity between the alloys in group 26  $G_1$  and those in  $G_2$ ,  $G_3$ , and  $G_4$  is significantly dissimilar. In addition, a small group of alloys (Fig. 3 a, gray 27 <sup>28</sup> region) is approximately different from all the others and 29 can be considered as outliers. The remaining alloys are 30 not assigned to any group to have confidence in the clus-<sup>31</sup> tering analysis results.

To evaluate the validity of the analysis process quan-32 1 mated. The prediction error for the Co-based alloys is  $_{33}$  titatively, we trained the regression models for  $T_C$  us-<sup>2</sup> critically reduced when some data of the other Co-based  $_{34}$  ing data from each of the four groups  $G_1, G_2, G_3$ , and  $_3$  alloys is included (Fig. 2 b). This observation supports  $_{35}$   $G_4$ . Moreover, we monitored their prediction accuracy 4 the hypothesis that the underlying mechanisms are dif- 36 on these groups. The confusion matrix summarizing the <sup>5</sup> ferent between the Co-based alloys and alloys of other  $_{37}$  correlation between the observed and predicted  $T_C$  by <sup>6</sup> transition metals. This facilitates the use of the eRSM <sup>38</sup> the four learned regression models is shown in Fig. 4.



FIG. 5. (a,d) Heatmaps illustrating the similarity matrices  $M_{quaternary}^{Mag}$  (a) and  $M_{quaternary}^{T_C}$  (d) extracted from datasets  $\mathcal{D}_{quaternary}^{Mag}$  and  $\mathcal{D}_{quaternary}^{T_C}$ , focusing on mechanisms of magnetization and  $T_C$ , respectively. (b,e) The confusion matrix summarizes the differences between the magnetization (b) or  $T_C$  (e) mechanisms of alloys in extracted groups. (c,f) Visualization of quaternary alloys in the two-dimensional embedding spaces constructed by applying the T-distributed Stochastic Neighbor Embedding (t-SNE) to  $M_{quaternary}^{Mag}$  (c) and  $M_{quaternary}^{T_C}$  (f). Red, blue, and gray contours indicate gaussian models  $\hat{G}_1^{Mag}$  $(\hat{G}_1^{T_C}), \hat{G}_2^{Mag}$   $(\hat{G}_2^{T_C})$ , and  $\hat{G}_3^{Mag}$   $(\hat{G}_3^{T_C})$ , respectively, learned by using the Gaussian Mixture Models<sup>33</sup> in the embedding space focusing on mechanisms of magnetization  $(T_C)$ . In addition, red and blue points in sub-figures b and c (e and f) indicate the alloys in  $G_1^{Mag}$   $(G_1^{T_C})$  and  $G_2^{Mag}$   $(G_2^{T_C})$ , respectively.

<sup>1</sup> The diagonal plots illustrate the cross-validation results <sup>18</sup> tion coefficient of 0.95 (Fig. 4, triangle scatters). This <sup>2</sup> of the models learned from the four groups of alloys. The <sup>19</sup> result is consistent with the observation of the previous <sup>3</sup> off-diagonal plot shows the correlation between the ob-<sup>20</sup> research<sup>30</sup>, when considering all binary alloys of transi-4 served  $T_C$  and the predictions made by the model learned 21 tion metals and rare earth metals in  $\mathcal{D}_{binary}$ ; the range  $\tau G_1, G_2, G_3$ , and  $G_4$ , respectively, dissimilarity between 24 alloys in  $G_1$  are Co-based alloys with high Curie tems the five groups, and intra-group dissimilarity of the alloys  $_{25}$  peratures ( $T_C > 600K$ ). By contrast, most of the other  $\circ$  considered as outliers. This indicates that the obtained  $_{26}$  Co-based alloys in  $\mathcal{D}_{binary}$  have lower Curie temperatures <sup>10</sup> results suggest that the physical mechanisms of alloys in <sup>27</sup>  $(T_C < 500K)$  and are assigned to  $G_2$ ,  $G_3$ , and  $G_4$ . These 11  $G_1$  may be different from those of the alloys in  $G_2$ ,  $G_3$ , 28 results are consistent with the observation that the re- $_{12}$  and  $G_4$ . Nonetheless, it is difficult to determine the dif- $_{29}$  gression model for Fe-, Mn-, and Ni-based alloys tends to 13 ferences between the mechanisms of the  $T_C$  of alloys in 30 underestimate the  $T_C$  of the Co-based alloys with high 14  $G_2, G_3, \text{ and } G_4$ .

Moreover, considering the alloys in  $G_1$ , there is a strong 32 alloys (Fig. 2 a). 16 linear correlation between  $T_C$  and the concentration of 33 In addition, we examine the behavior of eRSM on

 $_{5}$  from the alloys of the other groups. The obtained results  $_{22}$  of  $T_{C}$  is found to be correlated with the composition ra-6 confirm the intragroup similarity of the alloys in groups 23 tio of the transition metals. Furthermore, 13 of the 17  $_{31}$   $T_C$  and overestimates the  $T_C$  of the remaining Co-based

17 transition metals in the alloys with a Pearson correla- 34 toy datasets synthesized with outliers or multiple mech-



FIG. 6. Prediction accuracies for magnetization (a, b) and Curie temperature (c, d) of the alloys with 10-fold crossvalidations. Prediction validation results with single gaussian process regression models for magnetization and Curie temperature are shown in sub-figures (a) and (c), respectively. Prediction validation results with mixtures of expert models for magnetization and Curie temperature are shown in subfigures (b) and (d), respectively. Blue and white circles indicate magnetic alloys (finite magnetization) and non-magnetic alloys (zero magnetization), respectively.

1 anisms to assess the efficiency of this similarity measure. <sup>2</sup> Detailed results of these experiments are summarized in <sup>3</sup> the Supplementary Section II. Briefly, the eRSM demon-<sup>4</sup> strates that it can effectively assess the similarity between 5 the data instances and use the similarity for detecting 6 outliers and a mixture of mechanisms.

## 7 C. Assessment of the similarity between guaternary 8 high-entropy alloys based on mechanisms of magnetization

The effectiveness of the eRSM in detecting outliers and 9 10 identifying mixture mechanisms in the material dataset has been shown in the previous experiment. In the next 11 two experiments, we show the potential of applying the 12 measured similarity to design descriptors for materials. 13

Considering this experiment, we subsequently apply 14 the eRSM to assess the similarities between 990 quater-15 16 17 18 19 attempted to construct an optimal Gaussian process re-76 the alloys in groups  $G_1^{Mag}$  (red) and  $G_2^{Mag}$  (blue) when 20 gression model using the designed descriptors. The Gaus- 77 they form two separate regions with high density in the

<sup>21</sup> sian process can poorly regress the magnetization with an <sup>22</sup>  $R^2$  score of 0.75 and an MAE of 0.13 (T) in the ten-fold 23 cross-validation. The obtained results suggest that the <sup>24</sup> magnetization of these alloys may not be described by a <sup>25</sup> single model in the designed descriptor space. This in-26 dicates that the existence of outliers or mixture models 27 of the magnetization properties of these alloys in the de-28 scriptor space should be considered in the analysis of this <sup>29</sup> dataset.

Applying the eRSM, we obtain a similarity matrix <sup>14</sup>  $M_{quaternary}^{Mag}$  with two core groups of alloys denoted by <sup>25</sup>  $G_1^{Mag}$  and  $G_2^{Mag}$ , showing high intra-group similarities <sup>33</sup> and exceeding 0.5 (Fig. 5 a). Some of the alloys in  $G_1^{Mag}$ <sup>34</sup> are similar to those in  $G_{2_{-}}^{Mag}$ ; nonetheless, the rest show <sup>35</sup> apparent dissimilarities. Furthermore, one small group of alloys (Fig. 5 a, yellow region) showed dissimilari-36 37 ties with the others and could be considered as outliers. The remaining alloys in  $\mathcal{D}_{quaternary}^{Mag}$  do not exhibit apparent similarities with alloys in groups  $G_1^{Mag}$  and  $G_2^{Mag}$ . Therefore, they are not assigned to any group. 40

41 Similar to the previous session, to validate the obtained 42 results quantitatively, we trained three regression mod-43 els using data from each group,  $G_1^{Mag}$ ,  $G_2^{Mag}$ , and out-44 liers. We monitored the prediction accuracy of the three <sup>45</sup> learned regression models for data in all the groups. The confusion matrix summarizing the correlations between 46 47 the observed and predicted values of the target variable <sup>48</sup> using the learned regression models is shown in Fig. 5 (c). <sup>49</sup> The diagonal plots illustrate the *ten*-fold cross-validation <sup>50</sup> results of the models learned from these three groups of <sup>51</sup> alloys. The off-diagonal plot shows the correlation be-52 tween the observed magnetization and the predictions <sup>53</sup> made by the model learned from the alloys of the other 54 groups.

The obtained results confirm the intragroup similarity of the alloys in groups  $G_1^{Mag}$  and  $G_2^{Mag}$ , respectively, the 55 <sup>57</sup> dissimilarity between the two groups, and the intra-group 58 dissimilarity of the alloys considered as outliers. Specif-59 ically, we observe that group  $G_2^{Mag}$  consists of ferrimag-<sup>60</sup> netic alloys or alloys whose magnetization is relatively  $_{61}$  smaller (magnetization < 0.1 (T)) than the others in the <sup>62</sup> group  $G_1^{Mag}$ . In contrast, using the data in  $G_1^{Mag}$ , we <sup>63</sup> can construct a Gaussian process regression model with a high prediction accuracy with an  $R^2$  score of 0.992 and an MAE of 0.016 (T) in the ten-fold cross-validation. 65

66 Therefore, we can use the information of the con-67 stituent elements of each alloy to predict which group it belongs to in advance<sup>20</sup> and apply an appropriate regres-68 <sup>69</sup> sion model to improve prediction accuracy for the alloys. We combine the similarity measured by using the eRSM 70 71 with the Jaccard similarity coefficient<sup>34</sup> and apply the T-<sup>72</sup> distributed Stochastic Neighbor Embedding<sup>35</sup> (t-SNE) to nary high-entropy alloys comprising 14 transition met- 73 construct a two-dimensional embedding map (Fig. 5 c). als in the dataset  $\mathcal{D}_{quaternary}^{Mag}$  based on their magnetiza- 74 Details of the combination method are shown in Suppletion. To predict the magnetization of these alloys, we 75 mentary Section IV. As a result, we can easily distinguish

<sup>1</sup> embedding space. We apply a Gaussian mixture model<sup>33</sup> <sup>2</sup> (GMM) on the embedding space to identify groups and <sup>3</sup> calculate the probability of an alloy belonging to a par-4 ticular identified group. Alloys in different groups are <sup>5</sup> treated differently by using a mixture of experts<sup>36</sup> (MoE) α approach. Figure 6 (a-b) show a reduction of the pro-7 posed mixture of experts in MAE of 18% compared with **s** result of the single model, from 0.13 (T) to 0.11 (T). Fur-• ther analysis shows that applying the obtained similari-10 ties in MOE improves the prediction accuracy for mag-<sup>11</sup> netic alloys (Supplementary figure 7 a).

## 12 D. Assessment of the similarity between the quaternary 13 high-entropy alloys based on mechanisms of Curie 14 temperature

Considering this experiment, the target data are the 15 16 same as in the previous section  $(\mathcal{D}_{quaternary})$ ; however, 17 the physical property of interest is  $T_C$ . Å regression <sup>18</sup> model can be constructed using a Gaussian process. This 19 shows a rather high prediction accuracy in *ten*-fold cross-<sup>20</sup> validation with an  $R^2$  score of 0.85 and an MAE of 67  $_{21}$  (K). We also observe two distinguishable groups of qua-<sup>22</sup> ternary alloys in the dataset  $\mathcal{D}_{quaternary}^{T_C}$  when applying <sup>30</sup> tiveness of the data for detecting the mixture of multi-<sup>31</sup> the eRSM. Figure 5 (d) illustrates the similarity matrix <sup>32</sup>  $M_{quaternary}^{T_C}$  with two groups of alloys denoted as  $G_1^{T_C}$  <sup>33</sup> ods as in the previous experiment to construct a two-dimensional embedding man (Fig. 5 f) and then become <sup>25</sup> and  $G_2^{T_C}$ , showing high intra-group similarities and ex-26 ceeding 0.5. Some of the alloys in  $G_1^{T_C}$  are similar to  ${\scriptstyle {\bf 27}}$  those in  $G_2^{T_C}.$  Nonetheless, the others exhibit apparent 28 dissimilarities, which is consistent with the observation <sup>29</sup> of two high-density regions (red) in the embedding map 30 of  $M_{quaternary}^{T_C}$  (Fig. 5 e). Furthermore, a small group 31 of alloys (Fig. 5 d, yellow region) showed dissimilarities 32 with all the others and could be considered as outliers. The remaining alloys do not show apparent similarities with alloys in groups  $G_1^{T_C}$  and  $G_2^{T_C}$ ; thus, they are not 35 assigned to any group.

Following the same analysis procedure as in the previ-36 37 ous section, we trained regression models for Curie tem-<sup>47</sup> groups. We can also confirm the intra-group similarity of <sup>82</sup> underlying patterns. <sup>48</sup> the alloys in groups  $G_1^{T_C}$  and  $G_2^{T_C}$ , respectively, dissim-<sup>83</sup> Figure 7 shows that Fe and Co, which have a large spin 49



FIG. 7. Proportions of quaternary alloys containing Fe or Co in group  $G_1^{\overline{M}ag}$  (a) and  $\overline{G}_1^{T_C}$  (b).

<sup>55</sup> can construct a Gaussian process regression model with <sup>56</sup> a high prediction accuracy with an  $R^2$  score of 0.985 and an MAE of 19 (K) in the *ten*-fold cross-validation. 57

58 Therefore, we utilize the similarity information to de-<sup>59</sup> sign descriptors for quaternary alloys due to the effec-63 dimensional embedding map (Fig. 5 f) and then learn a 64 mixture of experts to predict Curie temperature of qua-<sup>65</sup> ternary alloys in the dataset  $\mathcal{D}_{quaternary}^{T_C}$ . The proposed <sup>66</sup> mixture of models exhibits higher prediction accuracy  $_{67}$  than the single model in 10-folds cross-validations (Fig. 68 6 c-d). The MAE of the proposed mixture of expert re-69 duces approximately 36%, from 67 (K) to 49 (K).

## 70 E. Discussion of the obtained similarities between materials 71 and the associated physical mechanisms

Regarding the experiments with the datasets 72 <sup>38</sup> perature using data from each of the three groups  $G_1^{T_C}$ , <sup>79</sup>  $\mathcal{D}_{quaternary}^{Mag}$  and  $\mathcal{D}_{quaternary}^{T_C}$  focusing on magnetiza-<sup>39</sup>  $G_2^{T_C}$ , and outliers and monitored their prediction accu-<sup>74</sup> tion or  $T_C$ , the datasets seem to be a self-evident 40 racy on these groups. Figure 5 (f) shows the confusion  $\tau_5$  example where magnetization and  $T_C$  are cases sensitive 41 matrix that summarizes the obtained results. The diag- 76 to finite or zero. As we can see from the results described 42 onal plots illustrate the ten-fold cross-validation results 77 above (Sections III C, III D, and Supplementary Section 43 of the models learned from these three groups of alloys. 78 VI), the prediction accuracy is low when considering a 44 The off-diagonal plot shows the correlation between the 79 single regression model for the entire dataset. In this 45 observed Curie temperature and the predictions made by section, we pay attention to the analysis of the extracted the regression model learned from the alloys of the other  $a_1$  alloys groups  $G_1^{Mag}$ ,  $G_2^{Mag}$ ,  $G_1^{T_C}$ , and  $G_2^{T_C}$  to identify

ilarity between the two groups, and intra-group dissimi- e4 moment, ferromagnetic interactions with many elements <sup>52</sup> harty between the two groups, and indra-group dissimi-<sup>53</sup> larity of the alloys considered as outliers. <sup>54</sup> Specifically, we observe that the Curie temperatures <sup>55</sup> of approximately all the alloys in group  $G_2^{T_C}$  have a low <sup>56</sup> elements comprising alloys in two groups  $G_1^{Mag}$  (a) and <sup>57</sup>  $G_1^{T_C}$ , which is 0 (K) or relatively smaller than that of the <sup>58</sup> other alloys. Furthermore, using the data in  $G_1^{T_C}$ , we <sup>59</sup> four constituent elements concerning the extracted four



FIG. 8. Effect of coexistence of the 14 transition metals on magnetization and Curie temperature mechanisms. Each pie chart results from quaternary alloys containing the respective element pair. They show the percentages of alloys that follow the magnetization mechanisms (lower-left triangle) and Curie temperature mechanisms (upper-right triangle), as extracted by the eRSM. Red and blue areas indicate the percentages of alloys whose magnetization and  $T_C$  are finite  $(G_1^{Mag} \text{ and } G_1^{T_C})$  and zero  $(G_2^{Mag} \text{ and } G_2^{T_C})$ , respectively. Yellow areas indicate the percentages of alloys that are detected as outliers. By contrast, gray regions indicate the fractions of alloys not assigned to the extracted groups.

1 groups  $G_1^{Mag}$ ,  $G_2^{Mag}$ ,  $G_1^{T_C}$ , and  $G_2^{T_C}$ , we observe that the 2 proportion of Fe-containing and Co-containing alloys in 3 two groups  $G_1^{Mag}$  (a) and  $G_1^{T_C}$  are significantly larger 4 than other groups (Fig. 8). Thus, the prediction mod-5 two groups  $G_1^{Mag}$  (b) and  $G_1^{T_C}$  are significantly larger 5 two groups  $G_1^{Mag}$  (c) and  $G_1^{T_C}$  are significantly larger 6 than other groups (Fig. 8). Thus, the prediction mod-7 the prediction mod-9 the pre

s els constructed from the data of the alloys in  $G_1^{Mag}$  or  $G_1^{T_C}$  are more suitable to predict magnetization or  $T_C$ , <sup>11</sup> which are not assigned to any of these mechanisms. Con-<sup>12</sup> versely, Mn-X alloys exhibit similar behavior as Fe-X and

10



FIG. 9. Correlation between magnetization (T) and Curie temperature (K) of quaternary alloys with non-zero magnetization and non-zero Curie temperature in datasets  $\mathcal{D}_{quaternary}^{Mag}$  and  $\mathcal{D}_{quaternary}^{T_C}$ . Marginal plots show histogram of the properties of the alloys.

 $_{3}$  categorized in the group  $G_{2}^{\hat{T}_{C}}$  of low  $T_{C}$  besides the other <sup>4</sup> groups. Especially among the Fe-X and Co-X alloys, the <sup>5</sup> percentage of Fe-Mn and Co-Mn alloys are considered as • outliers of the mechanisms extracted from  $G_1^{T_C}$  are rela-• tively higher, 55% and 43%, respectively (Fig. 8).

8 <sup>10</sup> absence of Mn. Figure 9 shows the correlation between <sup>65</sup> analysis of the detected subgroups improves the existing 11 magnetization and Curie temperature of 556 (56%) al- 66 knowledge of problems related to the applied datasets of 12 13 14 both  $T_C$  and magnetization are zero is 413 (42%), while 69 based binary alloys when using our method to a dataset 15 16 17 20 ate Curie temperatures. By contrast, the Mn-containing 75 terials with uncertainty, the method described herein is 21 alloys without Fe or Co have low Curie temperatures 76 expected to extract valuable information for describing 22 23 alloy groups do not offer any significant correlation be- 78 material datasets. tween magnetization and Curie temperature. However, 24 an apparent positive correlation between magnetization 25 and Curie temperature can be observed for the group of 79 SUPPLEMENTARY MATERIAL 26 Mn-free alloys. 27

28 29

31 netic moments divided by the unit volume. The local 32 magnetic moments are determined by the spin configura-<sup>33</sup> tions of atomic sites that stabilize the structure of alloys. Conversely, given a particular structure and spin con-34  $_{35}$  figuration, the  $T_C$  can be estimated from the spin-spin 36 exchange energy. First-principles calculations show that 37 early transition metals and late transition metals often have antiferromagnetic interactions $^{37}$ . This interaction 38 39 has also been confirmed in high-entropy alloys by using 40 automatic exhaustive calculations<sup>31</sup>. Mn lies between early and late transition metals; thus, the estimation 41 42 of the spin configuration (ferromagnetic or antiferromag-43 netic) in Mn-containing alloys should be cautiously considered in different situations, especially in high-entropy 44 alloys whose elements can stochastically exist at the same 45 atomic site. From this consideration, we can admit a hy-46 47 pothesis that the alloys containing Mn follow a different <sup>48</sup> rule for magnetization than those grouped into  $G_2^{Mag}$ . 49 Conversely, the alloys containing Mn may follow the same 50 rules for  $T_C$  as the alloys grouped into  $G_2^{T_C}$ , albeit with 51 a spin configuration that provides magnetization. The 52 details are beyond the scope of this paper and will not <sup>53</sup> be discussed here, but further analysis is promising.

### CONCLUSIONS 54 IV.

In this study, we developed a method that can be 55 1 Co-X when focusing on the magnetization mechanisms. 56 used to rationally transform material data from multi-<sup>2</sup> However, for the Curie temperature, the Mn-X alloys are <sup>57</sup> ple sources into evidence of similarities between materi-<sup>58</sup> als and combine the evidence to conclude the similarities 59 between materials. The extracted similarity-dissimilarity 60 information has significant potential for application in 61 subgroups discovery of materials. The effectiveness of 62 the eRSM in detecting homogenous subgroups of materi-For further investigation, we organized the raw data • als has been demonstrated by using two experiments on o of the quaternary alloys by focusing on the presence or 64 two datasets of magnetic materials. In addition, further loys with non-zero properties. The total number of data or magnetic materials. For example, we reveal the differinstances is 990, and the number of data instances where ences in the mechanisms of the Curie temperature of Cothere are twenty-one (2%) alloys with zero  $T_C$  but have  $\tau_0$  of 100 transition-rare earth metal binary alloys comprisfinite magnetization. We found that the alloys contain- 71 ing Ni, Mn, Co, and Fe. Moreover, we explored the ing all three elements, Mn, Fe, and Co, show high Curie 72 mechanisms of ferrimagnetic and low Curie temperature temperatures  $(T_C > 900 \ (K))$ . Conversely, the alloys 73 alloys from the magnetic dataset of calculated quatercontaining either pairs of Mn-Fe or Mn-Co show moder- 74 nary alloys. By measuring the similarity between ma- $(T_C < 250 \ (K))$ . Furthermore, the trends of these three  $\pi$  and interpreting the underlying physical mechanisms in

To interpret the results obtained, we considered a hy- so See supplementary materials for the following addipothesis of the origin of the observed data. The esti- at tional information: 1) Explanation of the formulation 30 mated magnetization is the sum of all the local mag- 32 modeling uncertainty, 2) Evaluation of the eRSM us-

1 ing the toy datasets, and 3) Features selection and pre- 57 <sup>12</sup>J. Bardeen, L. N. Cooper, and J. R. Schrieffer, "Theory of su-<sup>2</sup> analysis in the dataset of quaternary high-entropy alloys.

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## 12 DATA AVAILABILITY STATEMENT

Datasets related to this article are deposited to Zenodo 13 14 repositor  $v^{27}$ .

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