


Cite this: *RSC Adv.*, 2025, 15, 19126

Ternary materials discovery using human-in-the-loop generative machine learning†

Brandon Wilfong,^{ID} ^{ab} Alexander New,^{ID} ^c Gregory Bassen,^{ab} Wyatt Bustine,^b Michael J. Pekala,^c Nam Q. Le,^c Karun K. Rao,^c Elizabeth A. Pogue,^c Eddie Gienger,^c Michał J. Winiarski,^{ID} ^d Tyrel M. McQueen^{ID} ^{*abe} and Christopher D. Stiles^{*c}

Machine learning (ML) approaches to materials discovery are limited by data curation, availability, and bias. These issues can be addressed through the generation of new data points representing novel material compositions and/or structures. We demonstrate the implementation of this process to produce and subsequently determine the stability of novel materials using a generative ML model. Furthermore, we successfully synthesize two predicted materials, LiZn_2Pt and NiPt_2Ga , and use these predictions to extrapolate to other unreported ternary compounds in the Heusler family. Our work demonstrates and expands the use of generative ML models to successfully discover and synthesize novel materials. This has broad implications for material exploration by design, as previous ML approaches to materials discovery were biased by the limits of known phase spaces and experimentalist bias, and has the potential to enable inverse-design of materials with targeted properties.

Received 17th January 2025

Accepted 12th May 2025

DOI: 10.1039/d5ra00427f

rsc.li/rsc-advances

Recent advances in machine learning (ML) techniques for domain-specific learning tasks have led to an explosive adoption of data-driven scientific exploration in physics and chemistry.^{1–3} This is particularly true in materials development where the combinatorics of composition, stoichiometry, and

structure yield too many possible materials for the community to synthesize and characterize. As such, researchers have been applying ML techniques to this problem in attempts to predict novel materials with targeted properties, such as improved batteries, solar cells, semiconductors, and superconductors.^{4–8} One outstanding issue with the success of ML techniques in material science applications is the lack of high-quality data that is curated, easily ingestible, complete, and high fidelity.⁹ To address this, the coordination of materials databases have allowed the implementation of ML workflows into material science research.^{10–16} Although these databases contain hundreds of thousands of known and theoretical materials with corresponding structures and properties, the corpus of data is heavily biased toward explored phase spaces and realistically these databases contain only a small fraction of possible materials and their known properties.

To increase the success of ML-driven approaches, methods to improve materials data have been implemented to address two main issues: data completeness and data bias. One approach to address both issues simultaneously is to iteratively loop ML predictions with experimental validation thereby allowing the ML model predictions to continually evolve with the incorporation of new negative and positive data to the ML training sets.^{8,17,18} This “closed loop” treatment has recently been successful in the prediction and validation of new super-conductors and could be extrapolated to other property prediction problems.⁸ Although this approach does not require initial property labels for all materials of interest, the unlabeled, and therefore potentially interesting, material candidates must still come from the initial datasets. This restricts data de-biasing.

^aDepartment of Chemistry, Johns Hopkins University, 3400 N. Charles Street, Baltimore, 21218, Maryland, USA

^bInstitute for Quantum Matter, William H. Miller III Department of Physics and Astronomy, Johns Hopkins University, 3400 N. Charles Street, Baltimore, 21218, Maryland, USA. E-mail: mcqueen@jhu.edu

^cResearch and Exploratory Development Department, Johns Hopkins University Applied Physics Laboratory, 11100 Johns Hopkins Road, Laurel, 20723, Maryland, USA. E-mail: Christopher.Stiles@jhuapl.edu

^dFaculty of Applied Physics and Mathematics and Advanced Materials Center, Gdansk University of Technology, Narutowicza 11/12, 80-233 Gdansk, Poland

^eDepartment of Materials Science and Engineering, Johns Hopkins University, 3400 N. Charles Street, Baltimore, 21218, Maryland, USA

† Electronic supplementary information (ESI) available: All experimental, computational, and machine learning methods used in this work. Additional X-ray diffraction analysis of the LiZn_2Pt and LiZn_2Pd phases are provided in the context of order and disorder in the Heusler structure. X-ray diffraction and Rietveld refinement of as-synthesized NiPt_2Ga is presented. Microscopic analysis is performed using scanning electron microscopy (SEM) images and elemental analysis via energy dispersive spectroscopy (EDS) of as-recovered LiZn_2Pt , LiZn_2Pd , and NiPt_2Ga phases. Magnetization measurements for all three samples are presented with LiZn_2Pt and LiZn_2Pd measured down to 0.4 K in an attempt to observe superconductivity. Isothermal magnetization at various temperatures was measured for NiPt_2Ga to confirm the antiferromagnetic character of the sample up to 300 K. The representative crystal structures of the top 6 (excluding LiZn_2Pt) candidates from the PGCM model are shown. Band structures and DOS computations for LiZn_2Pt and LiZn_2Pd in the Cu_2MnAl -type Heusler structure and NiPt_2Ga in the $P4/mmm$ tetragonal Heusler are presented. See DOI: <https://doi.org/10.1039/d5ra00427f>



Therefore, a complementary strategy to expand the number of available materials, beyond what is available in current repositories, is the use of ML models or *ab initio* methods to generate novel materials and structures.^{19–27} These generative approaches can be paired with more traditional property-prediction ML models to target materials for specific design problems. To date, developing out-of-distribution material candidates with generative models and predicting properties with ML have been done independently.^{28–34} In this paper, we present the first successful use of a generative ML model to produce novel materials and structures, determine their thermodynamic stability, and subsequently experimentally synthesize and characterize these new materials in a single human-in-the-loop workflow. This study builds off of our prior work,³⁵ in which we analyzed the capabilities of the generative model we used in this paper and assessed. Here, we extend those results by computationally and experimentally characterizing specific generated materials of interest.

In particular, we use a state-of-the-art generative model, the PGCGM for the generation and prediction of novel materials.²³ The PGCGM is a Wasserstein GAN that can stochastically sample possible structures of ternary material systems given their constituent elements and space group.³⁶ In this study, we use the as-released PGCGM which was trained on data taken from Materials Project (MP), Open Quantum Materials Database (OQMD), and inorganic crystal structure database (ICSD). We randomly sample constituent element sets and space groups and then use the PGCGM to generate 27 116 material structures Fig. 1. As discussed in the PGCGM paper,²³ following generation, these structures were post-processed to merge together spatially-adjacent atoms of the same type to one crystallographic site.

Methods for theoretically assessing the range of possible structures a generative model can predict are limited. Such models can predict novel structures, but their predictions may lack diversity compared to their training data. Measuring the novelty and diversity of generated materials, especially in comparison to training data, remains a needed step in their use (see, *e.g.*, ref. 23 and 26). Averting pathologies such as mode collapse remains an active area of research.^{33,34,37} However, because the throughput of these methods is far higher than first-principles calculations or experimental synthesis and characterization, use of generative models for predicting structures allows for a faster sampling of unknown phase spaces.

Although the PGCGM is capable of generating large numbers of potential structures, there is no guarantee that the structures will be thermodynamically stable and thus synthesizable. Therefore, we also train a stability-prediction ML model to rapidly screen identified structures for stability (Fig. 1). Specifically, we construct a set of structures from MP with computationally-predicted decomposition enthalpy, previously identified by Bartel *et al.*³⁸ We use this data to train an ALIGNN model³⁹ to predict decomposition enthalpy relative to the convex hull based on phases in the ML. The decomposition energy is the same as the energy above convex hull, except for phases that lie directly on the convex hull. In those cases, the decomposition energy is the distance from the hypothetical convex hull formed by other phases. A structure is predicted to be stable if its predicted decomposition enthalpy is negative.

We use decomposition enthalpy as a metric for stability because decomposition enthalpy takes positive values for unstable compounds and negative values for stable compounds. In contrast, the commonly used energy above hull

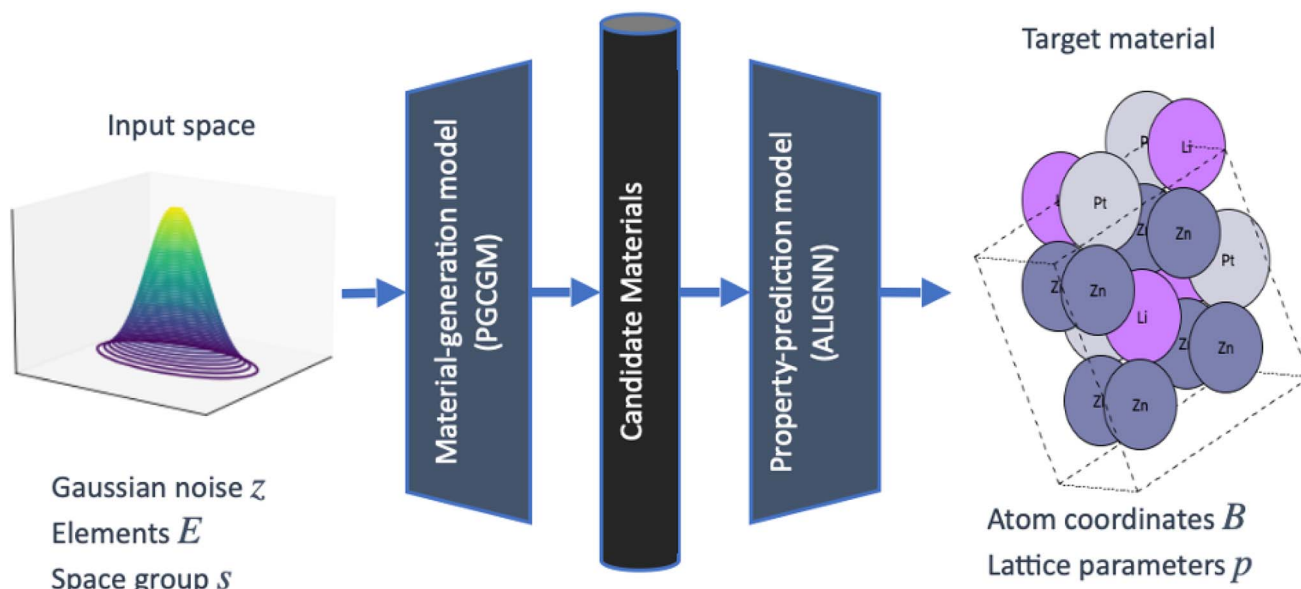


Fig. 1 The PGCGM is a GAN that maps a sample of random Gaussian noise z , three elements E , and a space group s to a ternary unit cell structure as characterized by its atom coordinates B and lattice parameters p . Varying z enables different structures with the same elements and space group to be generated. We generate 27 116 structures with the PGCGM and then use a GNN to predict the decomposition enthalpy of each structure.



takes a value of zero for all stable phases and does not indicate the degree of stability relative to competing phases. Thus, decomposition enthalpy provides more information about stability relative to other phases in the multicomponent system; its range of values also makes it more suitable for the loss functions used by regression-based ML methods. Outside of decomposition enthalpy and energy above hull, material stability can be characterized in other ways.^{40,41}

In general, this process can be adapted to include any secondary prediction model for thermodynamic stability and/or a desired property prediction such as superconductivity, elastic properties, and/or magnetism; this is the reason for including the general notation of property-prediction model in the workflow shown in Fig. 1.

After screening generated structures for stability, 2652 have a predicted decomposition enthalpy less than 0.1 eV per atom, 281 are less than 0.01 eV per atom, and 195 are less than 0 eV per atom. Domain expertise is further used to down-select materials based on factors such as oxidation states of constituent elements, coordination of atoms in the generated structure, and feasibility of synthesis. This step is currently required due to the large number of prediction candidates from the PGCGM process; future work and advances in synthesizability prediction would be required to incorporate this subject matter expertise in an automated way into the prediction process. Table 1 shows select PGCGM-generated materials and their ALIGNN-predicted decomposition enthalpies (E_d). The full list of structures and predicted properties is in the ESI.† Sampled PGCGM structures may be unique and strictly theoretical, or they may reflect the theoretical structures already listed in the training data. LiZn_2Pt with space group $Fm\bar{3}m$ is found in the subset of MP that PGCGM was trained on (mp-867251), and in the subset of OQMD PGCGM was trained on (OQMD IDs 1042055, 1042723, and 1047136) in the $P6_3/mmc$ space group. Even when the PGCGM predicts two compositions with the same stoichiometry and space group, the structures can still vary based on the precise positioning of atoms, which affects properties. The random sample z that the PGCGM takes as input enables generation of different structures with the same elements and space group.

In this work, LiZn_2Pt was selected as the most promising candidate from the initial round of material generation, energy classification, and down-selection due to its composition and

predicted structure. The selection of LiZn_2Pt was due to its second-lowest E_d and E_f amongst all candidates and that the predicted structure is a Heusler, a structure type that is ubiquitous in solid-state chemistry.^{42–44}

We considered several other material candidates as well. NaZn_2Pd and NaZn_2Pt appear in the list of PGCGM candidates (with predicted E_d values of -0.014375 eV per atom and -0.045241 eV per atom, respectively). Despite these negative predicted decomposition enthalpies, we were unable to successfully synthesize either. Thus, we also attempted to synthesize LiZn_2Pd ; this composition was targeted because there are no experimental reports of Li–Zn–Pd ternary phases, although it was not a material appearing in the PGCGM prediction list.

The X-ray powder diffraction patterns and subsequent Rietveld using the PGCGM-generated Cu_2MnAl -type Heusler ($Fm\bar{3}m$) structure are presented in ESI.† Importantly, the Heusler family has a range of related structures determined by the amount of order or disorder in the Heusler sublattices.^{43–45} Disorder in Heusler structures can be difficult to determine without synchrotron X-ray or neutron diffraction, but a qualitative approach to the X-ray diffraction offers some insight and is comprehensively shown in ESI† to justify the successful synthesis of LiZn_2Pt (and LiZn_2Pd) in the same structure as generated by the PGCGM. Overall, the successful synthesis of LiZn_2Pt was a proof-of-concept of the ability of the human-in-the-loop workflow to help discern which regions of phase space house stable phases that have not been previously experimentally realized.

Following the successful synthesis of LiZn_2Pt and LiZn_2Pd , several Heusler-type PGCGM predictions with negative E_f and E_d were targeted. To that end, a new ternary phase with stoichiometry NiPt_2Ga was successful. Fig. 2 shows the X-ray powder diffraction pattern and Rietveld refinement for NiPt_2Ga in the $P4/mmm$ tetragonal Heusler space group. Interestingly, this is

Table 1 The six PGCGM-generated structures and with the most negative (*i.e.*, stable) decomposition enthalpies (E_d in eV per atom) and NiPt_2Ga , as predicted by ALIGNN

Formula	Space group	Predicted E_d (eV per atom)
BaH_8Pt	$I4/mmm$	-0.1732166
LiZn_2Pt	$Fm\bar{3}m$	-0.1462675
HfH_{24}W	$Fm\bar{3}m$	-0.129171
Ba_3AsH_6	$R\bar{3}c$	-0.1043272
KPdF_6	$Fm\bar{3}m$	-0.0996605
RbAlS	$Immm$	-0.0987449
NiPt_2Ga	$Fm\bar{3}m$	-0.0070151

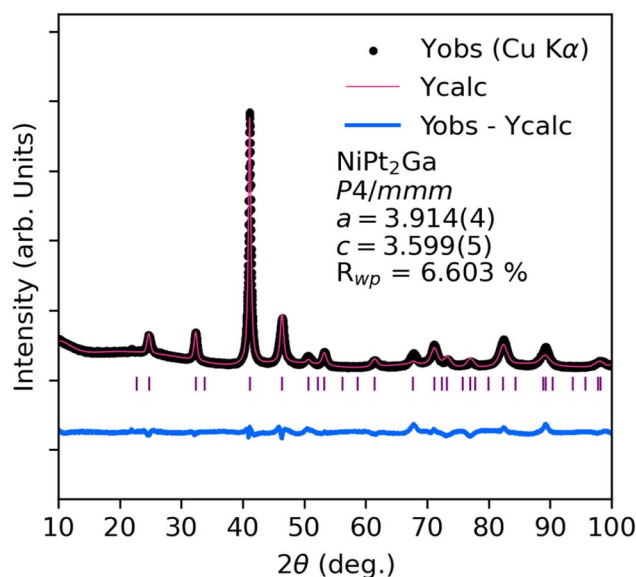


Fig. 2 Powder X-ray diffraction of phase pure NiPt_2Ga Rietveld refinement using the $P4/mmm$ tetragonal Heusler structure.



not the exact same structural prediction from the PGCGM which predicted a Cu_2MnAl -type cubic Heusler similar to LiZn_2Pt . These structures are obviously very closely related (ESI[†]), differing only by a tetragonal distortion. To confirm their proximity in energy, a comparison of DFT-computed energy for both structures was completed. The total energy for the PGCGM-predicted $Fm\bar{3}m$ structure and $P4/mmm$ differ by ≈ 0.15 eV per atom with $P4/mmm$ having lower overall energy (ESI[†]). Neither the $Fm\bar{3}m$ or $P4/mmm$ structures for NiPt_2Ga were present in the training data for the PGCGM – the $P4/mmm$ structure of NiPt_2Ga is present in OQMD (OQMD ID 1364774) but was not used to train the iteration of the PGCGM model used in this work. Thus, the successful synthesis of NiPt_2Ga takes the success of the PGCGM workflow beyond informing us to regions of phase space with unreported phases and structures that are likely to be stable, but enable us to synthesize unknown materials that exist beyond the database curation of the domain expertise.

These three materials are electronically quite unusual. They all have a valence electron count (VEC) of 33 for NiPt_2Ga and 35 electrons for $\text{LiZn}_2\text{Pt/Pd}$ – very high among known Heuslers (ESI[†]). Typically, this high VEC means that the electronic character of these Heuslers can be significantly different than the majority of Heuslers and enable access to unique regions of the band structure at the Fermi level. For example, the high VEC in LiZn_2Pt and LiZn_2Pd causes the Fermi level to sit directly at a band crossing between the L and Γ point which may be a source of unique transport properties. This band crossing is observed in Fig. 3 for NiPt_2Ga but due to its lower VEC, it is ≈ 1.5 eV above the Fermi level and possible exotic properties arising from this Fermiology is not easily accessible.

Here, we have worked with materials that, although electronically unusual, belong to the well-studied Heusler class. The

discovery of usable materials from truly novel classes is a challenge that still remains unsolved.^{26,46} Standard metrics like stable, unique, novel (SUN) assess only if a given structure is not in existing databases and do not check if the structure's materials class is novel.²⁶ As the field progresses, we recommend the use of distance-based measurements (whether using featurizations like Magpie⁴⁷ or ML model latent spaces) for determining how unlike generated materials are from known ones. This may provide additional mechanisms for going beyond current dataset limitations and mitigating dataset bias.

In general, the electronic properties of Heusler compounds are largely determined by their VEC and not their actual chemical composition, magnetism aside. For example, superconductivity in Heuslers is expected in the range of VEC of 26–29, with a peak in critical temperature (T_c) at VEC = 27.^{48,49} Measured magnetic susceptibility of NiPt_2Ga measured down to 1.8 K shows antiferromagnetic behavior which persists up to room temperature (ESI[†]) with isothermal magnetization at 300 K showing antiferromagnetic-like behavior. High temperature susceptibility is required to resolve the ordering temperature as NiPt_2Ga may be a candidate for study of metallic antiferromagnetism at room temperature. Magnetic susceptibility of LiZn_2Pt and LiZn_2Pd (ESI[†]) measured down to 0.4 K (for LiZn_2Pt and LiZn_2Pd) shows paramagnetic behavior and no anomalous magnetic transitions are revealed. Additionally, we performed a preliminary search of antiferromagnetic configurations with DFT (ESI[†]), none of which showed a significant decrease in energy from the ferromagnetic configuration. Overall, superconductivity is not observed in any of these compounds and in general, this is expected due to the high VEC outside the 26–29 range seemingly excluding superconductivity.

In conclusion, we have demonstrated the first successful use of a generative ML model to produce crystal structures, determine their stability through a secondary ML process, and experimentally verify these new structures. LiZn_2Pt was generated using the PGCGM model in the Cu_2MnAl -type Heusler structure and predicted to be stable. Subsequently, phase pure LiZn_2Pt was successfully synthesized. Due to its proximity and lack of reported ternaries LiZn_2Pd was also successfully synthesized. Following the successful synthesis of the aforementioned compounds, Heusler and Heusler-like PGCGM predicted structures were targeted leading to the successful synthesis of NiPt_2Ga in the tetragonal Heusler structure. This work has broad implications for material exploration by design as previous ML approaches to materials discovery were biased by the limits of known phase spaces and experimentalist bias. The ability to generate and synthesize novel materials that are structurally and compositionally unique enables inverse-design of materials with targeted properties.

Data availability

We used the pretrained PGCGM available at <https://github.com/MilesZhao/PGCGM/tree/main>, including its scripts for validating generated structures. The code for training and evaluating our ALIGNN model is hosted at <https://github.com/newalexander/generative-materials-discovery>. Other relevant

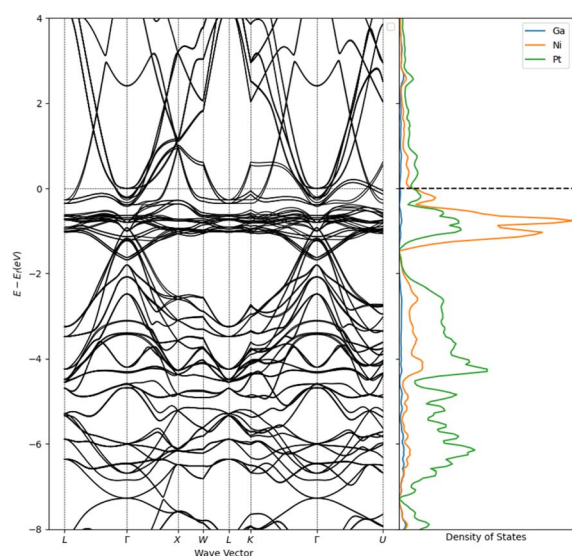


Fig. 3 DFT computed band structure of NiPt_2Ga in the tetragonal Heusler structure with lattice parameters determined from experimental data along high symmetry directions. DOS computed for total Ga, Ni, and Pt contributions.

data, including the MP structure IDs of the training data and trained ALIGNN model, as well as the structures of the generated materials and their ALIGNN-predicted decomposition enthalpies, are hosted on FigShare.⁵⁰

Author contributions

C. S., A. N., and T. M. M. contributed to the conception of the work. B. W., G. B., T. M. M., E. P., and E. G. contributed to the setup and design of experiments. B. W. and G. B. synthesized samples. A. N. created the ML model with some contributions from M. P. B. W., A. N., G. B., C. S., and T. M. M. heavily contributed to the writing and revising of the manuscript. B. W., G. B., W. B., and T. M. M. collected and analyzed experimental data. W. B., K. K. R., and N. Q. L. computed electronic structure calculations for the relevant compounds. M. W. contributed to the understanding of the electronic structure and bonding in these Heusler alloys.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors gratefully acknowledge internal financial support from the Johns Hopkins University Applied Physics Laboratory's Independent Research & Development (IR&D) Program for funding portions of this work. The MPMS3 system used for magnetic characterization was funded by the National Science Foundation, Division of Materials Research, Major Research Instrumentation Program, under Grant #1828490. Calculations were performed using computational resources of the Maryland Advanced Research Computing Center and the Advanced Research Computing at Hopkins (ARCH) Rockfish cluster.

Notes and references

- 1 K. T. Butler, D. W. Davies, H. Cartwright, O. Isayev and A. Walsh, *Nature*, 2018, **559**, 547–555.
- 2 G. Carleo, I. Cirac, K. Cranmer, L. Daudet, M. Schuld, N. Tishby, L. Vogt-Maranto and L. Zdeborová, *Rev. Mod. Phys.*, 2019, **91**, 045002.
- 3 G. E. Karniadakis, I. G. Kevrekidis, L. Lu, P. Perdikaris, S. Wang and L. Yang, *Nat. Rev. Phys.*, 2021, **3**, 422–440.
- 4 J. Im, S. Lee, T.-W. Ko, H. W. Kim, Y. Hyon and H. Chang, *npj Comput. Mater.*, 2019, **5**, 37.
- 5 Y. Liu, B. Guo, X. Zou, Y. Li and S. Shi, *Energy Storage Mater.*, 2020, **31**, 434–450.
- 6 V. Stanev, C. Oses, A. G. Kusne, E. Rodriguez, J. Paglione, S. Curtarolo and I. Takeuchi, *npj Comput. Mater.*, 2018, **4**, 29.
- 7 C. Qin, J. Liu, S. Ma, J. Du, G. Jiang and L. Zhao, *J. Mater. Chem. A*, 2024, **12**, 22689–22702.
- 8 E. A. Pogue, A. New, K. McElroy, N. Q. Le, M. J. Pekala, I. McCue, E. Gienger, J. Domenico, E. Hedrick, T. M. McQueen, B. Wilfong, C. D. Piatko, C. R. Ratto, A. Lennon, C. Chung, T. Montalbano, G. Bassen and C. D. Stiles, *npj Comput. Mater.*, 2023, **9**, 181.
- 9 L. Zhu, J. Zhou and Z. Sun, *J. Phys. Chem. Lett.*, 2022, **13**, 3965–3977.
- 10 A. Jain, S. P. Ong, G. Hautier, W. Chen, W. D. Richards, S. Dacek, S. Cholia, D. Gunter, D. Skinner, G. Ceder and K. A. Persson, *APL Mater.*, 2013, **1**, 011002.
- 11 K. Choudhary, K. F. Garrity, A. C. E. Reid, B. DeCost, A. J. Biacchi, A. R. Hight Walker, Z. Trautt, J. Hattrick-Simpers, A. G. Kusne, A. Centrone, A. Davydov, J. Jiang, R. Pachter, G. Cheon, E. Reed, A. Agrawal, X. Qian, V. Sharma, H. Zhuang, S. V. Kalinin, B. G. Sumpter, G. Pilania, P. Acar, S. Mandal, K. Haule, D. Vanderbilt, K. Rabe and F. Tavazza, *npj Comput. Mater.*, 2020, **6**, 173.
- 12 J. E. Saal, S. Kirklin, M. Aykol, B. Meredig and C. Wolverton, *JOM*, 2013, **65**, 1501–1509.
- 13 A. Belsky, M. Hellenbrandt, V. L. Karen and P. Luksch, *Acta Crystallogr., Sect. B*, 2002, **58**, 364–369.
- 14 A. O. Oliynyk, E. Antono, T. D. Sparks, L. Ghadbeigi, M. W. Gaultois, B. Meredig and A. Mar, *Chem. Mater.*, 2016, **28**, 7324–7331.
- 15 A. O. Oliynyk and A. Mar, *Acc. Chem. Res.*, 2018, **51**, 59–68.
- 16 J. Graser, S. K. Kauwe and T. D. Sparks, *Chem. Mater.*, 2018, **30**, 3601–3612.
- 17 A. G. Kusne, H. Yu, C. Wu, H. Zhang, J. Hattrick-Simpers, B. DeCost, S. Sarker, C. Oses, C. Toher, S. Curtarolo, A. V. Davydov, R. Agarwal, L. A. Bendersky, M. Li, A. Mehta and I. Takeuchi, *Nat. Commun.*, 2020, **11**, 5966.
- 18 Z. Rao, P.-Y. Tung, R. Xie, Y. Wei, H. Zhang, A. Ferrari, T. Klaver, F. Körmann, P. T. Sukumar, A. K. da Silva, Y. Chen, Z. Li, D. Ponge, J. Neugebauer, O. Gutfleisch, S. Bauer and D. Raabe, *Science*, 2022, **378**, 78–85.
- 19 A. P. Drozdov, P. P. Kong, V. S. Minkov, S. P. Besedin, M. A. Kuzovnikov, S. Mozaffari, L. Balicas, F. F. Balakirev, D. E. Graf, V. B. Prakapenka, E. Greenberg, D. A. Knyazev, M. Tkacz and M. I. Eremets, *Nature*, 2019, **569**, 528–531.
- 20 M. Somayazulu, M. Ahart, A. K. Mishra, Z. M. Geballe, M. Baldini, Y. Meng, V. V. Struzhkin and R. J. Hemley, *Phys. Rev. Lett.*, 2019, **122**, 027001.
- 21 D. V. Semenok, I. A. Kruglov, I. A. Savkin, A. G. Kvashnin and A. R. Oganov, *Curr. Opin. Solid State Mater. Sci.*, 2020, **24**, 100808.
- 22 M. Alverson, S. G. Baird, R. Murdock, E. S.-H. Ho, J. Johnson and T. D. Sparks, *Digital Discovery*, 2024, **3**, 62–80.
- 23 Y. Zhao, E. M. D. Siriwardane, Z. Wu, N. Fu, M. Al-Fahdi, M. Hu and J. Hu, *npj Comput. Mater.*, 2023, **9**, 38.
- 24 E. Kim and S. V. Dordevic, *J. Phys.: Condens. Matter*, 2023, **36**, 025702.
- 25 Z. Ren, S. I. P. Tian, J. Noh, F. Oviedo, G. Xing, J. Li, Q. Liang, R. Zhu, A. G. Aberle, S. Sun, X. Wang, Y. Liu, Q. Li, S. Jayavelu, K. Hippalgaonkar, Y. Jung and T. Buonassisi, *Matter*, 2022, **5**, 314–335.
- 26 C. Zeni, R. Pinsler, D. Züchner, A. Fowler, M. Horton, X. Fu, Z. Wang, A. Shysheya, J. Crabbé, S. Ueda, R. Sordillo, L. Sun, J. Smith, B. Nguyen, H. Schulz, S. Lewis, C.-W. Huang, Z. Lu, Y. Zhou, H. Yang, H. Hao, J. Li,



- C. Yang, W. Li, R. Tomioka and T. Xie, *Nature*, 2025, **639**(8055), 624–632.
- 27 E. T. Chenebuah, M. Nganbe and A. B. Tchagang, *npj Comput. Mater.*, 2024, **10**, 198.
- 28 C. W. Park and C. Wolverton, *Phys. Rev. Mater.*, 2020, **4**, 063801.
- 29 A. New, M. J. Pekala, N. Q. Le, J. Domenico, C. D. Piatko and C. D. Stiles, *ICML 2022 2nd AI for Science Workshop*, 2022.
- 30 R. E. A. Goodall and A. A. Lee, *Nat. Commun.*, 2020, **11**, 6280.
- 31 T. Xie and J. C. Grossman, *Phys. Rev. Lett.*, 2018, **120**, 145301.
- 32 T. Long, N. M. Fortunato, I. Opahle, Y. Zhang, I. Samathrakakis, C. Shen, O. Gutfleisch and H. Zhang, *npj Comput. Mater.*, 2021, **7**, 66.
- 33 Y. Zhao, M. Al-Fahdi, M. Hu, E. M. D. Siriwardane, Y. Song, A. Nasiri and J. Hu, *Adv. Sci.*, 2021, **8**, 2100566.
- 34 S. Kim, J. Noh, G. H. Gu, A. Aspuru-Guzik and Y. Jung, *ACS Cent. Sci.*, 2020, **6**, 1412–1420.
- 35 A. New, M. Pekala, E. A. Pogue, N. Q. Le, J. Domenico, C. D. Piatko and C. D. Stiles, *1st Workshop on the Synergy of Scientific and Machine Learning Modeling @ ICML2023*, 2023.
- 36 M. Arjovsky, S. Chintala and L. Bottou, *Proceedings of the 34th International Conference on Machine Learning*, 2017, pp. 214–223.
- 37 D. Saxena and J. Cao, *ACM Comput. Surv.*, 2021, **54**(3), 63.
- 38 C. J. Bartel, A. Trewartha, Q. Wang, A. Dunn, A. Jain and G. Ceder, *npj Comput. Mater.*, 2020, **6**, 97.
- 39 K. Choudhary and B. DeCost, *npj Comput. Mater.*, 2021, **7**, 185.
- 40 M. Aykol, S. S. Dwaraknath, W. Sun and K. A. Persson, *Sci. Adv.*, 2018, **4**, eaaq0148.
- 41 J. Jang, G. H. Gu, J. Noh, J. Kim and Y. Jung, *J. Am. Chem. Soc.*, 2020, **142**, 18836–18843.
- 42 J. K. Kawasaki, S. Chatterjee, P. C. Canfield and G. Editors, *MRS Bull.*, 2022, **47**, 555–558.
- 43 T. Graf, F. Casper, J. Winterlik, B. Balke, G. H. Fecher and C. Felser, *Z. Anorg. Allg. Chem.*, 2009, **635**, 976–981.
- 44 T. Graf, S. S. P. Parkin and C. Felser, *IEEE Trans. Magn.*, 2011, **47**, 367–373.
- 45 T. Graf, C. Felser and S. S. Parkin, *Prog. Solid State Chem.*, 2011, **39**, 1–50.
- 46 A. Merchant, S. Batzner, S. S. Schoenholz, M. Aykol, G. Cheon and E. D. Cubuk, *Nature*, 2023, **624**, 80–85.
- 47 L. Ward, A. Agrawal, A. Choudhary and C. Wolverton, *npj Comput. Mater.*, 2016, **2**, 16028.
- 48 T. Klimczuk, C. H. Wang, K. Gofryk, F. Ronning, J. Winterlik, G. H. Fecher, J.-C. Griveau, E. Colineau, C. Felser, J. D. Thompson, D. J. Safarik and R. J. Cava, *Phys. Rev. B*, 2012, **85**, 174505.
- 49 M. J. Winiarski, G. Kuderowicz, K. Górnicka, L. S. Litzbarski, K. Stolecka, B. Wiendlocha, R. J. Cava and T. Klimczuk, *Phys. Rev. B*, 2021, **103**, 214501.
- 50 A. New, Materials-Discovery, 2024, https://figshare.com/articles/dataset/materials-discovery_zip/25783221.

