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# Grouped representation of interatomic distances as a similarity measure for crystal structures†

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Determining how similar two materials are in terms of both atomic composition and crystallographic structure remains a challenge, the solution of which would enable generalised machine learning using crystal structure data. We demonstrate a new method of describing crystal structures based on interatomic distances, termed the Grouped Representation of Interatomic Distances (GRID). This fast to compute descriptor can equally be applied to crystalline or disordered materials, and encodes additional information beyond pairwise distances, such as coordination environments. Combined with earth mover's distance as a measure of similarity, we show that GRID is able to quantitatively compare materials involving both short- and long-range structural variation. Using this new material descriptor, we show that it can accurately predict bulk moduli using a simple nearest-neighbour model, and that the resulting similarity shows good generalisability across multiple materials properties.

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## Introduction

The exponential increase in crystal structures deposited in databases such as the inorganic crystal structure database (ICSD<sup>1</sup>) and Cambridge structural database (CSD<sup>2</sup>) continually expands our exploration of a multi-dimensional “structure-composition” (S-C) space. The arrangement of atoms and their types defined within this S-C space is intrinsically linked to all physical properties of a material; if two materials are proximate in S-C space, their properties are similar. For example, diamond and cubic boron nitride both adopt a cubic diamond structure, and carbon atoms are chemically similar to both boron and nitrogen. As a result, both materials are super-hard.<sup>3</sup> In contrast, diamond and graphite are compositionally identical but structurally distinct, giving very different properties for these polymorphs. Although this is a trivial example, the same approach could be applied to almost any problem. Although conceptually simple, the challenge of this idea lies in how to define S-C space so that proximity correctly reflects the physical property of interest. Indeed, some properties may only depend on some of the S-C dimensions. For instance, the crystal field splitting important for solid state phosphors is strongly dependent on the local bond lengths, angles and ligand species, but largely independent of longer-range crystal structure. In contrast, charge density waves (which are intertwined with superconductivity) are inherently a long-range effect. Existing

approaches to determine similarity between atomic structure tend to focus on either long-range similarity or short-range (local) environments. Long range methods are primarily concerned with comparing crystallographic symmetry and Wyckoff positions<sup>4,5</sup> and/or directly minimising the distance between periodic point sets through mapping.<sup>6–8</sup> In contrast, short-range approaches typically focus on matching local coordination environments and quantifying their similarity to different geometric environments,<sup>9</sup> and then combining all environments to determine overall crystal similarity.<sup>10</sup> Additionally, many of these algorithms neglect overall scale in order to group materials into structural prototypes, limiting their usefulness for properties that depend on lattice volume.‡ Although recent work has begun to address both long- and short-range similarity on an equal footing (for example ref. 10) it remains a significant challenge to quickly and accurately reflect the similarity within S-C space, whilst ensuring that small structure (or compositional) changes do not lead to discontinuous behaviour.

Machine learning (ML) offers a readily available framework to fit models within a multidimensional S-C space, but a definition of this space is still required. For crystalline materials in particular, this presents a problem. The standard way of representing structures based on a unit cell and atomic coordinates is not a suitable construction of S-C space; the infinite possible unit cell definitions for the same material result in a non-unique representation, and small atomic shifts can result in discontinuous changes in unit cell metric (*e.g.* by forming supercells). To overcome this limitation, it is common

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‡ Strictly speaking, the degree of similarity between two structures considering lattice scaling is a measure of mathematical congruence rather than similarity, however, we use the term similarity throughout this manuscript for simplicity.



to use crystal descriptors (also known as features) that represent a crystal structure in a unique way to allow direct comparisons. A range of descriptors exist for both organic and inorganic materials, and these typically focus on either the entire structure or on combining local structural features into materials' "fingerprints". A huge range of features are available (see for example ref. 11 or 12 for reviews) ranging from single-value metrics such as the Shannon entropy of a structure<sup>13</sup> or variance in bond lengths,<sup>14</sup> through vector descriptions such as the "bag of bonds",<sup>15</sup> all the way to multi-dimensional features based on polynomial expansions of radial or angular functions.<sup>16,17</sup> Features often focus on atom-centred functions (such as atomic fingerprints or smooth overlap of atomic positions, SOAP)<sup>18,19</sup> but for extended solids it is also possible to work with the Fourier-transformed atomic positions (*i.e.* diffraction-based).<sup>20</sup> Unfortunately, many existing approaches do not scale to different numbers of atoms in the unit cell, or the number of terms increases exponentially. More recently, successful results have been obtained using descriptions based on the bonding connectivity within materials, often incorporating additional atomic features.<sup>21–23</sup> Unfortunately, these approaches require non-standard ML architectures which are not suited to all problems. A common aspect of many existing approaches is that they treat structural and compositional information together. Separating them would allow a more flexible approach to constructing S-C space depending on the problem in hand, for instance emphasising structure over composition.

One of the simplest, least-parameterised features to describe atomic structure is simply the pairwise interatomic distances, or radial distribution function (RDF). While fast to compute, the RDF is an oversimplified crystal representation for ML as three structural dimensions are compressed to one. As such, related methods have been developed to include additional information, such as embedding atomic species information (partial RDF or the Coulomb matrix)<sup>24,25</sup> or bond angles (many-body tensor representation, MBTR).<sup>26</sup> Unfortunately, the size of both of these descriptors varies with the number of atomic species in a material, while established ML applications require a fixed descriptor size. Here we develop a method of encoding pairwise distance information called the Grouped Representation of Interatomic Distances (GRID). GRID encodes more information than RDF, whilst retaining its computational speed and insensitivity to the number of atoms present. GRID has been developed independently from a related approach (average minimum distance) which has recently been extended to pointwise distance distributions, and both have been proved to uniquely describe periodic point sets.<sup>27,28</sup> Because each unique distance appears twice in GRID, the asymmetric unit of a periodic structure can therefore be reconstructed from its GRID using a process of elimination, excluding chirality. To demonstrate the applicability of GRID to materials problems, we present its use in quantifying similarity between crystal structures as well as its effectiveness in predicting bulk modulus based on crystallographic structure.

## Methods

### Code

All data processing was performed using code written in Python 3.8. The most recent version of the code is freely available from <https://github.com/CumbyLab/gridrdf>, while a static version associated with this manuscript is available at DOI [10.5281/zenodo.7271754](https://doi.org/10.5281/zenodo.7271754). Crystal properties and structural data were collected and processed using tools within the pymatgen python package, including calculation of pairwise distances.<sup>29</sup> Machine-learning models were defined and optimised using scikit-learn.<sup>30</sup> Earth mover's distance (EMD) and cosine distance were calculated using `scipy.stats.wasserstein_distance` and `scipy.spatial.distance.cosine`, respectively.

### Bulk modulus dataset

Materials with associated bulk modulus ( $K$ ) calculations were extracted from the Materials Project (v2020.06) database using the materials API,<sup>31,32</sup> totalling 13 172 materials. Anomalous entries with  $K \leq 0$  were removed (12). To facilitate analysis, only entries with elements up to Bi (atomic number 83) exclusive of noble gas elements were kept. This means that, specifically to this dataset, entries with elements Ne, Kr and Ac–Pu were removed (788 materials). GRIDs were calculated to a maximum distance of 10 Å (see results for algorithm). Extremely low-density structures with fewer than 100 histogram 'groups' at 10 Å (194 materials) were omitted from analysis, leaving 12 178 materials in the analysed data. The distribution of the resulting bulk moduli range from 0–600 GPa, with most materials below 100 GPa (Fig. S1†).

### Machine learning model training

$k$ -Nearest neighbours ( $k$ NN) regression models<sup>33</sup> were implemented using the `KNeighborsRegressor` function in scikit-learn using pre-computed dissimilarity matrices. For simplification and to facilitate further analysis,  $k = 1$  was used; considering more than one neighbour ( $k > 1$ ) provided only a slight improvement in performance (Fig. S2†). A 5-fold random cross-validation implemented in scikit-learn was used to calculate the mean absolute error (MAE) as a function of the number of training samples. The  $k$  closest neighbours of each structure in the dataset were found using the  $n$ -smallest method implemented in Pandas and the pre-computed dissimilarity matrix.

Kernel ridge regression (KRR) models were implemented using the `KernelRidge` function in scikit-learn. Linear and radial basis function (RBF) kernels were tested, with 5-fold cross validation used to optimise the regularization and kernel width hyper-parameters in the logarithmic ranges  $10^0$ – $10^{-3}$  (4 steps) and  $10^{-2}$ – $10^2$  (50 steps), respectively. Minimum MAE was achieved with a linear kernel and regularization value of 1.

Visualisations using t-distributed stochastic neighbour embedding (t-SNE) were generated using scikit-learn, with a perplexity of 50.



## Composition similarity metric

The compositional similarity matrix was determined for 78 elemental pairs (atomic numbers to bismuth, excluding noble gases elements). Pairwise distances were computed based on previously reported similarity values ( $g_{AB}$ )<sup>34</sup> which were themselves computed from data-mining of literature crystal structures. To convert similarities to dissimilarities, we take the reciprocal logarithm of the original values,  $\text{diss}(A, B) = 1/\log_{10}(g_{AB} + 1)$ . While Hautier *et al.*<sup>34</sup> give similarity measures for multiple oxidation states for some elements, we have adopted the most commonly observed oxidation state for each element to avoid the need to estimate valence. To account for missing AB pairs in the original publication, we assign pairs containing Pm the corresponding Sm values, and otherwise set missing values to an arbitrary large dissimilarity ( $g_{AB} = 10^{-5}$ ) as a missing pair would suggest that substitution has not been experimentally observed. Given the overlap in data between the Materials Project and ICSD, we do not expect many materials to require these missing  $g_{AB}$  pairs. Fig. S3† shows the resulting compositional distance matrix, with elements arranged according to the Pettifor scale which places chemically similar elements near to each other.<sup>35</sup> Blocks of similar distance can clearly be observed for chemically related elements (*e.g.* rare-earths), as can the large distances between *e.g.* anions and cations. Note that Fig. S3† visualises  $\log(\text{diss}(A, B))$  to avoid the effect of extremely large values, whilst our compositional similarity calculations use  $\text{diss}(A, B)$ . Fig. S4† shows the distributions of EMD values for composition and structure, respectively. As only small values are useful in the  $k$ NN model, only the smallest EMD values (Fig. S4a and b†) and the average of 10 smallest EMD values (Fig. S4c and d†) are shown.

## Results and discussion

### GRID as a distance measure

The procedure to calculate the GRID is as follows: (1) compute all pairwise distances up to an arbitrary cutoff (here 10 Å) starting from each symmetry-unique atom ( $j$ ) within the unit cell, taking periodic boundary conditions into account; (2) rank the distances  $d_j = \{d_{1j}, \dots, d_{ij}, \dots\}$  in ascending order and then assign the  $i$ th-ranked value to the  $i$ th GRID 'group'; (3) to achieve a fixed-length descriptor, discretize the  $d_{ij}$  values of the  $i$ th group into a binned histogram, after smoothing them using a squared exponential kernel to avoid discontinuous jumps between bins caused by numerical or experimental error;

$$F_i(n) = \int_{(n-1)\Delta d}^{n\Delta d} \frac{\sum_j e^{-\frac{(x-d_{ij})^2}{2\sigma^2}}}{a} dx \quad (1)$$

where  $F_i(n)$  is the atom density in bin  $n$  (bin width =  $\Delta d$ , here 0.1 Å) normalised by the number of symmetry unique atoms  $a$ ,  $d_{ij}$  is the  $i$ th-ranked distance for atom  $j$  and  $\sigma$  is the standard deviation of the exponential smoothing applied to each distance (here 0.1 Å). Normalizing the resulting histogram enables comparison between structures with different numbers of atoms. The final GRID descriptor is formed by concatenating

the histograms of the first 100 groups, enabling numerical comparison between structures with different average density. Selecting the first 100 groups does not strictly enforce the 10 Å limit initially chosen, instead giving a skewed distribution of maximum distances peaking around 7.5 Å (Fig. S5†). This is sufficient to capture the structural features presented here, but longer distances could be considered at the expense of computational complexity.

As an illustrative example, Fig. 1 shows a simulated cubic perovskite  $ABX_3$ , and the first 8 groups of the corresponding GRID. From Fig. 1b, the coordination number of each atom within the perovskite is readily apparent; as the first two groups are identical but different to the 3rd group, one of the atoms (X) must have a coordination number of two. Similarly, the changes between 6th and 7th groups suggests a coordination number of six must be present (B). Whilst this sort of manual analysis becomes more challenging for complex or low-symmetry structures, the coordination number and local symmetry is inherently embedded within the GRID. This reduces the data loss in converting 3D coordinates into a 1D histogram compared with the RDF. It is worth noting, however, that summation of all GRID groups will recover the traditional RDF distribution.

As a demonstration of the effectiveness of the GRID in quantifying structural similarity, we have constructed three test

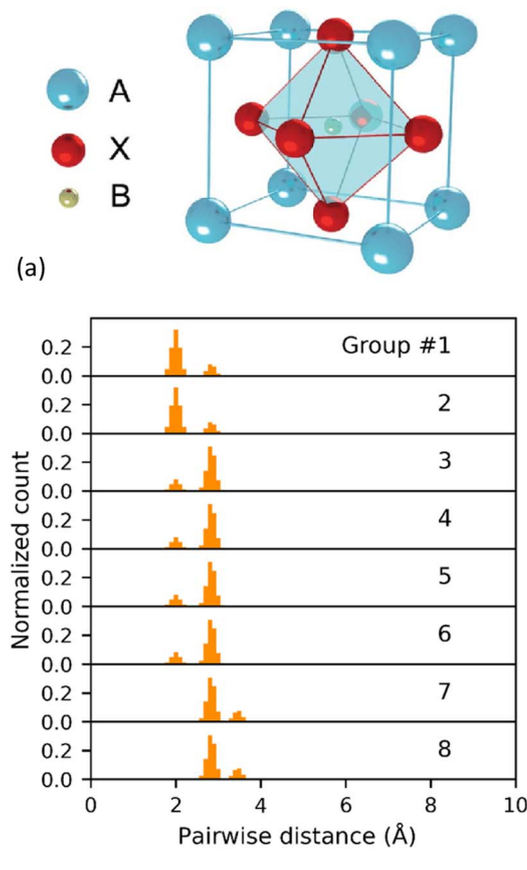


Fig. 1 (a) Crystal structure of a simulated cubic perovskite and (b) the first eight calculated GRID groups.



examples in order to investigate different aspects of structural similarity: (1) structures formed by the variable intergrowth of two sub-units; (2) changes due to atomic displacements such as at a phase transitions; and (3) the effect of lattice parameter variations. Between them these examples cover long-range structure, short-range variations and changes to overall symmetry, demonstrating the wide applicability of this descriptor to structural comparisons. While GRID focusses on the structural aspect of S-C space, we will show later in this manuscript that it can also be combined with a measure of compositional similarity.

Whilst the Euclidean distance is widely used to quantify difference between two multi-dimensional vectors, it is not well-suited to discrete distributions such as the GRID because of insensitivity to redistribution between bins. To illustrate this, the distribution  $X = [0, 1, 0, 0, 0]$  (with  $X$  representing the first five bins of an RDF) will have the same Euclidean distance to  $Y = [1, 0, 0, 0, 0]$  and  $Z = [0, 0, 0, 0, 1]$ , although  $X$  and  $Y$  are closer if viewed as a binned distribution. A more appropriate measure of distance for such distributions is the earth mover's distance (EMD). This metric (also known as the Wasserstein distance) can be considered as the minimum amount of 'work' needed to

transform one distribution into another, analogous to moving piles of earth (identical distributions have  $EMD = 0$ ). Such a metric has recently been suggested as a way to quantify compositional similarity in inorganic materials.<sup>36</sup> In this work, we have used EMD to compare between GRID distributions by computing the 1D pairwise ( $i$ th– $i$ th) distances, before finally computing the mean EMD across all GRID groups. Note, however, that a different weighting within the mean could be applied to favour short- or long-range similarity, or alternatively a 2D EMD could compare across all GRID groups at once, at the expense of computational time. This approach could be beneficial in situations where a property depends more strongly on short-range or long-range structural features, by biasing the overall EMD to emphasise short- or long-range similarity.

### Composite structures

The first example is based on the Ruddlesden–Popper (R–P) series  $Sr_{n+1}Ti_nO_{3n+1}$ , which can be considered as a composite structure of variable-width perovskite units (width determined by  $n$ ) separated by single rock-salt layers. Here we consider four structures (optimised using density functional theory, DFT)

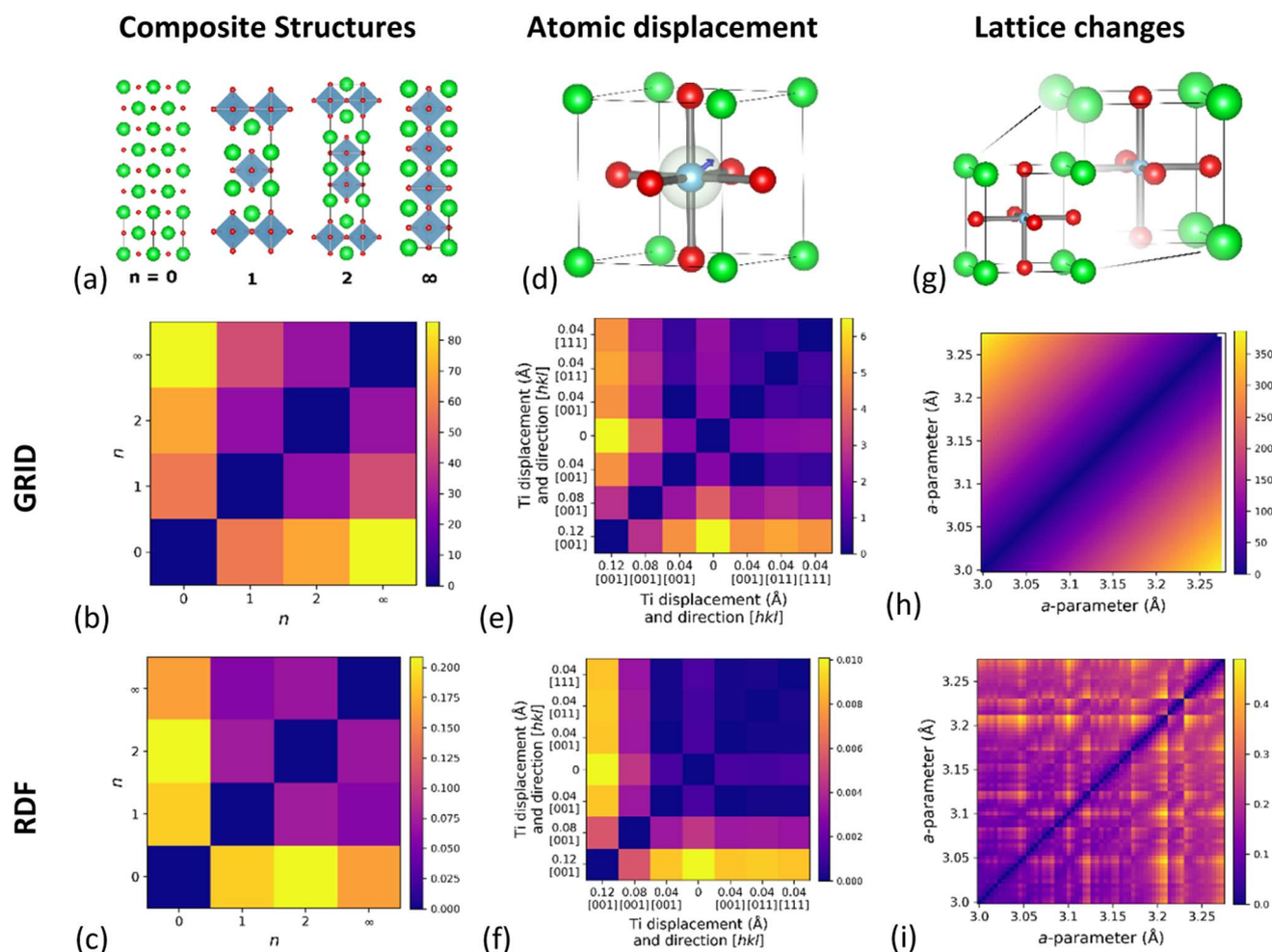


Fig. 2 Pairwise-comparison of EMD distances calculated using GRID and RDF for: (a–c) Ruddlesden–Popper phases  $Sr_{n+1}Ti_nO_{2n+1}$ ; (d–f) BaTiO<sub>3</sub> with varying Ti atomic displacements; (g–i) simulated cubic perovskite with linearly varying lattice parameters.



from the Materials Project (Fig. 2a): rock-salt SrO ( $n = 0$ , mp-2472), Sr<sub>2</sub>TiO<sub>4</sub> ( $n = 1$ , mp-5532), Sr<sub>3</sub>Ti<sub>2</sub>O<sub>7</sub> ( $n = 2$ , mp-3349) and perovskite SrTiO<sub>3</sub> ( $n = \infty$ , mp-5229). The results of GRID dissimilarity calculated using EMD (Fig. 2b) show that the distance from SrO increases with  $n$ . This agrees with the chemical intuition; as  $n$  increases the ratio of rock-salt to perovskite layers reduces. Importantly, SrO ( $n = 0$ ) and SrTiO<sub>3</sub> ( $n = \infty$ ) appear most dissimilar, as expected. In contrast, EMD calculated using the RDF (Fig. 2c) finds SrO and SrTiO<sub>3</sub> to be the most similar pair, against chemical intuition. We have also tested the commonly used cosine distance (Fig. S6†) and find that although the same broad trend is reproduced using the GRID and RDF, the quantitative discrimination between the different R-P phases is much reduced.

### Atomic displacements

Our second simulated example is based on idealized displacements of the B-site cation within a cubic perovskite, such as those important for ferroelectricity in BaTiO<sub>3</sub>.<sup>37</sup> The simulated dataset consists of six structures with varying B-site displacements, whilst artificially maintaining the same cubic unit cell metric to separate atomic from lattice effects (Table S1†). By applying B-site displacements of 0.04 Å along the [0, 0, 1], [1, 1, 0] and [1, 1, 1] crystallographic directions as well as varying the magnitude of the displacement along [0, 0, 1], we are able to compare the influence of B-site displacement on the dissimilarity measure. The results of pairwise EMD calculated using GRID (Fig. 2e) show that larger displacements do indeed give larger distance from the parent structure. We find that the displacement distance has a stronger effect than displacement direction on the similarity distance, which can be understood from the radial nature of GRID. There are, however, slight differences in EMD between the different displacement directions, showing that the GRID does encode some angular information for periodic structures. In comparison, equivalent EMD results using the RDF (Fig. 2f) show that while RDF captures the magnitude of atomic displacement, it does not distinguish displacement in different directions.

### Lattice expansion

A significant drawback to existing RDF-based descriptors is that two structures identical apart from isotropic lattice expansion will give completely different RDFs. To test this, we have simulated 61 cubic (space group  $Pm\bar{3}m$ ) perovskite structures with lattice constants ranging from 3 Å to 6 Å ( $\Delta a = 0.05$  Å), chosen to coincide the range of cubic perovskite lattice constant found in the Materials Project (3.05 Å–6.12 Å). Fig. 2h shows the pairwise EMD between structures for each lattice constant, and reveals a continuous increase with lattice constant difference. It is intuitive that as a structure expands it becomes less similar to the starting point, but this is in stark contrast with the RDF (Fig. 2i) which shows discrete jumps in EMD across the same range. We attribute this behaviour to the significant degeneracy of interatomic distances at large separations in the RDF, and the different rate at which these change with isotropic expansion. As these distances move into different RDF bins during expansion, the minimum EMD changes discontinuously due to

a change in EMD flow, as histogram ‘mass’ is preferentially moved to a different bin. A similar comparison using cosine distance (Fig. S6e and f†) reveals that cosine distance using GRID only gives sensible dissimilarities for numerically close lattice parameters, while cosine distance on the RDF shows even more discontinuities than EMD.

From these examples, it is clear that the GRID outperforms the RDF in all cases using either the EMD or cosine distance metrics, although the EMD gives more chemically intuitive dissimilarities. For this reason, the remaining results will focus on EMD as a useful measure of dissimilarity within S-C space.

### Predicting bulk modulus

To demonstrate the effectiveness of GRID in quantifying structural similarity with EMD, the following results demonstrate its use in predicting bulk modulus ( $K$ ) from crystal structures. Bulk modulus is an important parameter related to many technologically important properties such as lattice thermal conductivity and mechanical deformation. There are several existing literature reports of ML-based prediction of  $K$  (Table S2†) using different datasets, input features and machine learning algorithms. Comparison between these studies is challenging due to the different data sets and metrics used and whether the model is fitted on a linear or logarithmic scale, but the current state of the art methods using data from the Materials Project achieve a MAE of around 10 GPa (or 0.05 using  $\log_{10}(K/\text{GPa})$  as input).<sup>38,39</sup>

Using DFT-simulated bulk moduli obtained from the materials project (12 178 entries after cleaning, see Methods) we have computed the GRID for each material and determined the EMD between all structure pairs (the ‘dissimilarity matrix’). Such a dissimilarity matrix can be used directly to train a number of ML models; in this case we have used the simplest approach of interpolating between the values of the  $k$ -nearest neighbouring points ( $k$ NN). A  $k$ NN approach relies on the similarity matrix accurately representing proximity within S-C space, and also that the available data effectively sample this S-C space. For comparison, we have also trained a kernel ridge regression (KRR) model using a linear kernel (see Methods); for well-sampled data these two approaches should replicate each other.

Fig. 3 shows the MAE of bulk modulus prediction as a function of the training set size using both  $k$ NN and KRR. As expected the MAE decreases with greater sampling of S-C space, but importantly the  $k$ NN and KRR approaches give the same behaviour, *i.e.* the training and test data are sufficiently close that bulk modulus can be successfully predicted by the nearest neighbour. Notably,  $k$ NN actually out-performs KRR in most models, except when using only composition and training on fewer than 4000 samples. When using only composition as a feature, KRR is insensitive to the number of training samples, likely due to the short elemental vector (size of 78) compared to the large S-C space.

Based on GRID alone, the model approaches a minimum MAE of 33 GPa. Given that this model contains no composition information it is remarkably accurate, but has a fundamental limit; two materials with the same structure but different



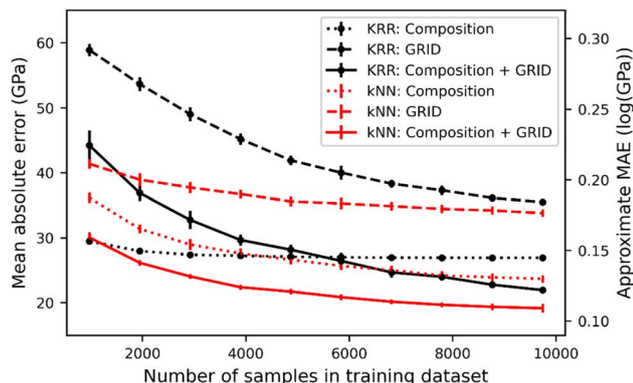


Fig. 3 Prediction error of bulk modulus as a function of training set size. The error bars are from 5-fold cross-validation as implemented in scikit-learn. KRR and kNN results are shown using colour and marker style, while different line styles correspond to different input features: composition (dots), GRID only (dash) and composition + GRID (solid).

composition will necessarily have the same predicted bulk modulus. It is therefore informative to consider how many materials within our data possess the same GRID; by comparing pairs of GRIDs we find that there are 279 non-unique GRIDs, shared across 582 materials (*i.e.* two or three materials have the same GRID). As an example,  $\text{ScGaNi}_2$  and  $\text{PtAlLi}_2$  (Fig. S7†) have identical GRIDs (EMD = 0) but have bulk moduli of 130 GPa and 88 GPa, respectively. Thus, structurally identical but compositionally distinct structures represent roughly 5% of the total dataset. It is worth noting that the number of structures with the same GRID will decrease with a greater distance cut off and/or smaller exponential smoothing of distances, at the expense of larger feature spaces (requiring more computational resources).

To address the absence of compositional information, we have therefore combined our GRID EMD with a similar compositional EMD, in a modified version of that demonstrated by Hargreaves *et al.*<sup>36</sup> Our method represents the normalised elemental fractions as a 78-element vector in atomic number order (considering elements up to Bi, but excluding noble gases); taking  $\text{SrTiO}_3$  as a representative example would give values of 0.6, 0.2 and 0.2 at the 7th, 19th and 34th elements in this vector, respectively. Rather than ordering this vector by Pettifor scale and computing EMD directly as in ref. 36, we instead introduce a pairwise dissimilarity metric (Fig. S3†) between elements based on the statistical likelihood of species occurring within the same crystal structure (see Methods).<sup>36</sup> The

advantage of this approach is that while the Pettifor scale assumes a constant distance between adjacent species, the substitutional (dis)similarity approach gives a more chemically meaningful metric. For example, the lanthanide series (La–Yb) covers a range of 14 steps on the Pettifor scale, while the dissimilarity approach gives a range of 0.4. In contrast, Na and F are adjacent on the Pettifor scale (one step), but have a dissimilarity of 4.38.

Using the composition information for each of the 12 178 data points, we have computed the pairwise EMD matrix of compositional distances; Table 1 gives the statistical results of models trained using composition alone as well as combined with GRID. Combining compositional and structural information gives the optimum bulk modulus prediction, achieving a minimum MAE of 18 GPa (Fig. 3). This approaches the state-of-the-art literature value ( $\sim 10$  GPa), but using a much simpler nearest neighbour model. Using EMD as a metric gives better predictions than cosine distance, particularly for composition comparisons.

The predicted and DFT-calculated values using combined GRID and composition distances are shown in Fig. 4. From this and the results in Table 1, it is apparent that while most

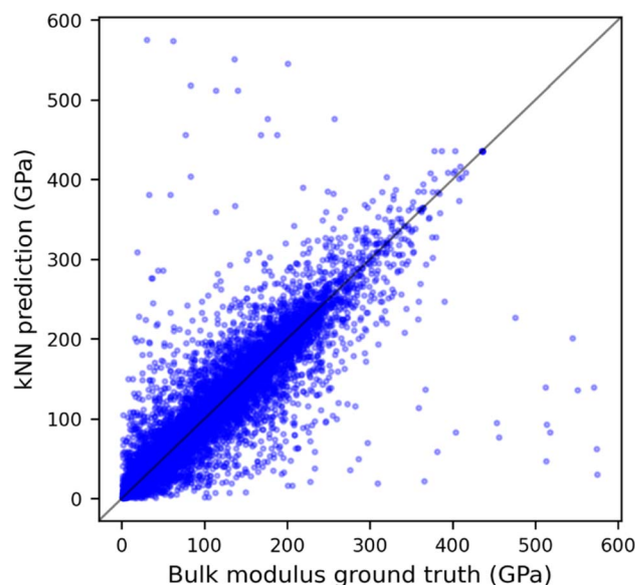


Fig. 4 Predicted bulk modulus vs. DFT-computed value for the whole dataset using kNN model and combined GRID and composition dissimilarity matrix.

Table 1 Prediction of bulk modulus using kNN and different dissimilarity measures (EMD is used except where cosine distance is stated). Statistics are computed using absolute errors (in GPa)

Similarity method	MAE	Standard deviation	Min	Median	Max
GRID – EMD	32.64	41.46	0	18	555
GRID – cosine	32.65	41.72	0	18	555
Composition – EMD	22.48	35.54	0	11	545
Composition – cosine	101.17	72.02	0	54	573
GRID + composition – EMD	18.39	30.29	0	9	545



predictions are close to the ground truth value, a few significant outliers give the resulting large MAEs; indeed, the median absolute error is only 9 GPa, comparable to the state-of-the-art model. Similar plots when fitting GRID or compositional similarity alone give a much broader spread around  $x = y$  (Fig. S8†). It might be assumed that these most erroneous points arise due to the small number of points with high bulk modulus, however, there seems to be little correlation between the prediction error for a material and the distance to its nearest neighbour (Fig. S9†). Interrogating this further, the maximum prediction error occurs for two anti-perovskite materials:  $\text{BiAsSr}_3$  ( $K_{\text{DFT}} = 575$  GPa, Fig. S10a†) and  $\text{BiPSr}_3$  ( $K_{\text{DFT}} = 30$  GPa, Fig. S10b†). These two structures are near-identical, and As and P are chemically very similar; it is unclear why these two (theoretical) materials should give such different bulk moduli, especially as the shear moduli are equivalent at 17 GPa and 22 GPa, respectively. This may point to irregularities within the bulk modulus data used to train the models, or alternatively suggest that accurate bulk modulus prediction requires information beyond structure and composition, such as detailed knowledge of the electron distribution.

## Generalisability

Used alone, composition information results in more accurate predictions of bulk modulus than structural information (GRID), but in both cases there are a significant number of materials where considering only one aspect of S-C space results in zero EMD but significant difference in bulk modulus (Fig. S7†). Both structure and composition are therefore critical to predict bulk modulus. Fig. 5 shows a two-dimensional representation of the high-dimensional feature space for each of the similarity measures described, using t-distributed stochastic network embedding (t-SNE). While these plots are a dramatic simplification of the high dimensional data, proximity within the 2D plane indicates proximity within S-C space (although the reverse is not necessarily true). Examining the bulk modulus data it is clear that clusters with similar bulk moduli appear for both GRID and composition features, and the combined descriptor shows particularly strong clustering of the low- $K$  materials.

The same t-SNE distributions can also be visualised with different material properties; in this case we have extracted shear moduli, formation energies and band gaps (Fig. 5). In all cases there is clustering of points with similar properties,

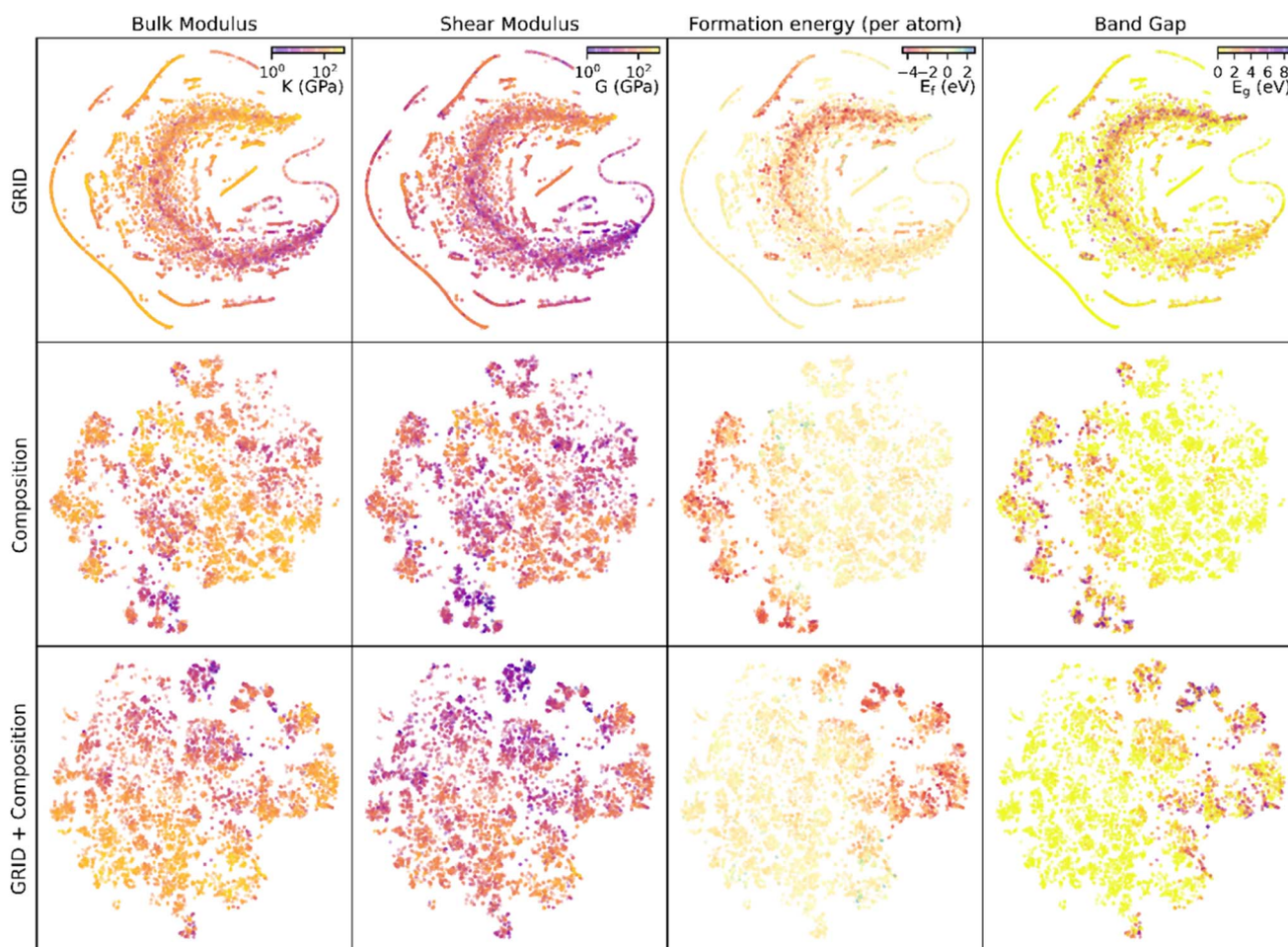


Fig. 5 t-SNE visualisations of structural, composition and combined descriptors, with points coloured by their physical properties. Clustering of points represents proximity within the high-dimensional space.



although the clusters occur in different points of the t-SNE plane and with different amounts of spread. This ability of GRID and compositional similarity to place materials with similar properties in close proximity shows that they are an accurate representation of the underlying S-C space. Using a similar  $k$ NN ( $k = 1$ ) model on each of these properties with no further optimisation yields predictions with a MAE (median AE in brackets) of: 15 (7) GPa for shear modulus prediction, 0.26 (0.12) eV for formation energy and 0.45 (0.0) eV for band gap. In all cases the median AE is lower than the mean, indicating that outlying data points are enlarging the MAE (note that the median AE of 0 eV for band gap prediction arises due to the significant number of metallic materials present in the data).

While GRID does not match the accuracy of graph-based neural networks when predicting properties from structure and composition, it is notable that it can achieve similar accuracy to other “traditional” machine learning algorithms (such as support-vector regression or random forests) using a model as simple as  $k$ NN (see Table S2† for details). We suggest that this reflects the way that GRID accurately reflects the underlying structural similarity, rather than relying on a high-dimensional model to recover that similarity. As such, we expect that the performance of GRID-based methods could be further improved, for example by weighting the contributions of different shells to the EMD, or by allowing comparisons between GRID shells (*e.g.* using two-dimensional EMD).

Unlike other existing representations of materials, GRID can not only be applied to periodic systems, but also those exhibiting disorder. This could include almost-periodic models such as structures with disordered occupations, but could equally be extended to truly amorphous materials as long as GRID shells could be calculated for each unique atom. The first challenge in this approach is finding a suitable structural model that could be used to generate a GRID representations, but this could be recovered from, for example, molecular dynamics simulations or reverse Monte Carlo fits to experimental data. The second challenge in applying this approach would be finding suitable datasets of disordered materials and their associated properties, such that a ML model could be trained. It is an open problem, but GRID gives the opportunity of comparing ordered and disordered materials using the same model.

## Conclusions

We have introduced the GRID as a more information-dense structure representation than the traditional RDF, but retaining its speed and relative simplicity. Combined with EMD as a distance measure between distributions, we find that we can quantify structural similarity in a range of materials-relevant problems in agreement with chemical intuition. We also introduce a modified approach to quantifying compositional similarity using EMD which gives a more accurate metric between different elements. Combining these two measures of similarity with bulk modulus data and an extremely simple  $k$ NN regression model results in prediction MAEs of 18 GPa, approaching the state of the art literature results; in fact the median AE of 9 GPa is directly comparable to recent studies.

More importantly, we find that the GRID/composition method described generalises well to other material properties, providing an accurate representation of the underlying structure-composition space that dictates observable behaviour. By extending these descriptors to more advanced, non-linear models, we expect that a wide range of physical properties could be modelled using a similar architecture.

## Data availability

The code for computing and analysing GRID representations can be found at <https://github.com/CumbyLab/gridrdf> with a static version associated with this manuscript available at Zenodo, DOI: [10.5281/zenodo.7271754](https://doi.org/10.5281/zenodo.7271754). The structural data used in this manuscript are publicly available from the Materials Project, <https://materialsproject.org/>.

## Author contributions

R. Z., S. S. and J. C. conceived the work and devised the methodology. R. Z. performed the data curation, software development and analysis/validation of results, with assistance from J. C. All authors contributed to the final manuscript.

## Conflicts of interest

There are no conflicts of interest to declare.

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