Probing atomic-scale symmetry breaking by rotationally invariant machine learning of multidimensional electron scattering

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The 4D scanning transmission electron microscopy (STEM) method maps the structure and functionality of solids on the atomic scale, yielding information-rich data sets describing the interatomic electric and magnetic fields, structural and electronic order parameters, and other symmetry breaking distortions. A critical bottleneck is the dearth of analytical tools that can reduce complex 4D-STEM data to physically relevant descriptors. We propose an approach for the systematic exploration of 4D-STEM data using rotationally invariant variational autoencoders (rVAE), which disentangle the general rotation of the object from other latent representations. The implementation of purely rotational rVAE is discussed as are applications to simulated data for graphene and zincblende structures. The rVAE analysis of experimental 4D-STEM data of defects in graphene is illustrated and compared to the classical center-of-mass analysis. This approach is universal for probing symmetry-breaking phenomena in complex systems and can be implemented for a broad range of diffraction methods.

ARTICLE

INTRODUCTION

Functionalities of materials including ferroics, superconductors, and charge density wave systems are governed by the physics of symmetry breaking phenomena. In systems with long-range discrete translation symmetries, these behaviors are readily amenable to neutron and X-ray scattering, providing insight into the minute details of atomic structure, electronic density distribution, and elastic and inelastic vibrational properties. In these systems, the long-range periodicity allows integrating the behaviors over multiple unit cells. Similar approaches can be extended to ordered 2D systems such as surfaces and interfaces, as accessed via low-energy electron diffraction or surface X-ray scattering. In systems with long-range symmetries, these behaviors are readily amenable to neutron and X-ray scattering, providing insight into the minute details of atomic structure, electronic density distribution, and elastic and inelastic vibrational properties. In these systems, the long-range periodicity allows integrating the behaviors over multiple unit cells.

However, this approach offers only limited applicability to materials such as nanoscale phase-separated oxides, ferroelectric relaxors, and morphotropic phase boundary systems, incommensurate charge- and spin density wave systems, and, more generally, systems with non-uniform ground states. Similarly, the local mechanisms describing the interplay between chemical disorder, including both lattice-preserving substitution and lattice breaking structural defects, and physical functionalities are often unknown. In all these cases, the lack of long-range translational symmetry limits the applicability of classical scattering techniques and requires the development of methods for probing correlated disorders.

At the same time, the last several years have seen an exponential growth of atomic-scale electron diffraction in scanning transmission electron microscopy (4D-STEM). The fast electrons in the electron probe are deflected by the electric field within the crystal. Negatively charged electrons are attracted to positively charged nuclei, which are screened by the surrounding electrons, meaning they contain sub-atomic scale components. This variation is most clearly seen in diffraction space, where the center-of-mass (COM) of the convergent beam electron diffraction (CBED) pattern is deflected toward the nuclei. Practically, the atomically sized focused electron beam is used to collect the local (2D) diffraction patterns over a dense spatial grid of (2D) points, producing the 4D-STEM data sets. A unique aspect of this method is that the size of the probe can be below the distance between the scatterers, resulting in very complex local diffraction patterns and encoding minute details of the local scattering potential.

Originally, 4D-STEM in its modern form was proposed by Rodenburg as an approach to achieve high spatial resolution, enabling a practical embodiment of the ptychographic idea of Hoppe. However, there were two main difficulties that prevented the widespread adoption of these methods. First, a practical problem was that CCD cameras were not fast or sensitive enough to keep up with the speed of the STEM probe, resulting in long acquisition times creating sample damage and stability problems. The second main problem was that the data sets were too large for existing computer infrastructure and the amount of computation required made it prohibitively expensive. Both of these difficulties have been addressed over the last 4–5 years. Modern computers and their associated storage and data-handling capabilities have improved dramatically in accordance with the well-known Moore’s law. Electron detection capabilities have grown both evolutionarily with incremental improvements in conventional designs and revolutionarily with the advent of direct-electron detectors.

Methods other than ptychography have been developed to analyze scanning nanodiffraction data. The position averaged CBED (PACBED) approach has been used primarily to determine specimen thickness. PACBED has recently been enhanced by the application of deep convolution neural networks to automatically analyze the data sets. Differential phase contrast (DPC) in the STEM was originally proposed in the early 1970s and was recently implemented using segmented detectors. The development of high-speed electron detectors has allowed DPC-STEM
to be readily applied. By determining the deflection of the COM of the CBED pattern as a function of probe position, insight can be gained about the local charge densities and fields or alternatively the electron scattering potential.

Despite these initial advances and the well-recognized promise of 4D-STEM for the sub-atomic scale exploration of materials properties, progress has been stymied by a lack of analysis tools to convert the 4D-STEM data sets into physically relevant parameters. The vast majority of the work presently relies on using a simple COM. Alternatively, a number of approaches using linear unsupervised dimensionality reduction methods such as principal component analysis (PCA) and non-negative matrix factorization (NMF) and clustering techniques have been explored and recently have become part of open-source platforms.

The applicability of linear separation methods for the analysis of 4D-STEM data sets is limited, stemming from the intrinsic symmetries of the atomic lattice. Linear unmixing methods such as PCA will separate Ronchigrams that differ by in-plane rotation only, creating multiple components describing rotational states of nominally identical objects. Similarly, conventional deep neural network architectures employing rigid convolutional layers combined with the distortions and deformations that are universally present in the imaging system and the mesoscale strain fields in the material will give rise to a very large number of weakly meaningful components that do not allow for the direct physical interpretation. Here, we propose an approach for the analysis of 4D-STEM data based on rotationally invariant autoencoders. In general, variational autoencoders (VAEs) are one of the primary classes of generative ML models that seek optimum representation of input high-dimensional data sets in terms of a small number of latent variables. More specifically, VAEs belong to a family of directed latent variable probabilistic models that can infer hidden structure in the underlying data. We assume that each observed data point, \( x_i \), is generated in a non-linear way by some latent variable, \( z_i \), and that the joint probability density of the generative model can be expressed as:

\[
p(x, z) = \prod_{i=1}^{N} p(x_i | z_i) p(z_i).
\]

where \( \theta \) is a global parameter that all datapoints depend on. In VAE, one introduces a variational family of distributions that approximate the true, but intractable posterior distribution, \( q_{\phi}(z|x) \approx p(z|x) \). The latent variable model is then learned by maximizing the evidence lower bound (ELBO) with respect to the model parameters, \( \theta \), and the variational parameters, \( \phi \), for any given datapoint \( x \). The nature of the VAEs dictate that both feature and target data are the encoded data sets.

RESULTS AND DISCUSSION

Application of rrVAE to simulated data

Figure 2 shows a plot of the simulated CBED patterns as a function of probe position for 60 kV electrons incident on graphene. An aberration-free probe with a 31 mrad probe forming aperture is used, which is chosen to be close to that used in the experiment. The CBED patterns are normalized by subtracting the mean CBED pattern as a function of probe position, insight can be gained about the local charge densities and fields or alternatively the electron scattering potential.

An aberration-free probe with a 31 mrad probe forming aperture is used, which is chosen to be close to that used in the experiment. The degree of deflection depends on the closeness of the probe to the atomic site and the electric fields of the other atoms, which leads to many CBED patterns with similar shapes but different rotations. It is this variation that is used in the CO method to reconstruct the electric fields and related quantities. Hence, the relevant question is whether rrVAE allows us to determine the same physical properties and perhaps provide additional insights in the structure of the 4D-STEM data sets.

For rrVAE training, it is important to have a consistent stopping criterion similar to most iterative processes. For the specific configuration used, the convergence of the rrVAE is examined in Supplementary Fig. S1, using the simulated dataset above. The training loss decreases rapidly at first and then gradually flattens
and reduces slowly in a monatomic fashion. While it might (naively) seem that more iterations would provide a better result, the results actually degrade if too many iterations are performed. In many cases, the latent variables appear closely related to the COM deflection map shown in Fig. 3d. In order to provide a robust measure of the correlation between the latent variables and the COM deflections, we use the Pearson correlation coefficient or the Pearson r factor that ranges in value between 1 and −1, with 1 being a perfect positive linear correlation and −1 being a perfect negative linear correlation. A value of zero represents no correlation. We will use the Pearson r factor to determine the number of iterations that provide the strongest correlation for each case we examine and present the corresponding results.

We investigate the application of 3D rrVAE to the simulated graphene data set in Fig. 3. The graphene unit cell used for the simulation is shown in Fig. 3b. The angle and magnitude of the COM deflection are shown in Fig. 3c and d, respectively. The COM magnitude plot has the expected distribution with minima on the atomic sites and the strongest deflections closest to the atoms. The rotation map in Fig. 3e illustrates the rotations of the CBED patterns about the atomic sites, albeit with reversed polarity. We used 1000 iterations in this case. The latent variable observed in Fig. 3e has a strong negative correlation with the COM magnitude plot has the expected distribution with minima on the atomic sites and the strongest deflections closest to the atoms. We will use the Pearson r factor to determine the number of iterations that provide the strongest correlation for each case we examine and present the corresponding results.

Application to simulated graphene CBED patterns. a Latent space for 3D rrVAE of simulated graphene CBED patterns for microscope operating a 60 kV with a 31 mrad probe forming aperture. b Model of the unit cell used for multi-slice simulation. c, d Angle and magnitude of the COM deflection calculated from the simulated CBED patterns, respectively. e Rotation map obtained from rrVAE analysis. Scale bar on (e) is 1 Å. f, g Two latent variable distributions (with Pearson r factor inset).

Fig. 3 Application to simulated graphene CBED patterns. a Latent space for 3D rrVAE of simulated graphene CBED patterns for microscope operating a 60 kV with a 31 mrad probe forming aperture. b Model of the unit cell used for multi-slice simulation. c, d Angle and magnitude of the COM deflection calculated from the simulated CBED patterns, respectively. e Rotation map obtained from rrVAE analysis. Scale bar on (e) is 1 Å. f, g Two latent variable distributions (with Pearson r factor inset).
experiment may need to be explored on a case by case basis. This rebinning should perhaps be best applied at the experimental level (on-chip binning usually results in faster possible readout-speeds, reducing the acquisition time). In addition, to reduce noise we applied PCA as implemented in the scikit-learn Python package28. An illustrative selection of PCA components from the analysis of experimental graphene CBED patterns are shown in Supplementary Fig. S4.

Using this approach, we applied the rrVAE algorithm to the experimental 4D-STEM data obtained from graphene. This is illustrated in Fig. 5 with the simultaneously acquired ADF-STEM image shown in Fig. 5b and the COM deflection angle and magnitude, calculated from the processed data, shown in Fig. 5c and d, respectively. The rotation plot produced by rrVAE is in phase with that derived from the experimental CBED patterns. The second latent variable shown in Fig. 5f shows a low correlation. The latent variable in Fig. 5g has a stronger correlation, but is still quite weak, which is most likely due to the noisy nature of the data. Increasing the number of latent dimensions to 5, as shown in Supplementary Fig. S5, does not provide more clarity, although the overall correlation is similar.

Figure 6 illustrates the effects of defects in graphene over two different length scales. Figure 6a–e shows the ADF-STEM, COM, rotation, and latent variables for graphene with a 3-fold Si impurity over a 1 nm by 1 nm field of view. The Si dopant is obvious in the ADF-STEM image but it is not strong in the COM map. The second latent variable is similar to that observed in the pure graphene case in Fig. 5f. If five latent variables are used, the degree of correlation is very much reduced, as shown in Supplemental Fig. S6. This is most likely due to the noise level. Interestingly, the position of the Si impurity is highlighted in Fig. S6c, suggesting a more careful analysis of the latent spaces may yield more than a COM analog.

Figure 6f–j shows a vacancy in graphene over a 1.5 nm by 1.5 nm field of view. The vacancy is clear in both the ADF-STEM image and COM map shown in Fig. 6f and g respectively. The second latent space, shown in Fig. 6j has a reasonable correlation with the COM map, but little can be seen in the first latent space or rotation map shown in Fig. 6i and h respectively. The expansion to 5 latent spaces, as shown in Supplementary Fig. S7, reduces the maximum correlation with the defect, which is clearly seen in only one space. In general, the presence of noise is better handled with fewer latent spaces.

To summarize, an approach for the analysis of local symmetry breaking via ML analysis of 4D-STEM images has been developed. The rotationally invariant variational autoencoder (rrVAE) approach enables the parsimonious representation of the 4D-STEM data in terms of a small number of latent variables including
the rotation angle. This approach allows the visualization of the structure of the 4D-STEM data sets in terms of a small number of compact maps, thus directly visualizing symmetry-breaking phenomena on the atomic level. While we have limited our examination to experimental parameters appropriate for our microscope, the methodology is applicable to a range of experimental accelerating voltages and aperture sizes, provided they are sufficient for atomic resolution. This approach is able to highlight both a single dopant atom and a single vacancy in monolayer graphene. Interestingly, this result not by examining the high-angle scattered intensity, but through probing the symmetry in the local scattering distribution. This distinction is important because several factors contribute to the ADF-STEM image intensity, making it difficult to distinguish things such as sample thickness changes or surface roughness from intrinsic effects. In the future this method should provide a route to probe defects in cases where there is a small (or no) atomic number difference and to identify visually distinct, but symmetry-related, anomalies. The proposed approach is expected to be universal for the analysis of hyperspectral imaging data sets containing multiple a priori unknown rotational variants. As such, it can be directly applied for a broad range of diffraction methods exploring the 2D diffraction spaces of system, including X-ray ptychography, EBSD, and more complex methods. Beyond exploratory image analysis, this approach provides a universal framework for probing symmetry-breaking phenomena in complex atomic and mesoscopic systems.

Fig. 6 Application to Experimental Graphene with defects. Graphene with a threefold Si impurity and 1 nm field of view after 90 iterations. a The simultaneously acquired ADF image, b The magnitude of the COM, c the rotation map from rrVAE and d, e the two latent variable distributions. The scale bar on (a) is 2 Å. Graphene with a vacancy and 1.5 nm field of view after 55 iterations. f The simultaneously acquired ADF image. g The magnitude of the COM, h the rotation map from rrVAE and i, j the two latent variable distributions. The scale bar on (f) is 5 Å.

METHODS
Materials
Atmospheric pressure chemical vapor deposition (AP-CVD) was used to grow graphene on Cu foil. Polymethyl methacrylate (PMMA) was spin coated on top of the graphene to protect the surface and form a mechanical stabilizer to facilitate the wet transfer to a TEM grid. The Cu foil was etched away in a bath of ammonium persulfate and deionized (DI) water and the graphene/PMMA stack was rinsed in DI water to remove residues. The graphene/PMMA stack was caught on a TEM grid and baked on a hot plate at 150 °C for 15 min. to promote adhesion of the graphene to the grid. After cooling, the grid was immersed in acetone to dissolve the PMMA and then dipped in isopropyl alcohol to remove the acetone and then dried in air. To remove residual hydrocarbon contaminants the sample was baked in an Ar-O2 atmosphere (10% O2) for 1.5 h at 500 °C. Prior to loading the sample into the STEM, the sample and holder cartridge were baked in a vacuum at 160 °C for 8 h.

4D STEM measurements
A Nion UltraSTEM 100 operated at 60 kV was used to acquire the experimental 4D STEM datasets. The CBED images were recorded with an optically coupled Hamamatsu Orca CMOS camera with a 2k by 2k pixel array. The camera was binned to 256 by 256 to increase read out speed. A nominal beam current of 60 pA and a nominal convergence angle of 31 mrad was used. The CBED patterns were acquired with a 7.5 ms dwell time and on a real space mesh of 64 by 64 for a total of 256 CBED patterns.

4D STEM simulation
All CBED patterns were calculated using a modified version of the µSTEM package. Graphene CBED simulations were carried out using the quantum excitation of phonons algorithm. For the simulations containing incoherence, temporal incoherence was added using weighted sum of defocus values over ±100 Å assuming a Gaussian energy distribution with a full width half maximum (FWHM) of 0.35 eV. Spatial incoherence was added using a weighted sum over CBED patterns and a Gaussian source size with a FWHM of 1.3 Å. Simulations for ZnS were done using the absorptive model and included a source size broadening with a FWHM of 0.75 Å. For the ZnS simulations the probe was focused into the midpoint of the crystal.

DATA AVAILABILITY
The datasets generated during and/or analyzed during the current study, along with the Jupyter notebooks performing the analysis are available at https://github.com/markpooley/rrVAE-for-4D-STEM.

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AUTHOR CONTRIBUTIONS

M.P.O. performed 4D-STEM simulations, applied machine learning analysis to data and simulations, produced the figures and co-wrote the paper. M.Z. implemented rotationally invariant VAE and co-wrote the paper. O.D. performed specimen prep, acquired experimental data sets and co-wrote the paper. A.R.L. assisted with data acquisition and paper writing. R.V. assisted with conceptual advances in the deep learning methods applied. S.V.K. proposed the concept, wrote the initial version of the analysis workflow and the first draft of the paper.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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