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A convolutional neural network for defect classification in Bragg coherent X-ray diffraction

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Coherent diffraction imaging enables the imaging of individual defects, such as dislocations or stacking faults, in materials. These defects and their surrounding elastic strain fields have a critical influence on the macroscopic properties and functionality of materials. However, their identification in Bragg coherent diffraction imaging remains a challenge and requires significant data mining. The ability to identify defects from the diffraction pattern alone would be a significant advantage when targeting specific defect types and accelerates experiment design and execution. Here, we exploit a computational tool based on a three-dimensional (3D) parametric atomistic model and a convolutional neural network to predict dislocations in a crystal from its 3D coherent diffraction pattern. Simulated diffraction patterns from several thousands of relaxed atomistic configurations of nanocrystals are used to train the neural network and to predict the presence or absence of dislocations as well as their type (screw or edge). Our study paves the way for defect-recognition in 3D coherent diffraction patterns for material science.

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INTRODUCTION

Defect detection and classification are important issues in material science, as defects strongly influence the properties of materials^{1–4}. Although metallurgy has long recognized the importance of defects for the macroscopic mechanical properties (e.g., such as enhanced yield strength of steel), their more widespread influence in other fields of material science is still lacking detailed understanding. Nevertheless, the concept of strain engineering in a vast variety of functional materials is attracting a lot of attention, opening great opportunities for the design and optimization of the mechanical, optical, electrical, or catalytic properties of materials via deliberate defect manipulation^{5–7}. Crystal defects of various nature and length scales are not always adverse but can instead activate specific functionalities, such as improving adsorption affinity or catalytic activity. For instance, twins and stacking faults can improve the catalytic efficiency of nanoparticles⁸, and more generally the strain generated by defects can affect the catalytic activity⁹. Similarly, the role of dislocations in battery performance has drawn the attention of scientists and could be a key point for further optimization⁶. This defect sensitivity might open new avenues to engineering the properties of nanostructures by introducing specific defects. In order to achieve this goal, it is important to detect and classify defects in nanomaterials to better understand their behaviors (nucleation, propagation, annihilation, defect-defect interaction).

Unlike perfect crystals that can be described as equilibrium structures, the physics and thermodynamics of defects is much harder to describe with the available theoretical tools. It is thus of greatest relevance to supply imaging techniques capable of delivering tomographic reconstructions of the crystal structure in the close environment of defects. Few experimental techniques can achieve this goal. Among them, transmission electron microscopy (TEM) is routinely used to image dislocations in real

space by selecting relevant diffraction vectors, according to established invisibility criteria¹⁰. It has atomic resolution and can directly image individual crystal defects. However, the technique is hindered by several constraints related to sample preparation. These constraints are relaxed for X-rays, which have a great potential to study defects in crystals. With the advent of new generation synchrotron sources with higher coherent flux, a very attractive technique to probe the microstructure of defects has emerged: coherent X-ray diffraction (CXD)^{11,12}. In Bragg geometry, it probes the local deviation from the perfect crystal lattice and is therefore highly sensitive to elastic strain¹³ and crystal defects such as stacking faults¹⁴ or dislocation loops¹⁵. In the past two decades, the technique has been turned into an imaging technique (Bragg Coherent Diffraction Imaging, BCDI), combining measurements of three-dimensional (3D) Coherent X-ray diffraction patterns (CXDPs) with phase retrieval algorithms^{16,17}, to obtain a spatial reconstruction of isolated nanoscale objects¹⁸. The technique has been used successfully to image the strain field in defective nanocrystals^{6,19} including for relatively complex defect configurations^{20,21}, but tends to fail for highly strained systems. In addition, phase retrieval algorithms are relatively slow, while a live evaluation of the data is often required during in situ and operando experiments. This is particularly true in the case of Bragg ptychography, which requires a considerable amount of data. There is therefore an interest in understanding CXDPs qualitatively and interpreting them directly in reciprocal space. Depending on their type and on the measured Bragg reflection, single-crystal defects have indeed a unique signature on CXDPs which enables their identification directly from the reciprocal space data²². For instance, a screw dislocation will lead to a ring-shaped Bragg diffraction signal, if the Burgers vector \mathbf{b} of the dislocation is parallel to the scattering vector at the measured Bragg position, \mathbf{g} .

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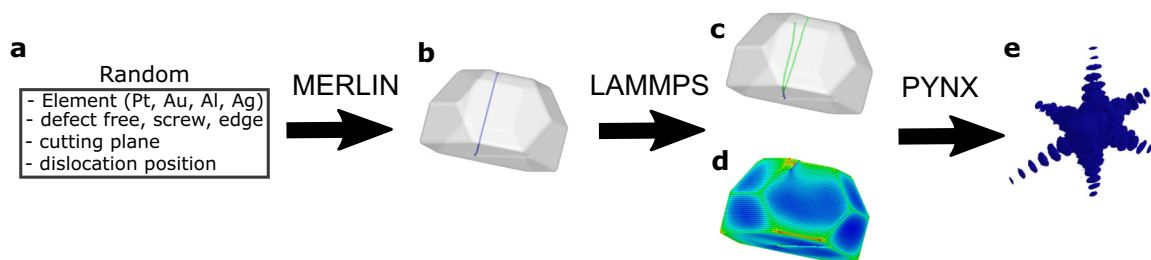


Fig. 1 Schematics of the framework for creating 3D simulated data sets. **a** A random element, class, crystal surfaces, and dislocation position are selected. **b** The corresponding crystal is constructed using MERLIN (Rodney (2000 2010). Merlin in a nutshell (unpublished)) and relaxed using LAMMPS³⁹. **c** The dislocation dissociates into Shockley partials and **d** a strain field builds-up in the crystal. **e** Finally, the corresponding 3D CXDP is calculated with PyNX⁴⁴.

For identifying defects, pattern classification^{23,24} and neural networks (NN) for fault detection²⁵ have been previously used, for example in diffraction phase microscopy. Deep learning has also been used successfully for optical surface-defect detection^{26–28} and for defect segmentation in scanning transmission electron microscopy²⁹. These methods are therefore relevant to detect and classify defects in CXDPs, which are very sensitive to the defect type. The need of extensive training sets and prior data with a different types of defects is one of the main difficulties to overcome with these computational methods. These requirements could potentially limit their performance and practical feasibility. However, with the exponential advancements in computational resources³⁰ and the possibility of ultra-fast atomistic relaxation and computation of diffraction patterns with massive parallelism or graphical processing units (GPUs), it is now straightforward to calculate the 3D CXDPs of single nanocrystals from their atomistic configurations. These configurations can be generated by varying the type and location of the crystal defects and then relaxed by energy minimization. The relaxation of the faulted crystal structure allows to model accurately the crystal defect and has been shown to have a large impact on CXDPs²², leading to a better agreement between the simulated 3D CXDPs and experimental measurements.

While models have been widely applied to generate 2D images, the generation of 3D structures is a nascent field. For example, a deep-learning NN model has been recently successfully developed for the classification of crystal structures from 2D diffraction maps of more than 100,000 simulated crystal structures³¹, but it has the drawback that the 2D diffraction fingerprint is not unique across space groups. Recently, several papers proposed to use deep-learning models trained on simulated CXDPs to perform phase retrieval^{32–36} which is commonly carried out using iterative algorithms. This demonstrates the emergence of deep learning in the field of CXD and BCDI.

In this work, we develop and train a 3D convolutional neural network (CNN), which aims to obtain a fast and precise defect classification in nanocrystals of common face-centered cubic (fcc) transition metals. The training data are generated from atomistic simulations that are representative of the physics of the material. Once trained, the network can predict dislocations on simulated and measured 3D CXDPs. The predictions are categorized in two (defect-free and single dislocation) or three (defect-free, single screw, and edge dislocations) classes. This work paves the way for automated defect detection and its reliable recognition from 3D CXDPs.

RESULTS AND DISCUSSION

Building the data sets

In order to build the data set required for training the neural network (NN), several material simulation tools were used. The data pipeline allows one to generate simulated CXDPs very close

to the ones obtained from Bragg CXD experiments. Figure 1 illustrates our approach for the creation of 3D CXDPs. The geometry considered in this study is derived from the Wulff construction, i.e., the equilibrium crystal shape of a free-standing crystallite obtained by Gibbs thermodynamic principle, which minimizes the total surface free energy associated to the crystal-medium interface³⁷. In order to take into account the presence of a solid-solid interface, i.e., the presence of an underlying substrate as in the experimental nanoparticles, the so-called Winterbottom shape, which can be described as a truncated Wulff construction, is employed³⁸. An example of a simulated crystal is shown in Fig. 1b–d. Only fcc transition metals are considered in this study (Al, Au, Ag, Pt), for which the Wulff/Winterbottom geometries mostly consist of $\{1\ 1\ 1\}$ and $\{1\ 0\ 0\}$ facets. The Winterbottom constructions are generated using the atomistic simulation code MERLIN (Rodney (2000 2010). Merlin in a nutshell (unpublished)), by creating a cube of atoms and cutting it along the $\langle 1\ 1\ 1 \rangle$ and $\langle 1\ 1\ 0 \rangle$ crystallographic directions, the position of the cut planes being defined by the ratio of the surface energies $\gamma_{111}/\gamma_{100}$ and $\gamma_{110}/\gamma_{100}$ of the material/potential of interest. The lattice orientations corresponding to the axes of the simulation cell are $x[1\ 0\ 0]$, $y[0\ 1\ 0]$, and $z[0\ 0\ 1]$ and are kept constant for all configurations. The interface plane is selected randomly among the eight possible $\{1\ 1\ 1\}$ planes, and is cut at a random position corresponding to 65–75% of the height of a free-standing Wulff particle.

Two crystal sizes are considered in this study, the small crystals consist of $40 \times 40 \times 40$ unit cells (Supplementary Fig. 1) while the large crystals are made up of $80 \times 80 \times 80$ unit cells (Supplementary Fig. 2). This corresponds to a size of $15 \times 15 \times (9–12)\text{nm}^3/100,000–140,000$ atoms for the small configurations, and $30 \times 30 \times (19–25)\text{nm}^3/800,000–950,000$ atoms for the large configurations, the height and number of atoms in the crystal depend on the distance of the interface plane with respect to the center of the particle, and on the lattice parameter of the element considered. For the purpose of this study, we focus on line defects, namely, edge and screw dislocations. A single dislocation and its corresponding displacement field (hypothesis of an isotropic and semi-infinite volume, see ref. 22) is introduced following two strategies. In the first type of configuration, hereafter referred to as CD, the dislocation is systematically introduced close to the center, within a range not exceeding 10% of the lateral size of the particle. In the second type of configuration, hereafter referred to as RPD, the position of the dislocation is completely random. The simulated dislocations have a Burgers vector of $\mathbf{b} = \frac{1}{2}[1\ \bar{1}\ 0]$ which is kept constant for all the configurations. This implies that the initial line directions are $\mathbf{t} = [1\ \bar{1}\ 0]$ and $\mathbf{t} = [1\ 1\ \bar{2}]$ for the screw and edge dislocations, respectively. If the Burgers vector and line direction of the dislocations are not varied, the random selection of the interface plane ensures that a large variety of orientations of the dislocation line with respect to the normal of the interface plane is available in the data set as shown in Supplementary Figs. 1 and 2.

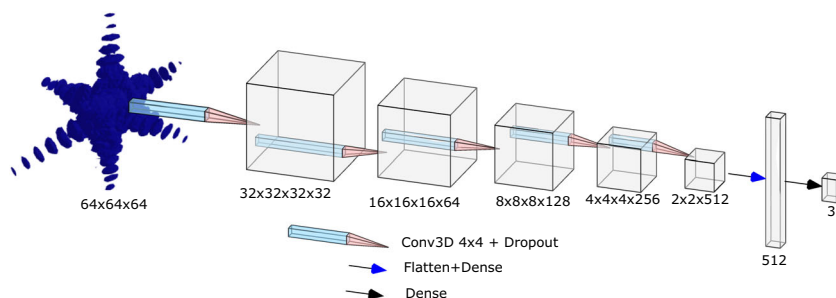


Fig. 2 Schematics of the neural network structure. The NN model consists of five convolution layers with Relu activation function. We use dropout⁴⁶ in all layers with a dropout rate of 0.2, to avoid overfitting. The model ends with 2 fully connected layers with a last softmax activation function. The model takes the 3D $64 \times 64 \times 64$ diffraction as input and predicts the probability for each class (defect-free, screw, edge).

Once the atomistic configurations are generated, the next step is to obtain accurate and realistic relaxed configurations that reproduce as faithfully as possible the displacement fields measured in the experimental particle. To do so, Molecular Statics simulations are carried out with the open-source Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS)³⁹. The interaction between atoms is modeled with different embedded-atom model (EAM) potentials that accurately reproduce elastic properties as well as surface and stacking fault energies, parameters that are essential to get an accurate description of the relaxed defects. For Al, Ag, Au, and Pt, we use the EAM potentials developed by Mishin et al.⁴⁰, Williams et al.⁴¹, Grochola et al.⁴², and Zhou et al.⁴³, respectively.

The crystals are relaxed at 0 K using a conjugate gradient algorithm. If the dislocations introduced close to the center of the nanocrystals are stabilized by the image forces during relaxation, one notable challenge is the tendency of the dislocations introduced close to a free surface to escape the crystal during the energy minimization. In order to prevent this phenomenon, energy tolerance is used as the main stopping criterion for energy minimization. The latter is defined as the energy change between two successive iterations divided by the total energy of the system, and is set to a value of 10^{-6} for the RPD configurations. This value is sufficiently high to ensure that the dislocations dissociate into Shockley partials and remain in the crystal at the end of the relaxation, as shown in Fig. 1c and Supplementary Figs. 1–3. The small number of minimization steps also prevents large rotations of the dislocations during the relaxation. It was indeed observed that edge dislocations are prone to rotate (thus becoming a mixed dislocation) during the energy minimization, especially when they are introduced in the vicinity of the free surfaces. Limiting the number of relaxation steps allows to retain the edge and screw character of the dislocation during the relaxation, even if dislocations very close to the free surfaces tend to have a mixed character as illustrated in Supplementary Figs. 1–3. Each data set typically contains 1000 relaxed configurations with one-third of defect-free nanocrystals, one third containing a relaxed screw dislocation, and the last third with a relaxed edge dislocation. The time required for the energy minimization of a full data set ranges between 10 and 25 min for the small crystal data set and 1 h 30 min and 4h for the large crystal data set.

The last step in the data set creation is the calculation of the three-dimensional CXDPs that are used as input data for our CNN. This is done by summing the amplitudes scattered by each atom with its phase factor, following a kinematic approximation:

$$I(\mathbf{q}) = \left| \sum_j f_j(\mathbf{q}) e^{-2\pi i \mathbf{q} \cdot \mathbf{r}_j} \right|^2, \quad (1)$$

where \mathbf{q} is the scattering vector, $f_j(\mathbf{q})$ and \mathbf{r}_j are, respectively, the atomic scattering factor and position of atom j . Note that the crystallographic convention is used in this manuscript, i.e., the 2π

factor is not included in q , which implies that a given q value corresponds to a real space distance d of $q = 1/d$. The computation is performed with a GPU using the PyNX⁴⁴ scattering package, which considerably speeds up the calculation of the CXDPs. Given a large number of atoms (10^5 – 10^6 atoms) and a large number of CXDPs that are generated for each data set (2000–15,000), the calculations are performed on $64 \times 64 \times 64$ reciprocal space points. The size of the 3D array is a trade-off between achieving a high enough resolution in the reciprocal space, which is required for an accurate comparison with the experimental CXDPs, and keeping the time required to generate the data set reasonable. Using a POWER9 machine, each CXDP is calculated in 0.25 s for the small configurations and 2 s for the large configuration. A data set containing 10,000 CXDPs is therefore typically generated in 40 min for the small nanocrystals and 6 h for the large ones.

In order to introduce enough variation in the data set and prevent overfitting of the model to the training set (Supplementary Fig. 4), each CXDP is rotated randomly around the chosen \mathbf{Q} vector, typically we consider 10 random orientations for each relaxed configuration. The reciprocal space sampling (δq) is also varied, which is equivalent to zooming around the Bragg reflection of interest (Supplementary Fig. 14). A low reciprocal space resolution (coarse sampling/large δq) can have detrimental effects on the accuracy of the network predictions (Supplementary Fig. 14c). To prevent this loss in accuracy, we typically selected δq values for which the oversampling ratio is consistent with the one used for experimental data. Note that even for the largest δq values, the oversampling criteria as defined by Sayre⁴⁵ are still fulfilled, as it is always the case for the experimental CXDPs. Since the simulated particles are significantly smaller than the experimental ones (typically by one order of magnitude), this also implies that a larger portion of the Brillouin zone is selected for the simulated particles. We will see in the following that this has little consequence for the accuracy of the network predictions. Before training the NN, the distribution of dislocation positions is typically estimated by comparing the maximum of the intensity scattered by the atomistic configurations in the data set with the maximum of the intensity scattered by a defect-free crystal with a similar number of atoms (Supplementary Fig. 15).

Convolutional neural network

The NN model architecture is displayed in Fig. 2. It takes as input the $64 \times 64 \times 64$ images of the CXDP intensity and encodes it through a series of convolution and fully connected layers. Dropout⁴⁶ is used in all layers with a dropout rate of 0.2, to avoid overfitting. This is a standard architecture, nevertheless, it already gives very accurate predictions on the simulated data set. Increasing the size of the model, adding extra layers, or increasing the number of filters in the convolution layers does not increase the model efficiency and even leads to an overfit of the training data set in some cases.

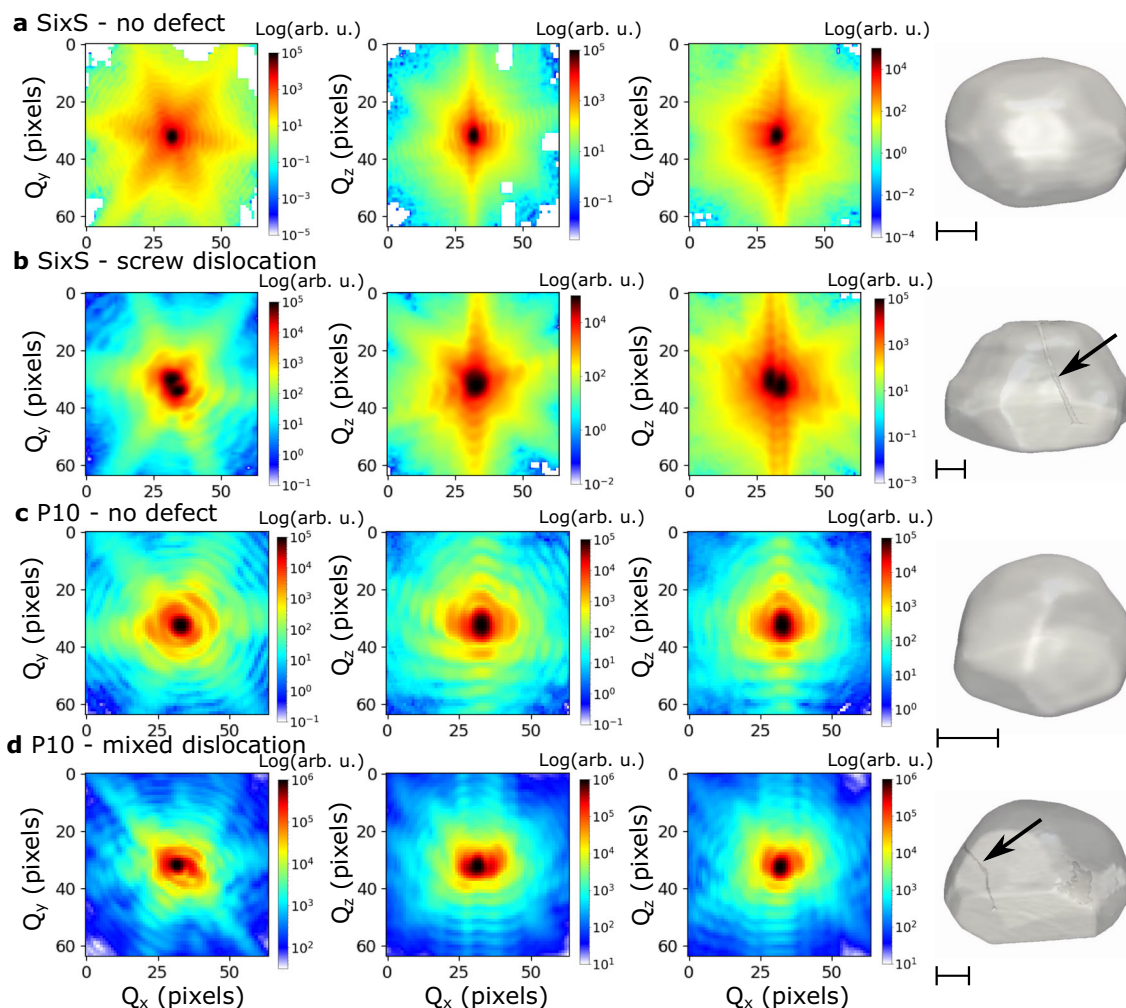


Fig. 3 CXDPs and reconstructed Bragg electron density from Pt NPs measured on the P10 and SixS beamlines. **a** Defect-free NP, SixS beamline **b** Defective NP, SixS beamline **c** Defect-free NP, P10 beamline **d** Defective NP, P10 beamline. The black arrows indicate the location of the dislocations. The scale bar indicates a length of 100 nm. The CXDPs are on a logarithmic scale to enhance the fringes visibility.

Training is performed using Adam optimization⁴⁷ with a learning rate of 10^{-3} and a batch size of 64. A large amount of 3D data sets are simulated. They systematically specify the correct output (defect class) for a given input (3D CXDP intensity), and minimize a categorical cross-entropy loss that quantifies the difference between the predicted and the correct class labels (defect-free, screw, and edge). Through this minimization, the weights (i.e., parameters) of the neural network are optimized to reduce the classification error. The weights of each convolutional and fully connected layers are initialized randomly. Moreover, the instances of the training data set are processed in a random order. Nonetheless, two independent trainings for a given data set a CNN always gives a very similar probability distribution as illustrated in Supplementary Fig. 16. The simulated data are split into training, validation, and test sets. The model fit is performed with the training set and stopped when the validation set accuracy reaches a maximum. The final model prediction on the test set containing 11,556 CXDPs calculated from 1284 atomistic configurations reaches a very high total accuracy score of 97.2%. In addition, the confusion matrix displayed in Supplementary Fig. 7 shows that almost all defect-free crystals are predicted. Most of the errors (4.7%) come from edge dislocations predicted as screw. Furthermore, as illustrated in Supplementary Figs. 11 and 12, a simpler two classes model (Supplementary Fig. 10) predicting either a defect-free or a defective crystal can reach an even higher

accuracy. From an occlusion sensitivity test⁴⁸ on a simulated CXDP shown in Supplementary Fig. 13, we demonstrate that the NN mainly uses the vicinity of the Bragg peak to make its prediction.

Validation on experimental data

The experimental data sets correspond to 3D reciprocal space maps obtained by measuring the Bragg CXDPs of Pt nanoparticles. Single particles were measured either at the SixS beamline of synchrotron SOLEIL (Orsay, France) or at the P10 beamline of synchrotron PETRA (Hamburg, Germany). The 3D Bragg CXDPs were collected at the asymmetrical $\bar{1}11$ Pt Bragg reflection at the SixS beamline or at the symmetrical (specular) 111 Pt Bragg reflection at the P10 beamline. The experimental reciprocal space data sets have been orthonormalized using the xrayutilities package⁴⁹. Figure 3 displays the CXDPs of the experimental data sets, as well as their reconstructed Bragg electron density using phase retrieval algorithms. Defect-free (Fig. 3a, c) as well as defective crystals (Fig. 3b, d) were measured. A closer look at Fig. 3b, d reveals the variety of dislocation configurations that are found in experimental nanocrystals. These dislocations were most likely nucleated during the growth of the nanoparticles, and did not escape during the annealing at 1100°C , suggesting that they are strongly pinned in the nanocrystal. For the SixS data, the screw dislocation is close to the center of the nanocrystal (Burgers vector of $\mathbf{b} = \frac{1}{2}[1\bar{1}0]$). On the other hand, the dislocation in the P10

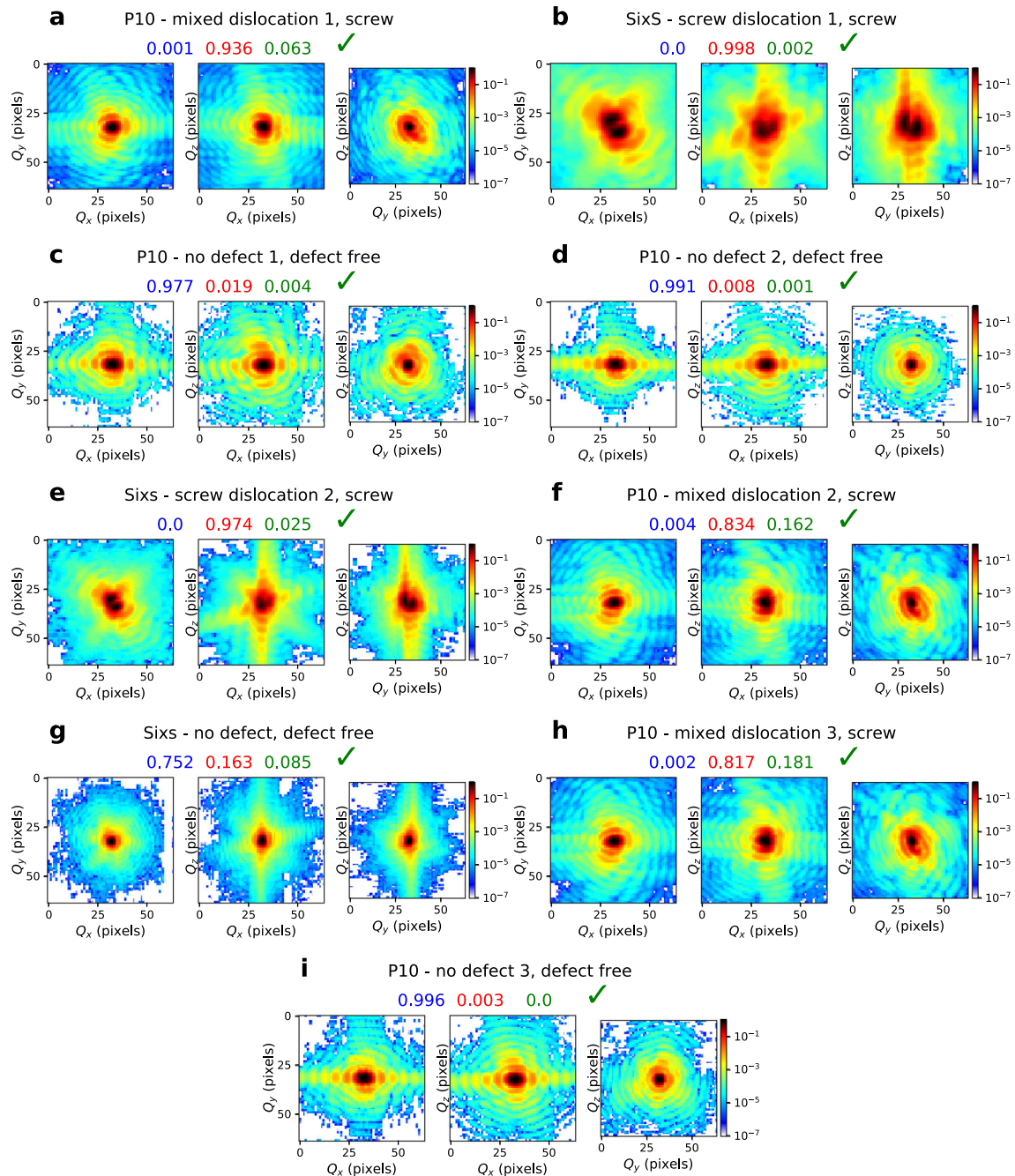


Fig. 4 Predictions from the best model on experimental data. For each example, the probabilities for the 3 classes are shown in the title where blue, red, and green correspond respectively to defect-free, screw, and edge class. Three cross-sections of the 3D CXDP are shown for each example. Using this NN model, all examples are correctly predicted (see green ticks). The CXDPs are on a logarithmic scale to enhance the fringes visibility.

defective nanocrystal is closer to the free surfaces. In addition, the dislocation line is not perfectly straight and parallel to the Burgers vectors ($\mathbf{b} = \frac{1}{2}[101]$). It can thus be described as a mixed dislocation with a dominant screw character.

In order to reinforce the agreement between the simulated and experimental data sets each diffraction measurement is pre-processed before computing the model prediction. The CXDP center of mass is placed at the center of the array, as it is also the case for the simulated data. Finally, the CXDP is normalized so that the maximum is equal to 1.

The results of our best NN model on the preprocessed CXDPs are displayed in Fig. 4 along with slices along Q_x , Q_y , and Q_z for

each experimental CXDP. Some crystals were measured several times under different experimental conditions (temperature, gas environment), for example, P10 (no defects 1, 2, and 3 in Fig. 4), allowing us to compare the model predictions for the same crystal but with slightly different CXDPs.

The performances of this model on experimental data are excellent, all the experimental examples being predicted in the correct class, and most of them with a very high probability (>95%). Although still very good, the predictions for the P10 data (mixed dislocation) are generally slightly worse with an accuracy ranging between 82% and 94%. This is not surprising given the mixed type of dislocation (with a dominant screw character),

Table 1. Predicted probabilities in % on the experimental data from several models trained with different simulated training data sets.

Experimental example	P10 mixed 1	SixS Screw 1	P10 no defect 1	P10 no defect 2	SixS Screw 2	P10 mixed 2	SixS no defect	P10 mixed 3	P10 no defect 3
Training data set									
Pt-unrelaxed small crystals CD	<i>p 99</i> <i>s 1</i> <i>e 0</i>	<i>p 0</i> <i>s 3</i> <i>e 97</i>	p 100 s 0 e 0	p 100 s 0 e 0	<i>p 0</i> <i>s 1</i> <i>e 99</i>	<i>p 97</i> <i>s 2</i> <i>e 1</i>	p 100 s 0 e 0	<i>p 100</i> <i>s 0</i> <i>e 0</i>	p 100 s 0 e 0
Pt-relaxed small crystals CD	<i>p 99</i> <i>s 1</i> <i>e 0</i>	p 0 s 100 e 0	p 100 s 0 e 0	p 100 s 0 e 0	p 0 s 69 e 31	<i>p 72</i> <i>s 18</i> <i>e 10</i>	p 72 s 16 e 12	<i>p 92</i> <i>s 6</i> <i>e 2</i>	p 100 s 0 e 0
Pt-relaxed big crystals CD	<i>p 99</i> <i>s 0</i> <i>e 1</i>	p 0 s 100 e 0	p 100 s 0 e 0	p 100 s 0 e 0	p 25 s 38 e 37	<i>p 99</i> <i>s 0</i> <i>e 1</i>	p 99 s 0 e 1	<i>p 99</i> <i>s 0</i> <i>e 1</i>	p 100 s 0 e 0
Multi-elements relaxed big crystals CD	<i>p 98</i> <i>s 0</i> <i>e 2</i>	p 0 s 100 e 0	p 100 s 0 e 0	p 100 s 0 e 0	p 0 s 98 e 2	<i>p 88</i> <i>s 5</i> <i>e 8</i>	p 99 s 0 e 1	<i>p 94</i> <i>s 2</i> <i>e 3</i>	p 100 s 0 e 0
Multi-elements relaxed big crystals RPD	p 9 s 53 e 38	<i>p 0</i> <i>s 36</i> <i>e 64</i>	p 100 s 0 e 0	p 100 s 0 e 0	<i>p 0</i> <i>s 8</i> <i>e 92</i>	p 0 s 58 e 42	p 99 s 1 e 0	p 0 s 63 e 37	p 100 s 0 e 0
Multi-elements-relaxed big crystals 75% CD, 25% RPD	p 0 s 94 e 6	p 0 s 100 e 0	p 98 s 2 e 0	p 99 s 1 e 0	<i>p 0</i> <i>s 97</i> <i>e 3</i>	p 0 s 84 e 16	p 75 s 16 e 9	p 0 s 82 e 18	p 100 s 0 e 0

In each cell, the prediction probability for the 3 classes (perfect: p, screw: s, edge: e) is shown in bold if the prediction is correct and in italics if it is wrong. CD and RPD stand for centered dislocations and random position dislocations, respectively.

which necessarily increases the probability of identifying the defect as an edge dislocation. Nonetheless, even if the dislocation is located close to a free surface and therefore induces weak distortions in the CXDP (Fig. 3d), our model still manages to identify this crystal as defective with almost a 100% probability. This demonstrates the robustness of the model trained on this data set, which can predict both centered and off-centered dislocations with very high accuracy.

The simulated training data set used to fit the NN model has a large influence on the accuracy of the predictions on experimental data. This data set must contain enough diversity, while sharing enough similarities with the experimental CXDPs. The predicted probabilities on experimental data for the same model architecture but different simulated training data sets are shown in Table 1. Six different simulated data sets have been trained: (1) single element (Pt) unrelaxed small crystals, 100% centered dislocations (CD), (2) relaxed Pt small crystals (100% CD), (3) relaxed Pt large crystals (100% CD), (4) relaxed large crystals with multiple elements (Au and Pt) (100% CD), (5) relaxed multi-elements large crystals with dislocations at random position (100% RPD) and (6) relaxed multi-elements large crystals with a mix of CD and RPD configurations (75% CD and 25% RPD). The first two rows of Table 1 emphasize the importance of accurately modeling the displacement field of the dislocations. Indeed, while these two models trained on relaxed and unrelaxed data sets predict accurately the defect-free configurations, they fail at identifying the mixed dislocation (P10 data). However, the model trained on the relaxed data set performs much better on the SixS-“screw” data, which is correctly identified as a screw dislocation (see also Supplementary Fig. 9). On the other hand, the size of the relaxed crystals does not have a major impact on the accuracy of the model (Table 1, second and third row), although the predictions of the models trained on the large configurations is slightly more accurate, in particular for defect-free configurations (Supplementary Tables 2 and 6).

The addition of several elements in the data set improves the accuracy of the predictions for the SixS data, but has no effect on the P10 data (Table 1, fourth row). Nonetheless, mixing several elements in the data set generally results in better model

predictions compared to the models based on single elements, in particular for the large crystal size (Supplementary Tables 5–8). The position of the dislocation also has a major impact on the model predictions. As seen from Table 1 (sixth row), introducing the dislocation at random positions, including positions close to the crystal-free surfaces, results in more accurate predictions for the P10 data. However, this improvement is at the expense of the predictions for the SixS data, which is correctly identified as a dislocation, but with an edge character instead of a screw. The predictions for the defect-free data are not affected and still excellent (see also Supplementary Fig. 8).

In order to obtain accurate predictions simultaneously for both P10-“mixed” and SixS-“screw” dislocations, one must increase the diversity in the training data set. This has been achieved by building a data set consisting of a mix of CD and RPD configurations (Table 1, seventh row). Training the CNN on this mixed data set significantly enhances the performances of the model and allows to predict correctly and with very high accuracy all the experimental examples.

We must emphasize that, despite the differences in the ability of the models to generalize to experimental data, the accuracy on the simulated test data for each training data set is always higher than 86% (Supplementary Tables 1 and 5). Our work illustrates the necessity of using a simulated trained data sets close to real structures: atomistically relaxed nanoparticles with accurate modeling of the dislocation displacement field, multiple atomic elements, and random location of the dislocations. It also demonstrates that a convolutional neural network can predict dislocations in a crystal from its 3D coherent diffraction pattern. Combined with the fast scanning capabilities of some synchrotron beamlines⁵⁰, this approach could be used to perform a fast screening of the nanocrystals on a sample of interest. This would allow to determine the proportion of defect-free nanocrystals as well as nanocrystals containing a specific type of crystal defect, and select the best candidate for a coherent diffraction imaging experiment. In addition, if the CNN was only tested on metallic fcc particles, we foresee that it could be extended to more complex systems like for instance multi-element particles.

From 3D coherent X-ray CXDPs, we used a convolutional neural network to predict defect classes. As a result, we obtain an automatic procedure for defect classification in fcc metals, which does not require any user manipulation, any intensive live data mining, and achieves high-accuracy classification even in the presence of defects close to the free surfaces of the nanocrystals. This tool can be exploited during experiment execution to provide rapid feedback to the investigator, enables one to identify on the fly target defect types present in individual nanocrystals, and furthers the possibility of unsupervised data collection, extremely relevant given the increases data rates expected at ever-improving facilities. Our study paves the way for defect recognition of three-dimensional structural data in big-data materials science.

METHODS

Training the network

We used the python deep-learning API Keras⁵¹ running the TensorFlow backend⁵² to build, develop and train our NN. The training was performed in parallel on two NVIDIA Tesla V100 GPUs and a POWER9 computer. We use a categorical cross-entropy loss function $L(y, \hat{y}) = -\frac{1}{B} \sum_{n=1}^B \sum_{c=1}^{N_c} y_{n,c} \log(\hat{y}_{n,c})$, where B is the batch size, N_c the number of classes, $y_{n,c} = 1$ for data element n if the true class is c and $y_{n,c} = 0$ otherwise. $\hat{y}_{n,c}$ is the predicted probability for class c . The simulated data set is divided into training, validation, and test, corresponding respectively to 85%, 10%, and 5% of the total data set. The model is trained with a learning rate of 10^{-3} and a batch size of 64 on the training set until the model accuracy calculated on the validation set reaches a plateau (Supplementary Figs. 5, 6, and 11). A typical training requires between 15 and 30 min depending on the data set (8–10 s per epoch and 100–200 epochs). Decreasing the learning rate and increasing the batch size do not further improve the model accuracy. Once trained, the model performance is evaluated on the test set and reaches a total accuracy >86% on the simulated data for all models presented in Table 1.

Sample growth

Pt nanocrystals were prepared by the solid-state dewetting of a 30-nm thin Pt film for 24 h at 1100 °C in air⁵³. The Pt film was deposited on α -Al₂O₃ (sapphire) with an electron beam evaporator. The Pt nanocrystals have their c -axis oriented along the [111] direction normal to the (0001) sapphire substrate. A standard photolithography method was employed to prepare a patterned layer of photoresist on sapphire prior to the electron beam evaporation of Pt. The lithographic processing route ensured that a number of dewetted Pt particles are well-separated from their neighbors and that only one crystallite is irradiated by the incoming X-ray beam. The particle size ranges from 100 to 700 nm.

DATA AVAILABILITY

The data supporting the findings of this work are available from the corresponding author on reasonable request.

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AUTHOR CONTRIBUTIONS

B.L. and M.D. performed the atomistic simulations. E.B. developed the convolutional neural network. M.-I.R. processed the experimental data. E.R. and E.A. prepared the samples. M.D., M.-I.R., S.L., A.R., and A.C. performed the experiments at the SixS beamline of synchrotron SOLEIL (experiment number: 20181680), and M.D., M.-I.R., S.L., and M.S. performed the experiments at the P10 beamline of synchrotron PETRA (experiment number: I-20180962). B.L., E.B., M.D., and M.-I.R. wrote the manuscript and all authors reviewed the manuscript. B.L., E.B., and M.D. contributed equally to this work and are considered “co-first authors” of this manuscript.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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