Segmentation of Crystal Defects via Local Analysis of Crystal Distortion

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Abstract: We propose a variational method to simultaneously detect dislocations and grain boundaries in an image of a crystal as well as the local crystal distortion. To this end we extract a distortion field $F$ from the image which is nearly curl-free except at dislocations and grain boundaries. The sparsity of the curl is promoted by an $L^1$-regularization of curl $F$ in the employed energy functional. The structure of the functional admits a fast and parallelizable minimization such that fairly large images can be analyzed in a few minutes.

1 INTRODUCTION

Imagine we are given a large transmission electron microscopy image or a simulated snapshot of a crystalline structure as in Fig. 1 left, and we want to detect all crystal defects, statistically analyze the size of defect-free regions, and evaluate correlations between crystal orientations in different regions, perhaps even for a whole time sequence of such images. Such tasks are of fundamental importance in materials science and physics to better understand the behavior and properties of crystalline materials such as metals, to validate mathematical models of such materials, to fit model parameters, or even to optimize material parameters. Often, the detection of crystal defects and crystal orientation is accomplished manually, e. g. (Wu and Voorhees, 2012), obviously with limited feasibility for larger images or sets of data.

We here devise a new automatic crystal segmentation method which yields the crystal defects, the local crystal orientation at each point, and the local crystal strain and which can be computed efficiently via a split Bregman iteration. The algorithm permits parallelization on a graphics processing unit (GPU) and thereby allows for the automatic analysis of images of a size as in Fig. 1 within few minutes.

We here restrict ourselves to two spatial dimensions which is sufficient for the classical application of examining microscope images. A generalization to three dimensions is straightforward.

The two subsequent sections provide the necessary background on crystal structures and introduce existing related methods before we present our model and algorithm in Sec. 2 and 3 as well as some numerical examples and a discussion in Sec. 4 and 5.

1.1 Grain Boundaries and Dislocations

We restrict our discussion to two dimensions. In a perfect atomic crystal, atoms are arranged in a Bravais lattice: the atom positions are given by $n_1a_1 + n_2a_2$ for two fixed vectors $a_1, a_2 \in \mathbb{R}^2$ and all integers $n_1, n_2$. Real materials form polycrystals, that is, crystalline regions (so-called grains) of different orientations, meeting at grain boundaries (Fig. 2 left). These grain boundaries are often denoted low or high angle boundaries, depending on the orientation mismatch between the adjacent grains.

Another type of crystal defect is a dislocation, where a line of atoms terminates inside a crystal (Fig. 2 middle). If we draw a closed curve (a so-called Burgers circuit) on the perfect lattice and map it onto the crystal around the dislocation (Fig. 2 right), a gap remains whose direction and length define the so-called Burgers vector. Low angle grain boundaries are typically composed of a sequence of dislocations (see e. g. Fig. 1).
Figure 1: Given a crystal image (left), our method computes a distortion map $F : \Omega \rightarrow \mathbb{R}^{2 \times 2}$ whose curl magnitude indicates crystal defects (second image; single spots are dislocations, dotted and continuous lines are low and high angle grain boundaries). Polar decomposition of $F$ yields the local crystal orientation (third image, $0 \leq \pi/3$), and $\text{dist}(F, \text{SO}(2))$ can be taken as a measure of crystal strain (right, $0 \leq 0.1$).

Physicists are interested for instance in how defects move and grain boundaries behave or in how the grain size distribution evolves over time. To this end crystal images are often analyzed manually.

1.2 Related Methods

Several efficient approaches for the automatic analysis of crystals do already exist, but they either only apply to molecular dynamics simulation results or they only provide a segmentation into different grains but ignore crystal strains or dislocations.

In (Berkels et al., 2008), the authors perform a piecewise constant Mumford–Shah segmentation of a crystal image $u : \Omega \rightarrow \mathbb{R}$ into disjoint regions $\Omega_i \subset \Omega$ of constant crystal orientation $\alpha_i \in [0, 2\pi)$, using a Chan–Vese level set approximation. The model can be augmented by a global deformation $\phi : \Omega \rightarrow \mathbb{R}^2$ describing the elastic crystal deformation in each region. This approach involves a large number of nonlinearly coupled degrees of freedom (the orientations $\alpha_i$, multiple level set functions, the deformation $\phi$) and thus is rather slow; also it concentrates on crystal grains and ignores the structure of their boundaries or single dislocations. Restricting to unstrained grains, the authors in (Boerdgen et al., 2010) were able to give a convex formulation of the above Mumford–Shah segmentation problem based on functional lifting (instead of using the orientation map $\alpha : \Omega \rightarrow [0, 2\pi)$ one rephrases the functional in terms of $I_\alpha : \Omega \times [0, 2\pi] \rightarrow [0, 1]$, $I_\alpha(x, \gamma) = 1$ if $\alpha(x) \geq \gamma$ and 0 else). They also propose a more efficient version in which the interfaces between the $\Omega_i$ are penalized according to the local angle difference, a method which can be improved further by taking into account the periodicity of the angle (Strekalovskiy and Cremers, 2011). A GPU implementation of this approach is fast, but needs lots of memory due to the lifted variable $I_\alpha$.

The method in (Singer and Singer, 2006) assumes a rather clean crystal image with a perfectly regular, unstrained grain interior. In that case the grain orientation can be found very rapidly by first convolving with a wavelet resembling a grid cell and then locally smoothing out the oscillatory pattern in the convolution. The resulting grey value indicates the average mismatch between the crystal and the wavelet and can thereby be identified with a crystal orientation. However, for slightly strained crystals this identification is erroneous; also one cannot distinguish between crystal orientations which equally mismatch the wavelet. The method seems to incidentally also identify dislocations by a different gray value.

A fast, non-variational dislocation detection method for three-dimensional molecular dynamics data is proposed in (Stukowski and Albe, 2010a). They first identify perfectly crystalline regions by a nearest neighbor analysis and then find shortest Burgers circuits around the non-perfect regions. This yields the Burgers vector and location of all dislocations. The method gains in robustness (but can no longer be used on the fly) by a technique of sweeping out tubes around the dislocation lines in 3D (Stukowski and Albe, 2010b). Additionally the authors identify and triangulate grain boundaries which are not composed of strings of dislocations. A similarly efficient method is introduced in (Begau et al., 2012). Like our approach, it is based on identifying the curl of the local crystal distortion, which in
a molecular dynamics simulation is readily obtained from the positions of the nearest neighbor atoms.

In contrast to the above approaches, our method simultaneously finds dislocations, grain boundaries, and crystal distortion, just based on a crystal image.

2 VARIATIONAL MODEL

In this section we explain how to identify crystal defects with curl concentrations of a distortion map which arises from interpreting an imperfect crystal as a deformed perfect crystal. We then introduce a simple energy functional whose minimization leads to the detection of such curl concentrations.

2.1 Defects as Curl Concentrations

Imagine that a domain $\Omega \subset \mathbb{R}^2$ is occupied by a perfect, unstrained crystal without defects. If this crystal is deformed according to a smooth deformation $\phi : \Omega \to \mathbb{R}^2$, which maps each position $x$ inside the crystal onto a new position $\phi(x)$, then the deformed crystal is still defect-free, though it might be strained. A polycrystal or a crystal with dislocations has a defect-free, albeit strained, crystal structure away from grain boundaries and dislocations. In those regions it can thus be interpreted as a smoothly deformed perfect crystal. Finding crystal defects hence amounts to the identification of places where such an interpretation is not possible. How to accomplish this task?

In their variational approach (Berkels et al., 2008), the authors attempt to directly find the global deformation $\phi$ together with a segmentation of the crystal into different regions across whose boundaries $\phi$ is discontinuous. These boundaries can then be interpreted as grain boundaries, separating defect-free crystal regions from each other. However, searching a joint global segmentation and global deformation is a highly nonlinear problem which makes the method rather slow. Instead, it would be advantageous to identify crystal defects based only on the local crystal structure.

Assume, at each point $x$ inside the crystal we know the local crystal distortion, that is, the local amount of rotation, stretching, and shearing (as compared to an unstrained perfect crystal). This distortion can be described by a linear map $F(x) \in \mathbb{R}^{2 \times 2}$. In case of a smoothly deformed perfect crystal, $F : \Omega \to \mathbb{R}^{2 \times 2}$ is given by the gradient of the crystal deformation $\phi$, $F = \nabla \phi = (\nabla \phi_1, \nabla \phi_2)^T$. In order to identify places where the crystal cannot be described by a smooth deformation of an unstrained perfect crystal, we now have to find those regions where $F$ cannot be interpreted as the gradient of a deformation, that is, where $F$ is not conservative. It is well-known that a tensor field $F$ is conservative if and only if it is curl-free,

$$\text{curl} F = (\partial_x F_{22} - \partial_y F_{12}, \partial_x F_{11} - \partial_y F_{21}) = 0.$$  

The curl-free regions thus represent the defect-free interior of crystal grains, while grain boundaries and dislocations are characterized by nonzero curl $\text{curl} F$.

In particular, if the boundary between two grains is sharp, $F$ is simply discontinuous there and thus has a measure-valued nonzero curl concentrated along the boundary, while real crystals typically exhibit a thin transition region across which the nonzero curl is spread out. Likewise, the curl of $F$ at a dislocation behaves like a $\delta$-peak. Its strength $\int \text{curl} F \, dx$ can be computed via a line integral around the dislocation,

$$\int_B \text{curl} F(x) \, dx = \int_{\partial B} F n^+ \, dx,$$

where $B$ is a ball around the dislocation and $n^+$ its unit outward normal rotated counterclockwise by $\frac{\pi}{2}$. The integrand describes how the curve $\partial B$ is locally stretched by the distortion $F$, and a nonzero integral value shows that if one traces the distorted curve one does not arrive at the starting point, but under- or over-shoots according to the dislocation’s Burgers vector. Indeed, $\int_{\partial B} F n^+ \, dx$ has the interpretation of the vector spanning the gap in Fig. 2 right.

Fig. 3 shows the curl concentrations in the scales pattern of a synthetic reptile skin. These are clearly located in places where an additional row of scales squeezes in between two rows. As discussed above, the shown direction of the curl is related to the Burgers vector and indicates the direction in which the additional row of scales is inserted. A red dot for instance implies that $\int_{\partial B} F n^+ \, dx$ points upwards, thus there is vertical dilation left and compression right of the dislocation so that additional scales are inserted left of it.
2.2 Variational Curl Segmentation

Given an image \( u : \Omega \rightarrow \mathbb{R} \) or some other measurement of a crystal structure, we would like to extract the distortion field \( F : \Omega \rightarrow \mathbb{R}^{2 \times 2} \), which we expect to have sparse curl, concentrated at dislocations and grain boundaries.

We shall follow the usual paradigm in variational image processing and devise an energy functional as the sum of a fidelity or fitting term \( E_{\text{fit}}[F] \), which penalizes a mismatch between \( F \) and the given data \( u \), and a regularization term, which incorporates a priori knowledge of the structure of \( F \) and correspondingly regularizes \( F \) in the case of noisy input data. For regularization we propose the \( L^1(\Omega) \)-norm of \( \text{curl} F \) since the \( L^1 \)-norm is expected to promote sparseness. Thus we arrive at an energy of the form

\[
E_{\text{fit}}[F] + \omega_1 \| \text{curl} F \|_1
\]

for some positive weight \( \omega_1 \). As for the fitting term, we restrict ourselves to images \( u \) as input data, in which case we can use the term proposed in (Boedgen et al., 2010),

\[
E_{\text{fit}}[F] = \int_{\Omega} \sum_k (u(x + F(x)v_k) - u(x))^2 \, dx,
\]

where \( v_1, \ldots, v_K \in \mathbb{R}^2 \) represent a stencil of lattice vectors such that in an image \( u \) of a perfect reference crystal we have \( u(x + v_k) = u(x) \) for all \( x \) and \( k = 1, \ldots, K \). The stencil elements \( v_k \) can for instance be chosen to agree with the \( K \)-fold rotational symmetry of the crystal by

\[
v_k = \ell (\cos(\frac{2\pi k}{K}), \sin(\frac{2\pi k}{K}))^T,
\]

where \( \ell \) represents the lattice spacing or characteristic period of the input image. In the case of a hexagonal crystal as in Fig. 1 we have \( K = 6 \).

Unfortunately, from an analytical viewpoint, the regularization \( \| \text{curl} F \|_1 \) in combination with a non-convex fitting term does not provide sufficient compactness to ensure existence of energy-minimizing tensor fields \( F \), even away from crystal defects, where a classical minimizer would be expected. Also, experimental results are improved by an additional smoothing term so that for our crystal segmentation problem we finally choose to minimize the energy

\[
E[F] = E_{\text{fit}}[F] + \omega_1 \| \text{curl} F \|_1 + \omega_2 \| \nabla F \|_2^2
\]

with weights \( \omega_1, \omega_2 > 0 \). For images with gray values between 0 and 255, the choice \((\omega_1, \omega_2) = (4000\ell, 8000\ell)\) seems to generally yield satisfactory results.

3 MINIMIZATION ALGORITHM

Here we describe the algorithm used to segment images based on the energy (2). We begin by describing some of the properties of the model and make the observation that we seek a local minimizer of (2). For this reason, a good initial guess for \( F \) is required. We propose one simple technique for initializing \( F \). Then we discuss the split Bregman formulation of (Goldstein and Osher, 2009) for problems with \( L^1 \)-type structure and indicate how we apply it to the proposed segmentation model.

3.1 Initialization of \( F \) as Rotations

The fitting term (1) makes the variational energy (2) nonconvex. It can be readily seen that the global minimizer of (2), \( F \equiv 0 \), is not of interest. Instead, we are interested in finding a local minimizer of (2) which is close to an appropriately-chosen initial guess \( F_0 \).

Two observations guide our choice of \( F_0 \): The images of interest consist of large defect-free, perfectly crystalline regions, and in those regions the local crystal distortion can be very well described by a mere rotation.

We choose \( F_0(x) \) at each \( x \in \Omega \) to be the rotation matrix

\[
F_0(x) = \left( \begin{array}{cc} \cos(\alpha(x)) & -\sin(\alpha(x)) \\ \sin(\alpha(x)) & \cos(\alpha(x)) \end{array} \right) = R_{\alpha(x)},
\]

parametrized by the angle \( \alpha(x) \) which approximately minimizes (1). To this end we consider a discrete set of \( p \) angles, \( \alpha_j = \frac{2\pi j}{p} \) with \( j = 0, \ldots, p - 1 \). At each pixel \( x \) we now identify that rotation \( R_{\alpha_j} \) which minimizes the integrand of (1) and either set \( F_0(x) = R_{\alpha_j} \) or \( F_0(x) = R_{\alpha^*} \) where \( \alpha^* \) minimizes the quadratic interpolation of \( \sum_k (u(x + R_{\alpha_j}v_k) - u(x))^2 \) at \( \alpha = \alpha_j, \alpha_{j+1} \) (the subscript \((j \pm 1)\) is to be interpreted mod \( p \)). In practice, we find it sufficient to choose \( p \sim 24/K \).

3.2 \( L^1 \)-Type Minimization

The split Bregman formulation of (Goldstein and Osher, 2009) applies the Bregman iteration idea proposed in (Bregman, 1967) and popularized by (Osher et al., 2005) to a convex energy composed of both \( L^1 \) and \( L^2 \)-type terms. The idea is to introduce a new variable \( \psi \) which corresponds to the \( L^1 \)-type term and then to add the appropriate constraint term to enforce this correspondence. Suppose \( E_{c}[v] = \| \zeta[v] \|_1 + H[v] \), with both \( \| \zeta[v] \|_1 \) and \( H[v] \) convex. This approach advocates solving

\[
\min_{v, \psi} \| \psi \|_1 + H[v] \quad \text{subject to} \quad \zeta[v] = \psi
\]
via Bregman iteration, which is in this context given by
\[
\{v^{k+1};\psi^{k+1}\} = \arg\min_{v,\psi} \|\psi\|^2 + H[v] + \lambda \|\psi - \zeta[v] - b^k\|^2_2, \tag{3}
\]
\[
b^{k+1} = b^k - \psi^{k+1} + \zeta[v^{k+1}]. \tag{4}
\]
The split Bregman simplification is to perform alternating minimization on \(v^{k+1}\) and \(\psi^{k+1}\), which allows for \(v^{k+1}\) to depend only on \(L^2\)-type terms, and gives an explicit form for \(\psi^{k+1}\):
\[
v^{k+1} = \arg\min_{v} H[v] + \lambda \|\psi - \zeta[v] - b^k\|^2_2, \tag{5}
\]
\[
\psi^{k+1} = \text{shrink}(\zeta[v^{k+1}] + b^k, \frac{\lambda}{2}), \tag{6}
\]
\[
b^{k+1} = b^k - \psi^{k+1} + \zeta[v^{k+1}], \tag{7}
\]
with
\[
\text{shrink}(z, \omega) := \frac{z}{\omega} \max(|z| - \omega, 0).
\]

Theorems proven in (Osher et al., 2005) and (Goldstein and Osher, 2009) guarantee that if the minimization problem (3) can be solved exactly, a minimizer of \(\|\psi\|_1 + H[v]\) is found which satisfies the constraint \(\psi = \zeta[v]\), i.e. one which solves the original minimization problem. The split Bregman simplification suggests that a single alternating minimization step for \(v^{k+1}\) and \(\psi^{k+1}\) is taken to approximate (3), however very good results and fast convergence are still observed in practice. The advantage is that the original energy, which had both \(L^1\)- and \(L^2\)-character, has now been reduced to a standard convex \(L^2\) minimization problem to advance \(v\) and to a pointwise and explicitly known update in the new variable \(\psi\).

The minimization problem considered here does not satisfy the hypothesis of the theorem, as \(E_{\text{in}}[F]\) is nonconvex, but the Bregman scheme appears to be very successful at locating a local minimum of our energy anyhow. We set
\[
H[F] = E_{\text{in}}[F] + \omega_2 \|\nabla F\|^2_2, \\
\zeta[F] = \omega_1 \text{curl}(F)
\]
and apply the update (5)-(7).

In order to perform the update (5), we apply the Fletcher–Reeves nonlinear conjugate gradient method with Armijo stepsize control, as Hessian-based methods are not expected to (and in practice do not) provide much benefit for the nonconvex fitting energy \(E_{\text{in}}\) which also requires interpolation of the image \(u\). We employed biquadratic interpolation which does not in general guarantee a continuous interpolation, much less one which is continuously twice differentiable.

4 NUMERICAL EXPERIMENTS

Fig. 1 already showed a result of our method, applied to a snapshot from a phase field crystal simulation (PFC, see e.g. (Elder and Grant, 2004)) with hexagonal crystals. Our method provides the crystal distortion field \(F\) at each pixel from which useful and intuitive output can readily be derived: The locations of high \(|\text{curl}F|\) represent crystal defects (either dislocations or grain boundaries, second image in Fig. 1), the rotation \(R\) of a polar decomposition \(F = HR\) (where \(H \in \mathbb{R}^{2\times2}\) is symmetric and \(R \in SO(2)\)) reveals the local crystal orientation and thus the extent of crystal grains (third image), and the distance of \(F\) to \(SO(2)\) has the interpretation of an elastic strain (right image). Together, these pieces of information fully describe the mesoscopic state of the polycrystal.

In the following sections we provide more examples of how to deal with large polycrystal images or crystals of different symmetries, and we report the typical computational effort.

4.1 Accounting for Point Groups

Especially in images of polycrystals there still remains an issue with the interpretation of our results: Any two matrices \(F_1, F_2\) should actually be interpreted as equivalent if they only differ by the point group \(P\) of the crystal (the set of all rotations and reflections that leave \(\{v_1, \ldots, v_K\}\) invariant). Indeed, if \(F_2 = F_1 P\) for a \(P \in P\), then the crystal distorted by \(F_2\) looks identical to the one distorted by \(F_1\). Our model does not automatically account for that fact and interprets \(F_1\) and \(F_2\) as different distortions, which can lead to spurious curl accumulations as shown in Fig. 4. As a remedy, we once initialize \(F\) with rotations between \([0, \frac{\pi}{3}]\) and once with rotations between \([\frac{\pi}{3}, \frac{2\pi}{3}]\). The result \(F_1\) from the first initialization typically shows spurious curl along boundaries of grains at an orientation close to 0 or \(\frac{\pi}{3}\), while the second result, \(F_2\), does so at grain orientations around
We use $F$ regions where $\mathbf{curl} F$ lies above a certain small threshold (e.g., convolving with a Gaussian $G$ of roughly two unit cell widths), then we pick the regions where $\mathbf{curl} F$ has higher values (blue region in right image). In this region we replace the first output by the second output, yielding the shown curl (right).

$\frac{\pi}{2}$ and $\frac{3\pi}{2}$. We now simply identify regions of spurious curl by comparing $|\mathbf{curl} F|$ in both results: in regions where $F_1$ has a significantly higher curl than $F_2$ we use $F_2$ instead of $F_1$. In detail, we first smooth $|\mathbf{curl} F_1|$ and $|\mathbf{curl} F_2|$ (e.g., convolving with a Gaussian $G$ of roughly two unit cell widths), then we pick the region where $G \ast |\mathbf{curl} F_1| - G \ast |\mathbf{curl} F_2|$ lies above a certain small threshold (in our simulations 0.01) and dilate this region by two unit cells. Within this region we consider $F_2$ as the correct distortion, outside $F_1$ (Fig. 5).

4.2 Coarsening Simulations

As a demonstration of this method, we segment a time series of crystals chosen from a PFC evolution. In Fig. 6, four time snapshots of the PFC evolution are considered (left to right). The resulting rotation map, local volume distortion map (measured as $\det F - 1$), elastic strain map, and curl magnitude map are displayed for each time point (top to bottom). Though each segmentation is performed independently, it can be seen that rotations are assigned in a consistent way. Volume distortion and strain are nonzero near grain boundaries and dislocations. High magnitude of curl ($|\mathbf{curl} F|$) is seen to denote grain boundaries and isolated dislocations, which can be easily distinguished from each other. Note that each dislocation creates a dipole structure in the volume distortion map, since the crystal is dilated on one side of the dislocation and compressed on the other. The orientation of this dipole can hence also be used as an indicator of the Burgers vector.

4.3 Strained Crystals

In contrast to the approaches from Sec. 1.2, our method directly provides the crystal strain, which can be of interest in particular for the analysis of non-equilibrated crystals. As a simple example, we performed a PFC simulation, where in regular time intervals we artificially shift a stripe of the simulated material horizontally by one pixel, thereby emulating the effect of a shearing force. The resulting strained crystals are shown in Fig. 7, once for a defect-free crystal and once for a crystal with three dislocations. The distribution of elastic strains indicates that the dislocations effectively reduce the strain in the bulk of the material (while there is a higher peak strain in close vicinity to the crystal defects).

4.4 Different Crystal Types

The method seems to work robustly on different types of input images, including real experimental images. Fig. 8 shows a HRTEM image of a nanocrystalline palladium thin film, courtesy of Nick Schryvers, from the work (Wang et al., 2012). The tensor field $F$ is robustly extracted, even though the coloring varies across the image, and $|\mathbf{curl} F|$ correctly identifies an isolated dislocation. The crystalline orientation obtained from polar decomposition of $F$ reveals that three distinct regions separated by twin boundaries are identified correctly despite no more than $10^\circ$ difference in orientation.

In Fig. 3 we used an even more challenging input image of a reptile skin, whose pattern is rather irregular compared to an atomic crystal. Nevertheless, using the six-point stencil of hexagonal symmetry, the locations as well as the types of the dislocations in the scales pattern are correctly identified. Note that the unit cells of the pattern vary strongly in size so that our initialization of $F$ with pure rotations seems much less appropriate, yet it obviously still suffices to achieve the desired local minimizer.

Lattices of different than six-fold rotational symmetry can be analyzed using an appropriately adapted stencil $v_1, \ldots, v_K$. Fig. 9 shows a snapshot of a PFC simulation with cubic lattice symmetry, for which an appropriate stencil is given by $v_1 = (\ell, 0)^T$, $v_2 = (0, \ell)^T$, $v_3 = (-\ell, 0)^T$, $v_4 = (0, -\ell)^T$ (where again, $\ell$ is the characteristic spacing). The underlying PFC model is a modification of the standard PFC model, presented in (Wu et al., 2010, Sec. VI) and originally proposed in (Lifshitz and Petrich, 1997) in the context of the Swift–Hohenberg equation. Isolated dislocations as well as low- and high-angle grain boundaries are properly identified. The local crystal orientation indicates that most grains have similar orientations, with atoms that in this particular simulation tend to align well with the computational grid.

4.5 Computational Effort

For images as in Fig. 1 we generally found twenty split Bregman steps to be sufficient, in each of which
Figure 6: Results of our algorithm applied to different snapshots of a phase field crystal (PFC) coarsening simulation. From top to bottom we show the time evolution of local crystal orientation (0 < \pi / 3), local volume distortion of the crystal measured as \text{det} F - 1, elastic strain measured as \text{dist}(F, \text{SO}(2)) (0 < 0.05), and \text{curl} F.

Figure 7: PFC simulation of a sheared crystal without (top) and with (bottom) defects, showing the crystal image (left), the elastic strain \text{dist}(F, \text{SO}(2)) (middle, 0 < 0.15), and \text{curl} F (right, \sim -0.2). The rightmost graph shows a histogram of \text{dist}(F, \text{SO}(2)). The crystal with defects has a higher peak strain, but lower overall strain.
we perform five nonlinear conjugate gradient steps for (5). The code is implemented in Matlab, using the included basic GPU support. Computation times on a Tesla S10 GPU are displayed in Tab. 1 for different image sizes. For an image with $2048^2$ pixels, the minimization takes slightly above six minutes of which the net time inside the split Bregman iteration (without screen output of the energy values) is roughly 82%. Most of the iteration time is spent in the nonlinear conjugate gradient steps which in turn spend more than half of their time interpolating the image $u$ to evaluate the fitting energy and its gradient. We observe a strongly sublinear scaling with image size due to the high GPU parallelization.

The bottleneck seems to lie in the evaluation of $u(x + F(x)v_k)$ at each pixel $x$ and for each stencil vector $v_k$ (computation times are reported for a six-point stencil). This image interpolation requires almost unstructured memory access which without code optimization is not very efficient on a GPU. Nevertheless a speedup can be expected from a non-Matlab implementation tailored to the GPU architecture.

Table 1: Rounded computation times of GPU implementation in Matlab for image in Fig. 1 and segments thereof.

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5 DISCUSSION

The proposed model is an efficient tool for the mesoscale analysis of polycrystals, an important task in materials science. It can analyze fairly large images in reasonable time, providing information about defect locations and local crystal distortion, from which different types of statistical data can readily be de-
rived. However, the simple Matlab implementation is limited in its applicability to image sizes up to around 2000^2 pixels, which can resolve roughly 5·10^4 atoms. With a tailored GPU implementation, this limit could probably be pushed back considerably.

Note that in some cases simpler methods are available. If the atom positions do not have to be extracted from images, but are known for instance from a molecular dynamics simulation, then the faster non-variational methods from Sec. 1.2 can be employed.

The advantage of our approach over these methods lies in the higher generality of the input data. Also, for images from phase field crystal (PFC) simulations, the PFC energy density already identifies defect locations (but not the local distortion) quite well for a range of model parameters. Furthermore, even though the local crystal orientation does indicate grain regions and can easily be obtained from our results, our method does not provide a sharp image segmentation into grains. The segmentation methods from Sec. 1.2 are better suited for such a purpose, however, they necessarily ignore single dislocations and details of the grain boundary structure. Overall, our method robustly yields more detailed and versatile information of the mesoscopic crystal structure (such as local strain or dislocation character) than previously existing methods, and it does so much faster than methods that try to extract a slightly lower level of detail from similarly general input data.

Finally, let us mention that our model is nonconvex and thus requires a sufficiently good initialization before minimization (which can be obtained at low cost in most cases). Also, the nonconvexity implies that the analytical convergence results for the split Bregman method cannot be relied on. Furthermore, a better mathematical understanding of the mutual influence and relative importance between the curl regularization and the smoothing of the distortion field might help to further improve the algorithm.

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REFERENCES