Quantification of 3D spatial correlations between state variables and distances to the grain boundary network in full-field crystal plasticity spectral method simulations

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Abstract

Deformation microstructure heterogeneities play a pivotal role during dislocation patterning and interface network restructuring. Thereby, they affect indirectly how the microstructure recrystallizes. Given this relevance, it has become common practice to study the evolution of deformation microstructure heterogeneities with 3D experiments and full-field crystal plasticity computer simulations including tools such as the spectral method. Quantifying material point to grain or phase boundary distances, though, is a practical challenge with spectral method crystal plasticity models because these discretize the material volume rather than mesh explicitly the grain and phase boundary interface network. This limitation calls for specific data post-processing methods to quantify the spatial correlations between state variable values at each material point and the points’ corresponding distance to the closest grain or phase boundary. This work contributes to the development of advanced such post-processing routines. Specifically, two grain reconstruction and three distancing methods are developed for solving above challenge. The individual strengths and limitations of these methods surplus the efficiency of their parallel implementation is assessed with an exemplary Düsseldorf Advanced Material Simulation Kit large scale crystal plasticity study. We apply the new
tool to assess the evolution of subtle stress and disorientation gradients towards grain boundaries.

Keywords: crystal plasticity, recrystallization, gradients, grain boundaries, data mining, Düsseldorf Advanced Material Simulation Kit, parallelization

(Some figures may appear in colour only in the online journal)

1. Introduction

Heterogeneities of constitutive and microstructural state variables, such as stress, strain, orientation, or dislocation density, play a pivotal role during metal forming and downstream annealing treatment [1–3]. These heterogeneities form as a consequence of local differences during dislocation self-organization [1, 4–7] or geometrical constraints which the crystal interfaces imprint on the individual deformation mechanisms [8]. The accumulation of misorientation and dislocation density at such interfaces [9–12], in turn, generates local conditions and gradients which facilitate the formation of dynamic or static recrystallization nuclei [13–17]. Therefore, considerable experimental effort [8, 18] has been devoted to the characterization of these deformation microstructure heterogeneities, both in 2D [19–24] and recently also 3D [25, 26]. Nowadays, it is common practice to investigate the evolution of the deforming microstructure with full-field 3D crystal plasticity representative volume element (RVE) computer modeling [27–29] and compare explicitly such results to experiments [18, 30–38].

Most commonly, such computer simulation studies report descriptive statistics of tensorial state variable values. These are either presented as histograms of state variable values or colorful renditions of the RVE domain or specific RVE sections. For characterizing how state variable values are distributed in the RVE volume, though, one should instead quantify the distance of each RVE material point to its closest interface point and evaluate the distributions of state variable values as a function of these distances.

We acknowledge that such quantification task faces challenges when one aims in addition for highest possible statistical significance and uses full-field spectral method models [38–45]: firstly, simulations for heterogeneity build-up demand for substantially finer spatially resolved RVE domains than are typically used for flow curve predictions. This increases numerical costs because statistical significance for texture and gradients demands to include not only as many grains but also an as fine as possible spatial resolution for each of them; all to reflect accurately that dislocation density and orientation gradients build three-dimensionally as a function of grain orientation and neighborhood [11]. For such costly simulations the key advantage of spectral method models with periodic boundary conditions are lower numerical costs than compared to similarly resolved finite element crystal plasticity models [44]. Nevertheless, this advantage comes with a second challenge: instead of probing explicit interface networks, now additional post-processing costs accrue to recover and characterize the interface network.

The requirement to analyze distances in the deformed configuration, i.e. distances as they are measured in an experiment, defines a third challenge: the successive shape distortion of the RVE domain has to be accounted for. This is challenging for periodic boundary conditions: ideally, the grains in contact with the RVE domain should get analyzed as well as to not make compromises in statistics. The grains are, though, frequently not only physically fragmented (in terms of their orientation) but also logically represented by a composite of
fragments on opposite sides of the RVE domain. Another challenge to master during any quantification of spatial correlations is to collect statistics with strongest significance. For simulations with several million material points and eventually dozens of strain steps this calls for efficiently parallelized post-processing tools to enable terabyte volume analysis productivity. As a fifth challenge, any post-processing of state-variable-to-interface-distance correlations for spectral method models calls for grain reconstruction methods because as the grains get deformed they accumulate internal misorientation.

At least for the grain reconstruction challenge, advanced post-processing tools were developed in the SEM/EBSD community [46–51]. Consequently, we assessed these [49, 52–54] for their potential and application to solve above reconstruction and distancing task. Not a single tool, though, combined grain reconstruction functionality for three-dimensionally distorted surplus periodically constrained point cloud data with documented application for multi-hundred million points.

This gap motivated our research which contributes methods that cope with all above listed challenges. Specifically, we implement statistical quantification methods which characterize how state variable values are distributed as a function of material point distance to grain boundaries. We implement two grain reconstruction methods and apply these in combination with three different distancing methods to compile a comparative assessment of long range spatial distributions for state variable values. The case study in this work focuses on single-material-point-resolved quantification of the Cauchy stresses and disorientation with respect to the mean orientation of the grain. These are exemplar quantities of frequent interest and controversy with respect to their tendency to form spatial gradients at interfaces.

Furthermore, we report how such methods can be implemented into an efficient strong and weak scaling tool supplementing the Düsseldorf Advanced Material Simulation Kit (DAMASK) [38]. Open source tools like this should encourage the crystal plasticity community to explore also their 3D RVE data in further quantitative detail.

2. Methods

2.1. Defining the data analysis task

This work presents a specific analysis workflow to post-process DAMASK [38, 43, 44, 55] simulation results. Two grain reconstruction, three material-point-to-interface distancing methods, and additional post-processing algorithms were developed. The workflow enables the quantification of spatial correlations between specific material point state variable values and each material points’ closest projected distance to an interface. The individual steps of the workflow are detailed in the appendix. The workflow was implemented into a parallelized original C/C++ program. Given that the entire source code is provided as open source, it suffices to report the key steps of the workflow and the strategies to their efficient implementation.

The workflow answers the following data mining question: How are state variable values distributed spatially as a function of the projected distance to the nearest interface for a given collection of strain steps? We define a strain step as a state variable value dataset which is probed from an RVE crystal plasticity simulation at a specific true strain value. From now on, it is assumed that this dataset contains results for every material point which supports the RVE domain.

Interfaces are either grain boundaries or phase boundaries. In this paper, we focus on single phase microstructures. The methods are formulated general enough to be applicable for
phase boundaries, provided their geometry is sampled by material points. We report 3D applications of the methods. The simplification to 2D is straightforward.

Under above provision the input data for each strain step consists of all $N$ material points. Identified as $p_n$, these define positions $x_i$ in the deformed configuration. The results are defined and analyzed in the laboratory Cartesian coordinate system $\mathbb{R}^3$. As such, all local displacements of a material point versus its initial position in the unloaded case are accounted for [43]. This makes the results comparable to the situation in experiments where a plastically deformed microstructure is probed typically by a posteriori sample preparation. For this coordinate system convention the initial grid of integration points becomes successively and irregularly distorted (figure 3(a)). This calls for the development of point cloud processing methods which back out the geometry of each grain and consider the fact that typical DAMASK simulations use full three-dimensional periodic boundary conditions.

Each material point $i$ has an associated set of state variable values $s_i$. Members of this state variable set are for instance the local deformation gradient $F_i$ or the orientation $q_i$. Although not all these variables are state variables in the microstructural sense, we have opted, for simplicity, to refer to these computed quantities as state variables.

Different strategies are in place in the crystal plasticity community to define an initial grid of integration points for sampling the microstructure in the RVE volume. Cubic grids are the most commonly employed strategy. Also full-field spectral method crystal plasticity simulation tools like DAMASK [38, 44] use such grid. This avoids explicit bookkeeping of the interface network geometry and thus cuts numerical costs. As a disadvantage, though, the interfaces have to be reconstructed through post-processing to enable a quantification of above distance correlations. Below workflow presents an original solution to solve this task; implemented as a supplemental DAMASK post-processing data mining tool, called damaskpdt for short. Here, the acronym pdt stands for post data treatment.

2.2. Pre-processing stage

The processing of the strain step data is distributed across the nodes of a cluster computer. Each process handles complete strain steps (at least one). The processes read independently all material point and respective state variable values from the simulation results file. In this work, data are parsed from the traditional binary container format file. An extension of the tool to use the HDF5 based container format in DAMASK [38, 45] remains as a task for the future.

After loading the strain step data, each process executes a pre-processing stage. This includes the calculation of the material point positions $x_i$ in the deformed configuration, the equivalent stress and strain tensors for each material point, and an RVE-averaging of these quantities using methods previously described [38, 43, 44, 55]. Next, three point clouds are defined for each strain step: the first point cloud is $\mathcal{P}^0_n$. It specifies the ensemble of the original $N$ material points in the deformed configuration. The second point cloud $\mathcal{P}^{0+\epsilon}_n$ specifies the ensemble $\mathcal{P}^0_n$ surplus all its 26 periodic image points inside a bounding box with thickness $\epsilon$ about the bounding box to $\mathcal{P}^0_n$. Here, $\epsilon$ defines a small fraction of the largest edge length of the box. The third point cloud $\mathcal{P}^1_n$ specifies all members of $\mathcal{P}^0_n$ surplus all their 26 periodic image points. Subsequent processing involves region queries on all three point clouds. Special data structures were used to accelerate these queries (section 2.5). The purpose of using multiple point clouds instead of always the complete set $\mathcal{P}^1_n$ is to optimize and reduce memory utilization. As an example, it suffices to work with $\mathcal{P}^{0+\epsilon}_n$ instead of $\mathcal{P}^1_n$ when querying neighboring material points at the RVE boundary within radii of at most $\epsilon$ during the
disorientation based distancing. Further details to the purpose of the point clouds are provided in the algorithms in the appendix.

2.3. Grain reconstruction

*Remark on classical grain reconstruction with DAMASK.* Attempts were made in the past to reconstruct the grains through application of cluster search algorithms on \( P^0_n \): the grains were built as a cluster of material points with the same texture ID within a critical distance and less than an *a priori* defined disorientation accumulated among each other. Unless a very large critical distance is chosen, though, this method cannot merge all periodic images of a grain, thus reducing the significance of the analysis. Large distances, though, will occasionally merge second- or even higher-order neighboring grains into a single object. Especially so when these neighbors have similar orientations.

Instead, two different approaches are proposed to reconstruct the grains. The methods borrow conceptually from achievements made in the SEM/EBSD community \([46, 49, 54]\) but add support for periodic boundary conditions and hierarchical software parallelization. Identifying the shape of the grains is a key step of every interface network reconstruction. With this process, material points get labeled as grains based on their orientation and point neighborhood. In what follows, two methods are specified for such labeling of the polycrystal:

*Modified grain reconstruction methods developed in this work.* The first method relabels the texture ID from the DAMASK microstructure synthesis step as the grain ID. This has been the common strategy so far when studying in-grain orientation gradient build-up: either explicitly \([9, 12]\) or implicitly by reporting orientation spread via pole figures \([11]\). This grain reconstruction method is tagged TEX in the results section.

The second method addresses the challenge that when grain fragmentation becomes significant, referring still to the initially instantiated grains is no longer correct. Instead, the individual grain fragments should be distinguished. Useful methods for this task meet three requirements: they work for irregular point cloud data, they can handle successfully three-dimensional periodic boundary conditions, and are sensitive to a gradual build-up of orientation gradients.

Graph clustering is one method which fulfills all above criteria. Application examples for characterizing microstructures with graph clustering methods, though, remain few. One is known in the SEM/EBSD community as fast multiscale clustering \([56, 57]\). Other examples, on which this work settles, are community detection algorithms \([58, 59]\). Such algorithms find frequent application in analyzing human communication and interaction patterns among individuals on social media platforms. This work uses the community detection method employed in \([59]\). Specifically, we use the open source implementation of the Louvain community detection method by Blondel *et al* \([60]\). This grain reconstruction method is tagged LOU in the results section.

*Extraction of a single periodic image per grain and bounding box ensemble.* After labeling each material point one can in principle compute the distance correlations. For arbitrarily deformed domains with periodic boundary conditions, though, practical challenges remain: first, macroscopic and microscopic shape distortions of the RVE domain walls have to be accounted for. Second, grains in contact with the RVE walls are fragmented into multiple pieces. Therefore, it is useful to merge the fragments into a single grain with simpler geometry. Placing it in a cuboidal box is even better because it decouples all downstream processing into grain-local operations.

Consequently, a geometry simplification step is executed. Therein, the individual periodic fragments are first fused using a DBScan \([61]\) clustering algorithm. Specifically, all grain
fragment neighbors within a threshold radius of $R = \sqrt{3}L_0$ with $L_0$ specifying the initial point-to-point distance are fused. Next, the closest replicate periodic image to the center of $\mathcal{P}_n^0$ is chosen and the associated material points are packed into a grain-local bounding box. Once boxed up, efficient multithreaded processing methods from the grain growth modeling community [62–64] speed up the grain-local analyses.

After the geometry simplification the global bounding box to all grains’ local bounding boxes are computed (figure 2(b)) and define a Rediscretization of this global bounding box. The actual Rediscretization, though, is applied inside each grain-local box only to improve efficiency. Once Rediscretized, the information contained in $\mathcal{P}_n^1$ is used to identify the nearest neighboring material point to each voxel. Cubic voxels of edge length $0.5L_0$ were used.

2.4. Distance quantification

**Disorientation based.** For this distancing method, tagged DIS in the results section, the challenge of grain reconstruction is ignored completely. Instead, position and orientation data for points in $\mathcal{P}_n^{0\rightarrow\varepsilon}$ are used. The quantity $\varepsilon$ defines a guard zone thickness that was chosen as 0.1 times the initial RVE domain edge length. The distance values are computed by finding for all material points in $\mathcal{P}_n^0$ all their respective neighbors within a region of radius $R = 0.1$, where $R$ is again in multiples of the initial RVE domain edge length.

Next, the identified neighbors are sorted in increasing order with respect to their Euclidean distance to the inspected material point. Finally, the distance value for the closest, above a critical threshold $\Theta_0$, disoriented pairing, is taken as the distancing result. In this work, a threshold of $\Theta_0 = 15^\circ$ was used. If no closest neighbor among the candidates was found, the material point made no contribution to the correlation statistics.

In a nutshell, the advantage of the DIS distancing method is that it requires no grain reconstruction as long as no disorientation-to-distance correlations are desired. The independence of each material point is another advantage which makes the calculations trivial parallel. Without taking the spatial arrangement of neighboring points into account, though, a disadvantage of DIS distancing is that only scalar distances are accessible.

**Signed distance/voxelization based.** Enabling the identification of normal distances to the grain boundary is the motivation for and the key strength of the second distancing method. It is tagged as SDF in the results section. The key idea is to evaluate signed distance level set functions in combination with latest method developments achieved for scalable grain coarsening simulations [64–67].

Level set function methods parameterize the position of interfaces implicitly as the isocontour of a real-valued level set function $\Phi(x)$ to an a priori defined iso-value. Defining a signed distance level set function (SDF) in combination with each boundary contour of a grain being defined as $\Gamma := \{x \in \Omega \mid \Phi(x) = 0\}$ with $\Gamma \subset \Omega \subset \mathbb{R}^3$ is a particular useful choice. Namely, this choice, i.e. $\Phi(x) \in \Omega \subset \mathbb{R}^3$, has the key advantage that $\|\nabla \Phi(x)\| = 1$ holds for most points $x \subset \mathbb{R}^3$ one practically encounters inside and outside the contour. Consequently, a consistent outer unit normal vector can be computed to each point on the contour by evaluating $n(x) = \frac{\nabla \Phi(x)}{\|\nabla \Phi(x)\|} \frac{\nabla \Phi(x)}{\|\nabla \Phi(x)\|}$ numerically. In effect, this allows for calculating arbitrarily projected distance vectors to the contour.

A two step procedure was employed to compute a signed distance function (SDF) for each grain. The procedure builds on the previously defined discretization of the bounding boxes about each merged grain. First, $\Phi(x)$ was initialized to positive values $(+h)$ for all voxel assigned to the grain under inspection and negative values $(-h)$ for points outside. The scalar $h$ was set equal to the cell width used for the global rediscretization of the domain. In a second step, the distance information was propagated with the fast sweeping method (FSM) [64, 68].
Also the third method, tagged VOR in the results section, delivers vectorial distances. Furthermore, it explicitly reconstructs a contour hull for each grain. The key idea of the method is to build up each grain from a collection of Voronoi polyhedra. These can be directly obtained for each grain from a Voronoi tessellation of the respective material point cloud. Formally, the VOR method is based on the approach of Bachmann et al [54]. We introduce two improvements, though: first, a generalization of the method for applications on distorted point clouds with periodic boundary conditions. Second, we introduce a hybrid-parallelized solution which executes substantially faster and might provide an avenue to explore for MTEX in the future as well. This parallelization alleviates much of the higher numerical costs, which the VOR method has in comparison to the DIS or SDF methods. VOR does not require a rediscretization of the bounding box to each grain but operates directly on the grain-local subset of material points from \( \mathbb{P}_n \).

In a first step, this local point cloud is extracted and each Voronoi cell computed. Each Voronoi cell gets labeled with the respective grain ID \( k \) and material point ID. Next, the algorithm identifies which of the cell facets have first-order neighboring Voronoi cells with a different associated grain ID than the currently processed. Such difference is a signature that the currently processed Voronoi cell makes contact with the grain boundary.

The resulting collection of all such facets of Voronoi cells about points tagged with the grain ID under inspection is used to construct the shape of the exterior contour hull (figure 1(b)). The contour hull extraction algorithm is executed in parallel for each grain \( k \).

The Voro++ tessellation library [69] was used to process the individual grain-local tessellations. Specifically, we re-implemented the library wrapping approach of Peterka et al [70] around calls to this library. To avoid truncating Voronoi cells of the inspected grain, the individual grain-local bounding boxes were enlarged by a guard zone of three times the initial point-to-point distance. Furthermore, possible Voronoi cell contact with the local bounding

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**Figure 1.** (a) Compositing Voronoi cells into a larger polyhedron can result in a polyhedron with concave and convex sections. This demands additional care when computing projected distances to the contour hull facets. (b) An exemplar result of the tessellation based contour hull extraction algorithm. A random subset of outer unit normal vectors is shown.
box domain walls was detected using Voro++ library functionalities. These steps cured any possibly truncated Voronoi cell, eventually through a recomputation after fattening the guard zone locally.

Once the exterior contour hull has been defined, the material point to contour hull distances can be computed. Given that projections on the polyhedron facets can be used, the VOR methods computes projected distances. The implementation is tricky, though, in detail because consistence and speed are desired:

**Algorithm to compute vectorial distances within VOR.** Using the edge circulation conventions of the Voro++ library, first a consistent outer unit normal vector for each contour hull facet is computed. Next, the normal and geometry of the facet is evaluated to compute a consistent right-handed local Cartesian coordinate system for each contour hull facet. Such coordinate system can be defined because each Voronoi cell is a convex polyhedron [71]. By virtue of construction each facet is a convex polygon in 3D space that is reduced, thanks to the local coordinate system to two-dimensions. These local coordinate systems accelerate point to polygon inclusion tests.

It remains to compute the distances. First, the general approach used is explained. Second, we detail how the processing of the facets gets accelerated with bounded volume hierarchy (BVH) querying algorithms. The distances are Voronoi cell interior-point-to-exterior-contour-hull-projected point distances. Distances were evaluated for each interior point. It suffices to sketch the procedure for a single interior point. The main task is to probe which facet of the contour hull lays closest to the point. While probing the facets for the absolute shortest projected distance the algorithm works as follows: firstly, each interior point gets normal projected onto the facet plane.
Secondly, it is probed whether the projected point lays inside the facet polygon or on its contour respectively or outside. The distance value is eventually used to update the currently closest distance.

If, though, the projected point is outside the polygon, one proceeds by normal projecting the currently tested interior point on each edge segment of the polygon. If such a projected point lays on the edge, the computed distance eventually updates the current closest distance.

If none of these projections map on a polygon edge segment, we evaluate the Euclidean distance of the currently tested interior point to each vertex of the polygon and update eventually thereby also the current closest distance. Once finished with the vertices, the algorithm proceeds with the next facet of the contour hull until no further facets remain for processing.

To the best of our knowledge advanced procedures like above are necessary to handle cases where, due to the possible non-convexity of the composed contour hull, a consistent normal distance is not immediately defined. In fact, such cases are frequently possible. One particular simple example is illustrated with the grayed-out region in figure 1(a). Evidently, there is not necessarily a projected point on a contour facet for every position inside an arbitrarily shaped non-convex polygon (here for the brightest yellowish region). At least not if one projects exclusively parallel to the direction of the facets’ outer unit normals.

While getting access to a defined contour hull for each grain surplus projected distances is a clear advantage, the VOR method has also disadvantages: most importantly, higher numerical costs. More subtle, the edge-on compositing of a contour hull from a collection of polygons faces discontinuities of the curvature at the polygon edges. These can be cured, though, with state of the art SEM/EBSD grain reconstruction software [49, 52]. This is another possible avenue to explore and use for reconstruction of the interface network in DAMASK in the future.

A strategy for accelerating the distance computations. Without additional tricks, the algorithm defines a quadratic time costly computational geometry task—for each interior point each contour facet needs inspection. Therefore, a pruning strategy was designed to reduce the total number of facets tested. The key idea is to use the shorter and shorter identified distance values to occasionally requery which contour hull facets require testing. Therefore, a BVH was built from the contour hull facets [72–74]. This utility data structure organizes the locations of the facet polygons into spatial regions, such that with narrowing down the search fewer and fewer candidates need testing.

First, we probe an arbitrary candidate and query the BVH for the resulting distance value. This value is used for querying the BVH to narrow down which candidate facets to probe next. Thereafter, the algorithm continues iteratively: each time the distance value gets lower than 0.9 of the last query, the candidate list gets updated until finding the absolute shortest distance per interior point.

Distilling spatial distributions of disorientation and stress. The results of each damaskpdt tool run are two ensembles of Cauchy stress tensor/distance and orientation quaternion/distance value pairs per material point. These data were finally processed with MATLAB (v2017a)/MTEX [75, 76] texture toolbox (v5.0.3) scripts to obtain spatial distributions of stress and disorientation.

A two stepped protocol was executed to quantify the disorientation of each material point to the respective mean orientation of the reconstructed grain: first, a mean orientation was computed for each grain. Documented methods for inferential quaternion statistics [77] were used for this task. Secondly, these mean orientations were evaluated against the individual orientation of each material point.
The so generated value/distance pairs were binned into distance bins on the interval [0.0, 24.0] using 0.2 step. Length units are reported in multiples of the initial [100]-direction point-to-point distance between neighboring material points \(L_0\). Quantile values of the resulting sub-distributions were quantified with MATLAB. Grain size distributions report how many material points were assigned per grain.

2.5. Parallel implementation and orchestrated hierarchical data placement

Above workflow was implemented into a hybrid-parallelized tool. Inherent parallelism in the data mining task was exploited: for the reason that each strain step is independent, we distribute the processing of the strain step ensemble via process data parallelism. Specifically, calls to the message passing interface (MPI) library \([78, 79]\) were implemented. Strain steps were distributed across the MPI processes in round-robin fashion. In addition, the processing of each strain step was accelerated via Open Multi-Processing (OpenMP) \([64, 80–85]\). Instead of using the commonly employed referencing of data items via global arrays, we actively enforce a partitioning of the material point positions and state variable values. Specifically, all derived quantities are split into thread-local data chunks. These chunks are placed preferentially in those memory locations with a fast connection to the executing computing core. This facilitates faster reading from memory through closer packing of data and less cache coherency traffic. We admit that our implementation currently uses a static load partitioning scheme whereby load imbalances may be stronger than for a dynamic workload distributing. This is a possible disadvantage to investigate in the future.

To efficiently execute all linear algebra operations of the workflow, fast Fourier transforms, singular value decomposition, and eigenvalue decomposition functionalities of the Intel Math Kernel Library (IMKL) were used.

The subsequent grain segmentation and characterization of the spatial descriptive statistics involves range querying tasks. Specifically, they demand the identification of all neighboring points inside a spherical volume of radius \(R\). Using 3D box binning with cubic buckets of width \(R\) for \(P_{0}^{+}\) and \(P_{0}^{-}\) enabled constant time complex queries. A disadvantage of this technique compared to other established fast position querying techniques \([72, 86, 87]\) is the larger memory footprint. This detail remains as future work for a possible code optimization aiming at a lower memory consumption.

2.6. Verification and validation simulation setup

**Microstructure instantiation.** We applied above methods for studying the evolution of the spatial distribution of stresses and disorientation in full-field crystal plasticity simulations. Simulations were executed with the DAMASK spectral solver \([38, 44, 55]\) and used the DAMASK Poisson–Voronoi tessellation microstructure synthesis and orientation sampling routines. Specifically, a 3D face-centered cubic single phase polycrystal RVE was instantiated. Full three-dimensional periodic boundary conditions were used. The RVE domain contained 500 grains. These were discretized by \(256^3\) grid points, i.e. material points. In effect, each grain has an average spherical equivalent radius of 20 material points. Grain orientations were sampled randomly and assigned in a spatially uncorrelated manner.

**Distance verification.** To the best of our knowledge there is no analytical solution which quantifies the distribution of shortest projected distances for an arbitrarily distorted point cloud to a given grain boundary interface network. As such, the verification of above methods is challenging. For Poisson–Voronoi microstructures, though, one can find exact linear algebra solutions for a given point cloud and evaluate them numerically. By virtue of
construction the above defined microstructure is a discrete approximation of a specific Poisson–Voronoi tessellation. Consequently, we verified our implementation as follows. First, the center positions of above grains were taken to define the corresponding fully-periodic Voronoi tessellation in continuous space. Second, the shortest projected distance to all facets of the enclosing Voronoi cell was probed for each material point. The resulting distances were binned using the same protocols.

Constitutive model. A phenomenological crystal plasticity model [88] was used. The tensor of elastic constants was parameterized based on experimental data for aluminum [89]. The tensor components were $C_{11} = 106.75$ GPa, $C_{12} = 60.41$ GPa, and $C_{44} = 28.34$ GPa, respectively. Dislocation slip on the 12 primary slip systems was assumed as the exclusive deformation mechanism. The phenomenological model was parameterized with a reference strain rate of $\dot{\gamma}_0 = 0.001$ s$^{-1}$ and a stress exponent of $n_{\text{slip}} = 20$. Stress parameters of this constitutive model were set to $\tau_0 = 31$ MPa, $\tau_{\text{sat}} = 63$ MPa, the hardening parameters were assumed as $a_{\text{slip}} = 2.25$ and $h_0 = 75$ MPa, respectively. Isothermal conditions and initially stress free grains were assumed.

Load case. The RVE was subjected to uni-axial compression along the $z$-axis, simulated by a deformation gradient rate $\dot{F}$ (equation (1)) with setting Piola–Kirchhoff $P$ stress boundary conditions (equation (1)). Stars identify unconstrained values.

$$
\dot{F} = \begin{pmatrix} * & 0 & 0 \\ 0 & * & 0 \\ 0 & 0 & -0.1 \end{pmatrix} \text{s}^{-1} \quad P = \begin{pmatrix} 0 & * & * \\ * & 0 & * \\ * & * & * \end{pmatrix} \text{Pa}.
$$

The resulting RVE domain is visualized in figure 3(a) exemplified for the deformed configuration in the lab coordinate system at the final equivalent von Mises strain $\varepsilon_{\text{eq}} = 0.21$. 

![Figure 3](image-url)
Simulation settings and monitoring. Default settings of the DAMASK spectral solver were used. At least 4 and at most 40 iterations per strain step were allowed. The simulation was executed on an in-house workstation (table 1) using it exclusively and employing one MPI process with 36 OpenMP threads without pinning the threads. Microstructure snapshots were written at 28 selected strain values which the black dots in figure 3(b) detail. The following quantities were logged for each material point at each strain step point: the volume, the deformation gradient $F$, its plastic $F_p$ and elastic part $F_e$, the first Piola–Kirchhoff stress tensor $P$, orientation quaternion $q$, and a reference ID, the so-called texture ID $T_i$. The latter integer specifies the discrete orientation ID of each material point. The resulting binary spectralOut file occupied 162 GB disk space.

Simulation data post-processing. Post-processing was executed on TALOS, a cluster computer (table 1). Different levels of hybrid, i.e. process and thread parallelization were benchmarked. Computing nodes were used exclusively and threads pinned (OMP_PLACES=cores). Explicit calls to the MPI_Wtime and omp_get_wtime functions were used to monitor how much time the individual workflow steps took. I/O and non-I/O operations were distinguished. The main virtual and resident set size memory consumption was probed at the node level by parsing on-the-fly from the /proc/self/stat system file.

The strong multithreading speed of the tool was benchmarked with runs on a single TALOS node. Specifically, quasi-sequential runs with one process spawning one thread were compared to repetitive runs of the same study with one process spawning 40 threads. The multinode weak scalability was probed by comparing the single node results with runs employing 28 MPI processes each of which spawning 40 threads. I/O operations were performed using a GPFS parallel file system built on top of 12 logical disks. Files were striped across a RAID6 array with 10 disks each (in 8+P+Q configuration). The low level stripe size was 4 MB. Supplemental MTEX post-processing was performed sequentially using a desktop PC. Given that the processing consumed only a few core hours in total, these analyses were not benchmarked.

2.7. Software details

DAMASK (git commit ID: v2.0.1-992-g20d8133) was compiled with GNU v8.2 using O2 optimization. The program was linked against the PETSc v3.11.0 numerical and the MPICH v3.3 MPI libraries. The damaskpdt post-processing tool (git commit ID: caea9-c52848809e0635ba7d0af313a592e3d0dd) was compiled with the Intel Compiler and Performance Suite (v2018.4) using O3 Skylake optimization. The program was linked against the corresponding IMKL and Intel MPI versions. Additionally, Boost (v1.66) [90] and the Voro++ (v0.4.6) [69] libraries were linked to the tool.
3. Results

3.1. Verification of the distance computations

First, we verify the distribution of distances through comparison to solutions for the initial configuration. Figure 4(a) reports the distances in multiples of $L_0$, i.e. the point-to-point distance in the unloaded configuration. Like at yield, in this configuration, the shape of the

![Figure 4](image-url)

**Figure 4.** Cauchy stress spatial distributions as a function of the distance to the nearest boundary, here exemplified for the largest stress component $\sigma_{33}$ and comparing specific quantiles of the distance and the state variable distributions. Theory details the results for the analysis of the distances of the voxel center to the Voronoi cell facets in continuous space. Abbreviations discern the distancing methods: disorientation (DIS), signed distance/voxelization (SDF), or tessellation (VOR) based, the grain reconstruction method (texture index TEX or Louvain LOU based), and whether large disorientations within a cluster were strongly ($SP, k = 1000$) or weakly ($WP, k = 75$) penalized. Grayed-out regions detail where numerical effects of the spectral method are expected strongest—either due to numerical effects or finite counting.
RVE is undistorted. Exemplified for the method combination DIS/TEX the histograms show quantization effects inasmuch as all the possible voxel-to-voxel center distances are recovered ($L_0$, $\sqrt{2}L_0$, and $\sqrt{3}L_0$). This approximates well the analytically accurate distribution of the actual normal distances of the voxel center to the Voronoi facets in continuous space. The figure substantiates that without injecting additional local grain boundary normal information, our methods are voxel accurate only. Given that a fixed number of material points is probed, a larger number of cases with shorter distances implies finding a lower number of the largest distances in the RVE than for the continuous space solution.

3.2. Quantifying the spatial distribution of stresses towards grain boundaries

Figure 4 summarizes the results from applying above methods to quantify how stresses are distributed in the RVE volume with respect to the distance of each material point to the nearest grain boundary. Using the two developed grain reconstruction methods in combination with the three distancing methods allowed for rigorous and comparative analyses on the same dataset. The results document the statistics at final strain when the stress distributions are broadest and highest in terms of absolute values. All methods identify with very similar significance and quantile values that the average $\sigma_{33}$ compressive Cauchy stress is 140 MPa. Only an at most 10% absolute variation from this average value is observed when probing deeper into the grain interior. Distances larger than approximately 15 distance units from the boundary should not be interpreted because the corresponding numerical support is finite counting limited, i.e. only very few points belong to these distance bins given the average grain radius of 20 voxels.

Comparing the distributions for each distance bin and method combination (figure 4(a)) conveys that all methods capture the flattening of the grains in the compression direction. Consequently, higher counts for the high distance bin are observed for the strain step at yield versus the one at final strain.

The results for the lower distance classes document a rigorous quantification of the numerical effects inherent to spectral based methods in general, and those specific for the DAMASK spectral solver: stresses at material points in the vicinity of numerical discontinuities, here the grain boundaries, show consistently different mean stresses (figures 4(b)/(c)) (10%–14%). Finding this difference consistently for material points at the discontinuities suggests that this is attributable to the Gibbs phenomenon [91, 92]. Latter is known as a systematic intensity overshooting in frequency space when attempting to Fourier expand at discontinuities.

3.3. Quantifying grain orientation spread accumulation towards boundaries

Motivation for this second application exercise comes from frequent literature findings which reported that material volume in the vicinity of grain [93] and phase boundaries [94] is differently disoriented than volume in the grain interior. Possible presence of such localized volume with different orientation has been identified as one reason why static and dynamic recrystallization nuclei [95] originate frequently from the faces or junctions of the grain boundary network [16, 17, 96].

The results in figure 5 deliver quantitative evidence that spatial gradients of disorientation are detectable with all grain reconstruction and distancing methods: material points in the proximity of a grain boundary show on average larger disorientation to the mean orientation of the grain than do points in the grain interior. It can be excluded that these differences are the result of averaging orientations from multiple material points, as it is the case when measuring e.g. kernel average orientation.
misorientations (KAM) at boundaries using the SEM/EBSD technique. In fact, when quantifying KAM values a kernel with multiple neighboring material points is evaluated. Probing with the kernel would contribute high disorientation values only if insufficiently strong criteria are set with respect to how much disorientation noise is allowed for a given kernel. In this work, though, no kernel was used. Instead, the disorientation values were computed independently to only one value for each material point—the respective grain mean orientation.

It can also be excluded that the observed gradients are random spatial correlations for the following reasoning: if a grain contains multiple material points in different orientations in the fundamental zone, it is expected that non-directionally correlated point-to-point disorientations are measured if one eventually compares the disorientation for two arbitrarily picked material points. If the orientation variation is strong and the grain small, i.e. the grain has few points support, eventually larger disorientations are measured on average. However, this scatter should also be correlated if there are orientation gradients with a strong component normal to the interface within the grain. Vice versa this scatter should remain practically uncorrelated if such gradients are absent or the grain contains practically uncorrelated small regions with spurious higher disorientation.

Given the strength of the gradient and mean disorientation value, we conclude that the grains are in an incipient stage of fragmentation. They have not yet accumulated localized regions of significant point-to-point disorientation in the high-angle boundary regime ($15^\circ$), except for a minority population of material points in an at most 2 pixel wide zone at the boundaries. Qualitatively, their higher disorientation is expected as also the absolute Cauchy stress values are slightly higher in this zone (figure 4).

There are two additional observations to make with practical significance for characterizing orientation gradients: first, only the methods with normal distancing capability
(SDF, VOR) report similar gradient slope ($\approx 4^\circ / \text{15keV}$). Compared to disorientation based distancing, the slope is moderately lower. Second, disorientation gradient characterization with the graph clustering method shows a strong parameter sensitivity (figure 5(b)).

4. Discussion

4.1. Comparing the two grain reconstruction methods

We identified that one practical challenge of using the graph clustering method for grain reconstructing is the strong parameter sensitivity. Therefore, it was worth studying the grain size distributions (figure 6(a)) and consistency with which individual grains were (re-)
identifiable (figure 6(b)) as a function of the penalization parameter $K_L$ (algorithm B). Figure 6 summarizes the key findings.

The larger the penalization parameter $K_L$ is set, the stronger every accumulation of high disorientation between nodes of the same community gets penalized. In effect, more grains with lower in-grain orientation variation are found on average for a stronger penalization ($K_L = 1000$).

Neither the values for a low nor for a high penalization can avoid systematic challenges when using the reconstructed grain ensemble for characterizing the distance correlations. In fact, if the penalization is weak, the algorithm reconstructs different grains and comes not even close to the initially synthesized number. Figure 6(b) proofs that this is a systematic consequence of the method’s tendency to merge neighboring grains with low disorientation at low $K_L$ values.

In effect, the mean orientation of many grain pairs is an average of at least two orientation ensembles with low intra ensemble but possible noticeable inter ensemble disorientation. With respect to spatial disorientation gradients this has two consequences: first, such gradients are flatter and second they are shifted to larger disorientation on average. This explains why the gradients for weakly penalized graph clustering (figure 5(b)) are shallower.

One could avoid the possible bias introduced by the systematic grain merging through applying a stronger penalization, such that eventually the initial grains are re-identified (figure 6(a)). Figure 6(a) shows the detrimental effect of such procedure when applying it to analyze the (highly) deformed configurations: we observe oversegmentation with an associated qualitative change of the grain size distribution from unimodal to bimodal. A detailed inspection of the respective material points identified that this spreading into eventually bimodality is a consequence of the fact that the strongest disoriented material points at the grain boundary get now segmented into a decoration of the boundaries by very small grains. These findings have a methodological and a practical implication.

The methodological implication is that graph clustering based grain reconstructions should always be backed up by rigorous quantitative parameter sensitivity studies. One strategy could be to report always unnormalized distributions of grain sizes as a function of the segmentation parameter. Otherwise, different physical conclusions might be drawn even though one uses the same method. One example is exemplified in figure 5(b): disorientation accumulates at boundaries or not.

The practical implication reads as follows: grain fragmentation should be better characterized ideally via the evolution of the grain boundary network rather than to continue insisting logically on the initial grain as the decisive object. In fact, not the homogeneous regions within a deformed grain but the heterogeneities are the key microstructure locations and first descriptors to investigate where and how annealing microstructure evolution initiates.

4.2. Implications of numerics-induced scatter for non-local and coupled crystal plasticity/ interface migration spectral method models

Evincing such numerics-induced scatter suggests to apply above quantification protocol in the future regularly. Especially, it should be applied in so-called non-local crystal plasticity models. These predict the local material point values via evaluating a kernel of neighboring material points. Examples are continuum scale crystal plasticity solvers with incrementally and locally coupled dislocation flux sub-models, such as the one from [31, 97] or full-field crystal plasticity models with incremental coupling to phase-field solvers [35].
One should always start inspecting such solvers from an in-depth monitoring of the state variable values and the non-local flux terms for each material point as a function of strain. Methods, such as the ones developed in this paper, could be used to assess via such monitoring in how far numerical noise remains uncorrelated and low to avoid a systematic biasing of the results at larger strains.

Another example in which rigorous quantitative monitoring of state variable values is useful are incrementally coupled grain boundary migration/crystal plasticity models. These typically evolve microstructures along complex transients. As such, any permeation of correlated numerical noise into the flux or interface migration terms should be reduced as best as possible to keep rigorously controlled conditions, physical accuracy, and precision.

Knowledge in the discrete dislocation dynamics community teaches us that uncontrolled stress spikes should better remain numerically controlled, for instance by supplementing the integration with some time trajectory before making choices [98, 99]. For these future challenges the work delivers an approach based on which to build further tools for assessing the numerical and physical quality of such predictions.

4.3. Comparing computational efficiency

Strong scaling multithreading performance. Figure 7 summarizes wall clock timings to document the tools’ strong scaling efficient multithreading capabilities. Furthermore, the figure documents key computing time contributions for all combinations of above post-processing methods applied to the same data, using the same computer, and exclusive execution. The left column of each column pair shows the sequential execution time. The results identify that SDF, in combination with either of the two grain reconstruction methods, is the least costly combination. Results for a single strain step and sequential
execution are available after 2 h approximately. Given the capability to identify normal
distances is another argument to choose the SDF method over disorientation based distanceing.

Figure 7 documents that all methods concentrate at least 87.5%–99.3% of the total execution
time in at most three algorithmic code sections. For the disorientation based distanceing these steps are the querying of neighboring points and computing point-to-point
disorientation angles. For the signed distanceing method it is the identification of periodic
images, volume rediscretization, and the remapping of every voxel to a closest unique
material point. For tessellation based distanceing the evaluation of the projected normal distances for each material point demands contour facet querying.

In this work, all these most costly steps are parallelizable. Thus, specific care was taken
to implement them in parallel. In addition, data placement was orchestrated such that GB-
sized data chunks were placed close in the memory of the executing core taking into account
the memory hierarchy of the cluster computer.

In effect, this speeds up the execution of every method combination by 13.6–24.4 times
the sequential baseline. With 40 OpenMP threads required to achieve this, a strong scaling
efficiency between 61% and 34% is documented. The main obstacle to achieve higher effi-
ciency in this study was load imbalance. Such was strongest for the tessellation (VOR)
method because grains were distributed to the threads statically in round-robin fashion. When
building tessellations, however, the grain size critically affects how many facets the contour
hull contains; and thus how dissimilar the distanceing costs are for different grains. In light of
this, the distributing of only 500 grains across 40 threads will equip each thread with possibly
too few grains to compensate for the large variety of grain sizes. In effect, the cost and facet
count in the contour hull processing stage differs.

A possible improvement for the future is to change the implementation and employ
dynamic work scheduling via for instance OpenMP tasking. This, though, demands a com-
promise: the current round-robin distributing procedure allows to place the grains in con-
trolled memory locations [82]. Dynamic scheduling, in turn, will likely keep the cores more
frequently busy but at the cost of typically more memory traffic.

Another potential improvement of this work, with immediate practical benefit, is to
distribute also the graph clustering algorithm for which CPU- and GPU-parallelized com-
munity detection solvers have been developed in the past [100–103].

The results document additional practical improvements in which the present work
advances the field: the first is improved speed, when compared to hitherto reported values for
DAMASK post-processing [38]. This was achieved with sequential optimization of the file
accessing strategy surplus employing parallelized I/O to cut I/O costs by at least an order of
magnitude.

As an additional strategy to improve sequential performance, state of the art linear
algebra libraries were used to compute point-wise tensor quantities. Such improvement is of
practical use not only for executing numerics within DAMASK when solving an RVE but
also for post-processing flow curves. As the second improvement, this work details how to
combine these strategies with multithreaded execution surplus trivial data parallel processing
of strain steps on multiple nodes of a cluster computer.

Having the possibility to construct contour hulls to each grain in the deformed con-
furation is the third improvement. On the one hand because it allows to quantify a volume for
each grain not only by counting the number of material points it contains but by accumulating
the volume of its supporting Voronoi cells. The resulting polygon mesh allows for volumetric
rendering of the grain as a polyhedron, as exemplified in figure 1(b). These polyhedra may be
useful to allow further analyses using other microstructure characterization tools like
DREAM3D [53] or QUBE [52].
Hybrid execution performance. During hybrid execution every MPI process evaluates a different strain step and spawns an own group of threads. Data are read independently from a priori known sections of the DAMASK results file. In effect, this enabled to evaluate all 28 strain steps at once using 1120 cores. Figure 8 summarizes the key results for this hybrid execution mode. Specifically, two sets of common analysis cases were probed: a flow curve extraction versus analyses of aforementioned state variable values to boundary distance correlations, six of them in total.

There are two key findings of practical importance. First, hybrid processing enables to cut wall clock time by an additional order of magnitude. In effect, the evaluation of a flow curve, which took damaskpdt 3.2 h sequentially, was solvable in 76 s when using parallelism. Similarly, distancing results for all strain steps were analyzed after 1.5 h when using tessellation (VOR), the most costly method. Distances were even faster accessible, namely in 8 min, using the signed distance method.

The practical benefit of these findings is put in perspective by the following observation. While the SDF method was already finished for the entire ensemble, the classical DAMASK post-processing routines had not even completed the processing of the flow curve for the first strain step.

The second key finding is that parallel performance gains are limited. Respective arguments for the multithreaded analyses were already identified (figure 7). Not parallelizing all post-processing computational steps is another apparent reason for limited scalability [104]. The most significant obstacle to cut execution time further, though, are microstructure-induced differences. Specifically, differences which are caused by interface networks with dissimilar shape and distribution of facet area across the strain step ensemble. Consequently, the individual algorithms show different solving time resulting in workload differences across the strain step ensemble.

Two observations are important to add here. First, these inter strain step work load differences would remain even if the multithreaded execution of each strain step gets better...
work load partitioned. Second, it is in fact the successive reduction of the relative numerical complexity for a strain step with increasing strain which limits primarily the hybrid performance of MPI parallelization. Provided there were more strain steps to process very likely even more processes could be employed. This is one of the key advantages to our true scientific computing performance solution for post-processing DAMASK simulations.

One example is visible for signed distance based distancing. Given that the RVE shape change results in a moderate increase in the total bounding box volume, more voxel have to be probed for the higher than the lower strain steps. Another, by far stronger, example for microstructure induced work load imbalances is visible for tessellation based distancing (figure 8).

When using the strongly penalized grain reconstruction, more grains are detected at higher strains. Therefore, also more boundary area is generated on average, as a consequence of which the material points lay closer to a boundary, providing for faster distancing operations. In addition also different contour hull shape and facet compositions are generated. This modifies the composition of the individual bounded volume hierarchies, and thus their individual response and pruning effectiveness during querying them.

It is useful to compare the computational costs of post-processing to the actual simulation costs. The DAMASK deformation simulation occupied 36 cores for 354 h wall clock time, i.e. 12 744 core hours were spent in total. The post-processing of the entire strain step ensemble using the most costly distancing method kept 1120 cores busy for 4822 s wall clock time, i.e. 1500 core hours in total, or at most 11.8% of the simulation costs.

Lastly, the memory consumption of the simulation and post-processing should be reported. DAMASK allocated a total of 230 GB virtual memory, i.e. 14.3 kB per material point. Post-processing demands less virtual memory per process when evaluating the flow curve only (21 GB). However, virtual memory consumption peaks at 134 GB for the most costly strain step and using the VOR/LOU distancing method. One reason for the larger memory consumption are the storage demands for position values for a portion of the 26 periodic images to all material points. Another reason is that Voronoi polyhedra and their facets have to be stored for tessellation based grain reconstruction.

Another contribution is due to the conservative strategy we used to implement the querying structures. This could be optimized in the future with potential for a memory footprint reduction of at least a factor two.

5. Conclusions

A set of strong and weak scaling post-processing methods were developed to quantify the accumulation of state variable values at grain boundaries in 3D full-field spectral method crystal plasticity simulations. Exemplified for the DAMASK spectral solver and a phenomenological constitutive crystal plasticity model the results substantiate.

- Voxel accurate methods were developed which reconstruct successfully the grain boundaries and operate on arbitrarily deformed and periodically confined RVE domains with respect to the deformed configuration. This allows for direct comparison to experiments and rigorous numerical assessment of the spectral method.
- Long range gradients of material point disorientation, with higher values to the boundary than in the grain interior, were quantified. They are a signature of incipient grain fragmentation.
As an alternative method to capture the individual fragments, also a graph clustering grain reconstruction method was assessed. Compared to the classical method of assigning grain IDs based on the initial conditions, though, the quantification of the gradients is very parameter dependent.

Three methods were detailed to quantify the distance of a material point to the boundary. All performance critical computational tasks were implemented distributed and shared memory parallel. Therewith, processing the entire set of 28 strain steps from a 256³ RVE cube DAMASK simulation, worth 162 GB, was possible in 470 s wall clock time using 28 processes each spawning 40 threads. Compared to exclusive multithreaded (11 063 s) or sequential execution (150 000 s), this hybrid approach allowed for 320 times faster post-processing. The efficiency could be improved further if heterogeneous work load across the individual strain step data, which is method dependent, gets better balanced and remaining sequential code portions parallelized.

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Data and code availability

The source code of the tool and its supplementary MATLAB scripts with which we post-processed all results are open source software. This software, all DAMASK post-processing parameter settings files including the SLURM batch system submission scripts are available for download [105]. The entire repository of compressed post-processed data occupies 844 GB. It is available from the authors upon serious request.

Work distribution

MK designed and implemented the tool surplus performed and post-processed the simulations. MK and FR wrote the manuscript and discussed the results on a regular basis.

Appendix A. Detailed specification of the post-processing workflow

Algorithmic details. In what follows, the individual steps of the post-processing workflow are detailed. For all steps with a larger than linear time complexity $O(N^2N_0N_0, \ldots = N^2)$, additional details are commented. In every step, $N$ denotes the total number of unique material points supporting the RVE. Indices $n$ specify the strain step. Three point cloud sets are distinguished: the first contains the $N$ unique material points of the original domain $\mathcal{P}_n^0 \in \mathbb{R}^3$. The second, $\mathcal{P}_{n+1}^0$, contains $\mathcal{P}_n^0$ surplus all periodic images of $\mathcal{P}_n^0$ which lay inside a cuboidal bounding box about a cuboidal tight bounding box to $\mathcal{P}_n^0$. The outer bounding box
is fattened by $\epsilon = 0.1$ in length units of the initial RVE domain. The third point cloud set, $\mathcal{P}_n^1$ contains $N_n^1$ points, i.e. all members of $\mathcal{P}_n^0$ plus all their 26 periodic copies. The members $i$ of the material point ensemble $\mathcal{P}_n^0$ have positions $p_{i,n}^0$, which are defined by their initial locations in the RVE in the reference configuration $[38, 44]$ plus a deformation-induced displacement which maps into the deformed configuration in the laboratory coordinate system $\Delta x_{i,n}^q$. Every material point has associated state variable values $\{s_{i,n}\}$. In addition, a reference ID and periodic image variant ID is stored to track which periodic image points are assigned to which unique material point of $\mathcal{P}_n^0$. This allows for a dereferencing of state variable values $s_{i,n}$ rather than duplicating them. IMKL specifies in which steps the Intel Math Kernel Library is used.

**Algorithm 1.** Post-processing workflow.

1: procedure DAMASKPDT
2: I/O parse parameter and initialize MPI
3: I/O parse spectralOut file layout to identify content
4: Partition $N_n$ strain increments round-robin on MPI processes
5: for $\epsilon_{n} ^{-1} \rightarrow N_n$ do
6: Process parallelized MPI I/O spectralOut file reading
7: $\triangleright$ State variable values $s_{i,n}$ per point $x, F, P, P, q$
8: Spatial distributing of material point cloud to OpenMP threads
9: RVE averaging $F, P, m, c, x$ via IMKL
10: Threaded displacements $\Delta x_{i,n}^q$ via IMKL $\triangleright O(N \log(N))$
11: Compute periodic images $\mathcal{P}_n^{q+1}$
12: Build a spatial index $B_{n+1}^q$ that partitions $\mathcal{P}_n^{q+1}$
13: Threaded grain reconstruction
14: if Initial assignment/texture ID based then
15: RECONSTRUCT_GRAINS_TEXTUREID (see B)
16: if Community detection based then
17: RECONSTRUCT_GRAINS_LOUVAIN (see B)
18: GRAIN_GEOMETRY_SIMPLIFICATION (see B)
19: Threaded state variable/distance to boundary quantification
20: if Disorientation thresholding based scalar distances then
21: ANALYZE_DIS (see C)
22: if Signed distance/voxelization based normal distances then
23: ANALYZE_SDF (see C)
24: if Tessellation based contour hull normal distances then
25: ANALYZE_VOR (see D)
26: MPI parallel I/O binary results $(d, \sigma, q)_{n}^{-1}$ voxel or $i$
27: MPI aggregate flow curve data $(\epsilon_{M}, \sigma_{M})_{n}$
28: I/O flow curve, internal profiling results, finalize MPI and exit

**Appendix B. Detailed specification of the grain reconstruction methods**

*Texture ID based grain reconstruction.* $T_i$ specifies the initial texture ID assigned during microstructure synthesis DAMASK [38, 44].
Algorithm 2. Texture based grain reconstruction.

1: procedure RECONSTRUCT_GRAINS_TEXTUREID
2: for $i \leftarrow 1, N$ do
3: Assign point $i$ a grain ID $\mathcal{G}_{i,n} = T_i$

Graph clustering based grain reconstruction $N_{mv}$ specifies the number of executed node relabeling operations in the current iteration step, $Q$ the Newman–Girvan modularity [58] using a critical quality value of $Q_c = 0.01$. $K_L$ specifies the penalization strength, i.e. whether a weak $K_L = 75$ or a strong $K_L = 1000$ penalization was used. $||q_{i,j,n}||$ denotes the rotation angle argument of the disorientation quaternion. The search radius during graph edge construction was chosen as $R_{clv} = 256$ distance units.

Algorithm 3. Louvain community based grain reconstruction.

1: procedure RECONSTRUCT_GRAINS_LOUVAIN
2: for $i \leftarrow 1, N$ do
3: Find all neighboring points $p_{i,n} \in \mathcal{P}_{n_0} \cup \{||x_{i,n} - x_{j,n}|| \leq R_{clv}^n\}$
4: Compute disorientation angle $q_{i,j,n}$
5: Compute edge weights $w_{i,j,n} = \exp(K_L \cdot (||q_{i,j,n}|| - 1))$
6: Add one graph edge with weight $w_{i,j,n}$ for every position pair $p_{i,n}, p_{j,n}$
7: do
8: Iterative graph clustering according to [59, 60, 106]
9: while $N_{mv} > 0$ and $Q > Q_c$
10: for $i \leftarrow 1, N$ do
11: Assign $i$ the final label of the top-level community as grain ID

Grain geometry simplification. $N_{k,n}^0$ and $N_{k,n}^1$, respectively are the number of material points supporting the $k$th grain in $\mathcal{P}_{n_0}$ and $\mathcal{P}_{n_1}$ respectively. $A_{k,n}$ specifies a tight axis-aligned bounding box about the positions $N_{k,n}^0$. $A_n$ is the tight cuboidal global bounding box which contains all $A_{k,n}$. In this work a cell size of half the normalized initial material point point-to-point distance was used, i.e. $d_{cell} = \frac{1}{\sqrt{356}}$.

Algorithm 4. Threaded extraction of a single periodic image per grain.

1: procedure GRAIN_GEOMETRY_SIMPLIFICATION
2: Re-organize members of $\mathcal{P}_{n_0}$ into disjoint sub-sets $\mathcal{P}_{k,n}^0$ per grain $k$ ▷ In what follows, multithreaded processing $\forall k$ grains
3: Threaded memory initialization
4: Threaded periodic images $\mathcal{P}_{k,n}^1$ and bounding boxes $A_k$
5: Threaded building of sparse spatial index $\mathcal{B}_{k,n}^1$ via $B_k^1$
6: Threaded merging of possible grain fragments via DBScan [61] $\mathcal{O}(N_k \log(N_k))$
7: Threaded picking of one representative per grain $\hat{A}_{k,n}$
8: Identify global bounding box $A_n$ enclosing all $\hat{A}_{k,n}$
9: Define a global voxelization $L_n$ of volume $A_n$ with cell size $d_{cell}$
10: Fuse members of $\mathcal{B}_{k,n}^1$ into one global spatial index $B_n^1$
Appendix C. Detailed specification of the distancing methods

Disorientation based distancing. A disorientation angle of $\Theta = 15^\circ$ was used for thresholding.

Algorithm 5. Disorientation thresholding based scalar distances.

1: procedure ANALYZE_DIS
2: for $i \leftarrow 1, N$ do \Comment{Multithreaded, dynamic scheduling}
3: Find all $N_j$ neighbors $P_{ij}^{0 \rightarrow} \in P_n^{0 \rightarrow} \mid \|x_{i} - x_{j}\| \in R_n^3$
4: Sort by distances $d_{i,j,n} := \|x_{i,n} - x_{j,n}\|$ \Comment{$O(N_j \log(N_j))$}
5: for $j \leftarrow 1, N_j$ do
6: Compute disorientation angle $\Theta_{i,j,n}$
7: if $\Theta_{i,j,n} \geq \Theta_i$ then
8: Report $(d_{i,j,n}, s_{i,n})$
9: break \Comment{ignore $i$ if no pair found}

Signed distance/voxelization based normal distances. Abbreviations denote a SDF and the FSM [64, 68].

Algorithm 6. Signed distance/voxelization based normal distances.

1: procedure ANALYZE_SDF
2: Threaded voxelizing of $k$ according to $L_n, d_{cell}$ using $B_k^n$
3: Specifically, $\forall$ voxel $\in B_k^n, \text{identify closest member of } P_n$
4: Threaded initialization of SDF $\Phi_k(x)$
5: Threaded spreading of SDF via FSM $\Phi_{k,n}(x)$ as in [64]
6: Threaded report $(\Phi_k(x), s_{j,n}) \forall$ voxel with $\Phi_{k,n}(x) \geq 0$

Appendix D. Tessellation based contour hull normal distances

$\hat{A}_{k,\alpha}$ specifies an axis-aligned bounding box about the identified single periodic image of grain $k$. The box is fattened by a guard zone of thickness $\frac{3}{325}$ of the normalized initial material point-to-point distance. The fattening assures that the volume about each material point belonging to the grain $k$ can be tessellated and no Voronoi cell gets arbitrarily cut off by the bounding box walls. The query structure of the Voro++ library was configured to include five points per spatial bin on average. For each grain $k$ interior points are distinguished from exterior points with respect to their logical identity when building the exterior contour hull of the grain. Specifically, interior points $ip$ originate from a material point in $P_n$ with grain ID $k$. BVH is short for bounded volume hierarchy. $N_{ip}^k$ denotes the number of exterior facets identified for each grain $k$. $N_{ip}^k$ is the number of interior points per grain $k$. Exterior facets to a Voronoi cell of an interior point are all facets to the first order neighboring cells of an exterior point if any such exist.
Algorithm 7. Tessellation based contour hull normal distances.

1: procedure ANALYZE_VOR \(\triangleright \forall k\)
2: Threaded pulling of all points \(\hat{A}_k\) from \(B^n_k\)
3: Threaded mapping periodic image point to unique material point
4: Threaded Voro++ build spatial index to accelerate tessellator
5: Threaded Voro++ tessellate all Voronoi cells for points
6: Threaded Voro++ exterior facets, consistent outer unit normal
7: procedure IDENTIFY_NORMAL_DISTANCES
8: Threaded construction of a BVH with all facet polygons
9: \[\text{for } ip \leftarrow 1, N^n_k \text{ do}\]
10: \[\text{Pool all } nf \text{ facets } f \text{ of the contour as initial candidates}\]
11: \[\text{while } f < nf \text{ do}\]
12: \[\text{Normal projection interior point } x_{in} \text{ on facet plane } x_{proj}^f\]
13: \[\text{if } x_{proj} \text{ covered by facet polygon } = \text{ false then}\]
14: \[\text{for } e \leftarrow 1, N^e_f \text{ do } \triangleright \text{Circulate facet edges } e \text{ of } f\]
15: \[\text{Normal projection } x_{in} \text{ on } e\]
16: \[\text{if } x_{proj} \text{ on the edge } = \text{ true then}\]
17: \[\text{Eventually update current shortest distance}\]
18: \[\text{Eventually re-query facet candidate pool}\]
19: \[\text{for } v \leftarrow 1, N^v_f \text{ do } \triangleright \text{Circulate facet vertices } v \text{ of } f\]
20: \[\text{Compute distance } d_{in,v} = \left\| x_{in} - x_v \right\|\]
21: \[\text{Eventually update current shortest distance}\]
22: \[\text{Eventually re-query facet candidate pool}\]
23: \[\text{else}\]
24: \[\text{Eventually update current shortest distance}\]
25: \[\text{Eventually re-query facet candidate pool}\]
26: Threaded report value pairs \((d_{ip,n}, s_{ip,n}) \forall ip\)

ORCID IDs

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