Identifying Structure–Property Relationships Through DREAM.3D Representative Volume Elements and DAMASK Crystal Plasticity Simulations: An Integrated Computational Materials Engineering Approach

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Predicting, understanding, and controlling the mechanical behavior is the most important task when designing structural materials. Modern alloy systems—in which multiple deformation mechanisms, phases, and defects are introduced to overcome the inverse strength–ductility relationship—give rise to multiple possibilities for modifying the deformation behavior, rendering traditional, exclusively experimentally-based alloy development workflows inappropriate. For fast and efficient alloy design, it is therefore desirable to predict the mechanical performance of candidate alloys by simulation studies to replace time- and resource-consuming mechanical tests. Simulation tools suitable for this task need to correctly predict the mechanical behavior in dependence of alloy composition, microstructure, texture, phase fractions, and processing history. Here, an integrated computational materials engineering approach based on the open source software packages DREAM.3D and DAMASK (Düsseldorf Advanced Materials Simulation Kit) that enables such virtual material development is presented. More specific, our approach consists of the following three steps: (1) acquire statistical quantities that describe a microstructure, (2) build a representative volume element based on these quantities employing DREAM.3D, and (3) evaluate the representative volume using a predictive crystal plasticity material model provided by DAMASK. Exemplarily, these steps are here conducted for a high-manganese steel.

INTRODUCTION

Controlling the mechanical behavior is the key task when developing materials for structural applications. Replacing mechanical tests by simulation studies to evaluate the mechanical performance of candidate alloys is highly desirable as it enables a significant reduction of resource allocation in the alloy design process. However, in order to get reliable results, the simulation tool needs to correctly predict the mechanical behavior in dependence of alloy composition, microstructure, and texture.

In this study, an integrated computational materials engineering (ICME) approach that enables such virtual material development is presented. It is based on the DREAM.3D1 and DAMASK2 (Düsseldorf Advanced Materials Simulation Kit) software packages and consists of the following three steps:

1. Acquire statistical quantities that describe a microstructure.
2. Build a representative volume element (RVE) based on these quantities using DREAM.3D.
3. Evaluate the RVE using a predictive crystal plasticity material model implemented in the DAMASK framework.

Exemplarily, these steps are here conducted for a high-manganese steel (HMnS). Like other modern high-performance alloys, HMnS combine multiple deformation mechanisms to overcome the inverse strength–ductility relationship. Besides dislocation glide, both, transformation-induced plasticity (TRIP) and twinning-induced plasticity (TWIP), serve as additional deformation mechanisms. From an engineering point of view, additional deformation mechanisms create the challenge that they are influenced by several microstructural and environmental parameters such as grain size, texture, chemical composition, temperature, strain rate, and the nonlinear interactions between them. These complex interactions and influencing factors can significantly hamper accelerated alloy design, particularly when using the material under complex loading and strain path conditions such as commonly encountered in engineering sheet-forming applications. Hence, for complex engineering materials, it is indispensable to evaluate the performance of potential microstructures by fast and reliable simulations before conducting mechanical tests and even before synthesizing prototype alloys. With such simulation studies, a suitable regime in the huge parameter space can be located beforehand in order to conduct classical alloy prototyping and mechanical tests only for target-oriented validation of the computed domain.

In the proof-of-concept study presented here, statistical microstructural quantities are retrieved from an existing and experimentally well-characterized material. This allows for a comparison of calculated results with experimental data to evaluate the capabilities of the approach. However, as outlined below, using a predictive crystal plasticity model and synthetic microstructure generation raises the opportunity to use the procedure to also investigate new materials with the aim of forecasting suitable microstructures for different loading states.

The study is organized as follows. First, to give the reader a background of the investigated model alloy, a concise synopsis on HMnS is provided. Next, we explain how the microstructure features obtained from the experimental characterization are translated into appropriate statistical quantities. Then, it is discussed how these microstructure measures are subsequently used for the generation of representative microstructures using DREAM.3D. After that, we explain the simulation details of the crystal plasticity model implemented into the DAMASK software package. Finally, after presentation of the simulation results, we draw conclusions and provide an outlook on how to further improve and apply the methodology.

MODEL MATERIAL

The concept of HMnS is based on stabilizing the face-centered cubic (fcc) austenite phase. This is usually accomplished by adding a high amount of manganese (15–30 wt.%). Small proportions of carbon (0.05–1.00 wt.%), aluminium (0.0–3.0 wt.%) and/or silicon (0.0–3.0 wt.%) can be added for tuning the stacking fault energy and the oxidation layers. HMnS are considered to be part of the second generation of advanced high-strength steels (AHSS). The active deformation mechanism(s) in HMnS mainly depend(s) on the stacking fault energy (SFE), which is normally below 20 mJ m\(^{-2}\) for TRIP steels and in the range between 20 mJ m\(^{-2}\) and 40 mJ m\(^{-2}\) for TWIP steels. In HMnS with SFE values above 45 mJ m\(^{-2}\), dislocation glide dominates plastic deformation, whereas TRIP and TWIP effects are suppressed. The selected TWIP steel has a composition of 22.5Mn-1.2Al-0.3C wt.%. Its SFE was determined by a subregular solution thermodynamic model as approximately 25 mJ m\(^{-2}\). Hence, deformation twinning is expected to be the only active deformation mechanism besides dislocation glide. Since the material has been extensively investigated in previous works, details on the production and post-processing procedure are not repeated here.

INTEGRATED COMPUTATIONAL MATERIALS ENGINEERING PROCEDURE

Experimental Characterization

From the initial material in hot-rolled condition, six different microstructural states have been produced by imposing different (thermo-) mechanical treatments. First, the material was cold-rolled to 30%, 40%, and 50% thickness reduction. From these three states, recrystallized samples have been produced by subsequent annealing at 700\(^\circ\)C for 15 min after 30% reduction and for 10 min after 40% and 50% reductions.

Details on sample preparation for the following electron backscatter diffraction (EBSD) and x-ray diffraction analyses used for the material characterization can be found in an already published study.

A LEO 1530 field emission gun scanning electron microscope operated at 20 kV accelerating voltage and a working distance of 10 mm was used for EBSD analyses. EBSD mappings were generated with a step size of 0.28μm. The HKL Channel 5 software was utilized for data post-processing and the removal of wild spikes and non-indexed points, taking at least five neighbor points into account. Based on EBSD measurements, grain size distributions of the cold-rolled and subsequently recrystallized state were retrieved. In contrast, the shape and the mean grain size of the cold-rolled samples were estimated from measurements on the material in the hot-rolled state. EBSD band contrast maps of
the 40% cold-rolled samples before and after recrystallization are shown exemplarily in Fig. 1. The typical grain morphology of cold-rolled material, i.e. grains elongated along the rolling direction (RD) and compressed along the normal direction (ND), is shown in Fig. 1a. The smaller, equiaxed grains of the recrystallized sample are displayed in Fig. 1b.

The crystallographic texture was characterized by means of x-ray pole figure measurements. Three incomplete (0° to 85°) pole figures—\{111\}, \{200\}, and \{220\}—were acquired at the mid-thickness layer of the sheet on a Bruker D8 Advance diffractometer, equipped with a HI-STAR area detector, operating at 30 kV and 25 mA using filtered iron radiation and polycapillary focusing optics. The orientation distribution functions (ODFs) were calculated in the MATLAB-based MTEX package.\(^\text{20,21}\) \(\varphi_2 = 45^\circ\) sections of the ODF of the six investigated states are shown in Fig. 2. The corresponding legend of the ideal texture components is given elsewhere.\(^\text{22}\) With increasing rolling degree, the texture transformed gradually from Cu-type to Brass-type, as indicated by the decreasing \{112\}(111) Cu texture component and the more pronounced \(\alpha\)-fiber (\langle110\rangle\parallel ND) with a spread towards the \{552\}(115) CuT texture component. After recrystallization, the rolling texture was retained but significantly weakened in intensity as a result of the oriented nucleation and the formation of recrystallization twins.\(^\text{23,24}\)

The obtained grain size distribution data as well as the orientation information obtained from the experimental ODFs are then used as input to generate the RVEs as outlined in the following.

**Representative Volume Element Generation Using DREAM.3D**

The DREAM.3D software\(^\text{1}\) was used to generate the RVEs using statistics extracted from the experimental data following a procedure outlined elsewhere.\(^\text{25,26}\) DREAM.3D has been extensively used to
characterize experimentally acquired microstructures and generate microstructure-based models for subsequent simulations.\textsuperscript{27–32} Since it produces 3D virtual volumes, the experimentally obtained data were extrapolated from 2D to 3D based on some underlying assumptions. First, the diameter of the sphere-shaped grains in the original (hot-rolled) state was determined experimentally. These spheres were then assumed to deform isomorphically during 30\%, 40\%, and 50\% thickness reduction to yield the grain morphology of the solely cold-rolled states. In contrast, the grains in the cold-rolled and subsequently recrystallized states were assumed to be nearly equiaxed, i.e. the measured diameters ($d$) along RD, ND, and transverse direction (TD) are similar ($0.8 < d_{ND} / d_{RD} < d_{TD} / d_{RD} < 1.0$). The average grain diameters for each state were then determined by calculating the mean lineal intercept in 2D and extrapolating to 3D by the relationship $d_{3D} = 1.5d_{2D}$. The crystallographic texture in each state was used directly from the calculated ODFs. The microstructure was simplified in such a way that a single orientation was assigned to each grain and no internal orientation gradients exist for either cold-rolled nor cold-rolled and recrystallized states. It is important to note that the constructed volume elements are microstructurally representative, and multiple instantiations will produce volumes with similar microstructural characteristics. However, the volumes can be termed properly representative only if multiple instantiations yield similar simulated properties (Fig. 3; Table I).

### Crystal Plasticity Simulation Using DAMASK

We used a dislocation-based crystal plasticity constitutive model from the DAMASK package suitable for capturing the mechanical characteristics of the investigated material class.\textsuperscript{8} This constitutive formulations introduced in DAMASK for the prediction of the mechanical response of TWIP and TRIP steels is capable of reproducing the active deformation modes depending on the SFE of the material. It is important to realize that the SFE can be directly computed from ab initio simulations. Hence, a bridge from the atomistic scale to the component scale can be established, and the model is predictive with respect to the chemical composition. As can be seen from Tables II and III, almost all other parameters of the crystal plasticity model also have a clear physical meaning, physics-related upper and lower bounds, and can be determined from small-scale simulations or microstructure characteristics.

The same material parameters as used in a previous study,\textsuperscript{15} have been applied with the exception of the grain size $d$ and initial values for the edge dislocation density $\rho_e$, the dipole dislocation density $\rho_d$, and the twinned volume fraction $\eta_t$.

#### Table I. Grain morphology properties of the generated representative volume elements

| State     | Size/$\mu$m$^3$ | Grain count | Shape    |
|-----------|-----------------|-------------|----------|
| 30\% CR   | 640.0           | 4086        | Ellipsoidal |
| 30\% CR + RX | 128.0         | 3859        | Spherical   |
| 40\% CR   | 640.0           | 4041        | Ellipsoidal |
| 40\% CR + RX | 64.0          | 4585        | Spherical   |
| 50\% CR   | 640.0           | 4044        | Ellipsoidal |
| 50\% CR + RX | 51.2          | 4664        | Spherical   |

For the cold-rolled and recrystallized samples, the average grain size diameter is determined from the RVE characteristics given in Table I: first, the average grain volume is computed as the RVE size divided by number of grains, followed by the calculation of the diameter of a sphere with this volume.
For the cold-rolled states, ellipsoidal grain shapes are assumed which result from the rolling of the initially spherical grains with diameter \( d = 27 \mu m \). The shortest axis of the ellipsoid (along ND) \( d_{ND} = 27 \mu m / \varepsilon^{2} / C^{2} \) is taken as the limiting size that determines the mean free path for dislocation glide. The initial values of \( q_{0e} \), \( q_{0d} \), and \( f_{0t} \) are derived from the results of a plane strain compression simulation by calculating the average over all 12 slip and 12 twin systems at the respective deformation level, i.e. neglecting any partitioning to specific systems.

Simulations are performed using a spectral method coupled with DAMASK. The RVEs are subjected to uniaxial tension at a loading rate of \( 1 \times 10^{-3} \text{s}^{-1} \). The microstructures representing the cold-rolled states are loaded along all three directions (RD, ND, TD) to investigate the anisotropy introduced by the preceding deformation. As no such anisotropy is expected for their recrystallized counterparts, owing to the weak crystallographic textures and the absence of grain shape effects, those are loaded only along RD. The final strain levels have been adjusted to the experimentally obtained values for the cold-rolled states and are set to a true strain of \( \varepsilon = 0.16 \) for simulation in cold-rolled and recrystallized conditions.

**SIMULATION RESULTS OBTAINED BY DAMASK**

Stress–strain curves of the tensile tests conducted along RD are shown in Fig. 4 for all investigated material states. The significant hardening introduced by the cold rolling is clearly captured by the model, with the yield point of the sample with 50% thickness reduction being more than 1.0 GPa higher than for the states with preceding recrystallization heat treatment. Moreover, the effect of the different grain sizes among the cold-rolled and recrystallized conditions can also be seen, i.e. the 30% cold-rolled and recrystallized sample hardens less than the 50% cold-rolled and recrystallized sample. Comparing these results with the experimental results (lines marked by dots), a good qualitative agreement in the elastic–plastic deformation regimes, i.e. before damage is expected to set in, can be observed. Also, the evolution of the twin volume fraction \( f_{t} \)

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Table II. Values of the parameters and their symbols used in the crystal plasticity model for all simulations

| Parameter | Symbol | Value | Unit |
|-----------|--------|-------|------|
| Burgers vector magnitude | \( b_{s} \) | \( 2.56 \times 10^{-10} \) | m |
| Activation energy slip | \( Q_{s} \) | \( 1.5 \times 10^{-19} \) | J |
| Activation energy climb | \( Q_{c} \) | \( 2.0 \times 10^{-19} \) | J |
| Activation volume climb | \( V_{c} \) | \( 1.7 \times 10^{-29} \) | m³ |
| Obstacle profile top | \( p \) | 1.0 | |
| Obstacle profile bottom | \( q \) | 1.0 | |
| Mean free path in multiples of dislocation spacing | \( i_{s} \) | 45.0 | |
| Mean free path in multiples of twin spacing | \( i_{t} \) | 1.0 | |
| Self-diffusion prefactor | \( D_{0} \) | \( 4.0 \times 10^{-5} \) | m²s⁻¹ |
| Minimum dipole spacing | \( C_{anni} \) | \( 8.96 \times 10^{-28} \) | m |
| Velocity prefactor | \( v_{0} \) | \( 1.0 \times 10^{4} \) | ms⁻¹ |
| Solution strengthening | \( s_{sol} \) | \( 2.0 \times 10^{7} \) | Pa |
| Burgers vector | \( b_{t} \) | \( 1.47 \times 10^{-10} \) | m |
| Nucleus width | \( L_{t} \) | \( 2.56 \times 10^{-7} \) | m |
| Avg. twin thickness | \( t_{t} \) | \( 5.0 \times 10^{-8} \) | m |
| Activation volume cross-slip | \( V_{cs} \) | \( 5.0 \times 10^{-32} \) | m³ |
| Profile width exponent | \( A \) | 13.96 | |
| SFE | | 25.0 | mJm⁻² |

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aInteraction coefficients for hardening are obtained by discrete dislocation dynamics simulations.33
follows the experimental observations: essentially, there is no twinning activity at the simulated deformation levels. The yield stress values for the cold-rolled and recrystallized state are, however, lower by approximately 180 MPa and the hardening rate is underestimated.

When analyzing the stress–strain curves predicted for tensile tests along the three different loading directions given in Fig. 5, it can be seen that a significant anisotropy is introduced in the cold-rolled state. This behavior is especially pronounced for 50% thickness reduction. Except for the 30% cold-rolled and recrystallized state, where only small anisotropy is observed, loading along RD gives the softest response.

Given the good qualitative agreement between the experimentally observed and simulated average responses, the simulation results can be used to study the spatial stress and strain distributions. With typical experimental equipment, these quantities cannot be readily determined at all (in the case of stress), or the acquisition requires significant efforts (in the case of strain, where, e.g., digital image correlation techniques are needed). The local distributions of equivalent true strain ($\varepsilon_{eq}$) and stress ($\sigma_{eq}$) at the final loading state, i.e., at $\varepsilon = 0.16$ for the 30% cold-rolled and recrystallized sample are shown in Fig. 6. We observe a significant and broad range in the distribution of stress and strain—not only among different grains but also within individual grains. Even though a quantitative agreement to the experimental stress–strain response cannot be achieved without tedious model parameter fitting, the quantitative analysis of the stress and strain distribution gives helpful insights for the development of improved alloys.

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**Table III. Values of the parameters (grain size: $d$; edge dislocation density $\rho_e^0$; dipole dislocation density $\rho_d^0$; twinned volume fraction $f_t^0$) used in the crystal plasticity model adjusted for specific material states**

| State             | $d/\mu m$ | $\rho_e^0 / m^2$ | $\rho_d^0 / m^2$ | $f_t^0$ |
|-------------------|-----------|------------------|------------------|---------|
| 30% CR            | 28.35     | $4.5 \times 10^{13}$ | $8.1 \times 10^{13}$ | 0.0033  |
| 30% CR + RX       | 10.04     | $1.0 \times 10^{12}$ | 1.0              | 0.0     |
| 40% CR            | 24.29     | $7.0 \times 10^{13}$ | $1.4 \times 10^{14}$ | 0.0039  |
| 40% CR + RX       | 4.78      | $1.0 \times 10^{12}$ | 1.0              | 0.0     |
| 50% CR            | 20.25     | $9.2 \times 10^{13}$ | $1.7 \times 10^{14}$ | 0.0041  |
| 50% CR + RX       | 3.80      | $1.0 \times 10^{12}$ | 1.0              | 0.0     |

CR cold-rolled, RX recrystallized.
CONCLUSION AND OUTLOOK

The presented ICME approach can assist, accelerate, and guide the design of new alloys and suitable processing pathways. Based on statistical descriptions of microstructures, RVEs are created on the basis of which the material’s performance is evaluated. The use of DREAM.3D enables microstructure RVE generation at high fidelity, incorporating statistical features which can be evaluated using the spectral solver and physically-based crystal plasticity laws for specific materials\(^8\),\(^{41,42}\) available in DAMASK. Such a strictly microstructure-oriented ICME approach enables the drastic reduction in the number of experiments typically required in alloy, microstructure, and process development.

To further strengthen the link between measured microstructural features and generated RVEs, additional statistical quantities can be taken into account. Features that might have an influence on the mechanical behavior could be the in-grain orientation distribution (especially for the cold-rolled states), the grain-to-grain misorientation in the form of a misorientation distribution function,\(^{43}\) and the exact grain shape. While incorporating rules to construct microstructures based on these additional pieces of information is a challenging task by itself, for a comparison with experimental results, data acquisition efforts will also increase. For both crystallographic features, the experimental characterization would need to be expanded to obtain not only global orientation information but also spatially resolved and neighborhood-sensitive quantities\(^1\). Similarly, measurement complexity for a more exact grain shape determination requires statistically relevant 3D information, e.g., from EBSD measurements conducted on mutually orthogonal surfaces.

The employed crystal plasticity model has revealed its predictive capabilities for different chemical compositions and various temperatures.\(^3\) Most of the parameters used in this model have a physical meaning and can be obtained from small-scale simulations. However, two weak points in the model can be identified that are deemed to impede a quantitative agreement with experimental results. First, the grain size is only considered as a fixed average value to limit the mean free path for dislocation slip and does not affect the local hardening based on dislocation density gradients. Out of the various approaches of taking gradients into account, the physically most sound approaches are based on the flux of dislocations.\(^{42,44}\) While, on the one hand, the associated computational costs prohibit the use of such constitutive models for large-scale simulations as presented here, on the other hand the effect on macroscopic properties is probably not that pronounced.\(^{45}\) Secondly, solid solution strengthening is also not implemented in a physics-based way and its temperature dependence is neglected in the current approach.\(^8\)

Another area of current research where progress will increase the quality of the results is the coupling of damage and crystal plasticity models. As damage is the dominating mechanism for the highly cold-rolled states, their behavior cannot be captured correctly within the current simulation framework. Nevertheless, current approaches of implementing damage models into DAMASK\(^{46,47}\) will allow the tackling of this challenge in the near future.

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