Correlation between Taylor Model Prediction and Transmission Electron Microscopy-Based Microstructural Investigations of Quasi-In Situ Tensile Deformation of Additively Manufactured FeCo Alloy

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Submitted: 20 May 2021 / Accepted: 11 June 2021 / Published online: 3 August 2021

Within this research, the multiscale microstructural evolution before and after the tensile test of a FeCo alloy is addressed. X-ray μ-computer tomography (CT), electron backscattered diffraction (EBSD), and transmission electron microscopy (TEM) are employed to determine the microstructure on different length scales. Microstructural evolution is studied by performing EBSD of the same area before and after the tensile test. As a result, \(\{001\}||TD\), \(\{011\}||TD\) are hard orientations and \(\{111\}||TD\) is soft orientation for deformation accommodation. It is not possible to predict the deformation of a single grain with the Taylor model. However, the Taylor model accurately predicts the orientation of all grains after deformation. \{123\} \(\langle111\rangle\) is the most active slip system, and \{112\} \(\langle111\rangle\) is the least active slip system. Both EBSD micrographs show grain subdivision after tensile testing. TEM images show the formation of dislocation cells. Correlative HRTEM images show unresolved lattice fringes at dislocation cell boundaries, whereas resolved lattice fringes are observed at dislocation cell interior. Since Schmid’s law is unable to predict the deformation behavior of grains, the boundary slip transmission accurately predicts the grain deformation behavior.

Keywords EBSD, FeCo alloy, quasi-in situ tensile test, Taylor model, TEM

1. Introduction

Soft magnetic materials are important components of electric machine devices requiring high flux density. Among soft magnetic materials, equiatomic FeCo alloys possess the highest Curie temperature and good permeability combined with highest saturation magnetization. These alloys also reveal a soft magnetic materials, equiatomic FeCo alloys possess the electric machine devices requiring high flux density. Among previous study (Ref5), this paper characterizes the microstructure and voids) showing dislocation accommodation and also impeding dislocation motion (Ref5). Thus, following the previous study (Ref5), this paper characterizes the microstructure on multiple length scales (from μm to sub-nm level) to understand the deformation accommodation mechanism of the AM FeCo alloy using a quasi-in situ approach. A detailed multiscale microstructural characterization enables to observe deformation behavior from millimeter to nanometer lengths. Therefore, the internal porosity is studied by x-ray microtomography (μ-CT) on the millimeter scale. Subsequently, the grain rotation of the same region before and after tensile testing is examined via electron backscattering diffraction (EBSD) at the submicron scale. In this regard, the grain rotation behavior is examined using the Schmid factor and grain boundary slip transmission factor. The grain boundary shape change upon deformation is analyzed using fractal behavior. The Taylor model is used to explain the overall texture of the same sample region before and after the tensile test. Here, the focus is to identify active slip systems and to explain the overall deformation behavior. Further, the evolution of the deformation substructure is addressed by transmission electron microscopy.

The online version contains supplementary material available at https://doi.org/10.1007/s11665-021-06065-9.

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(TEM) and correlative high-resolution transmission electron microscopy (HRTEM) on the nanometer level. The dislocation density is also calculated from the deformation substructure with the help of energy-filtered transmission electron microscopy (EFTEM).

2. Experimental Procedure

2.1 Sample Preparation

Spherical pre-alloyed FeCo powder containing 50 wt.% Co is produced by gas atomization. The average d50 (diameter of 50% particles) is 38 μm. All FeCo specimens are produced by LBM (SLM250HL machine, SLM Solutions Group AG, Lübeck, Germany) applying a 400 W ytterbium continuous laser with a spot size of 70 μm. Argon gas is supplied during LBM to prevent oxidation, and the building platform is heated to 200 °C to reduce internal stress. Based on our previous parameter optimization study (Ref 8), a laser power of 270 W, a laser with a spot size of 70 μm, a scan speed of 700 mm/s, a hatch distance of 0.11 mm, and a powder layer thickness of 0.05 mm are set as process parameters.

2.2 Tensile Testing

Tensile samples with the gauge section parallel to the build direction (BD) are prepared by LBM. The dimension of the gauge section is 10 mm (length) × 3 mm (width) × 1.5 mm (thickness). Tensile testing (MTS tabletop system 858) is conducted with the tensile direction (TD) parallel to BD. A strain rate of 10⁻³ s⁻¹ is imposed during tensile testing, and the strain is measured with an extensometer.

2.3 μ-CT

To determine the internal porosity, μ-CT (Skyscan 1275, Bruker) is performed on the gauge section before the tensile tests. For this, the μ-CT is operated at 100 kV and 10 W with a tungsten target and copper filter (1 mm). During μ-CT, a sample rotation of 360° is imposed in a step size of 0.2°. A sample volume of 20.2 mm³ is analyzed by μ-CT. For μ-CT data post-processing, a 7 μm voxel size is used.

2.4 EBSD Sample Preparation and Investigation

Before performing the tensile tests, all samples are polished using SiC abrasive paper. The gauge section is further electropolished (Struers Lectropol-5) with an electrolyte containing 590 mL ethanol, 330 mL butoxyethanol, and 78 mL perchloric acid for acquiring EBSD maps. Operating conditions during electropolishing are 20 V and 1 °C for 60 s.

Before performing the tensile tests, the gauge sections of the electropolished samples are examined via optical microscopy (Zeiss Axiophot). Furthermore, EBSD maps are captured in the same part of the gauge section before and after the tensile tests. For this, EBSD is performed in a scanning electron microscope (SEM, Zeiss Ultra Plus) equipped with an EBSD detector (EDAX). The SEM is operated at 13.5 mm working distance and 20 kV accelerating voltage; the EBSD maps are collected at 200 nm step size and 77 pA probe current.

2.5 EBSD Data Analysis

The EBSD file is exported from TSL OIM 7.1 software to MTEX 5.4.0 software (Ref 9). The EBSD data analysis is done using MTEX 5.4.0 (Ref 9). A misorientation angle larger than 15° is defined as a high angle grain boundary. Finally, the inverse pole figure (IPF) map along TD is plotted. Based on this, the Schmid factor (SF) of grains is calculated considering {110} {111}, {112} {111} and {123} {111} slip systems. Moreover, the Taylor model is applied to predict the deformation behavior during tensile testing. A strain tensor [1, 0, 0; 0, −0.5, 0; 0, 0, −0.5] is imposed for calculation in the Taylor model.

The grain boundary shape before and after the tensile test is characterized by the fractal number. The fractal number is calculated using the multiple iterations box-counting methods. In the box-counting method, the boxes of size r are drawn on the micrograph and the number of boxes containing the grain boundaries is counted. The size of boxes is changed in each iteration, and the number of boxes containing the grain boundaries is counted again. The fractal number is calculated using Eq 1:

\[ N = r^{-D} \]  

where \( N \) is the number of boxes, \( r \) is the size of the box, and \( D \) is the fractal number (Ref 10).

2.6 TEM Sample Preparation and Investigation

After the tensile tests, the gauge section of each sample is cut applying the Struers Secotom-5. For TEM, a disk with a diameter of 3 mm is prepared and subsequently polished to a thickness of \( \approx 100 \mu m \). For preparing the TEM sample, electropolishing is done using a 5% perchloric acid-ethanol solution in Struers Tenupol-5 at a voltage of 21 V, a current of 30 mA, and a temperature of \(-21 \) °C.

Subsequently, TEM is performed using the JEOL JEM-ARM 200F which is equipped with a cold field emission gun and a CEOS hexapole Cs corrector. An emission current of 15 μA is used for sample imaging. A double tilt sample holder is used during all TEM investigations, and HRTEM images are also acquired.

Additionally, EFTEM images are captured to measure the average sample thickness. For capturing the EFTEM images, an energy filter of width 10 eV is selected. An EFTEM image is taken without any image filter. Another EFTEM image is taken with the energy filter placed around the zero-loss peak. The sample thickness \( t \) is calculated using Eq 2:

\[ t = \lambda \ln(I/I_0) \]  

where \( \lambda \) is mean free path, \( I_0 \) is the intensity with the energy filter, and \( I \) is the intensity without the energy filter. The mean free path is calculated from the chemical composition using the Iakoubovskii model (Ref 11). The model is implemented using the Digital Micrograph script (Ref 12).

To calculate the dislocation density, lines are drawn on each TEM micrograph. Afterward, the dislocation density is calculated based on Eq 3:

\[ \rho = \frac{2N}{Lt} \]  

where \( N \) is the number of boxes, \( t \) is the thickness of the sample, \( l \) is the free path, and \( I \) is the intensity without the energy filter.
where $\rho$ is the dislocation density ($m^{-2}$), $N$ is the number of intersection points, $L$ is the total length of all the lines ($m$), and $t$ is the sample thickness ($m$) (Ref 13).

3. Results

3.1 Optical Microscopy and $\mu$-CT

Figure 1(a) shows a schematic of a tensile sample. The gauge section is highlighted in yellow. The optical micrograph of the representative gauge section in Fig. 1(b) proves an almost defect-free sample surface, which is supported by the $\mu$-CT image in Fig. 1(c) showing the gauge section of the same specimen. Only little internal porosity is observed. However, the total porosity is determined to be 0.1%. Figure 1(d) shows a plot of the pore diameter versus sphericity of pores revealed from the $\mu$-CT data. Based on the literature (Ref 14), the pores are classified into gas porosity or incomplete fusion porosity. Most of the pores are spherical (sphericity $\approx 0.9$) and are formed due to gas entrapment during LBM. During LBM, the melt pool is at a higher temperature resulting in high solubility of gases. Due to very fast cooling during LBM, there is insufficient time for the dissolved gases to escape the melt pool (Ref 15). This results in gas entrapment.

3.2 Tensile Test

Figure 2 summarizes the engineering stress–strain curve and strain hardening rate curve for the FeCo specimen. Based on the engineering stress–strain curve in Fig. 2(a), the yield strength is 315 MPa, the ultimate tensile strength (UTS) is 738 MPa, and the total elongation is 34%. The engineering stress–strain curve shows only a little elastic linear behavior, followed by a sharp increase in stress until UTS. Subsequently, after UTS a gradual decrease in stress is observed due to strain localization. The strain hardening rate curve calculated from the true stress–strain curve is shown in Fig. 2(b). Stage 1 is characterized by a rapid decrease in strain hardening rate due to the onset of dynamic recovery. Stage 2 shows a slower decrease in strain hardening rate due to the onset of slip-on different slip systems.
systems. Stage 3 which represents the onset of failure due to strain localization is not shown here.

3.3 EBSD Investigations

Figure 3(a) and (b) shows the EBSD map of the same area before and after the tensile test. The microstructures in Fig. 3 (a) and (b) are acquired from an area away from the necking region. Almost all grains are elongated along with the BD/TD. The average grain size before and after tensile testing is 3.7 μm ± 2.9 μm and 3.7 μm ± 3.1 μm, respectively. Upon plotting the grain orientations on IPF (Supplementary Figure S1), the [111][TD orientations have the lowest SF, whereas [001][TD orientations exhibit the highest SF. Therefore, [001][TD orientated grains are anticipated to undergo deformation. However, experimental data show that [001][TD and [011][TD oriented grains are hard orientation. These orientations do not undergo much rotation and are resistant to deformation accommodation.

During fractal analysis, a linear relationship between the box size and the number of boxes is observed (Supplementary Figure S2). This is due to the fractal nature of the microstructure. From the slope of the curve, the fractal number before and after tensile testing is 1.84 indicating no change in fractal behavior. As there is no change in the fractal behavior, no significant grain boundary shape change is expected for the overall microstructure. But there may be grain boundary shape change on a local scale.

Figure 3(a) displays a [001][TD oriented “grain 1”. Correspondingly, Fig. 3(c) indicates “grain 1” to have a high SF value (> 0.48). However, no significant grain rotation is observed for “grain 1” in Fig. 3(b). Therefore, Schmid’s law is inapplicable to explain the deformation of this grain. Figure 3(a) also highlights “grain 2” with a [1 24][TD orientation. According to Fig. 3(c), this should have a high SF value (> 0.46), too. “Grain 2” undergoes significant rotation upon deformation. Therefore, Schmid’s law is applicable to explain its deformation. Consequently, Schmid’s law is capable to explain the deformation behavior of only some specific grains (here: grain 2).

Also, Fig. 3(d) presents a map with a slip transmission factor (Ref 16) across boundaries. Here, the slip transmission factor quantifies grain boundary ability to transmit slip between two neighboring grains. The slip transmission factor measures the relative orientation of slip planes and slip directions for slip transmission between two neighboring grains. A slip transmission factor value equal to one means no grain boundary resistance to slip transmission. Slip transmission factor (Ref 17) is calculated using Eq 4:

\[ m' = \cos \phi \cdot \cos \kappa \]  

(Eq 4)

where \( \phi \) is the angle between slip plane normals, \( \kappa \) is the angle between slip directions, and \( m' \) is the slip transmission factor.

In Fig. 3(d), “grain 1” has boundaries with low slip transmission (≈ 0.2), whereas “grain 2” has boundaries with high slip transmission (≈ 0.9). Hence, “grain 2” is more susceptible to deformation accommodation compared to “grain 1”. Based on this, boundary slip transmission is capable to explain the deformation behavior of both grains.

3.4 Taylor Model Simulation

It is well known that the deformation behavior of grain is influenced by its neighboring grains. To restrict the number of neighboring grains, the highlighted grain (dotted rectangle in Fig. 3(a) and (b)) is chosen for local analysis. This grain is completely circumscribed within a single grain. Figure 4(a) depicts the EBSD map of the highlighted grain (dotted rectangle in Fig. 3(a) and (b)) before and after tensile testing. A fragmentation of grain orientations (IPF blue dots, Fig. 4(a)) is observed after deformation. Here, the Taylor model is applied to predict grain orientation. As the EBSD observation area is away from the neck section, the deformation level is assumed to be a uniform elongation strain for the Taylor model. It is stated that the Taylor model does not predict the orientation of the individual grain accurately. According to the IPF in Fig. 4(a), a rotation of ≈11° is expected [00 1̅] between the initial orientation and the prediction based on the Taylor model. However, after the tensile test a rotation of ≈6° [1 3̅2] occurred.

Applying the Taylor model to predict the global orientation of all grains in Fig. 3(a) leads to the results summarized in Fig. 4(b). The superimposed IPF in Fig. 4(b) presents all grain orientations before (black dots) and after tensile tests (red dots) as well as the prediction of the Taylor model (blue dots).
Besides, the contour IPF plot shows a clustering of orientations to [056]||TD according to the Taylor model (Supplementary Figure S3). From the experiments, a formation of strong [146]||TD and [023]||TD orientations is stated (Supplementary Figure S4). These orientations are close to [056]||TD which is predicted by the Taylor model. A (101) texture is reported to develop during tensile testing of bcc crystals (Ref 18) and also for FeCo alloy (Ref 5).

3.5 TEM Investigations

Figure 5 shows TEM and HRTEM images from the gauge section after deformation. In Fig. 5(a), yellow arrows indicate grain boundaries. The formation of dislocation cells (red arrows) is observed. The dislocation cell boundaries are not sharp. The size of the dislocation cells is not uniform and varies between \( \approx 130 \) and \( \approx 360 \) nm. The magnified images of dislocation cells from a different region are shown in Supplementary Figure S5. The size of the dislocation cells here is between \( \approx 140 \) nm to \( \approx 400 \) nm. This supports the results from the previous EBSD measurements regarding grain fragmentation (Fig. 3b). The local average thickness of the TEM sample is calculated as \( \approx 51 \) nm (Supplementary Figure S6). The dislocation density in Fig. 5(a) calculated according to Eq 2 is \( 8.5 \times 10^{13} \) m\(^{-2} \). In this regard, our previous study shows a dislocation density in the as-built FeCo alloy sample to be \( 2.2 \times 10^{13} \) m\(^{-2} \) (Ref 8). Thus, an increase in dislocation density is observed upon the tensile test. Deformation-induced substructure formation is not reported for FeCo alloy in previous literature. Previous studies have reported the development of subgrains in FeCo alloy upon cold rolling and annealing. In this regard, for FeCo1.8V0.3Nb alloy, the size of the subgrains is reported to be 100 nm and 150 nm for different annealing treatments (Ref 19). However, for FeCo2V alloy the subgrain size is 1 \( \mu \)m after annealing (Ref 20).

Figure 5(b), (c) and (d) represents the correlative HRTEM images presenting the local lattice structure. Figure 5(b) is taken from the interior of a dislocation cell (Fig. 5a). Figure 5(c) and (d) is taken from a region close to a dislocation cell boundary (Fig. 5a). The inset fast Fourier transform (FFT) pattern in Fig. 5(b) shows FeCo [012] zone axis. The lattice spacing of (121) planes measured from the FFT pattern is \( \approx 0.11 \) nm. Figure 5(c) shows the lattice fringes due to (121) planes with a spacing of \( \approx 0.11 \) nm. Due to the presence of lattice strain near the dislocation cell boundary, the contrast from the lattice fringes is not clear as highlighted by white arrows (Fig. 5c).
The lattice fringes are due to (110) planes with a spacing of \( \approx 0.20 \text{ nm} \) (Supplementary Figure S7). The inset FFT pattern in Fig. 5(d) shows the FeCo \([-\overline{1}3]\) zone axis. The presence of areas with unresolved lattice fringes is highlighted by white arrows (Fig. 5d). These areas are due to lattice strain or lens aberrations. Lattice strains are attributed to the presence of dislocations due to grain subdivision. Extensively increased dislocation density, grain rotation, and subgrain formation are observed in a tensile tested FeCo alloy sample (Ref 5).

4. Discussions

The inability of the Taylor model to accurately predict the orientations of one single grain upon deformation is due to its assumption. The main assumption is that each grain experiences the same amount of external macroscopic strain. This may not be true on a microscopic level of a single grain. Furthermore, interaction effects between grains are disregarded.
This leads to inaccurate predictions of the orientations in the deformed state. However, the Taylor model predicts the texture evolution of all grains in the global microstructure accurately. Based on the coefficient of the Taylor model, the activity of three slip systems of the FeCo alloy is presented in Table 1. It is observed that all three slip systems are activated in varying amounts. \{123\} \{111\} is the most active and \{112\} \{111\} the least active slip system. In this regard, Fig. 6 shows the sample surface of the tensile deformed sample. The presence of slip bands and wavy slip lines is seen. This is an indication of cross slip due to the activation of multiple slip systems. Moreover, upon tensile deformation orientation change within grains is highlighted by black arrows in Fig. 3(b). This is due to grain fragmentation due to the action of different slip system on different parts of the grain. Qualitatively, stage 2 of the strain hardening curve also indicates the activation of multiple slip systems. The high dislocation density may be also due to

**Table 1** Activity of slip systems of FeCo alloy based on the deformation in tensile tests

| Slip system     | Relative activity |
|-----------------|-------------------|
| \{110\} \{111\} | 7.6               |
| \{112\} \{111\} | 5.6               |
| \{123\} \{111\} | 14.2              |
multiple slip system activation. Previous studies reported only the activation of the \{110\} \langle 111 \rangle slip system (Ref 1). It is stated that the \{110\} \langle 111 \rangle slip system provides five independent slip systems and thereby satisfies the von Mises criteria for polycrystalline plasticity. However, in the current analysis, the five independent slip systems are not activated only by the \{110\} \langle 111 \rangle slip system. Therefore, the activation of more slip systems is required during the deformation.

5. Conclusions

Based on the study, the following conclusions can be drawn:

1. Grain fragmentation is observed after tensile testing due to the activation of multiple slip systems. Grains with [001]||TD and [011]||TD orientations are observed to be resistant to deformation accommodation. No significant change in grain boundary shape (fractal number) is observed before and after the tensile test.
2. Schmid’s law is not applicable to explain the deformation behavior of all grain orientations. However, the grain boundary slip transmission is suitable to explain the global deformation behavior of all grain orientations.
3. The Taylor model does not accurately predict the orientation for one single grain upon deformation. However, the model predicts the orientation of all grains globally upon deformation with high accuracy. The \{123\} \langle 111 \rangle slip system is the most activated one, whereas the \{112\} \langle 111 \rangle slip system is the least activated during tensile testing.
4. Based on the EBSD micrographs, a grain subdivision after tensile testing is suggested. Here, TEM micrographs support the occurrence of dislocation cells. The dislocation density in the deformed sample is \(8.5 \times 10^{13} \text{ m}^{-2}\). Correlative HRTEM images show the unresolved lattice fringes near the dislocation cell boundary and resolved lattice fringes in the dislocation cell interior.

Acknowledgments

The authors would like to thank Anja Puda and Thomas Janzen for their support in sample preparation and execution of the hardness tests and EBSD analysis. Dr.-Ing Anatoli Andreiev and Dennis Mileage are acknowledged for performing \(\mu\)-CT measurements and tensile tests.

Funding

Open Access funding enabled and organized by Projekt DEAL.

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Data Availability

The raw/processed data required to reproduce these findings cannot be shared at this time as the data also form part of an ongoing study.

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