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The evolution with strain of the stored energy in different texture components of cold-rolled IF steel revealed by high resolution X-ray diffraction

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ABSTRACT

During the deformation of low carbon steel by cold-rolling, dislocations are created and stored in grains depending on local crystallographic orientation, deformation, and deformation gradient. Orientation dependent dislocation densities have been estimated from the broadening of X-ray diffraction lines measured on a synchrotron beamline. Different cold-rolling levels (from 30% to 95% thickness reduction) have been considered. It is shown that the present measurements are consistent with the hypothesis of the sole consideration of screw dislocations for the analysis of the data. The presented evolutions show that the dislocation density first increases within the α fiber (={hkl}<110>) and then within the γ fiber (={111}<uvv>). A comparison with EBSD measurements is done and confirms that the storage of dislocations during the deformation process is orientation dependent and that this dependence is correlated to the cold-rolling level. If we assume that this dislocation density acts as a driving force during recrystallization, these observations can explain the fact that the recrystallization mechanisms are generally different after moderate or large strains.

1. Introduction

Polycrystalline metals, when subjected to a heat treatment after cold deformation, usually undergo a drastic microstructural transformation by the so-called static recrystallization process, during which new and dislocation-free grains are formed within the deformed microstructure [1]. This induces in the material a strong modification of the crystallographic texture and in turn of the physical and mechanical properties. This texture evolution is accompanied by a release of the energy stored in the material during plastic deformation, which is known to act as a driving force for nucleation and/or growth of new grains [1]. It is thus important, in order to understand, model and control the texture evolution during annealing to be able to quantitatively characterize this stored energy, and especially to be able to assess its evolution with strain and its dependence with crystallographic orientation [2].

Indeed, it is quite common to relate this stored energy to the production and organization of dislocations during deformation. If some basic linear relationships between stored energy and dislocation density can thus be found in the literature [3–5] (see below), the way these dislocations are locally organized is also well documented and shown to strongly depend on the orientation of the grain. A number of papers relate e.g. experimental data obtained through Transmission Electron Microscopy (TEM) [5–7] or Electron Back Scattering Diffraction within a Scanning Electron Microscope (SEM/EBSD) [8-10], which allow their authors to distinguish various types of dislocation arrangements as a function of strain, strain mode, material and grain orientation. In all investigated polycrystalline metals deformed up to large strains at room temperature, it is generally recognized that a high proportion of stored dislocations are organized into "patterns" within individual grains, composed of elements such as dislocation cells or walls. Simultaneously, some more isolated dislocations are also observed, which are either randomly distributed inside the cells (between the dislocation walls), or more localized close to the grain boundaries to accommodate the strain incompatibilities between neighboring grains. And classically, the total dislocation density into one grain ρ_{tot} can thus be decomposed as [4,11]

$$\rho_{tot} = \rho_{\rm w} + \rho_{\rm c} + \rho_{\rm GB} \tag{1a}$$

where ρ_w is the density of the dislocations constituting the cell walls, ρ_c the density of the dislocations randomly distributed within the cells and ρ_{GB} the density of the dislocations present at grain boundaries. As the dislocations at grain boundaries as well as the one composing the cell walls involve misorientations, both can be grouped under the classical expression of geometrically necessary dislocations

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(GNDs) [12,13], whereas the randomly distributed dislocations are called statistically stored dislocations (SSD). Eq. (1a) can then be rewritten more simply as

$$\rho_{tot} = \rho_{GND} + \rho_{SSD}. \tag{1b}$$

In order to measure these various dislocation densities. TEM. SEM/EBSD and X-ray Diffraction (XRD) are classically used. However, these techniques do not necessarily allow to distinguish all various dislocation types and thus to measure all terms in Eq. (1b). With TEM or SEM/EBSD, ρ_w or ρ_{GB} is easily deduced from local misorientation measurements. These measurements are more rarely completed by the assessment of ρ_{c} , which involves tedious and long counting procedures [4,6]. With XRD, which is a more statistical method, the analysis of the shape of a Bragg peak (and especially the broadening of the peak) measured inside a grain or a texture component is used to assess one "average" dislocation density, usually estimated to be the sole density of the isolated dislocations ρ_c [11,14–16]. The main argument advanced to eliminate any influence of the dislocation boundaries observed in highly deformed materials on peak broadening is of different types. It is most often claimed that essentially statistical dislocations (SSD) can affect peak broadening: this is for example an implicit assumption of Wilkens model [17]. More rarely, it is argued that the density ρ_c is by far the greatest (by a factor of 10) and that the other two can thus be neglected [11]. This last argument was advanced for measurements performed in steel. However, most of TEM observations performed on Al [6], Cu [4] or steels [5] do find the opposite result, *i.e.* show that ρ_w/ρ_c varies between 3 and 10 (instead of 1/10 in [11]). Additionally, some recent work aiming at comparing both SEM and X-ray methods conclude to a final ratio ρ_w/ρ_c of 1 [18].

Once the various dislocation densities have been evaluated, several equations are used to derive the associated stored energy; for example, if only GNDs are considered, the associated stored energy is usually expressed as [4]

$$E_{\rm s} = \frac{\mu b^2}{2} \rho_{\rm GND} \tag{2}$$

where ρ_{GND} is calculated from the local misorientation measured by TEM or EBSD through a linear relationship. In Eq. (2), μ is the shear modulus and *b* the norm of the Burgers vector of the dislocations. As for measurements performed by X-ray diffraction, the expression used to evaluate the stored energy is in this case [19]

$$E_{\rm s} = \frac{\alpha \mu b^2}{2} \rho_{\rm XRD} \ln \frac{R_e}{b} \tag{3}$$

where ρ_{XRD} and R_e are derived from the peaks measured by XRD (see below). This expression has been shown by Wilkens [17,19] to be valid, not only for randomly distributed dislocations, but also for what Wilkens calls restrictedly random distributions of dislocations, which can be seen in a way as random distributions by parts. In Eq. (3), there is also an extra parameter α which depends on the character (edge, screw or mixed) of the considered dislocations (see Section 3).

For steels, the first XRD measurements performed after rolling by Every and Hatherley [14] established the following hierarchy for the stored energy (E_s) as a function of fiber orientations {hkl}³

$$E_{s}\{200\} \le E_{s}\{211\} \le E_{s}\{222\} \le E_{s}\{110\}.$$
(4)

These authors explicitly mention that, to their opinion, only the randomly distributed dislocations are taken into consideration in their measurements. This hierarchy was since then never contradicted by more recent measurements, but the published quantitative values are relatively scattered: Borbely et al. [11] estimate the ratio E_s {222}/ E_s (200} to 2.8 after 48% rolling strain on UHP (ultra high purity) iron and to 3.6 after 88% rolling whereas a factor of 1.8 is reported by Castelnau et al. [20] for an IF (Interstitial Free) steel after 90% rolling. Using neutron diffraction and a quite different method for deriving the stored energy directly from the diffraction peaks, Rajmohan et al. [21] also found a factor close to 2 for this ratio after 80% rolling. It is worth mentioning though that, apart for differences in the used X-ray or neutrons devices or in the investigated steels, the methods used for the instrumental correction and analysis of the data may differ strongly from one publication to another.

Thus, the aims of the present paper are: (i) to present some recent data concerning the quantitative assessment of dislocation densities by XRD as a function of orientation and strain in a low carbon steel deformed in rolling, (ii) to discuss in some details the link between the obtained values and the various dislocation densities introduced above and to propose an analysis method for XRD data, (iii) to validate this method by some complementary EBSD measurements and (iv) based on all these experimental data, to propose a general scenario of microstructural evolution during rolling as a function of strain and grain orientation and to show that it is already consistent with some published data concerning nucleation and growth of new grains during annealing of low carbon steels, after various amounts of strain.

The present paper is thus organized as follows: the material, the used experimental techniques and the characterization of the crystallographic texture are first described in Section 2, the method used for the analysis of the XRD data is then detailed and justified in Section 3, the data obtained after various deformation levels, as well as some complementary EBSD measurements are then presented in Section 4; and the comparison of all these data allows us to discuss the validity of our approach before drawing some conclusions in Section 5.

2. Material and experimental procedures

2.1. Material and thermomechanical processing

The investigated material is an Interstitial Free Steel (IF steel), i.e. a low carbon steel often used in industry for automotive applications. The alloying elements present in the material are given in Table 1: in the present case, all interstitial dissolved nitrogen and carbon atoms are captured by titanium mainly. The deep-drawing properties of this steel depend on its final crystallographic texture, resulting from the overall thermomechanical processing route, composed of a first hot rolling step, followed by an intense cold rolling step and a subsequent controlled heat treatment designed to achieve complete recrystallization (not studied in the present paper, but during which the basic nucleation and growth mechanisms will be directly affected by the distribution of stored energy which is studied here). In the present case, the selected IF steel has reached a final thickness of 3 mm and an average grain size of 20 µm after hot rolling. It has then been deformed by cold-rolling in several steps, in a laboratory rolling mill up to a final reduction of 93%, which corresponds to a final thickness of 0.23 mm.

2.2. Microstructure and texture examination after cold deformation

The microstructures of the material have been analyzed after various rolling strains (between 15% and 93%) by Electron Back Scattering

Table 1

Chemical composition of the investigated IF steel: concentration of the alloying elements in $10^{-3}\,\rm wt\%$

| С | Mn | S | Al | Ν | Р | Cu | Ti | Nb |
|-----|-------|-----|----|-----|------|-----|------|-----|
| 2.8 | 154.2 | 7.8 | 31 | 2.2 | 14.5 | 6.3 | 49.1 | 2.8 |

³ Here {hkl} represents the ensemble of orientations having a {hkl} plane parallel to the rolling plane. See Section 2 for more details about the definitions of orientations and orientation fibers.

Diffraction (EBSD) on a Cambridge S360 scanning electron microscope equipped with the automatic Orientation Imaging Microscopy (OIM) software from Tex-SEM laboratories Inc. for data acquisition. The HKL software from Channel 5 has also been used for part of the data analysis. For EBSD observations, the samples were mechanically polished with silicon carbide paper down to grade 4000, then electro-polished in the Struers A2 solution (composed of perchloric acid and butoxyethanol).

The crystallographic textures of these samples have been measured in laboratory by XRD on an Inel four circle diffractometer in Bragg Brentano geometry. The main purpose of these measurements is to be able to select the diffracting positions for the subsequent high resolution X-ray measurements (Section 2.3). From the three {110}, {200} and {211} experimentally determined pole figures, the Orientation Distribution Function (ODF) has then been calculated with the Labotex[©] Software, which is based on the ADC (Arbitrarily Defined Cells) iterative discrete method, which allows to describe the texture as a discrete set of orientations and associated volume fractions [22]. For the result analysis and for the selection of the orientations to be studied in high resolution X-Ray diffraction, we restricted ourselves to the main orientations which are usually found after rolling and annealing, and which are all located within the $\varphi_2 = 45^\circ$ section of the Euler space⁴ (see Fig. 1). Orientations A, B and C are located on the so-called α fiber = {hkl}<110>,⁵ designating the group of orientations presenting a <110> direction parallel to the rolling direction. Orientations C and F are located on the γ fiber = {111}<uvw> designating the group of orientations having a {111} plane parallel to the rolling plane. The orientations C' and F' are symmetrically equivalent orientations of C and F respectively which are statistically as equally present as C and F, because of the orthotropic symmetry of the rolling process. The orientations D and E (which are sometimes present after very large strains and may thus play a role during recrystallization) are slightly shifted from the γ fiber (from 6 and 11° respectively).

The texture evolution during cold rolling is presented in Fig. 2. In the initial state, which corresponds to hot rolling, the texture is quite scattered as a result of a complex process involving hot deformation, recrystallization and phase transformation. Then, during cold deformation, a classical rolling texture is formed, which consists in a partial α fiber ranging from $\{001\} < 110 >$ towards $\{111\} < 110 >$ and a complete γ fiber. It is seen in this figure that the exact position of the absolute maximum in this section evolves with strain and seems to be close to the E orientation, intermediate between the B and F orientations, after large strains. To be more precise, the percentages of the main orientations have been calculated from the ODFs (with a spread of $\pm 7.5^{\circ}$, classically selected for moderate to high values of strain, as in the present work). Results are reported in Table 2 and graphically presented in Fig. 3 as a function of strain. It is clear from the figure, that the B and E orientations become the strongest after 60% strain, to the expense of the A orientation. The γ fiber (C and F orientations) is only slightly reinforced during rolling (up to 75% rolling), whereas the D component remains at the lowest level, very close to that of the neighboring F orientation. However, it is important to note that, due to the selected spread, the orientations F and D may significantly overlap.

2.3. Acquisition of high resolution diffraction data

2.3.1. Experimental setup

As the analysis of the XRD data for stored energy is based on peak broadening, it is important first to eliminate any influence of the instrumental device on it. In the present study, high resolution data were

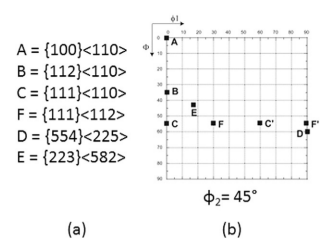


Fig. 1. (a) Selected orientations for XRD measurements and (b) their representation in the Euler section at $\varphi_2 = 45^{\circ}$.

obtained at BM02 beamline (7-circles diffractometer of the Synchrotron ESRF, Grenoble, France) at an energy E of 20 keV, thanks to (i) the very small beam divergence and bandwidth ($\Delta E/E$ about 10⁻⁴), (ii) the use of an analyzer crystal in the diffracted beam, and (iii) a solid state detector for photon counting. It is worth mentioning that such a resolution cannot be reached using laboratory goniometers unless the acquisition time becomes excessively large. With this energy, the attenuation length is ~50 μ m for iron, which is thus much larger than the grain size (which is about $15 \,\mu\text{m}$) even for the less deformed sample [10]. Moreover, the beam cross section was selected to be 3 mm² in order to scan a large number of grains simultaneously. It is a difficult task to evaluate the exact number of grains that have been scanned simultaneously, but we estimate that it must be at least several thousand for each specific orientation for all specimens; this guarantees our data to be statistically representative for the texture component as a whole. Additionally, the diffracting volume varies with the volume fraction of the scanned orientation; this feature may have an influence of the profiles, especially in terms of overlapping of various orientations. For each measured θ -2 θ line profile, 310 data points were acquired (each at different θ values), with ~10 points describing the peak itself and ~300 points describing the intensity decay in the peak tails, important for line profile analysis. This discretization was selected such that the complete measurement scan always corresponds to 30 times the full width at half maximum (FWHM) of the profile. As a result, the whole amplitude of the 2θ scans was equal to 0.9° for the less deformed specimens but increased to 2.9° for the most deformed ones due to the broadening of the Bragg peaks with strain.

A reference powder of CeO_2 (Reference NIST SRM674a) was first used to characterize the resolution of the setup at BM02 (so-called Caglioti curve), and an almost constant and low value (~0.03°) of the Integral Breadth (IB) of the peak has been recorded in the investigated 2 θ range (see Fig. 4), as typically found for synchrotron setups. The IB measured on two deformed samples is also provided for comparison, showing that, for highly deformed specimens, peak broadening is mostly due to the lattice distortion by dislocations, and that the instrumental broadening coming from the optical aberrations can be neglected in our analysis. In fact, the treatment of the peaks has been performed with and without subtracting this instrumental broadening in the worst cases, and it was concluded that this correction has only little influence on the final data.

2.3.2. Typical high resolution XRD data

The experimental set-up parameters were then selected to analyze the six specific crystallographic orientations indicated in Fig. 1 separately, *i.e.* to minimize a possible overlap between several of the investigated orientations (this is hardly possible though for the F and D

⁴ The Euler space is the commonly used orientation space in which orientation distribution functions can be represented and in which each orientation is represented by 3 Euler angles (φ_1 , Φ , φ_2) which vary between 0° and 90° because of the orthotropic symmetry. ⁵ The metallumical potntion in terms of MUL is W = 0.000

⁵ The metallurgical notation in terms of Miller indices {hkl}<uvw> is used throughout this paper to describe orientations and orientations fibers; {hkl} refers to the Miller indices of the rolling plane and <uvw> are the ones of the rolling direction.

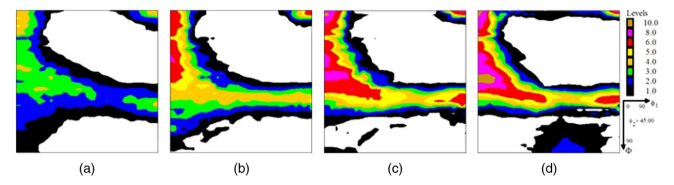


Fig. 2. Texture evolution ($\varphi_2 = 45^\circ$ section) during rolling: (a) initial state, after hot rolling, (b) after 29%, (c) 61%, (d) 76% (Labotex® software). Intensities are in m.r.d. (multiples of random density).

orientations which are separated by 6° only). For all orientations, diffracted intensity was measured along the diffracting vector (θ -2 θ scans) on {110}, {200}, {211}, {220}, {310} and {400} diffracting planes according to theoretical positions on the pole figures, which were selected as follows: first, the positions of all possible poles (namely the $\{110\}, \{200\}, \{211\}$ and $\{310\}$ poles) of the 6 selected orientations (including all orthotropically symmetrical orientations) were plotted. These are presented in Fig. 5. As it can be seen, the number of poles is quite large on the {211} and {310} pole figures, due to the multiplicity of these families of crystallographic planes. Then, for each orientation and each diffracting plane family, one pole has been selected which respects the double constraint of being as close as possible to the center of the pole figure (in order to minimize goniometer rotations and to maximize the penetrating depth of the X-ray beam) and as far as possible from the poles of the other orientations. The selected positions have been highlighted in red in Fig. 5 for the 6 orientations within one guarter of each pole figure, in which all other labels have been erased for the sake of clarity. It is seen that the selected points do avoid any overlapping of orientations for the {200} and {110} poles, but that it is not the case for the C orientation on the {310} pole figure (where it is very close to an F pole), and not the case either for the B (close to E) and C (close to F) orientations on the {211} pole figure. We can thus expect a higher degree of overlapping for the {211} measurements.

Also, since XRD only provides an average measurement along the fiber determined by the orientation of the diffraction vector, the corresponding data can additionally contain some information coming from minor texture components of the fiber (which form at least 25% of the overall texture, see Table 2). In order to verify that their contribution to the measured intensity is negligible, two ODFs have been calculated for each strain level, by considering either the sole 6 orientations and associated volume fractions (see Table 2) or all remaining orientations composing the rest and the background. Then, the corresponding pole figures have been plotted. One example of such a calculation is shown in Fig. 6 for the 29% cold rolled texture and {200} and {110} pole figures: it is clear that the orientations which compose the rest and the background, which represent ~40% of the texture in the selected example, are quite scattered and as a

Table 2

Calculated volumes fractions of the 6 selected orientations; the spread around each orientation is set to \pm 7.5° and the background to 10%.

| | Initial | 29% | 61% | 76% | 88% |
|------------|---------|-------|-------|-------|-------|
| А | 4.35 | 9.69 | 10.48 | 10.07 | 9.13 |
| В | 5.45 | 8.72 | 12.28 | 12.65 | 12.65 |
| С | 4.15 | 8.13 | 9.16 | 9.6 | 8.04 |
| F | 3.53 | 6.63 | 7.4 | 8.15 | 7.65 |
| D | 4.05 | 6.94 | 7.15 | 7.55 | 6.5 |
| E | 7.2 | 13.18 | 15.36 | 16.84 | 15.88 |
| Rest | 61.27 | 36.7 | 28.16 | 25.15 | 30.75 |
| Background | 10 | 10 | 10 | 10 | 10 |

consequence, their contribution to the measured peaks can be considered to be negligible.

Some examples of Bragg peaks are now shown in Fig. 7: the peaks have been normalized and centered, and are presented in two different scales, in order to emphasize the main characteristics of these peaks. The normalization is rendered necessary by the fact that the counting parameters are not always the same, since they depend on the actual spread of the scanned peak as well as on the intensity of the diffracting orientation. It is worth mentioning though that the signal to noise ratio varies between 10³ in the worst case to 10⁴ in most of the cases, which is remarkably high and typical of synchrotron measurements. When the peaks are plotted in a logarithmic scale, it becomes obvious that the tails are entirely captured (the lowest values are very close to the background value) and that these parts present a very small scatter.

It is also worth noting that, in the present case, all measured Bragg peaks are remarkably symmetrical (see Fig. 7). This could be an indication of a low level of long-range internal stresses associated with the cell-wall dislocation structure at the level of the texture component since this is not the case when measurements are performed in single crystals [20,23–27]. However, in highly deformed grains, peak asymmetry can also be due to the polarization of the dislocation microstructure, *i.e.* the fact that when dislocation boundaries are formed during plastic strain, the dislocations of a certain sign can be stopped on one side of the boundary and those of opposite sign on the other side of the boundary. This polarity may give rise to an asymmetry of the slip resistance [28]. We can thus probably anticipate that, in the present case, the investigated microstructures (measured at the level of one texture

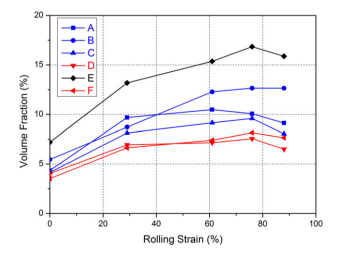


Fig. 3. Calculated volume fractions of the various texture components as a function or rolling strain.

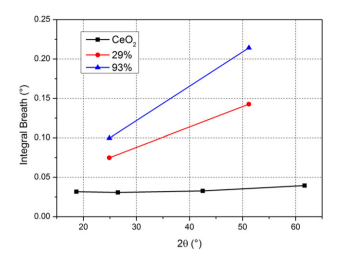


Fig. 4. Values of integral breadth for different analyzed planes for the reference powder CeO_2 ({220}, {400}, {620} and {840} planes), the less (29% strain) and the most deformed samples (93% strain) ({200} and {400} planes). The measurements are performed for specimen orientations corresponding to the center of the pole figures, which always correspond to a high texture intensity for the deformed samples.

component) are not strongly polarized. In other words, even if the microstructure is composed of dislocation cells, these must be relatively homogeneous in size and more or less randomly oriented within the investigated volume (which is composed of several grains of the same orientation). This can be due to the fact that the fragmentation of the

grains during deformation is due to the simultaneous effect of the increase of the dislocation content within the dislocation walls and of subsequent different reorientations during strain of the various scanned subgrains. In that context, we can then assume that the concept of a restrictedly random distribution of dislocations introduced by Wilkens [17] may provide an adapted model for our microstructures, and this justifies a priori the use of Eq. (3) for the stored energy, as well as the Fourier analysis presented below.

We can further argue that this conclusion has an additional consequence on the various dislocation densities present in the material (Eqs. (1a) and (1b)). Considering that our measurements correspond to several grains associated with the same orientation, the total dislocation density given by Eq. (1a) can be simplified to $\rho_{tot} = \rho_w + \rho_c$, the third term ρ_{GB} – estimated to be much smaller than the other two, see *e.g.* [11,29] – being included into the ρ_w term as a contribution to the so-called GND density (see Introduction section). Now, two different assumptions can be made before analyzing the XRD peaks:

(i) Either these XRD measurements give access to ρ_c only, but the absence of internal stresses implies then that [23,24]

$$\rho_{\rm c} = \rho_{\rm w} \tag{5}$$

and the measured density can then simply be multiplied by a factor of 2 in order to get the complete dislocation density ρ_{tot} .

(ii) Or, both cell and wall (sub-grain or grain boundaries) contribute to the measurements, again in more or less equal proportions because of the observed absence of internal stresses, and we measure directly ρ_{tot} by XRD in that case.

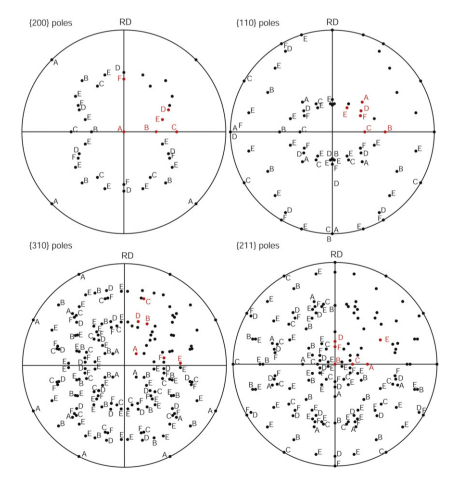


Fig. 5. {200}, {110}, {310} and {211} pole figures for the 6 individual A to F orientations including the orthotropic symmetry. On one quarter, the labels have been erased and the positions selected for high resolution X-ray diffraction have been highlighted in red.

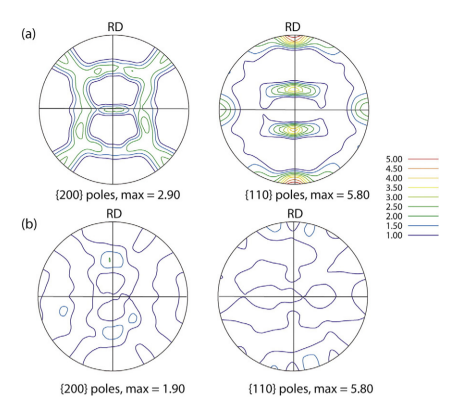


Fig. 6. (200) and (110) pole figures calculated, for the 29% strain case, from the set of orientations constituting the distribution around the 6 A to F orientations (a) and from the set of orientations constituting the remaining part of the texture and the background (b).

This must be kept in mind when comparing measured values to published ones. But in both cases, it will be possible to quantitatively compare values obtained by XRD in different orientations, as long as the relative contribution of the walls and the cells is indeed comparable. It is interesting to note that in recent papers relating measurements or calculations of dislocation densities from XRD peaks, *e.g.* [30–33], the question of the possible influence of the wall dislocations into the evaluation of the average dislocation density is never addressed. Although – to the authors' knowledge – this fact has never been either proved or inferred, it seems that, in all mentioned cases, it is explicitly assumed that indeed $\rho_{tot} = \rho_w + \rho_c$ is evaluated in the case of asymmetrical peaks. In the present work, we will simply call the measured density ρ_{XRD} and try to deduce from our analysis whether we are closer to hypothesis (i) $\rho_{XRD} = \rho_c$ or (ii) $\rho_{XRD} = \rho_{tot}$.

3. Analysis method for high resolution diffraction data

3.1. The used line broadening model

Our investigated material being very low alloyed and highly deformed, we first assume that both broadening and shape of the diffraction peaks may be affected only by the size of the coherent domains and the presence of dislocations. We thus can make use of the modified Warren Averbach (mWA) method introduced by Wilkens and further developed by Ungar and coworkers [24–26,33] to analyze the XRD data. The principle is the following: Bragg diffraction is sensitive to the position of atoms in the specimen. Dislocations distort the crystal lattice, creating a displacement field of atoms which signature is contained (in a statistical way) in the shape of any Bragg peak. The mWA method provides a (relatively simple) way to extract the dislocation density from

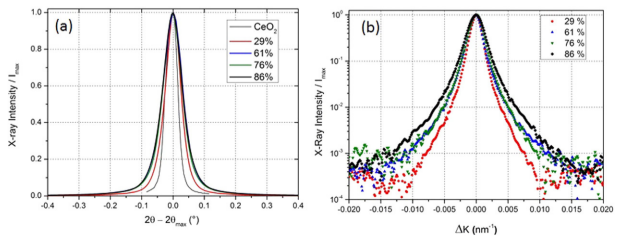


Fig. 7. Examples of {200} peaks measured within the B orientation after various rolling strains, compared to the {400} peak for the CeO₂ reference powder. The measured intensities are normalized by the maximum one. In Fig. 7b, $K = 2 \sin \theta / \lambda$.

the peak shape. Some other statistical parameters of the dislocation structure (for example polarization) can also be estimated using higher order moments of the dislocation peak, *e.g.* describing peak asymmetry. We do not go that far in this work since data treatment can become sensitive. The measured Bragg peaks $I(2\theta)$ have to be expressed in the crystal reciprocal lattice space $I(\mathbf{K})$, with \mathbf{K} being the diffraction vector. According to the mWA model, the Fourier Transform A(L) of $I(\mathbf{K})$, for small values of L, reads

$$\ln|A(L)| = \ln|A^{s}(L)| - \frac{\pi b^{2}}{2} \rho_{XRD}L^{2} \ln\left(\frac{R_{e}}{L}\right)K^{2}\overline{C} + Qb^{2}L^{4} \ln\left(\frac{R_{2}}{L}\right) \ln\left(\frac{R_{3}}{L}\right)\left(K^{4}\overline{C}^{2}\right).$$
(6)

Here, *L* corresponds to a physical length in the crystal in the direction of the diffraction vector, *K* is the norm of the diffraction vector **K** (*K* = 2 sin θ/λ , with λ the wavelength), ρ_{XRD} is the dislocation density, and R_e an outer cut-off radius for the dislocations ($R_e \approx 1/\sqrt{\rho_{XRD}}$). In the last term of Eq. (6), *Q*, R_2 , and R_3 are additional material parameters. This Fourier transform is indeed linked to the various moments of the atomic displacement field within the investigated volume. This formula has been obtained by doing some assumption on the dislocation structure, *e.g.* all dislocation segments are considered to be straight.

The first term A^s expresses the broadening due to the small size of the diffracting subgrains, and which becomes important essentially for sizes smaller than ~ 100 nm. The dimensionless coefficient \overline{C} is the average contrast factor of the dislocations present within the material, which accounts for the anisotropy of the deformation field created by the dislocations [34]. This factor can be calculated, as long as some hypotheses are made about the number of slip systems populated by dislocations and the possible dislocation types present in the material (see below).

3.2. The influence of the number of considered slip systems

In the literature, the following standard assumptions can be found:

- (i) It is often considered that all possible slip systems are equally populated and that it is not possible to really distinguish between edge and screw type; in that case, half of the dislocations are considered of edge type and the other half are screws. This approximation is often shown to be consistent with the peak broadening analysis [30,35,36];
- (ii) Crystal plasticity is sometimes used to reduce the number of populated systems; this approach is most often shown to improve [11,16,37] or more rarely to deteriorate the quality of the fitting procedure (exposed below) [38]; it is sometimes completed by somewhat arbitrary assumptions on the dislocation character [11];
- (iii) All possible slip systems are taken into account and assumed to be equally populated, but the character (screw, edge or mixed) is not a priori set, but deduced from a minimization process [31].

Ideally, it seems tempting to use a polycrystalline model to get a more precise idea on the actual number of active systems and to deduce the dislocation character from the peak analysis. However, since most of the models, used up to now, do not take explicitly into account the building up and the recovery of dislocation microstructures, there is no reason why the number of predicted active systems should reflect the number of experimentally observed populated systems. Another alternative consists in using a DDD (discrete dislocation dynamics) code to evaluate the number of populated systems and type of dislocations. This has been done recently to assess the validity of the Wilkens model [32]. However, if this procedure is convenient for small strains in single crystals, it cannot be applied anymore to the case of strongly deformed polycrystals, due to the tremendous effects of stress and strain localization associated with the mechanical interaction between neighbor grains. To our opinion, taking thus all systems equally populated (as in point (i) above) seems to be closer to the reality in this case, since it integrates the total deformation history and associated changes of active systems.

3.3. The influence of the dislocation character

As for the dislocation character, it is not at all easy to identify it experimentally (see e.g. [39]). In that case, the adjustment procedure seems tempting. But this does not seem appropriate in the case of bcc metals, in which the screw dislocations are known to glide much slower than the edge ones [39-41]. As a consequence, mainly screw types remain in the material after deformation. We are thus going to make use of this observation and assume for simplification that all dislocations present in our material are of screw character. During the Fourier analysis, this assumption will also be compared to the classical half screw/half edge assumption. The corresponding average (on all systems) contrast factors are listed in Table 3 for the two different hypotheses (50% screw + 50% edge or 100% screw). Note that if the values of the $K^2\overline{C}$ coefficients are not so different between the two cases, the points corresponding to the various diffracting planes will not be placed in the same order in the plot of the Fourier coefficients as a function of $K^2\overline{C}$ (Eq. (6)) and this will of course strongly affect the quality of the fit. An example of such a fit by Eq. (6) of the Fourier coefficients of the peaks measured in the F orientation after 76% strain is shown in Fig. 8. assuming that all dislocations are screw. The plotted curves are obtained by a least square fit on the coefficients obtained on various peaks and for various L values, through a homemade software. This procedure allows the determination of all material parameters composing Eq. (6). The very good quality of the fit justifies a posteriori the assumption made on dislocation character and renders useless any additional minimization procedure, which would most probably lead to a very similar result, *i.e.* a stronger screw character (see Section 3.2).

Also, in this example as in all other treated cases, imposing 50% screw and 50% edge dislocations leads to a much poorer fit of the experimental data by Eq. (6) (not shown here). It is also worth mentioning though that two peaks, namely the {211} and {400}, have been systematically taken out from the present analysis. Indeed, for the {211} measurements, it has been seen previously that it was difficult in the present case to get data corresponding solely to one main texture component. As for the {400} measurements, due to the much larger value of $K^2\overline{C}$ (see Table 3), the quality of the adjustment is slightly reduced when those peaks are taken into account.

4. Presentation and analysis of the experimental results

4.1. Assessment of the XRD dislocation density

The results of the X-ray analysis described in Section 3 by considering only screw dislocations are now presented in Table 4 and Fig. 9. The evolution of the dislocation density is quite different for the α and γ fiber orientations. In the first deformation steps, the dislocation density increases mainly in the α orientations (at least up to 20% for the B

Table 3

Calculated averaged contrast factors for all investigated planes with the 2 selected hypotheses.

| | | {110} | {200} | {211} | {220} | {310} | {400} |
|--------------|-------------------|--------|---------|---------|---------|---------|---------|
| 1/2 Edge 1/2 | \overline{C} | 0.1414 | 0.2736 | 0.1416 | 0.1414 | 0.2260 | 0.2736 |
| Screw | $K^2\overline{C}$ | 3.4421 | 13.3199 | 10.3405 | 13.7685 | 27.5086 | 53.2798 |
| 100% Screw | \overline{C} | 0.104 | 0.3055 | 0.104 | 0.104 | 0.2329 | 0.3055 |
| | $K^2\overline{C}$ | 2.5317 | 14.8730 | 7.5966 | 10.1268 | 28.3485 | 59.4919 |

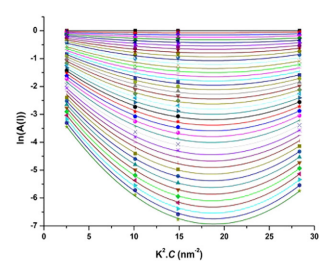


Fig. 8. Application of the mWA method on a specimen cold-rolled to 76% for the F orientation ({111}<112>). Here, Eq. (6) nicely matches the Fourier transform of {110}, {220}, {200} and {310} planes, up to L = 66.3 nm (in this figure, L decreases from the top (L = 0 nm) to the bottom (L = 66.3 nm) with a step of Δ L = 1.7 nm).

component but possibly up to a larger strain like *e.g.* around 40%, and up to 60% for the A and C orientations), while it is more or less constant in the other orientations. After 60% strain, the dislocation density increases strongly in the F orientation while it decreases strongly in all α orientations. Additionally, it is seen that the dislocation density is more or less constant in the D and E orientations which are considered as orientations intermediate between α and γ fibers. At this stage, we can also say that the present data are only partly consistent with previous ones, as mentioned in Section 1: in the present work, the observed hierarchy between various texture components is clearly dependent on the strain level, and possibly strongly affected by the initial texture.

As these orientations become more or less stabilized with strain, it is tempting to conclude at this stage that the observed different evolutions are the consequence of different hardening and softening mechanisms which can take place within the different orientations during deformation; these mechanisms are not only dependent on the number and nature of active slip systems, but also on the degree of reorientation which will affect in turn the fragmentation of the grains [42], and thus the various dislocation densities listed in Section 2.

4.2. Assessment of the reorientation capacity of the various orientations

In order to visualize the possible degree of reorientation of each fragmented grain, a simple polycrystalline model can be used to calculate the reorientation field for each point of the Euler space. This has been done here with a viscoplastic Taylor model, which assumes that all grains of a polycrystalline sample are subjected to the same imposed strain rate tensor, and which takes into account 12 {110>111> and 12 {112>111> slip systems with the same reference shear stress [43]. Then, the Euler space (here, only the $\varphi_2 = 45^\circ$ section) is discretized into regularly spaced orientations, and one single plane strain compression step is imposed to each of them. The assessment of the slip rate on all active slip systems enables then to calculate the three independent

Table 4

Measured dislocation densities (m⁻²) using Eq. (6), considering only screw dislocations.

| Orientation | 29% | 61% | 76% | 88% |
|-------------|----------|----------|----------|----------|
| А | 2.59E+15 | 7.75E+15 | 4.10E+15 | 3.67E+15 |
| В | 6.38E+15 | 3.81E+15 | 1.35E+15 | 1.56E+15 |
| С | 3.18E+15 | 6.99E+15 | 4.28E+15 | 1.91E+15 |
| F | 3.73E+15 | 2.44E+15 | 7.25E+15 | 6.70E+15 |
| D | 3.33E+15 | 2.94E+15 | 2.88E+15 | 2.49E+15 |
| E | 2.48E+15 | 2.52E+15 | 2.24E+15 | 2.16E+15 |

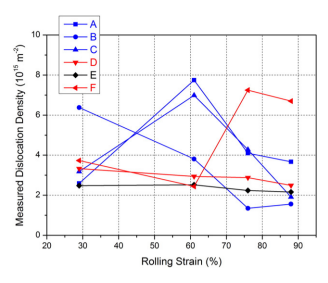


Fig. 9. Dislocation densities extracted from the peak analysis.

components of the lattice spin tensor Ω_{23} , Ω_{13} , and Ω_{12} . The components of orientation change can then be deduced from the following evolution laws for the 3 Euler angles [44]:

$$\dot{\varphi}_{1} = -\Omega_{12} - \dot{\varphi}_{2} \cos \Phi$$

$$\dot{\Phi}_{=} -\Omega_{23} \cos \varphi_{1} - \Omega_{31} \sin \varphi_{1}$$

$$\dot{\varphi}_{2} = -\Omega_{23} \frac{\sin \varphi_{1}}{\sin \phi} + \Omega_{31} \frac{\cos \varphi_{1}}{\sin \phi}.$$
(7)

From these equations, the three increments $\Delta \varphi_1$, $\Delta \Phi$, $\Delta \varphi_2$ can then be easily deduced. In Fig. 10, each considered orientation is represented by a dot whose size is proportional to $\Delta \varphi_2$ (related to reorientation outside the $\varphi_2 = 45^\circ$ section), and associated with a small line whose components are proportional to $\Delta \varphi_1$ and $\Delta \Phi$, and which represents reorientation inside the $\varphi_2 = 45^\circ$ section. It is interesting to note that this figure confirms the fact that the orientations A and B are stable, whereas a general reorientation of orientations from F to B through

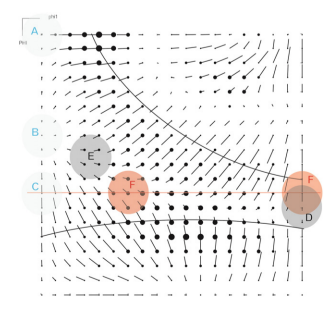


Fig. 10. Reorientation map calculated with the Taylor model. The large circles represent the spread around each studied orientation used for percentage calculations (see Table 2). The two black lines delimitate the zone occupied by the initial texture of the material (see Fig. 2).

E is predicted by such a model (with a main rotation around TD). The C orientation tends also to reorient towards B through a rotation around RD. The area occupied on this section by the initial texture (see Fig. 2) is also identified in this figure (by the two extreme lines), which enables to understand why the E orientation remains highly populated up to large strains while the B orientation is reinforced. It is also worth noting that as soon as a dislocation substructure is built up inside a given grain, if the associated orientation is unstable, the various subgrains are generally subjected to slightly different reorientation rates, and this can further enhance the grain fragmentation and modify the dislocation density within the subgrain boundaries [42].

4.3. Assessment of the fragmentation from the EBSD measurements

This fragmentation can easily be visualized by EBSD. Some microstructures measured on the RD–ND plane are shown in Fig. 11. It is clear that some fragmentation takes place in some grains at the very early stage of deformation, but that it is difficult to distinguish different behaviors in α and γ grains, especially at large strains.

In order to go further, some microstructural parameters have been extracted from these maps. The first one is the average intercept along the RD or the ND. This intercept is defined here as the distance along a line parallel to RD or ND between two boundaries associated with misorientations larger than 10° or 1°. This parameter allows estimating the grain size even when the material is highly deformed. It can then be calculated for a given map along several lines parallel to RD or ND in such a way to have at least one measurement per grain, and then simply averaged. These different intercepts are represented in Fig. 12 for the two selected limiting misorientation values. As the misorientation limit of 10° is quite close to the generally accepted limit between grain boundaries and subgrain boundaries, it is obvious that the observed evolution is indeed associated with grain fragmentation (since, without fragmentation, the grain size and thus the intercept value should constantly increase along RD, as a result of plane strain compression; this is indeed the case below 15% strain). It is also clear that there is a transition between two different regimes between 30% and 50% strain: fragmentation gradually increases within the grains up to 40% strain where it reaches the limiting value of 10° and then gradually saturates, especially after 60% strain; this could be due to the combined effect of fragmentation and elongation of the grains together with the recovery of the subgrain and grain boundaries (which will directly affect the density of GNDs). As for the intercept calculated with the misorientation value of 1°, which directly characterize the subgrain boundaries or dislocation cells, a similar observation of a general saturation after 60% can also be drawn.

To further quantify this subgrain boundary dislocation density, one second parameter can be extracted from the EBSD map, *i.e.* the local intragranular misorientation inside grains belonging to specific texture components (fiber or individual orientation). It is calculated here as the so-called kernel average misorientation (KAM), *i.e.* the average misorientation between a point on the measurement grid and its nearest

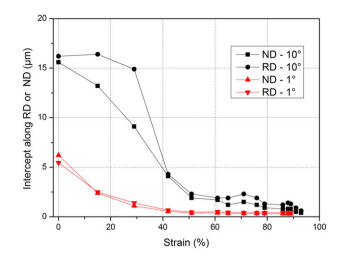


Fig. 12. Evolution of the intercept along RD or ND with strain, for two different misorientation values (10° or 1°).

neighbors. To ensure that all considered points belong to the same grain, misorientations larger than 5° have been excluded from this average. For one given texture component, a frequency histogram can then be plotted, and the peak value, *i.e.* the misorientation value associated with the maximum of the histogram can then be extracted. This parameter, characterizing the intragranular misorientation, is plotted in Fig. 13 as a function of strain for α and γ grains. Again, a transition between two regimes is visible around 60%: intragranular misorientations increase first slowly similarly in all grains, whereas it increases more significantly in the γ grains than in the α grains after 60%.

4.4. What happens during rolling?

It is worth noting first that when we measure a diffraction peak within a given orientation during strain, we are in a way in an Eulerian scheme: we focus on one orientation but not on specific grains. In other words, we examine here three different types of grains: (i) some grains associated with stable orientations from the beginning of the deformation, (ii) some grains which are unstable at the beginning of the strain path and which will always remain unstable because the strain level is not high enough or (iii) some unstable grains which will finally reach some stable orientations. Thus, the observed behavior of one orientation is indeed the collective behavior of various grains, whose initial orientation was not necessarily the same [44]. Also, if some papers relate the hardening behavior of single crystals (e.g. [45]), the hardening/softening behavior of specific orientations within a polycrystalline sample is rarely documented in the literature, and mostly associated with moderate strains only [46]. Furthermore, the previous modeling efforts clearly demonstrate that the results are indeed strongly dependent on the selected polycrystalline model [47,48].

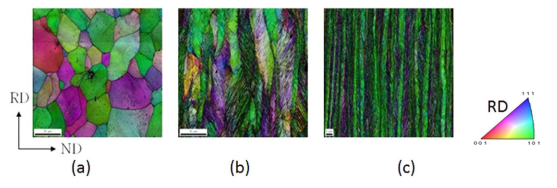


Fig. 11. EBSD maps measured on the transverse section of the cold-rolled material after (a) 15%, (b) 61% and (c) 88% rolling strains. Maps of 100 µm × 100 µm, step size of 0.2 µm.

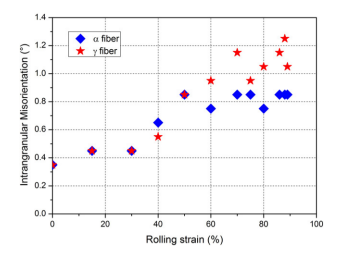


Fig. 13. Evolution of the intragranular misorientation measured by EBSD (calculated as the misorientation value at the peak of the Kernel histogram for EBSD maps of 100 μ m by 100 μ m measured with a step size of 0.2 μ m) in α and γ fiber grains as a function of strain.

Having said that, can we still try to explain the observed evolutions in terms of GND and SSD densities? It seems difficult at this stage to separate the two, although some experimental facts are worth recalling. Indeed, it was already qualitatively observed that, during complex strain paths, these two dislocation populations did have different evolutions within different orientations [44]: both ρ_c and ρ_w were observed to (i) first increase and then decrease with strain within stable orientations (first ρ_c and then ρ_w), (ii) remain quite low within unstable orientations and (iii) to increase again in stabilized and fragmented orientations.

Based on these previous observations and on the simultaneous inspection of both X-ray and EBSD data as well as Taylor calculations, we can then propose the following scenario for the microstructural evolution taking place in various grains during rolling:

From 0 to 40% strain. All grains harden and slightly fragment (see Fig. 13), but the dislocation density ρ_{XRD} increases mainly in the α fiber, and especially in the stable A and B orientations. Those, as well as the E orientation are strongly reinforced because of the texture evolution (see Fig. 8). But as the A and B orientations are stable, the fragmentation of these grains is only due to the building up of a dislocation substructure (both ρ_c and ρ_w increase significantly, as in [44]) whereas fragmentation is enhanced by (or even mostly due to) reorientation in the E orientation and thus presumably associated with a lower ρ_w dislocation density since reorientation favors the absorption of dislocations within cell walls (it is also the case in the C, F and D orientations, which are all unstable at this stage). We can thus conclude at this point, that both cell and wall dislocations present in orientations A and B are indeed taken into account in our X-ray analysis (otherwise, ρ_{XRD} would not increase so significantly in A and B, compared to the other orientations).

From 40 to 60% strain. The same scenario continues, but the level of dislocation density has now reached a saturation value in B; as a result, recovery of the microstructure takes place within this orientation provoking a strong decrease of both ρ_c and ρ_w . The dislocation density increases mostly in A but also in the C orientation, whose fragmentation due to reorientation is now reduced (the overall texture intensity increases slightly).

From 60 to 80% strain. The saturation dislocation density value is reached in A and C, and thus recovery of the microstructure takes also place in these orientations, leading to a strong decrease of the overall dislocation density ρ_{XRD} together with a stagnation of the

intragranular misorientation. Because of texture evolution, the spread of orientation becomes now reduced around F (see Fig. 2), and we observe an enhancement of intragranular fragmentation in this orientation (see also Fig. 13), associated with a strong increase of the dislocation density ρ_{XRD} . These observations are similar to those performed in the B orientation at low strains, and we can thus conclude that both ρ_c and ρ_w increase significantly at this stage in the F orientation, leading in turn to a strong increase of ρ_{XRD} . After 80% strain. Only the stable B orientation is still reinforced, and the overall reorientation towards this final orientation is slower and slower. Since the saturation value has also been reached in the F orientation, recovery becomes active in this orientation leading to a slight decrease of the dislocation density.

Also, we can notice that the ratio of ρ_{XRD} {222}/ ρ_{XRD} {200} (or more precisely ρ_{XRD} (F)/ ρ_{XRD} (A) in the present investigation) evolves here from about 0.5 after 50% strain to about 2.0 after 90% strain (unlike in the previously reported analyses, where this ratio was always claimed to be larger than 1, see Section 1). The initial hot rolling texture, mainly composed of components close to the α fiber, and its subsequent evolution during cold rolling could be largely responsible for this result as proposed above.

If the observed evolution of the dislocation density can be quite surprising, especially the fact that we observe strong decrease of ρ_{XRD} within some orientations for various levels of strain, it is nevertheless consistent with previous studies: for example, based on the scenario initially proposed by Ridha and Hutchinson [49] reporting an enhanced recovery within Cube oriented grains deformed in plane strain compression for copper, a hardening/softening law was recently proposed [48] which allows to account for this mechanism, and which is completely consistent with prior X-ray ρ_{XRD} measurements [50]. More recently, a strong decrease of the dislocation density has also been observed in various fcc metals subjected to severe plastic strain at room temperature [51,52].

It is finally worth noting that if the value of the outer cut-off radius R_{e} , extracted from the peak analysis also evolves with strain, the evolution of the stored energy due to the dislocations (Eq. (3)) is exactly the same as the one of the dislocation density shown in Fig. 9. If we recall that, during annealing, nucleation of recrystallization always starts in the areas associated with maximum stored energy (or equivalently dislocation density) (see *e.g.* [53]), we can thus conclude that the present data are completely consistent with the general observation of the initiation of recrystallization of steels within the α fiber after moderate strains and within the γ fiber after larger strains (see *e.g.* [54]), even though the actual nucleation mechanisms may be radically different in both cases (*e.g.* strain induced boundary migration at low strains *versus* intergranular nucleation at higher strains).

5. Conclusions

In this work, dislocation densities have been estimated from X-ray diffraction measurements performed in several texture components of a rolling IF steel sheet, after several strain levels, using synchrotron radiation. The proposed analysis is based on the Fourier analysis of the peaks, by considering that all possible slip systems are equally populated but that all remaining dislocations are of screw character.

The main findings of our work are the following:

- The measured peaks have been shown to be highly symmetrical for all investigated orientations; to our opinion, this implies that the scanned dislocation microstructures are not polarized, which could suggest that the measurements are statistically representative of orthotropic texture components;
- The assumption of only screw dislocations has been shown to provide a very good fit for all investigated texture components;

- The evolution of the dislocation density varies differently with strain from one texture component to another; it increases in the α fiber during the first deformation steps, before increasing within the γ fiber; this can be partly explained by simultaneously looking at the texture evolution;
- With the aid of additional EBSD observations and simple calculations, it is proposed that the measured dislocation densities do include the contribution of both cells and walls (SSD and GND).

In order to go further, these measurements should be repeated in a material presenting a stronger texture, enabling to differentiate between various components without any ambiguity.

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