The effect of stress triaxiality on the phase transformation in transformation induced plasticity steels: Experimental investigation and modelling the transformation kinetics

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A R T I C L E   I N F O

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A B S T R A C T

In situ multiaxial loading during neutron diffraction tests were undertaken on a low-alloyed Quenched and Partitioning (Q&P) Transformation Induced Plasticity (TRIP) Bainitic Ferrite (TBF) steel with dispersed austenite particles. The effect of stress triaxiality on the evolution of the deformation-induced martensite is investigated under uniaxial- and equibiaxial-tension as well as tension/compression with a ratio of –1:6. It is shown that transformation is not a monotonic function of stress triaxiality; the amount of deformation-induced martensite is similar under uniaxial and equibiaxial tension but it is significantly smaller under tension/compression. The transformation kinetics are modeled using a recently developed kinetic model that accounts for the stress state and the stability and size of the austenite particles. The larger austenite particles transform first and the mean volume of the austenite particles decreases with increasing strain; the decreasing austenite particle size impedes the phase transformation as the deformation proceeds. It is concluded that stress triaxiality alone cannot account for the differences in the transformation kinetics between different loading states and that the number of potential nucleation sites depends on the stress state.

1. Introduction

The transformation induced plasticity (TRIP)-assisted steels are a grade of low-alloyed steels that have been widely used in the automotive industry. They feature multiphase microstructures consisting of ferrite, bainite, martensite with body-centered cubic (bcc) crystal structure, and dispersed particles of metastable austenite with face-centered cubic (fcc) crystal structure. When subjected to mechanical loading, the retained austenite transforms into martensite. The shape and volume changes accompanying this transformation cause local plastic deformation in the surrounding ferrite grains, which increases the steel strength while retaining its high ductility [1,2]. The stability of austenite does not only depend on the stacking fault energy (SFE) via chemical composition and temperature [3–6], but also on the grain size [7–10], and the stress state [11–14]. The conventional TRIP steels with polygonal ferrite are well known for their good combination of high tensile strength and high elongation [15]. However, these steels exhibit moderate bendability, flangeability and edge formability for applications that require high localized strain accommodation. Quenched and Partitioning (Q&P) steels [16] are quenched below the martensite start (Ms) temperature and kept at the quenching temperature or reheated above the Ms temperature in order to temper the martensitic matrix. Isothermal bainitic transformation of TRIP Bainitic Ferritic (TBF) steels is undertaken above the Ms temperature, resulting in a microstructure that consists of a bainitic matrix and dispersed particles of retained austenite [17,18]. In order to avoid the presence of polygonal ferrite in the microstructure, full austenitization and a critical cooling rate are required for TBF and Q&P steels. The replacement of the polygonal ferrite matrix, as present

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in conventional TRIP steels, by bainite and/or tempered martensite, diminishes hardness differences among the respective phases. This results in outstanding edge formability and stretch-flangeability. Moreover, the metastable retained austenite particles ensure high elongations during conventional deep drawing processes [19].

The mechanical behavior of these materials is complex and macrosopic plasticity models need to account for the hardening due to plastic deformation of the phases and to incorporate kinetic models for the deformation-induced martensitic transformation [20–22]. Several transformation kinetic models have been developed. The kinetic model of Olson and Cohen [23] was the first to take into account the physical mechanisms of martensitic nucleation induced by plastic strain in austenitic steels. Olson and Cohen [23] concluded that shear band intersections are the potential nucleation sites for the formation of martensite and the rate of shear band formation is influenced by the composition and temperature through the SFE. This model has been incorporated in continuum plasticity constitutive models that describe the kinetics of phase transformation and the mechanical behavior of austenitic stainless steels subjected to uniaxial tensile loading [23–26].

Recent experimental studies on austenitic steels subjected to multiaxial loading [27–29] have shown that the loading state significantly affects the TRIP effect. The combination of crystallographic texture and loading direction defines whether e-martensite will form in low SFE steels, which is an intermediate phase for α'-martensite [27] or whether deformation twins will form in medium SFE steels, which suppress the formation of α'-martensite [29]. Therefore, a key element for the validation of continuum plasticity models is their ability to model the kinetics of martensite formation under multiaxial loading states which occur during operation or metal forming processes. The classical Olson–Cohen model has been modified to account for the stress state dependence of the martensitic transformation [30–32]. In most of these studies, the triaxiality factor Σ, defined as the ratio of the hydrostatic stress p to the von Mises equivalent stress σ_{eq}, is used as a fit parameter affecting the kinetics of the martensitic transformation. The hydrostatic stress p and the von Mises equivalent stress σ_{eq} are defined as

\[
p = \frac{\sigma_{11} + \sigma_{22} + \sigma_{33}}{3}, \quad \sigma_{eq} = \sqrt{\frac{1}{2} \sum_{i=1}^{3} \sum_{j=1}^{3} \sigma_{ij} \delta_{ij}},
\]

\[\sigma_{ij} = \sigma_{0i} - p \delta_{ij}\]

where \(\sigma_{0i}\) and \(\sigma_{ij}\) are the components of the stress and deviatoric stress tensors respectively and \(\delta_{ij}\) is the Kronecker delta (\(\delta_{ij} = 1\) if \(i = j\), \(\delta_{ij} = 0\) if \(i \neq j\)). However, there is experimental evidence that the stress triaxiality alone cannot account completely for the dependence of phase transformation on the stress state (e.g., in 201 and 301 L austenitic steels [13,27] and in TRIP780 steel [33]). Beese et al. [13] introduced an additional stress state parameter, the Lode angle θ, in the kinetic model of Santacreau et al. [34] to account for the complex dependence of martensite evolution on the stress state. The Lode angle θ determines the direction of the stress deviator on the so-called “II-plane” and is defined from the relationship (e.g., Hill [35], Nayak and Zienkiewicz [36])

\[\sin 3 \theta = -\frac{27}{2} \frac{J_2}{\sigma_{eq}^3} \left( -\frac{\pi}{2} \leq 3 \theta \leq \frac{\pi}{2} \right)\]

where \(J_2\) is the determinant of the stress deviator. This model was further developed to describe the deformation of TRIP-assisted steels at non-ambient temperatures by introducing a dependency of the martensitic transformation on temperature [33]. Most recently, a new variation of the classical Olson-Cohen model was coupled with crystal plasticity to model the mechanical behavior of metastable austenitic steels under different stress states [28]. This coupled kinetic model and crystal plasticity framework accounts for i) the effect of stress state on the nucleation of martensite, ii) the evolution of the deformation texture, and iii) the SFE.

Haidemenopoulos et al. [37] developed recently a kinetic model for the martensitic transformation in TRIP-assisted steels with a relatively small fraction of metastable austenite in the form of dispersed particles in a ferrite/bainite/martensite matrix. This model describes two modes of transformation, i.e. the stress-assisted and strain-induced transformation and it is based on the modification of the nucleation site potency distribution by the applied stress and plastic strain. The model can account for the effect of austenite size, chemical composition, and stress state on the stability of austenite. It accounts for the potency of martensite nucleation in pre-existing nucleation sites (defects) or in newly created nucleation sites by deformation (formation of new defects by plasticity). The kinetic model has been applied to experimental results under uniaxial tensile loading, however, it has never been validated by experimental results under multiaxial loading (i.e. using loading states with Σ values other than 0.33).

In this work the kinetic model by Haidemenopoulos et al. [37] is evaluated, for the first time, under multiaxial loading in order to understand the mechanisms that control the TRIP effect under multiaxial loading. So far multiaxial loading experiments in TRIP-assisted steels have been only performed under uniaxial loading, shear, torsion or a combination of all [38,39] or punch tests [12]. The neutron diffractometer POLDI (Pulse OverLap DIffractometer) at the at Swiss spallation neutron source SINQ is equipped with a novel multiaxial loading rig which is capable of applying proportional and non-proportional biaxial loads [40,41]. In this study we report on a series of experiments undertaken for validating the kinetic model by Haidemenopoulos et al. [37] and understanding the mechanisms that control the deformation-induced phase transformations under different loading states (i.e. varying values of Σ). In addition, synchrotron X-ray diffraction is performed utilizing a miniaturized multiaxial machine for quantitatively assessing the evolution of austenite size with deformation. The evolving austenite size is used as model parameter and it is seen that it affects the transformation kinetics significantly.

2. Materials and experiments

The material was supplied by voestalpine AG, Austria as hot-rolled sheets with a thickness of 4.35 mm. The chemical composition is given in Table 1. The mechanical response of the material upon monotonic ex situ uniaxial deformation of a dogbone sample is shown in Fig. 1.

Cruciform-shaped samples with reduced thickness at the center were designed with the aid of finite element simulations, the schematic of the cruciform is shown in Fig. 2. The special shape and the thickness reduction in the middle of the cruciform is tailored to the mechanical properties of the specific material and it is a result of a trade-off between optimizing the volume of material for sufficient neutron diffraction statistics and the possibility to reach relatively high plastic strain at the center of the cruciform prior to fracture at the cross arms.

Let \(\sigma_1\) and \(\sigma_2\) be the stresses applied macroscopically on the cruciform specimens. Three different loading combinations were tested:

1) \(\sigma_1 = 0\) (uniaxial tension = UN),

| C    | Si  | Mn  | P   | S   | Al  |
|------|-----|-----|-----|-----|-----|
| 0.203| 1.46| 2.48| 0.011| 0.0003 | 0.058 |
The corresponding macroscopic values, i.e., the values corresponding to \( \sigma_1 \) and \( \sigma_2 \), of the stress triaxiality \( \Sigma = \frac{\sigma_1}{\sigma_0} \) and Lode angle \( \theta \) are:

- ii) \( \Sigma = 1 \) (equibiaxial tension = EQ),
- iii) \( \Sigma = \frac{1}{2} \) (biaxial tension/compression = TC).

The applied macroscopic loads \( \sigma_1 \) and \( \sigma_2 \) are increased proportionally from zero. The simulations show that uniform deformation is concentrated at the center of the cruciform, as shown in Fig. 3c. It is interesting to point out that the calculated values of \( \Sigma \) and \( \theta \) at the center of the specimens are different from the “macroscopic” values given above. In particular, the actual values at the center of the specimens are calculated as \( \Sigma = 0.18, \theta = -16^\circ \) for the TC loading, whereas for the EQ loading the calculated value of \( \Sigma \) agrees with the theoretical. Since the loading combination for the TC loading is close to uniaxial tension, the difference between the calculated and the macroscopic values can be attributed to the “ring” and “Poisson” effects that have been reported when cruciform specimens are loaded in uniaxial tension [41–43]. Such effects are not important when biaxial macroscopic loads are applied, and this explains the agreement between the calculated and macroscopic values for \( \Sigma \) and \( \theta \) under EQ loading. The \( \Sigma \) and \( \theta \) values remain essentially constant after the specimen deforms plastically at the cruciform center.

The in situ deformation and neutron diffraction experiments were undertaken on the POLDI instrument at the Swiss Spallation source SINQ, Switzerland, using the biaxial machine described in Refs. [40,41]. The biaxial deformation system is equipped with a 2-camera digital image correlation (DIC) system (GOM, Aramis 5 M) for measuring the in-plane macroscopic strain at the center of the cruciform, which is sprayed with a black/white pattern for tracking the displacement. The neutron diffraction measurements were carried out in load control mode after interrupting the loading at predefined force intervals and holding the displacement. The loading rate was 40 N s\(^{-1}\) for UN, 40 N s\(^{-1}\) and -6.7 N s\(^{-1}\) in tension and compression.

![Fig. 1. Experimental true stress-strain plot for the hot-rolled Q&P/TBF steel.](image1)

![Fig. 2. Schematic of the cruciform geometry.](image2)
respective for TC. Evaluating the stress state at the center of a cruciform is not as straightforward as for dogbone specimens as the cruciform samples do not have a well-defined cross-section; while experiments can only capture the applied force along each axis and strain from DIC. Hence, FE simulations are utilized for predicting the (equivalent) strain curve under EQ and TC loading as shown in Fig. 4. It is seen that the FE prediction for the TC loading, using data from UN tests, is very good since TC with a −1:6 ratio is a loading state that is not significantly different than UN. However, the FE prediction is not as good as the experimental data for EQ due to mechanical anisotropy of the material along the two loading axes.

An example of neutron diffraction patterns before and after deformation is shown in Fig. 5a. The neutron diffraction data were reduced and fitted using the open-source software Mantid [44]. The obtained results were analyzed in terms of the evolution of the integrated intensity of austenite and the lower bainite/tempered martensite matrix. Since neutron diffraction cannot distinguish between the different phases with bcc crystal structures, the fitted martensite fraction corresponds to the volume fraction of deformation-induced martensite with respect to the initial volume fraction of austenite, which is calculated as

\[
f_m = 1 - \frac{f_{\text{bcc}}}{f_{\text{fcc}}} = \frac{I_{\text{fcc}}^{0}}{I_{\text{fcc}}^{0} + I_{\text{bcc}}^{0}},
\]

where \( I_{\text{fcc}}^{0} \) and \( I_{\text{bcc}}^{0} \) are the integrated intensities of the fcc reflections before deformation and during deformation respectively, and \( I_{\text{fcc}} \) and \( I_{\text{bcc}} \) are the integrated intensities of the bcc reflections before deformation and during deformation respectively. Hence, \( f_m \) varies from 0 (before loading) to 1 (when austenite tranforms completely).

In situ synchrotron X-ray diffraction under equibiaxial loading was performed in transmission mode at 20 keV on the MS beamline of the Swiss Light Source using the biaxial machine described in Ref. [45]. A miniaturized cruciform sample with 250 μm thickness in the arms and 80 μm thickness in the center was loaded until fracture with a displacement rate of 0.1 μm s⁻¹, while continuous diffraction measurements were undertaken. The Mythen detector of the MS station was used to record diffraction data parallel to one loading axis, a X-ray diffraction pattern is shown in Fig. 5b. The data were fitted using the open-access software WinPlotr [46]. The peak position and the full-width at half-maximum were used for assessing the evolution of the peak broadening as a function of the lattice strain. According to Williamson and Hall [47], a plot of the integral breadth \( \beta_{hkl} \) versus \( \sin(\theta_{hkl})/\lambda \), where \( \theta_{hkl} \) is the diffraction angle and \( \lambda \) is the X-ray wavelength, produces a straight line: the intercept with the ordinate and the slope of this straight line yields values for the crystallite size and the microstrain, respectively. The intercept is inversely proportional to the crystallite size \( D \). Due to lower resolution such investigation is not possible with the data obtained through neutron diffraction. The elastic lattice strain \( \varepsilon_{\text{el}} \) is determined by the relative change of the interplanar lattice spacing \( d_{hkl} \) of a specific family of planes \( \{hkl\} \) with respect to \( d_{hkl}^{0} \), which is the initial value prior to deformation:

\[
\varepsilon_{\text{el}} = \frac{d_{hkl} - d_{hkl}^{0}}{d_{hkl}^{0}}
\]

Electron backscatter diffraction (EBSD) was carried out on the as-received material. The sample was ground with 1200 grit SiC paper
and then electropolished for 10 s with a 16:3:1 (by volume) ethanol, glycerol, and perchloric acid solution using 52 V current. A field emission gun scanning electron microscope (FEG SEM) Zeiss ULTRA 55 equipped with EDAX Hikari Camera operated at 20 kV in high current mode with 120 μm aperture was used. The EBSD raw data were post-processed using the commercial EDAX OIM Analysis 7.3 software.

3. The kinetic model

The kinetic model of Haidemenopoulos et al. [37] determines the evolution of martensite volume fraction during deformation-induced martensitic transformation of dispersed austenite particles in low-alloy TRIP-aided steels. The kinetic model is based on the Olson-Cohen [23] theory of heterogeneous nucleation of martensite, which is described by a site potency distribution function. The model thus considers that the material contains a certain number of nucleation sites from which a deformation-induced martensite fraction will transform into martensite. The overall model accounts for the influence of austenite grain size and stress-induced (by plastic deformation) contributions. Additionally, the model accounts for the influence of austenite grain size and stress triaxiality on the martensitic transformation. The overall deformation-induced martensite fraction $f_m(\varepsilon)$ is given by:

$$f_m(\varepsilon) = 1 - \exp[-v_p N_s(\varepsilon)]$$

where $\varepsilon$ is the equivalent plastic strain, $v_p$ the mean volume of austenite particles, and $N_s$ the number of the operational nucleation sites for martensite (i.e. the sites with sufficient potency to nucleate martensite). The term $N_s$ consists of the operational sites under the applied stress (stress-assisted nucleation), $N_{s}^0$, and the additional sites produced by plastic deformation, $N_{s}^p(\varepsilon)$:

$$N_s(\varepsilon) = N_{s}^0 + N_{s}^p(\varepsilon)$$

where $N_{s}^0 = N_{s}^{00} \exp(-\alpha_s n' \varepsilon)$

$$N_{s}^p(\varepsilon) = N_{s}^{p0} \exp(-\alpha_p n' \varepsilon)$$

where $N_{s}^{00}$ is the number of pre-existing nucleation sites, $\alpha_s$ is a shape factor constant for the stress-modified potency distribution, and $n'$ is the critical embryo thickness given by:

$$n' = \frac{2\gamma_f}{\rho(\Delta G_h + \Delta G_e(\Theta) + E_{str} + W_f)}$$

where $2\gamma_f$ is the fault/matrix interfacial energy, $\rho$ is the density of atoms in the fault plane (in moles/unit area on a crystal plane), $\Delta G_h$ is the chemical driving force for the martensitic transformation, $E_{str}$ is the elastic strain energy associated with the distortions in the fault interface, $W_f$ is the frictional work of the interfacial motion. The mechanical contribution to the chemical driving force, $\Delta G_e(\Theta)$ depends on the von Mises equivalents stress $\sigma_e$ and on the stress triaxiality $\Sigma$:

$$\Delta G_e(\sigma_e, \Sigma) = - (0.725 \sigma_e + 0.3206 \sigma^2)$$

where $\sigma_e$ is the yield stress of the material. Based on the model of Haidemenopoulos et al. [37], as the triaxiality factor increases, the critical size for martensite nucleation $n'$ decreases and martensite formation becomes easier.

The number of operational nucleation sites upon plastic deformation $N_{s}^p(\varepsilon)$ is given by:

$$N_{s}^p(\varepsilon) = N_{s}^{p0} \exp(-\alpha_p n' \varepsilon)$$

where $N_{s}^{p0}$ is the number of pre-existing nucleation sites, $\alpha_p$ is a shape factor constant for the strain-modified potency distribution, and $n'$ is the critical embryo thickness.

4. Results

4.1. Crystallite size distribution

Fig. 6 shows the intercept of the Williamson-Hall (W-H) plot as a function of the lattice plane strain of the (111) lattice plane family for the in situ biaxial and synchrotron X-ray diffraction test. The evolution of the intercept of the W-H plot with deformation gives a good qualitative estimate of the relative evolution of the crystallite size. During elastic deformation, the full-width-at-half-maximum (FWHM) of the (111) lattice planes families is nearly constant and the W-H plot intercept remains relatively unaffected. At approximately 2250 με lattice strain of
the (111) lattice plane family, plastic deformation takes place and the peak width starts to increase and, at the same time, the W–H plot intercept starts to increase, indicating a decrease of the mean crystallite size of austenite. By the end of the test the intercept value becomes 3 times larger, indicating a decrease of the crystallite size in the order of approximately 75%. The decreasing austenite size can be i) due to earlier transformation of the larger austenite grains, thereby reducing the average size of the ones that did not transform, ii) by the partial transformation of some austenite grains, which splits the austenite grains into smaller subgrains, or iii) by plastic deformation, which introduces intragranular misorientation within the initially (relatively) large austenite grains. A similar decrease of austenite particle size and mean crystallite size during plastic deformation under UN deformation was observed in a recent work by Haidemenopoulos et al. in a low-alloy TRIP steel with magnetic force microscopy (MFM) and X-ray diffraction (XRD) [48]. The application of the W–H plot in Ref. [48] showed that the mean crystallite size reduces by 75% after deformation, meanwhile MFM showed that the austenite grain size reduces by 50%. Based on the findings in Ref. [48] and due to the agreement of the W–H results in the current work, we assume that the austenite grain, regardless the loading state, reduces by 50% by the end of each loading test. The evolution of the mean austenite size must be considered when using the model, since it affects the transformation kinetics. After the material fails, a small amount of austenite is still detectable (cf. Fig. 5a and b), such small fraction is beyond the detection limit of neutron diffraction.

The orientation map with inverse pole figure (IPF) coloring shown in Fig. 7 indicates a relatively mild crystallographic texture of the as-received material. The phase map of the initial microstructure consists of a lower bainite and tempered martensite matrix, both with bcc crystal structure, and dispersed austenite particles with a mean diameter of approximately 0.35 μm and fcc crystal structure; this is in good agreement the W–H analysis, which suggests a mean crystallite size of 0.37 μm, assuming that the grains are equiaxed and the crystallite size represents the diameter of a circle, the area of which is the same as the grain projection in 2D, in the undeformed material. Fig. 6 suggests the decreasing austenite grain size is a linear function of plastic strain, which is also confirmed in Ref. [48]. Hence, the mean austenite particle diameter, for each loading state, can be given by:

$$D_A(\varepsilon_p) = D_A^0 - \frac{D_A^0}{2\varepsilon_{max}}\varepsilon_p$$

where $D_A^0$ is the diameter of the austenite particles before deformation, i.e., $D_A^0 = 0.35$ μm. The volume of the austenite particles $v_p$, is then

Fig. 6. Plot of the W–H plot intercept and the FWHM of the (111) lattice plane family as a function of the lattice strain of the (111) lattice plane family. The black dashed lines are guides for the eye. The blue dashed line indicates the point at which both FWHM and W–H intercept start to change indicating yielding of the material. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

Fig. 7. (a) Orientation map with IPF coloring of the undeformed material in the out-of-plane direction. (b) Phase map showing the initial microstructure consisting of a bcc matrix (mixture of lower bainitic/tempered martensite) shown with red and dispersed austenite particles with fcc crystal structure shown with green. The mean austenite particle diameter is 0.35 μm. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

Fig. 8. The evolution of the mean austenite volume with plastic strain for TC, EQ and UN, used for as $v_p$ in Eq. (3).
calculated by the evolution of the volume of a sphere with a diameter of \( D_0(\sigma_f) \). The evolution of \( v_p \) is plotted in Fig. 8 for the three loading states considered. The relationship in Eq. (10) is used in the kinetic model to describe the evolution of the mean austenite particle volume in Eq. (3).

The evolution of the deformation-induced martensite fraction with equivalent plastic strain is shown in Fig. 9 (f-e curves). The phase transformation under TC is significantly delayed compared to the UN and EQ loadings, which exhibit similar behavior. The latter is in very good agreement with results from uniaxial and punch tests undertaken on TRIP-assisted steels by Jacques et al. [12] where it was shown that uniaxial and equibiaxial loading result in a similar evolution of martensite phase fraction.

The f-e curves in Fig. 9 do not show a monotonic dependence on stress triaxiality, \( \Sigma \), which is implied by the kinetic models in Ref. [30, 37]. It appears that the dependence of the phase transformation on the stress state is more complicated and may involve a strong dependence on parameters such as the different martensite nucleation mechanisms.

### 4.2. Model fitting

The fitted kinetic model is able to describe the stress state dependence of the martensitic transformation, as shown in Fig. 9. The model was fitted to the experimental data for all three tests simultaneously with \( N^0 \) and \( N \) as fit parameters. The possibility of \( \theta \)-dependence is not considered here; a detailed investigation of this subject is now underway. The rest of the model parameters were fixed. The fit parameters of the model are summarized in Table 2. The number of pre-existing nucleation sites, \( N^0 \), is a material property, which is not related to the loading state. Therefore, it was fitted and assumed to be independent of the loading state. The critical value of the embryo thickness, \( n^* \), depends on triaxiality; therefore different \( n^* \) values were used for each loading state. The parameter \( n^* \) was calculated using Eq. (6), taking the parameter \( \Delta G_m \) equal to \(-2500 \text{ J/mol}\), which is the median value used in Refs. [37]; fluctuations of \pm 1000 J/mol do not affect the fit significantly. The parameter \( \Delta G_m \) was set equal to \(-423, -510 \) and \(-451 \text{ J/mol MPa for TC, EQ, and UN respectively} \), which is approximately the yield stress of the material, and the appropriate value of \( \Sigma \). The parameter \( \sigma_0 \) was set to 500 J/mol from Ref. [50] and \( W_f \) was calculated using the expression given in Refs. [37], which accounts for the chemical composition of the material. The maximum number of additional nucleation sites, \( N \), was fitted and assumed to depend on the stress state. The constants \( \alpha_0, \sigma_0, \) and \( \kappa \) where set to the values given in Ref. [37], and \( m \) was fitted as a common constant for all loading states.

### 5. Discussion of fitting results

Haidemenopoulos et al. [37] have demonstrated that the effect of triaxiality on the transformation kinetics is moderate, when the rest of the fitting parameters between different loading states are the same. However, the experimentally obtained evolution of the martensite fraction, in the present study, indicates that there is a stronger effect of the loading state. The obtained value for the number of pre-existing
nucleation sites, $N_0^p$, is in agreement (within the order of magnitude) with the previous fitted number of pre-existing nucleation sites under uniaxial loading state [37]. The number of additional nucleation sites that are formed by deformation is different for the different loading states, i.e., EQ loading produces slightly more nucleation sites than UN, and both significantly more than TC. This suggests that the extent of martensite formation depends strongly on the number of nucleation sites for martensite and it is less dependent on the loading state (via the triaxiality factor and hence via $ΔG_m$). This is in good agreement with previous experimental studies that show that the mechanisms of nucleation of martensite are strongly affected by the loading state, the texture of the austenite and the SFE [27,51–53].

Haedemenopoulos et al. [37] have demonstrated, in theory, the influence of austenite grain size on the stabilization of austenite and on the suppression of deformation-induced martensite. In the present study, the evolution of the mean austenite volume is considered for fitting the kinetic model to the experimental data. Fig. 10 shows the predictions of the model when the evolution of the mean austenite volume is considered for fitting the kinetic model when the evolution of the mean austenite volume is considered for fitting the kinetic model. The raw/processed data required to reproduce these findings cannot be shared at this time as the data also forms part of an ongoing study.

6. Conclusions

Utilizing cruciform-shaped samples and in situ multiaxial deformation on the neutron diffraction instrument POLDI at SINQ, we follow the TRIP effect in a low-alloy Q&P/TBF deformed under different loading states, namely uniaxial (UN), equibiaxial (EQ), and a non-proportional tension/compression (TC) loading. The results indicate that the loading state influences the martensitic transformation; however, it does not change monotonically with triaxiality ($Σ$) as suggested in previously developed kinetic models: TC loading exhibits the slowest martensite formation kinetics ($Σ$=0.18), whereas UN ($Σ$=0.33) and EQ ($Σ$=0.67) are nearly similar. The experimental results are used for validating a recently developed kinetic model, which accounts for the effect of austenite size and stress triaxiality. It is observed that the effect of triaxiality on the transformation kinetics is moderate; however, in order to describe the observed differences, the model parameters that describe the martensite nucleation sites need to be fitted separately for different loading states. This result is confirmed by previous observations, where the martensite nucleation is dependent on the loading state. Hence, it is concluded that transformation kinetics are dictated strongly by the number of nucleation sites of martensite formation, rather than the growth of martensite (via the triaxiality factor). In situ synchrotron X-ray diffraction under biaxial loading indicates that the austenite grain size decreases with deformation; the observed evolution of austenite size is used in the kinetic model. The decreasing austenite size plays important role on the transformation kinetics, as it slows down the transformation.

Data availability

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

CRediT authorship contribution statement

E. Polatidis: Conceptualization, Data curation, Formal analysis, Writing - original draft. G.N. Haedemenopoulos: Conceptualization, Methodology, Writing - review & editing. D. Krizan: Resources, Writing - review & editing. N. Aravas: Conceptualization, Methodology, Formal analysis, Writing - review & editing. A. Mertens: Resources, Writing - review & editing. M. Smid: Investigation, Writing - review & editing. I. Papadioti: Formal analysis. N. Casati: Investigation. S. Van Petegem: Investigation, Writing - review & editing. H. Van Swygenhoven: Funding acquisition, Writing - review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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