


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## Journal Article

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# Room temperature recovery of cryogenically deformed aluminium alloys

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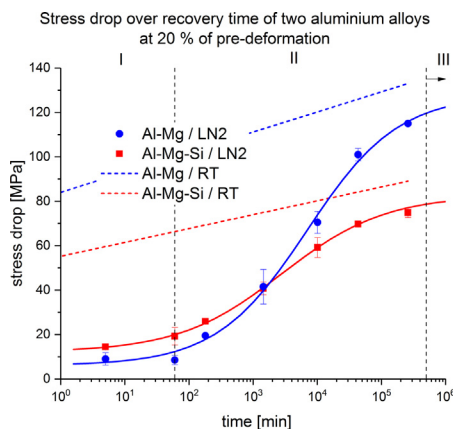
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## HIGHLIGHTS

- Room temperature recovery after cryogenic deformation can be divided into three different mechanistic phases
- These can be distinguished by dislocation rearrangement, dislocation annihilation and a change in strength (stress drop)
- Room temperature deformation directly after cryogenic deformation can result in immediate fracture

## GRAPHICAL ABSTRACT



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## ABSTRACT

Increased formability of aluminium alloys has been demonstrated via cryogenic deformation. In previous studies, the microstructures of samples deformed at low temperatures were analysed after reheating to room temperature (RT) and storage. However, after heating the dislocation structure and density of the deformed material do not reflect the cryogenic situation. In this work, we investigate the evolution of flow stress during recovery in Al-Mg and Al-Mg-Si alloys. We examine the RT recovery behaviour of samples pre-strained at 77 K to different strain levels, and evaluate the structural stability upon subsequent deformation. We also study microstructural evolution via in-situ synchrotron X-ray diffraction, starting from initial conditions at cryogenic temperatures to long-term RT-recovery. Recovery of cryogenically deformed samples at RT results in reduction of the flow stress, in dependence on RT storage. The recovery process can be divided into three distinct sections, each based on a different mechanism characterized by either the arranging or the annihilation of dislocations. Subsequent further straining at room temperature after cryogenic forming also generates plastic instabilities and premature fracture due to unfavourable hardening and recovery assisted softening interplay.

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## 1. Introduction

In the past few years the automotive industry has answered growing demand for lightweight materials to increase fuel efficiency and therefore diminish CO<sub>2</sub> emissions. By using aluminium sheets in car bodies, up to 20% of total car weight can be reduced. [1]. Currently, however, various different alloys for inner and outer panels are being used which weaken the efficiency of the recycling route [2]. Here the main hindrances to a unialloy system with medium to high strength are surface quality and ductility. Production of complex structures often requires high forming limits. Al-Mg based alloys exhibit good formability, but hardening via heat treatment procedures is not possible [3]. In addition, low surface quality prevents the usage of Al-Mg based systems for exterior application or visible parts [4]. Interaction between the magnesium solute atoms and dislocations causes Lüders yielding or PLC effects (also known as dynamic strain aging) [5,6]. Both generate an uneven surface, visible in flame-shaped elevations (stretcher strain markings) or bands. Al-Mg alloys are therefore used mainly for structural or inner parts. In contrast, Al-Mg-Si alloys display better surface quality and higher strength values due to possible precipitation hardening upon paint bake treatment [7,8]. In the automotive industry this artificial aging treatment is applied after the last deformation step (i. e. deep drawing), simultaneously to coat drying and hardening. The main disadvantage of Al-Mg-Si alloys lies in their lower level of formability compared to Al-Mg alloys. This is also a disadvantage of aluminium alloys compared to steels in general [1,9]. A possible way to overcome this is to shift the forming process to higher temperatures which increases the forming limit of the alloys [10] and prevents potential PLC effects [11]. However, this can cause negative effects such as unpredicted thinning and necking, and lower age-hardening potential [12–15].

Recent studies show the advantages of deformation at sub-zero temperatures [16–24]. Schneider et al. [18] described an increase in uniform elongation of 75% for the Al-Mg alloy EN AW 5182 when the temperature is lowered to 77 K. In a multiaxial stress state, realized by limiting dome height experiments, a significant improvement was also achieved. Cryogenic temperatures also suppress the PLC effect thus preventing an uneven surface due to band formation and generate higher surface quality [25]. This makes it appropriate for the automotive industry to deploy Al-Mg alloys in outer panels, and renders a unialloy system possible.

Although the effect of an extended plastic deformation regime was first examined by the National Advisory Committee for Aeronautics (NASA) in 1950 [26], the mechanism behind it is still difficult to investigate due to the limits of the characterisation methods applied. Unfortunately, the directly accessible fracture surface provides few insights into the underlying deformation mechanism and microstructure [27]. Microstructural characterisation of low-temperature strained samples was typically performed after reheating to room temperature (RT). This is certainly due to the fact that sample preparation under low temperatures is hardly possible and cold insertion and measurement of the samples in the electron microscope is only feasible with special setups. Electron microscopy studies were conducted [16,17] via deformation at low temperature; heating to RT for sample preparation; and storage and evaluation of the microstructure at RT. Such measurements do not provide a full picture of the deformation state at low temperatures. Hence, most research into cryogenic deformation has been conducted on samples where the microstructure was recovered to a potentially unknown extent. Moreover, another aspect is the industrial application of cryogenic deformation. Therefore it is important to understand the effects of recovery after cryogenic deformation to be able to determine final component properties. It is also crucial to understand the influence of cryogenic deformation on further RT deformation, as this is relevant in industrial post-processing.

Recovery is characterized by dislocation rearrangement and annihilation of defects to reduce the total energy of the system after

deformation. Only at certain temperatures could the barrier for this processes be overcome, leading to climb and cross slip of dislocation [28–33]. Recovery due to dislocation movement is sensitive to stacking fault energy (SFE). Because the SFE of pure aluminium is high, with values of up to 250 mJ/m<sup>2</sup> [34,35] depending on the measurement method, there is a strong propensity for recovery and initial recovery processes will start even at sub-zero temperatures [28,30,36]. Hence, heating cryogenically deformed material to RT triggers recovery, and the dislocation structure of the deformed material changes.

The aim of this study is to determine the recovery of the lattice defects and the flow stresses after deformation at 77 K and RT storage in order to draw conclusions about the dislocation arrangement at the cryogenic state. Deeper insight into the mechanisms of cryogenic deformation will be possible by considering recovery for previous and future microstructural investigations. Moreover, the industrial process of cryogenic forming has not yet been fully developed and the influence of recovery on subsequent RT post-processing (e.g. multi-step forming) or final component properties will be revealed.

## 2. Experimental

### 2.1. Pre-deformation and tensile tests

Specimens were produced from aluminium sheets EN AW 5182 O and EN AW 6016 T4 with a thickness of 1.2 mm. The chemical composition of the alloys is listed in Table 1.

Pre-deformation and tensile tests were conducted using a Zwick/Roell system with a maximum of 100 kN loading. The sample geometry was in accordance with EN ISO 6892-3 [37] with a width of 12.5 mm and a reduced section of 57 mm. All samples were mill-cut from sheets perpendicular to the rolling direction. An extensometer (Sander 20–10 O) with a gauge length of 20 mm was used to measure the change in length. The tensile tests were carried out at RT (~296 K) and under liquid nitrogen (LN<sub>2</sub>) at 77 K at a strain rate of 0.008 s<sup>-1</sup> (exceptions are specifically mentioned). The strain rate was chosen in accordance with EN ISO 6892-3 [37], but no strain rate change was made during the tensile tests, as this was not possible due to the measurement setup. During deformation at cryogenic temperatures, the reduced section of the specimens and the extensometer were kept in a Dewar filled with liquid nitrogen. The experimental setup of the cryogenic tensile tests is shown in Fig. 1. For temperature control, thermocouples were attached to the sample and inserted in the liquid nitrogen pool. At the same time, the selected strain rate prevented temperature increase in deformation zones, as enough time for heat dissipation is available. Therefore, the sample temperature was between 77 K and 78 K during the entire measurement.

Two sets of measurements were conducted. The measurements sets are simply interrupted tensile tests, which were RT-recovered at different times in the course of the interruption. Both started with pre-deformation at liquid nitrogen to 10%, 20% or 30% strain. Subsequently, the specimens were heated and stored in a Memmert IPP 400 incubator at a constant 25 °C for 5 min to up to 6 months (recovery times were chosen on a logarithmic basis). To ensure fast heating and a measurable initial time for recovery, the samples were unloaded and detached at cryogenic temperatures, and rapidly heated by immersion in ethanol at RT. After different recovery times, tensile tests at 77 K or RT were performed. Each measurement was repeated three times; the error bars refer to the standard deviation. In Table 2, the two different

**Table 1**  
Chemical composition in wt% of EN AW 5182 and EN AW 6016.

Alloy	Mg	Si	Mn	Fe
EN AW 5182 O	4.7	0.10	0.42	0.20
EN AW 6016 T4	1.1	0.35	0.08	0.15

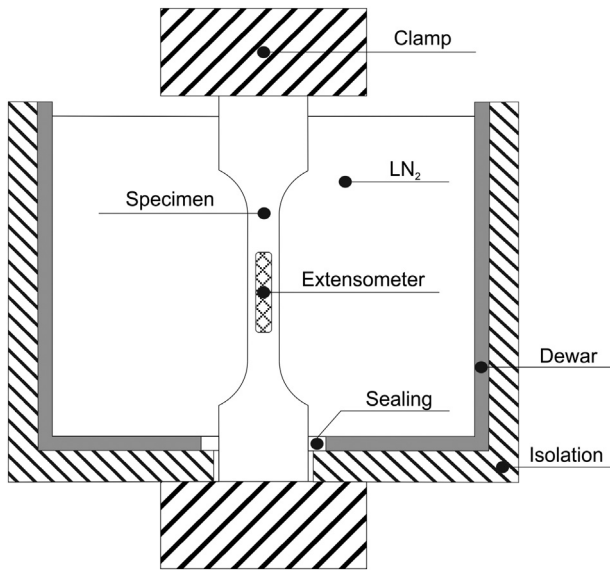


Fig. 1. Diagrammatic sketch of the experimental setup of the cryogenic tensile tests.

measurement sets are listed. All information on investigations involving final deformation until fracture at 77 K are given in Sections 3.2 and 4.1, "Recovery after cryogenic deformation". Samples involving final deformation at RT are summarized in Sections 3.3 and 4.2, "RT deformation after cryogenic deformation". For comparison purposes we have also carried out a third set of measurements (not listed in Table 2). These tests were done only at RT and consisted of tensile pre-deformations, recovery of unloaded specimens for 6 months, and subsequent continuation of tensile tests.

To get a better overview of the recovery processes, the stress drop between the stress level of pre-deformation and the flow onset at further straining were evaluated. Because various irregularities occur at the flow onset, the engineering stress/strain curves were transformed into true stress/strain curves. The flow stress was set in relation to the dislocation density through the Taylor eq. (1) [38]. Here,  $\sigma_0$  is the initial yield stress,  $\alpha$  a material constant,  $\mu$  the shear modulus,  $M$  the Taylor factor,  $b$  the Burgers vector and  $\rho$  the dislocation density.

$$\sigma = \sigma_0 + \alpha \mu M b \sqrt{\rho} \quad (1)$$

In Eq. (1) the dislocation density is dependent on the strain. The evolution of the dislocation density during deformation is given by the Kocks-Mecking model in Eq. (2) [39,40] and consists of two parts. The term  $k_1 \sqrt{\rho}$  represents the dislocation generation based on plastic deformation and the term  $k_2 \rho$  the dynamic dislocation annihilation.

$$\frac{d\rho}{d\varepsilon} = k_1 \sqrt{\rho} - k_2 \rho \quad (2)$$

By means of the KM model, the flow curves were approximated and the difference in stress between pre-straining and the calculated flow onset after recovery evaluated. All parameters used in the modelling are listed in Table 3.

Table 2  
The two different sets of measurements and their experimental parameters.

Pre-deformation [%]	Recovery time at RT	Further straining until fracture
10, 20 and 30 at LN <sub>2</sub>	5 min, 1 h, 3 h, 1 d, 1 w, 1 m, 6 m	LN <sub>2</sub>
10, 20 and 30 at LN <sub>2</sub>	5 min, 1 h, 3 h, 1 d, 1 w	RT

## 2.2. Synchrotron measurements

Synchrotron measurements were performed using a fixed photon energy of 60 keV on the Petra III beamline P02.1 at DESY (Hamburg). The beam size was  $200 \times 200 \mu\text{m}$  and diffraction patterns were recorded by a 2-dimensional Perkin-Elmer XRD1621 detector. The wavelength and sample to detector distance were calibrated using LaB<sub>6</sub> (NIST660b) and CeO<sub>2</sub> (NIST674b) standards, respectively. Similar to the tensile tests, a set of cryogenic preloaded samples was measured during heating and storage at RT. Therefore, tensile test samples of sheet material were deformed to 10%, 20% and 30% of strain under LN<sub>2</sub>. After unloading, sample preparation and mounting to the temperature stage were carried out in a Linkam furnace under LN<sub>2</sub> using a solid state sample holder designed for relaxation studies [44] to create the unique possibility to study recovery processes from the moment of heating up. Diffraction patterns were recorded in-situ during heating from 77 K to RT and held at RT for up to 3 h to evaluate the recoverable fraction of dislocations upon heating and storage. Additionally, samples with a recovery time of 12 h, 24 h, 1 d, 1 week and 1 month were measured separately. Afterwards the diffraction patterns with respect to the azimuthal range were integrated using pyFAI-software [45]. The change in dislocation density was calculated by X-ray line profile analysis using Fityk [46] for the peak position evaluation and Convolutional Multiple Whole Profile fitting (CWMP) [47,48] with a combinatorial approach to achieve a high numerical stability [49,50].

## 3. Results

### 3.1. Stress-strain behaviour at RT and 77 K

Fig. 2 shows the stress-strain curves of alloys EN AW 6016 T4 and EN AW 5182 O at RT and 77 K, respectively. Particularly noteworthy is the strain hardening behaviour, which is strongly temperature-dependent in both alloys.

At the onset of plastic deformation the influence of the temperature on the flow stress is rather moderate (difference between RT and 77 K:  $\Delta\sigma \approx 30$  MPa for EN AW 6016 T4 and  $\Delta\sigma \approx 35$  MPa for EN AW 5182 O, mainly determined by the thermally activated hardening contributions of the obstacle strength of clusters in EN AW 6016 T4 [51,52], and solid solution hardening in EN AW 5182 O [53]). The temperature influence increases markedly with increasing strain. At 20% plastic strain, for example, the differences in flow stress are  $\approx 80$  MPa (EN AW 6016 T4) and  $\approx 100$  MPa (EN AW 5182 O), respectively. The hardening effect becomes particularly apparent in the plots in Fig. 2(b) and 1(d), where the slope of the curves provides information on the strain hardening rate. The low slopes at 77 K indicate that even at high strength levels considerable strain hardening occurs. It can therefore be concluded that the increase in dislocation density with plastic strain is more pronounced at 77 K and a higher density of lattice defects will be generated. Consequently, heating of samples that have been deformed at 77 K to RT may result in softening, via processes described in the next section.

Table 3  
Defined parameters for the calculation of the flow curves [41–43].

Parameter	Value	Unit
$\sigma_{0, 6016 LN_2}$	138	MPa
$\sigma_{0, 5182 LN_2}$	152	MPa
$\alpha$	0.6	–
$\mu$	$2.83 \cdot 10^4$	MPa
$M$	3.06	–
$b$	$2.8 \cdot 10^{-10}$	m
$\rho_0$	$1 \cdot 10^{-10}$	m <sup>-2</sup>

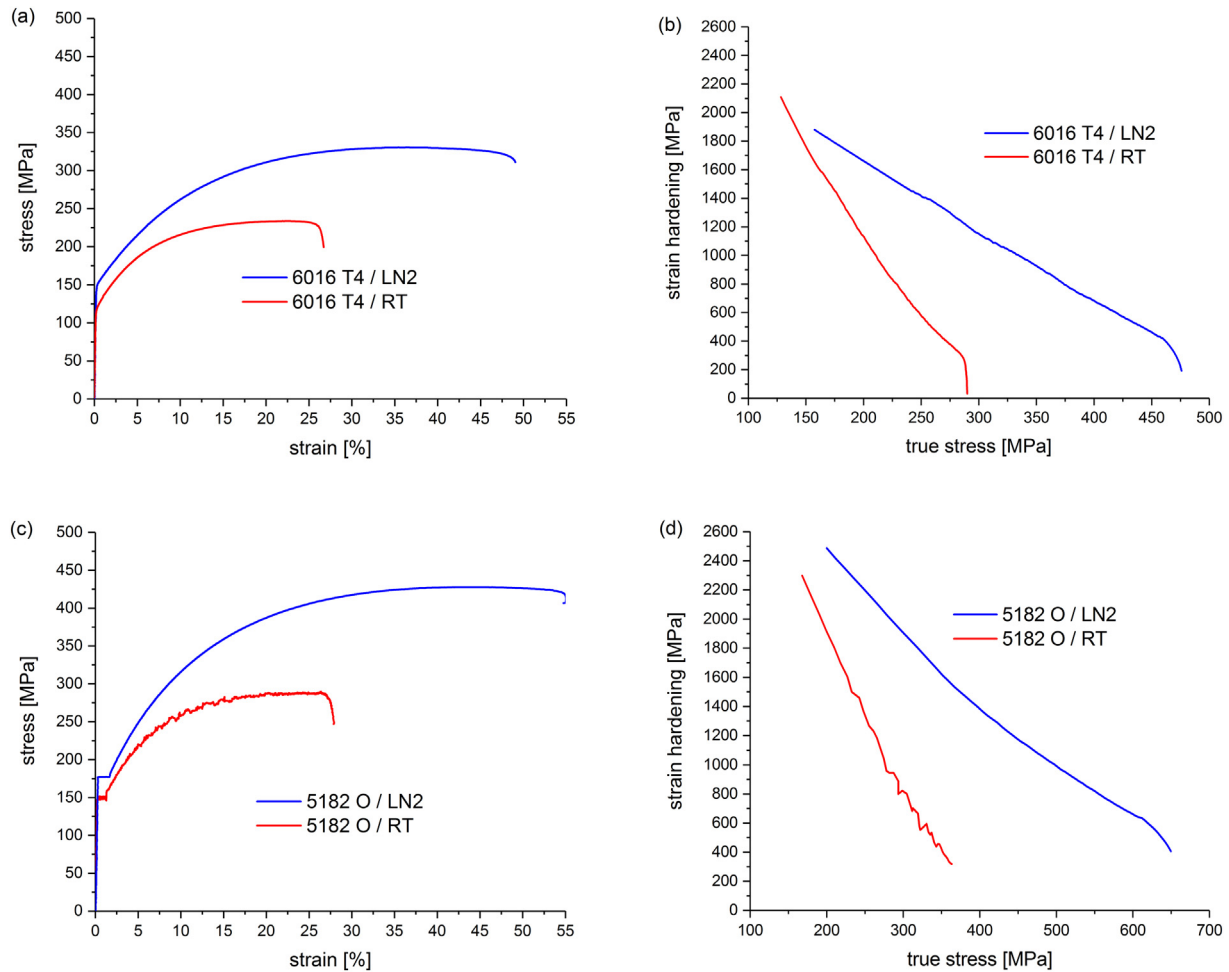


Fig. 2. Tensile test curves of alloys EN AW 6016 T4 and EN AW 5182 O at 296 K and 77 K (a,c) and corresponding strain hardening plots (b,d).

### 3.2. Recovery after cryogenic deformation

In Fig. 3 and Fig. 4 tensile test curves of alloy EN AW 6016 T4 and EN AW 5182 O at 77 K are shown where the tests were interrupted and the samples recovered at RT for a certain time. Each measurement consists of a pre-deformation at 77 K with subsequent unloading and RT recovery, and afterwards final straining until fracture at 77 K. Except for the curve for 5 min of intermediate RT recovery, all other curves are shifted vertically by  $-25$  MPa each to improve the visibility of the effects present. Note that the curves are displayed only to uniform elongation. For longer recovery times between pre-deformation and final deformation the yield stress of the second curve decreases. For short recovery times this drop is not as pronounced. However, the onset of flow at the second straining of the samples needs attention. Two cases were observed: a pronounced plateau-like area with constant or slightly negative slope and a yield point followed by a stress decrease. The evolution of the onset flow is best illustrated by alloy EN AW 6016 T4. The tensile test curves with 10% pre-strain in Fig. 3(a) show no plateau but only a stress drop after the yield point. At 20% pre-deformation (Fig. 3(b)) the plateau occurs between 5 min and 3 h, followed by stress decrease. Similar to this, Fig. 3(c) with 30% pre-straining displays a wide plateau at 5 min, even with a drop in stress compared to the pre-straining curve. After 1 day, the plateau has disappeared. A similar sequence is visible for alloy EN AW 5182 O in Fig. 4. In every pre-deformation state a plateau with a negative slope appears within 3 h of storage at RT, followed by a yield point with a subsequent stress drop. At 10% and 20% pre-deformation a slight stress overshoot also

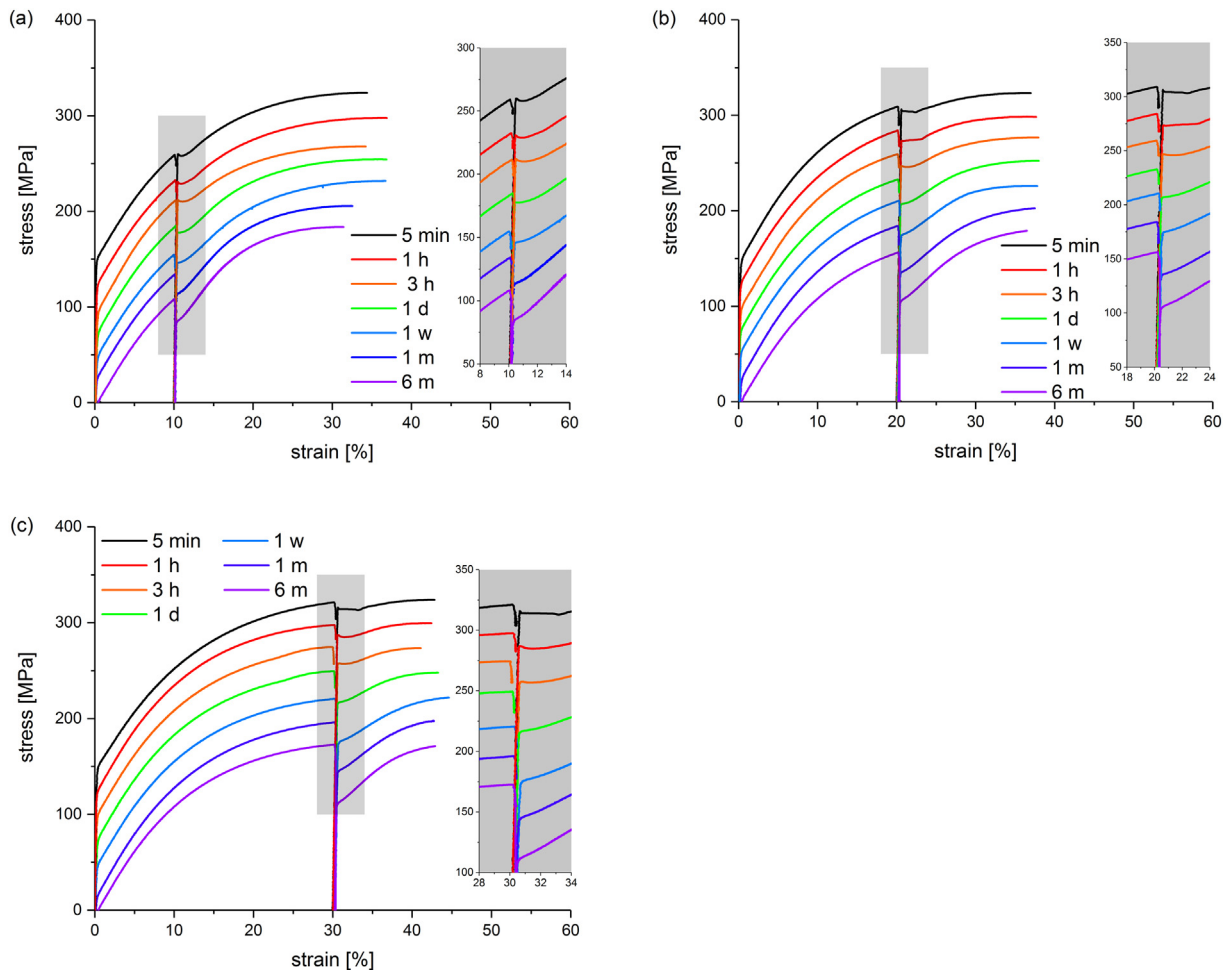
appears. Where there is very short RT recovery this can happen to EN AW 5182 O with a slightly higher flow onset than when pre-deformation is stopped.

As seen in Fig. 3 and Fig. 4 it is difficult to access the actual stress drop due to stress plateaus and yield points followed by a stress decrease evolution. For a more detailed evaluation and to bypass uncertainties in the flow onset, the stress drop after recovery was determined by back extrapolation of the yield point from the flow curve with the KM model. A more detailed description can be found in the supplementary.

In Fig. 5 the stress drop derived upon RT recovery is displayed for alloys EN AW 6016 T4 and EN AW 5182 O. For guidance purposes, the inclined dashed lines of corresponding colour show the theoretically maximum achievable recovery, calculated by the difference in strength between flow curves deformed to identical strain at RT and 77 K. They consider also the RT recovery of RT-deformed samples, which is expressed in the slope of the lines. This slope was assumed to be linear logarithmic [54,55] and defined by the initial difference in strength between flow curves at RT and 77 K and after 6 months of recovery. Because 30% strain was not achievable at RT tensile testing due to failure at lower strain (see Fig. 2), the flow curves were extrapolated using the KM approach.

Both alloys in Fig. 5 display a minor stress drop after heating to RT at short times, before recovery accelerates. Past this onset, the kinetics changes at approximately 1 h for EN AW 6016 T4 and 5 h for EN AW 5182. Afterwards an increase in recovery generates a faster stress reduction until it slows down again. Following this decrease in recovery, the slope is similar to the stress drop of RT-deformed samples. The slowing





**Fig. 3.** Tensile test curves of alloy EN AW 6016 T4 after pre-straining at 77 K for (a) 10%, (b) 20% and (c) 30%, and intermediate recovery at RT for 5 min to 6 months and subsequent deformation at 77 K. All curves are displayed to uniform elongation and except 5 min are shifted by  $-25$  MPa for better visibility of the effects.

down of the recovery rate is only visible after 6 months or longer and is therefore at the limit of the observation period. These three different stages of recovery are separated by grey, vertical, dashed lines.

### 3.3. RT deformation after cryogenic deformation

In addition to the interrupted RT-recovered tensile tests at 77 K, a second experimental set was performed to gain knowledge of the remaining RT deformation behaviour after cryogenic pre-deformation. As illustrated in Fig. 6 and Fig. 7, samples of alloys EN AW 6016 T4 and EN AW 5182 O were pre-strained at 77 K, stored at RT for periods of 5 min to 1 week, and afterwards tensile tested at RT. The only difference to the previous measurements was that the final straining was carried out at RT and not at 77 K.

Fig. 6(a) displays this treatment after 10% of cryogenic pre-deformation for alloy EN AW 6016 T4. If the final tensile test at RT is performed shortly after heating (i.e. 5 min), limited elongation is observed. However, the elongation recovers after 1 h of RT storage. With longer RT recovery, minor changes in the flow onset are visible but the reachable elongation remains the same. In Fig. 6(b) and (c) with 20% and 30% of pre-deformation at 77 K, short recovery (up to 3 h) causes immediate fracture at further RT deformation. Interestingly, significant further deformation potential could be achieved after 1 day (20%) or 1 week (30%) of RT recovery.

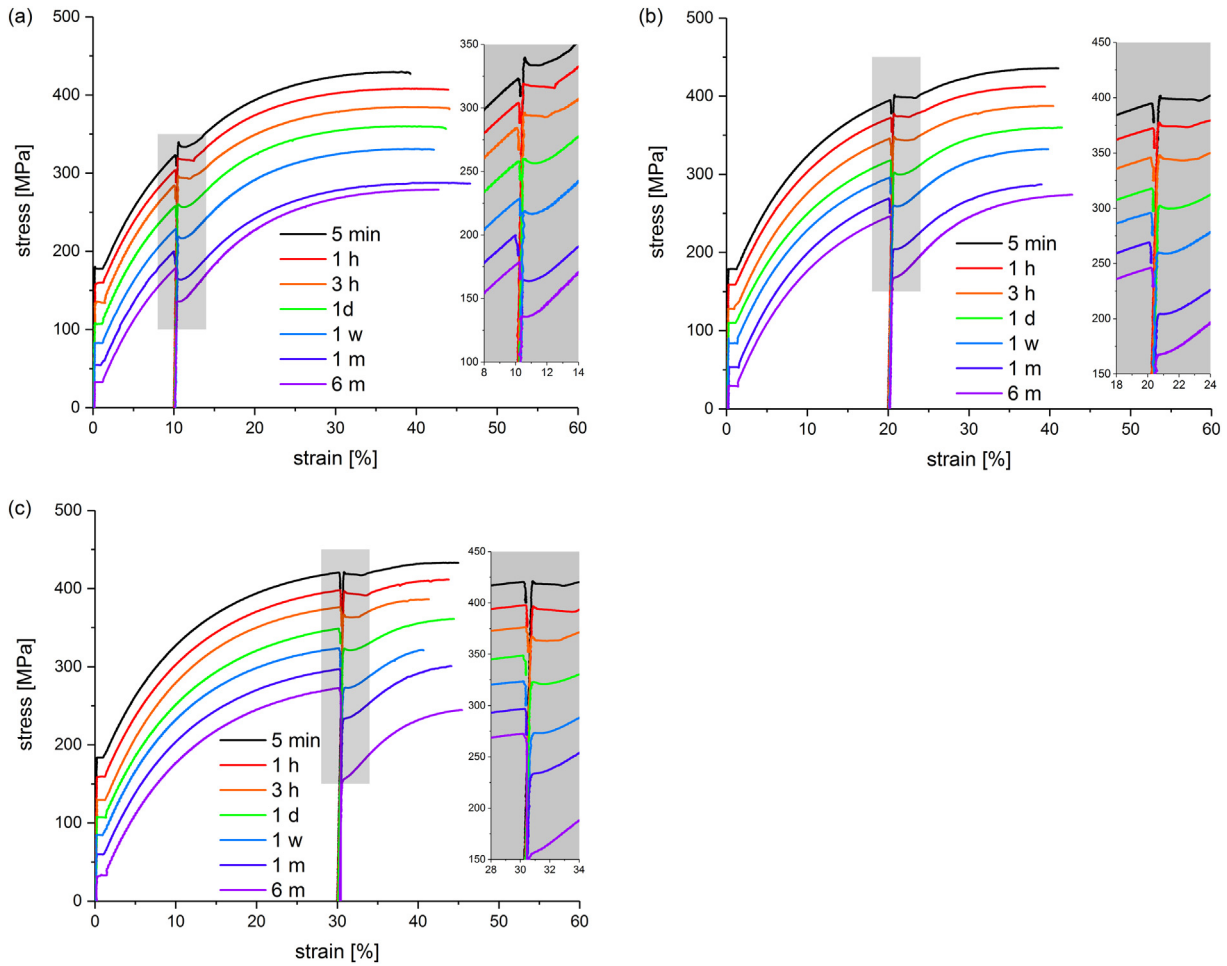
The behaviour of alloy EN AW 5182 O in Fig. 7 shows the same trend, but at RT jagged tensile test curves are obtained due to the known

interaction of magnesium in solid solution with the dislocations. Similarly to alloy EN AW 6016 T4 at higher pre-deformation, additional elongation is only possible after 1 day and 1 week of RT storage. However, even 5 min of recovery are sufficient to ensure further deformability in the 10% pre-deformed samples.

In Fig. 6(a) the 5 min RT recovery curve is of particular interest, as it is close to immediate fracture. To gain additional information the strain rate was varied from  $0.008$  s $^{-1}$  to  $0.08$  s $^{-1}$  and  $0.0008$  s $^{-1}$ . The results, illustrated in Fig. 8, reveal that at lower strain rates failure appears earlier, i.e. instantly after yielding. At the higher strain rate of  $0.08$  s $^{-1}$  a further 12% elongation can be applied, comparable to that applied to the specimens stored for 1 h at RT.

### 3.4. Dislocation densities

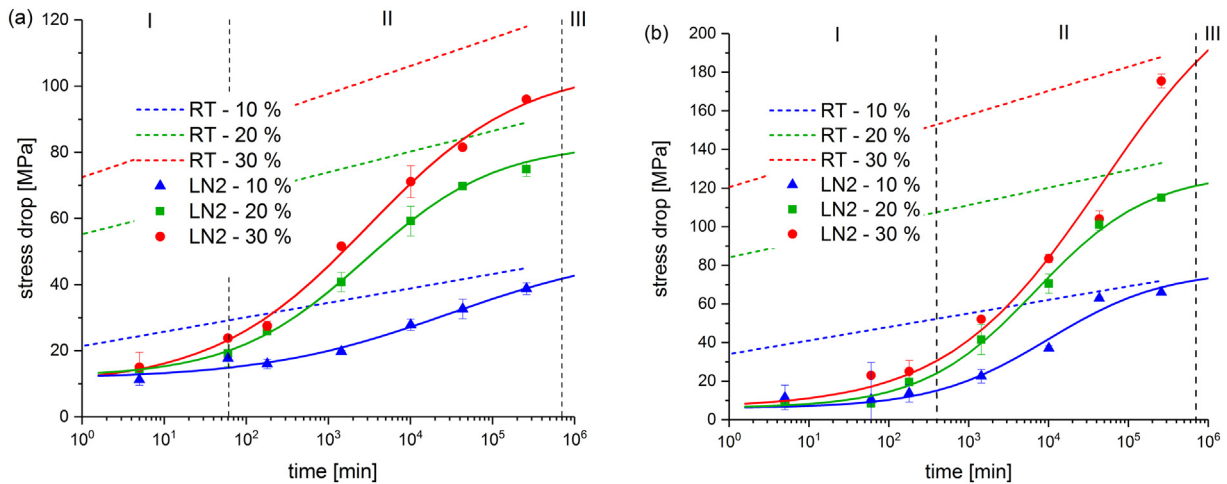
Fig. 9 and Fig. 10 compare the dislocation densities over RT recovery time of EN AW 6016 T4 and EN AW 5182 O after deformation at 77 K. The data was acquired by two different means. On the one hand the dislocation density was simply calculated from the tensile test curves in Fig. 3 and Fig. 4 by back extrapolation with the KM model (Fig. 9 (a) and Fig. 10(a)). The solid lines represent the dislocation density with recovery time after varied pre-deformation at 77 K. For comparison, the dashed lines show values for recovery after simple deformation at RT which were also deduced from tensile tests (compare with Fig. 5). On the other hand data from X-ray line profile analysis from synchrotron measurements are displayed (Fig. 9(b) and Fig. 10(b)).



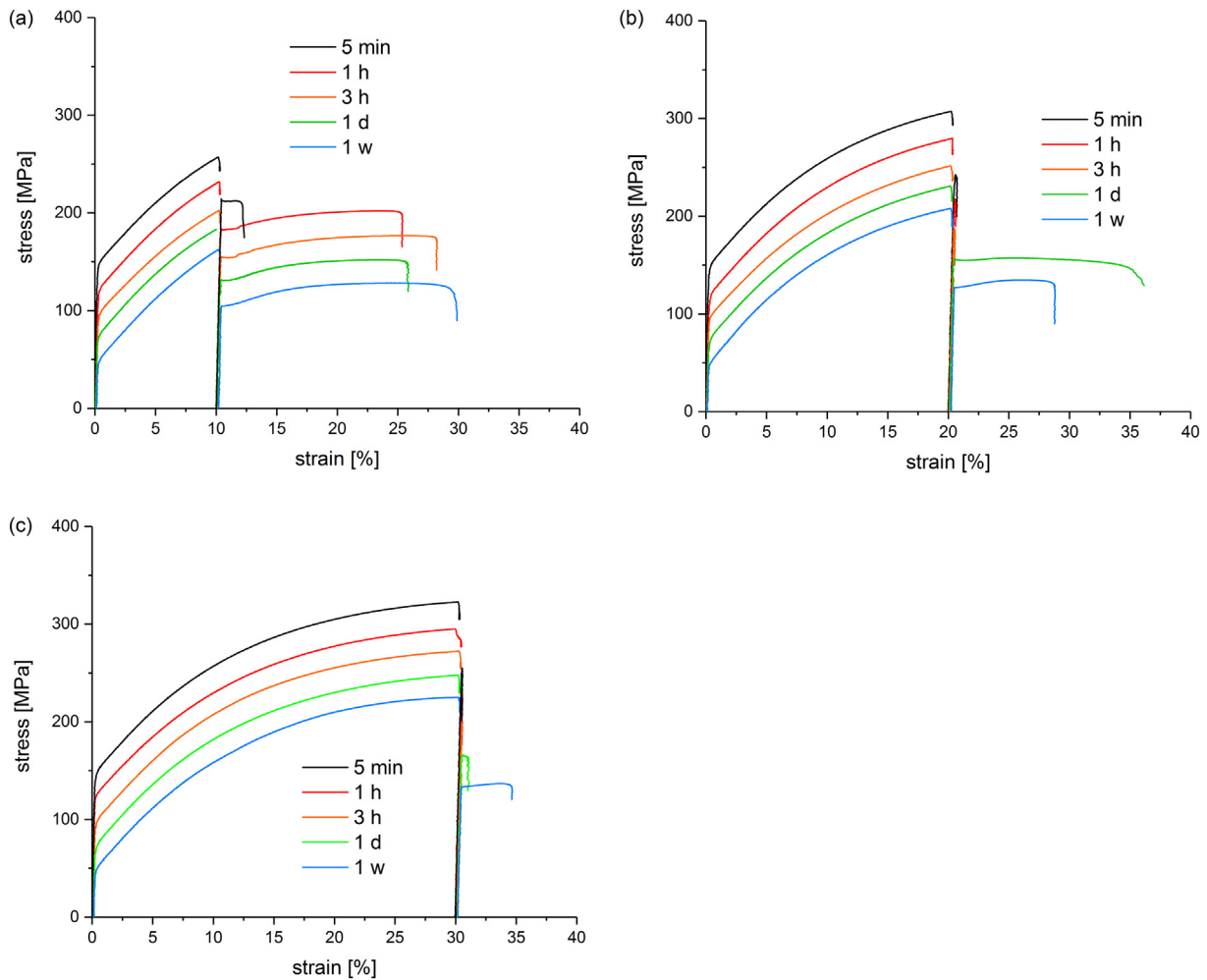
**Fig. 4.** Tensile test curves of alloy EN AW 5182 O after pre-straining at 77 K (a) 10%, (b) 20% and (c) 30% and intermediate recovery at RT for 5 min to 6 months and subsequent deformation at 77 K. All curves are displayed to uniform elongation and except 5 min are shifted by  $-25$  MPa for better visibility of the effects.

The results from the tensile tests predict an immediate minor decline in dislocation density for the samples pre-deformed at 77 K, followed by a rapid drop. After long recovery times of up to 6 months this drop in dislocation density is reduced. Compared to the synchrotron measurement in Fig. 9(b) and Fig. 10(b), the trend of decreasing dislocation density is the same, but tends to higher values. Another difference

is that the data from synchrotron measurements show no initial changes in dislocation density. This is even more pronounced for EN AW 5182 O (Fig. 10 (b)). Note that in contrast to the dislocation densities deduced from the tensile tests the synchrotron measurements also include the in-situ heating of the pre-strained samples from 77 K to 296 K.



**Fig. 5.** Stress drop after deformation at 77 K over RT recovery time for alloys (a) EN AW 6016 T4 and (b) EN AW 5182 O for different pre-straining.



**Fig. 6.** Tensile test curves of alloy EN AW 6016 T4 upon pre-straining at 77 K for (a) 10%, (b) 20% and (c) 30% and intermediate recovery at RT for periods of 5 min to 1 week and subsequent deformation at RT. All curves except 5 min are shifted by  $-25$  MPa for better visibility of the effects.

## 4. Discussion

We have probed the deformation and recovery behaviour of the two alloys EN AW 5182 O and EN AW 6016 T4 at RT and 77 K via tensile tests, measurements of the dislocation densities by X-ray line profile analysis from synchrotron, and a simple KM modelling approach. At low temperatures a lower annihilation of the dislocations occurs and strain hardening stays at higher levels. Heating samples to RT results in complex recovery phenomena which creates various types of distinct flow onset behaviour upon reloading, which will be discussed in Section 4.1.

Further RT deformation of cryogenically pre-deformed samples can even lead to direct failure. However this depends on the degree of cryogenic pre-deformation, type of alloy and strongly on the applied strain rate at RT. Mechanisms and technological implications behind this unexpected behaviour are discussed in Section 4.2.

### 4.1. Recovery after cryogenic deformation

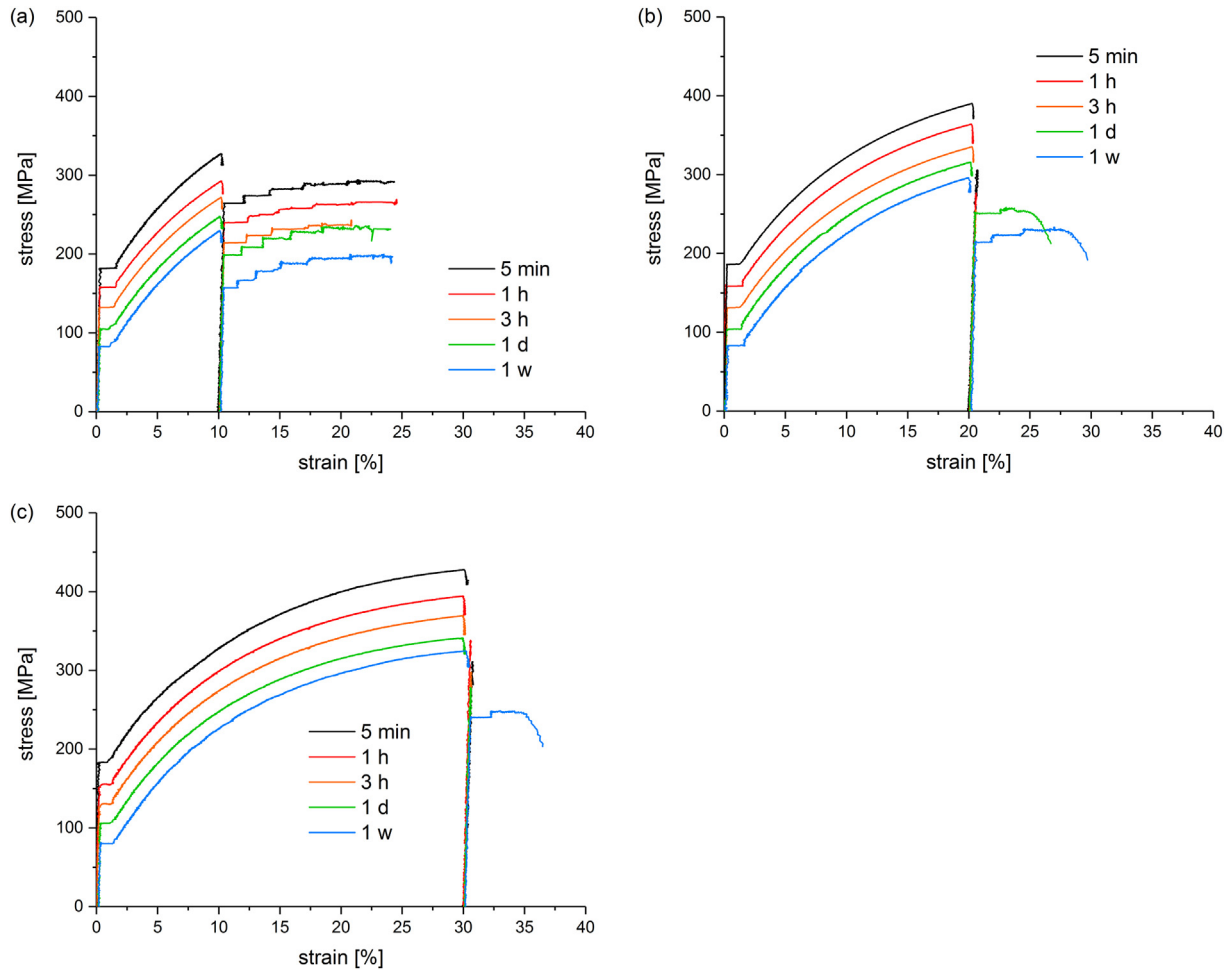
Fig. 5 provides the best overview of the processes during RT recovery of samples deformed at 77 K. Note that the shape of the stress drop due to RT recovery over recovery time in Fig. 5 cannot be modelled by introducing a simple static recovery term to the KM model for either of the two alloys. Such term for static recovery is usually impacted by vacancy-assisted dislocation climb and is reported to be negligible at temperatures below 300 K in aluminium alloys [41,56]. The curves

from Fig. 5 cannot even be reduced to a trivial linear logarithmic scale as typically seen for recovery after conventional RT deformation [54,55]. In summary, the observed recovery process can be divided into three stages. As described, both alloy systems initially show only a slight stress drop in the first few hours after heating up to RT (i), with a minor dependence to the applied pre-strain. After this phase the recovery increases rapidly (ii) until saturation starts (iii). The increase (ii) and the time to saturation (iii) depends strongly on the pre-deformation. A similar behaviour has also been observed after deformation during static recovery at elevated temperatures [57]. This was caused by recrystallization processes, which could not take place here. In the following the observed three stage behaviour is discussed.

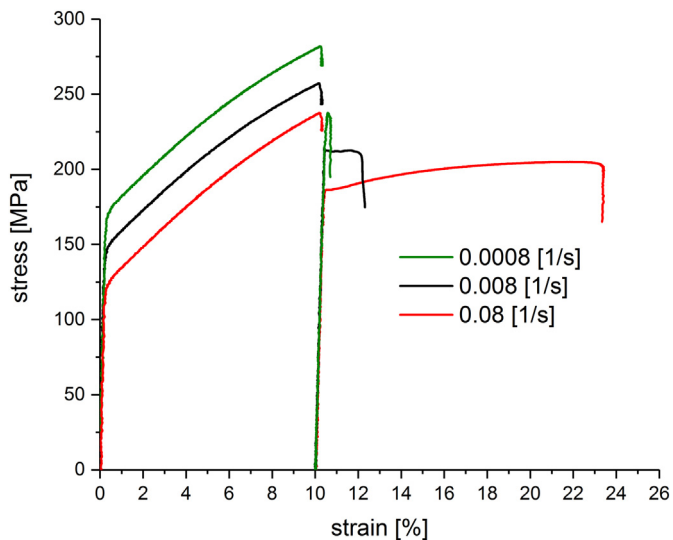
#### 4.1.1. Stage I

The first stage of recovery (i) starts directly after heating, when the tensile test curves in Fig. 3 and Fig. 4 show a change in the type of flow onset. The development of a yield point with stress decrease and stress plateaus indicate initial recovery processes. The plateaus cannot be ascribed to Lüders yielding caused by magnesium in solid solution for at least two reasons. Firstly, alloy EN AW 6016 T4 in natural aged condition contains a very low content of magnesium atoms in solid solution and also displays the respective behaviour. Although dissolution of Mg-Si clusters after extensive dislocation cutting may increase Mg in solid solution, the maximum possible amount for achieving Lüders yielding does not match the observations of several authors





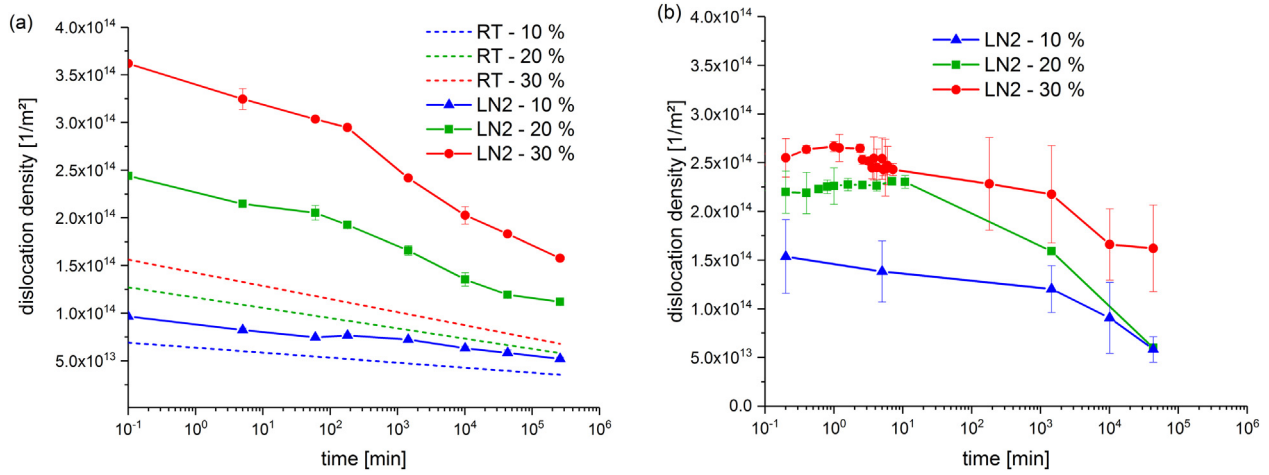
**Fig. 7.** Tensile test curves of alloy EN AW 5182 O upon pre-straining at 77 K for (a) 10%, (b) 20% and (c) 30% and intermediate recovery at RT for periods of 5 min to 1 week and subsequent deformation at RT. All curves except 5 min are shifted by  $-25$  MPa for better visibility of the effects.



**Fig. 8.** Influence of strain rate on tensile test curves of alloy EN AW 6016 T4 upon pre-straining at 77 K for 10%, intermediate recovery at RT for 5 min and subsequent deformation at RT. All curves except  $0.008 \text{ s}^{-1}$  strain rate are shifted by  $-25$  MPa for better visibility of the effects.

[5,6,58,59]. Secondly, these “stress plateaus” occur and also vanish in alloy EN AW 5182 O after longer recovery times and exhibit mostly negative slopes. We suggest that these observed stress-plateaus can be seen as dislocation rearrangement triggered by applying stress. This is supported by the fact that during long term recovery first rearrangement processes and also dislocation annihilation took already place, the plateaus do not occur anymore. A detailed explanation for this assumption is further given below on the discussion of strain hardening.

In alloy EN AW 5182 O a slight stress overshoot also appears, which even produces an increased flow onset after short RT recovery. This could be explained by the migration of vacancy-Mg to dislocations or the direct interaction of Mg [60] with dislocations generating a strong dislocation-solute interaction [61]. At RT deformation of this alloy, dynamic strain aging would occur, stemming from the interaction between dislocations and these complexes [62] or single Mg atoms [60]. In the case of interrupted deformation at 77 K, the Mg interactions with mobile dislocations appears during RT storage. Hence the stress overshoot is assumed to result from “frozen-in dynamic strain aging”. Simultaneously to the strong interaction between Mg and dislocations always recovery processes occur. However, at low pre-deformations and times, the recovery processes are less pronounced and a stress overshoot is clearly visible. With prolonged storage at RT, the recovery becomes more predominant and no increase in strength is visible. At 30% of pre-deformation the dislocation density is higher and hence the driving force for recovery. As a result, the recovery predominates and the stress overshoot is no longer noticeable.

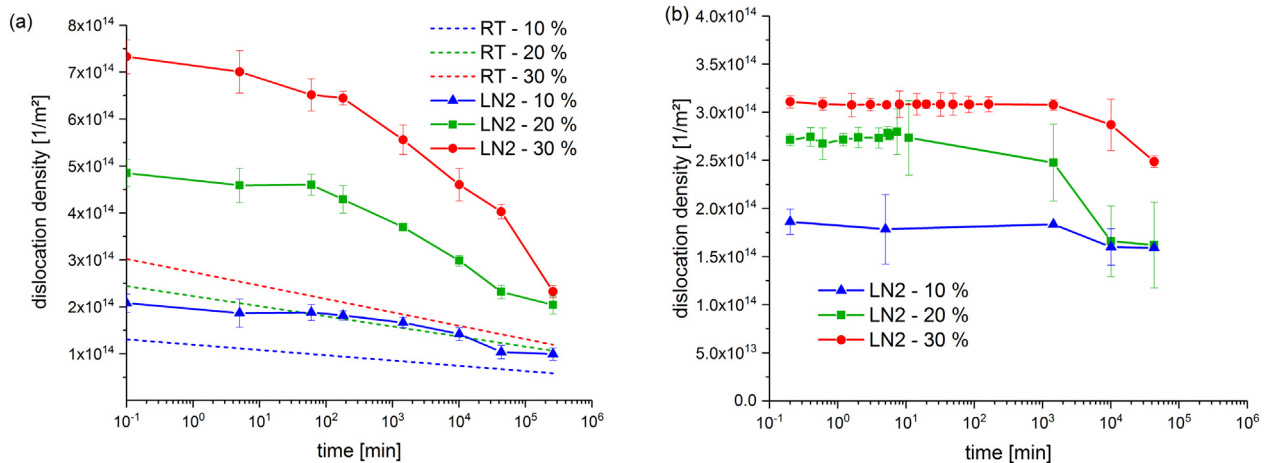


**Fig. 9.** Dislocation densities versus RT recovery time of alloy EN AW 6016 T4 pre-deformed at 77 K, derived by (a) back extrapolation from tensile tests and (b) from X-ray line profile analysis of synchrotron data.

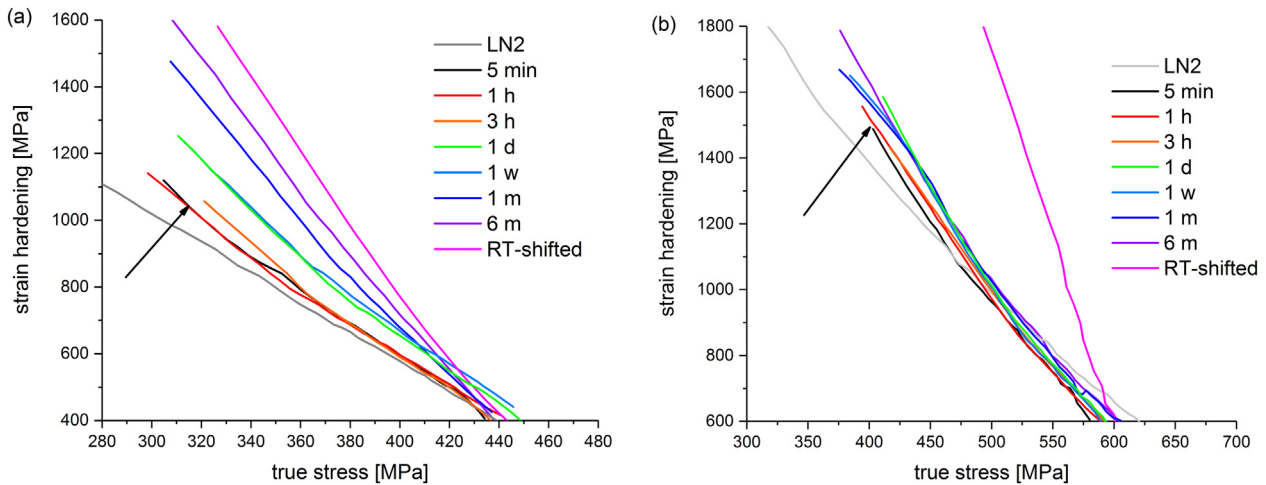
To gain a better insight into the flow onset phenomenon, we focus on the Kocks–Mecking plots of the deformation behaviour after RT recovery (Fig. 11). This allows an easier and clearer explanation of the strain hardening behaviour of the tensile test curves. Only at the beginning of straining in phase (i) of recovery the curves differ and display a higher strain hardening rate (steeper slope). Further on, the strain hardening rate in alloy EN AW 6016 T4 is equivalent in position (dislocation storage) and slope (dislocation annihilation) to the continuous tensile test under LN<sub>2</sub> without interruption. This can be explained as follows. During deformation, moving dislocations generate vacancies. They annihilate on slowed dislocations, which leads to joggling and local climb (pinning points). The dislocation in between bows out, building loops of different Burgers vectors and tangling in the end [55]. Short RT recovery afterwards leads to a relaxation and to unkinking and untangling of dislocations [63,64], also expressed by a yield phenomenon. Removing these bowed out extra line lengths is easy; only low driving forces are needed [65,66] and the subsequent strain hardening curve are left unchanged. Similar results were also found in pure aluminium cold worked at RT [33,67]. For short recovery times these studies showed that after an initial phase with very high strain hardening rates the curve will overlap with the original continuous straining at RT. Only at higher recovery times and higher temperatures is the RT behaviour

overcome. This initial phase is explained by the transformation of a recovery of loosely tangled dislocations to a cell or subgrain structure with regular dislocation networks and a decrease in the dislocation density, followed by subgrain coarsening and growth. Compared to our results, and taking much lower temperatures into account, this means that after an initial onset (arrow in Fig. 11(a)) the strain hardening curve follows the original tensile test without RT recovery. Therefore no significant annihilation of dislocations may occur, as revealed below. The onset is likely to appear due to a slight rearrangement of dislocations leading to untangling or sharpening of cell walls.

As previously mentioned, in stage (i) there is only a minor stress drop. In the Taylor equation (Eq. (1)) the average dislocation density is proportional to the stress drop. Hence, also the calculated dislocation density will only be minimally influenced (Fig. 9(a) and Fig. 10(a)). It is important to note that this simple approach provides no insights into dislocation arrangement, and a change in strength can only be expressed through a change in dislocation density. In contrast, our synchrotron measurements (Fig. 9(b) and Fig. 10(b)) show clearly no change in the dislocation densities during heating and the first minutes of RT storage. This is compatible with the above described mechanistic of the observed flow onset phenomenon, where at the onset of recovery only detangling of dislocations is assumed to occur.



**Fig. 10.** Dislocation densities versus RT recovery time of alloy EN AW 5182 O pre-deformed at 77 K, derived by (a) back extrapolation from tensile tests and (b) from X-ray line profile analysis of synchrotron data.



**Fig. 11.** Strain hardening curves versus true stress (KM plot) upon recovery of alloys (a) EN AW 6016 T4 and (b) EN AW 5182 O. Note that the RT curves are shifted to higher stress levels to still see the difference in between the LN<sub>2</sub> curves (compare to Fig. 2(b,d)). The arrow indicates the onset of the strain hardening curve, which differs from the LN<sub>2</sub> curve, after short times of recovery (stage (i)) of up to one day.

#### 4.1.2. Stage II

After stage (i) a rapid increase in recovery follows, which is referred to here as stage (ii) of recovery (Fig. 5). The change in stress drop is slightly dependent on the pre-deformation and is linked to the alloy system. Crucial for a suppressed start seems to be the dissolved magnesium in alloy EN AW 5182 O, which is known to be responsible for lower mobility of dislocations [68]. Accordingly, a higher permanent dislocation density is achieved by pre-deformation in EN AW 5182 O due to higher strain hardening (and obviously higher strength values). These aspects are assumed to generate a higher driving force and, after a longer period of moderate recovery, a sudden and stronger stress drop as seen in Fig. 5.

In stage (ii) of recovery a rapid decrease in dislocation density is calculated (Fig. 9(a) and Fig. 10(a)). This is more pronounced in alloy EN AW 5182 O. Comparing the synchrotron measurement to the results from the tensile tests in Fig. 9(b) and Fig. 10(b) provides only a rough insight into the trend of the evolution of dislocation densities, indicating a recovery process dominated by annihilation. The data from the synchrotron indicates a later onset of annihilation of dislocations as the calculations from the tensile tests. Although the absolute values are slightly shifted compared to the calculated dislocation densities, they match surprisingly well.

More insight into the dislocation arrangement is given by the strain hardening curves of the recovered samples in Fig. 11. Alloy EN AW 6016 T4 exhibits a rotation in the slope of the strain hardening curve. The slope slowly changes after one day of recovery to exhibit strain hardening characteristics which would be observed at RT deformation and therefore gives some insight into the dislocation structure involved. Please note that the RT curves are shifted along the abscissa to gain a more compact overview of the slopes. It should be noted that the shift in true stress does not affect the dislocation annihilation parameter. The behaviour of alloy EN AW 6016 T4 illustrated in Fig. 5 leads us to the conclusion that the recovery has progressed to a dislocation structure similar to that of RT-deformed samples, which is stable enough to determine the strain hardening rate during further deformation at 77 K. With the change in slope to RT work hardening behaviour with further deformation, the maximally achievable stress is also slightly reduced because with further deformation the strain hardening decreases significantly. Hence the uniform elongation is also decreased. In contrast, alloy EN AW 5182 O shows a much smaller change in strain hardening slope, which begins immediately after heating, but does not change with continuous recovery up to 6 months, and never approaches RT behaviour. As already mentioned, alloy EN AW 5182 O exhibits

reduced mobility of dislocations due to the high content of Mg [68], and therefore less stress drop during the first few hours, but this changes after longer storage time at RT. Nevertheless, this is not visible in the strain hardening characteristics, because the LN<sub>2</sub>-deformed samples do not approach the RT curve even with longer recovery times. With further straining the recovered dislocation structure is replaced immediately with further deformation at 77 K, also after 6 months. It may be assumed that either no stable dislocation structure is obtained after a longer recovery, as is the case with EN AW 6016 T4, or the RT curve has such a slope only due to the PLC effect, which is suppressed at lower temperatures.

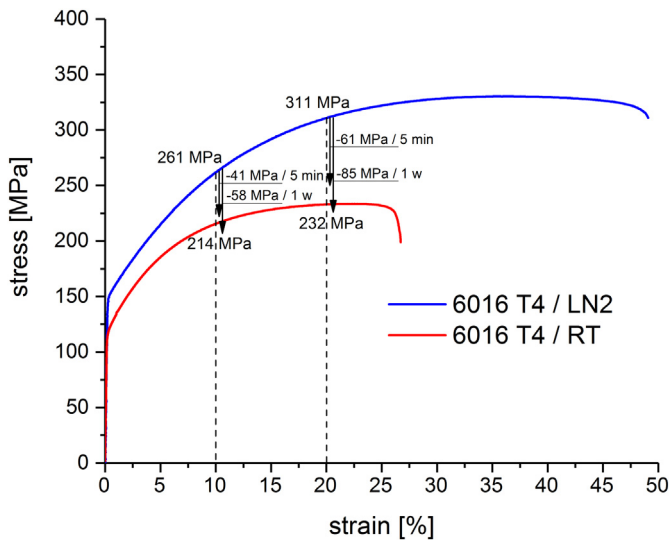
#### 4.1.3. Stage III

In stage three (iii) the further stress drop or dislocation reduction is comparable to RT-deformed and stored samples [55]. Because the defect density has rapidly decreased in the first two stages the driving force of the recovery decreases. The dislocation density approaches the density of RT-deformed structures (stress level difference to RT-deformed samples shown by the dashed lines in Fig. 5) and the stress drop curve turns in the direction of logarithmic behaviour [32,55,69].

#### 4.2. RT deformation after cryogenic deformation

In both alloys after 20% and 30% of pre-deformation at 77 K we observe immediate fracture at further straining at RT. Additional deformation is only possible after one day or one week of RT storage. Alloy EN AW 6016 T4 allows no direct post-straining at RT even with 10% pre-deformation at 77 K, because the sample displays no net strain hardening and fracture occurs after just 2% of plastic deformation. This information is especially important in industrial processes, as the cryogenically formed component is very susceptible to failure directly after heating. This effect may require consideration in further processing of components.

Fig. 12 provides an exemplary overview of the tensile test curves of alloy EN AW 6016 T4 at RT and 77 K with explicit values of 10% and 20% straining. The arrows indicate the yield point of RT tensile tests after five minutes and one week of RT recovery after deformation at 77 K. Only 10% pre-deformation at 77 K (261 MPa) generates higher strength values than the ultimate tensile strength at RT (232 MPa). Even if the thermally activated portion of the yield strength ( $\approx 30$  MPa) is subtracted, the stress level of the 77 K pre-deformed sample is still in the range of the ultimate tensile strength at RT. Shortly after pre-straining at 77 K almost no additional deformation at RT is possible;

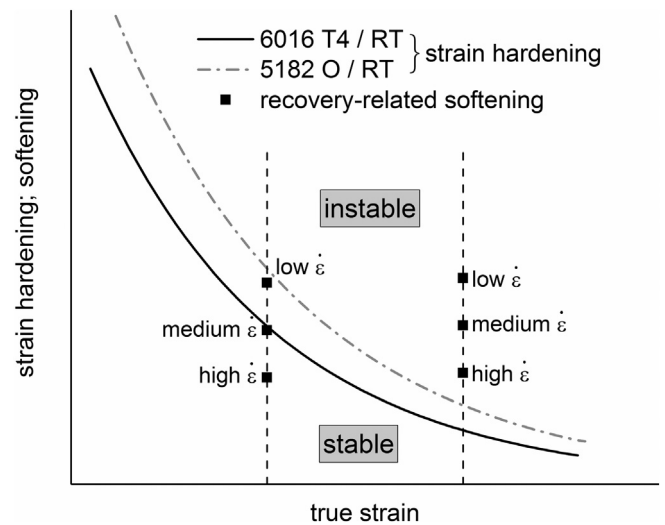


**Fig. 12.** Tensile test curve of alloy EN AW 6016 T4 at RT and at 77 K. The arrows indicate the yield strength of RT reloading after deformation at 77 K and 5 min and 1 w of RT recovery, respectively (RT yield strength of recovered samples).

this is shown in Fig. 6(a), where the sample fails after slight straining. If the pre-deformation is higher, the strength is far beyond the ultimate tensile strength of RT and only after short recovery times does immediate fracture at yielding occur (Fig. 6(b)). However, after 1 week of RT recovery the stress level is below the tensile test curve at RT (Fig. 12) and additional straining becomes possible. The same problem occurs in alloy EN AW 5182 O. The ultimate tensile strength at RT is reached after 10% pre-deformation, but the LN<sub>2</sub>-to-RT stress drop within the first 5 min of recovery is greater. Hence, immediate RT straining after 10% deformation at 77 K is possible.

We now consider the influence of the strain rate on the deformation or failure behaviour at post-cryogenic RT forming. In Fig. 6(a) the sample with 10% pre-deformation at 77 K and 5 min recovery at RT is already close to immediate fracture with roughly 2% plastic deformation and exhibits no net strain hardening. Varying the strain rate to lower and higher values reveals unusual behaviour, as shown in Fig. 8. At a low strain rate of  $0.0008 \text{ s}^{-1}$  the sample displays immediate fracture, but a high strain rate of  $0.08 \text{ s}^{-1}$  enables further plastic elongation. If softening outweighs strain hardening an unstable situation and failure will occur, otherwise further elongation could be realized.

The interplay between strain hardening and softening is schematically displayed in Fig. 13. The solid and dash-dot lines represent the strain hardening of alloys EN AW 6016 T4 and EN AW 5182 O at RT, which can be assumed to be independent of the strain rate [70,71]. The solid squares indicate the recovery-related softening which occurs during the time required to deform the sample by the amount  $\Delta\varphi$  in the RT tensile test. If a low strain rate is applied a longer test time is required to achieve the plastic deformation  $\Delta\varphi$ , which accompanies increased softening. Such a situation will result in immediate fracture. The opposite is true for high strain rates, where only a little softening can take place because of limited time. In this case, hardening outweighs softening. Finally, in Fig. 13 all forming procedures with a softening component above the strain hardening curve will result in an unstable situation, i.e. catastrophic failure, while softening conditions below the strain hardening curve could generate maintained forming capacity. In Fig. 13 the softening points for 20% pre-deformation at 77 K are shifted to slightly higher values compared to those of 10% pre-deformation. This takes into account the higher recovery/softening rate of the 20%-deformed structure (compare the LN<sub>2</sub>-to-RT stress drop values in Fig. 6 and Fig. 7). Thus, higher deformed structures are more susceptible to failure during RT deformation for two reasons: strain hardening



**Fig. 13.** Schematic illustration of strain hardening and softening over strain for the alloys tested. Strain hardening (solid and dash-dot lines) and recovery-related softening occurring during post-cryogenic RT deformation at different strain rates (points). Stable situations are below, unstable above the strain hardening lines.

decreases, and softening increases. For alloy EN AW 5182 O the curve displays a higher strain hardening level, i.e. it is shifted upwards compared to EN AW 6016 T4 in Fig. 13. Consequently, this alloy is slightly less susceptible to sudden failure at RT forming after cryogenic pre-deformation.

## 5. Conclusion

This work investigates the room temperature recovery behaviour of cryogenically deformed Al-Mg and Al-Mg-Si alloys and thus clarifies to what extent the room temperature storage influences the mechanical and microstructural properties of the samples. Tensile test curves interrupted and recovered at room temperature show that the recovery process can be divided into three stages.

- Initial recovery processes occur due to untangling of dislocations and dislocation rearrangement (stage (i) of recovery). The length of this phase is dependent on the alloying system and the amount of pre-deformation.
- In the second stage of recovery (ii), a major stress drop and the main reduction of dislocation density occurs. In EN AW 6016 T4 this is accompanied by a rearrangement of dislocations, which is stable with further deformation.
- The third stage of recovery (iii) is marked by a lesser decline in strength and dislocation density which is comparable to room temperature recovery.

Room temperature tensile tests after cryogenic deformation can result in immediate fracture. Sudden failure occurs mainly when the post-cryogenic RT deformation is performed soon after cryogenic forming and at low strain rates. In this case, recovery-related softening dominates strain hardening, which results in an undesirable unstable situation.

## 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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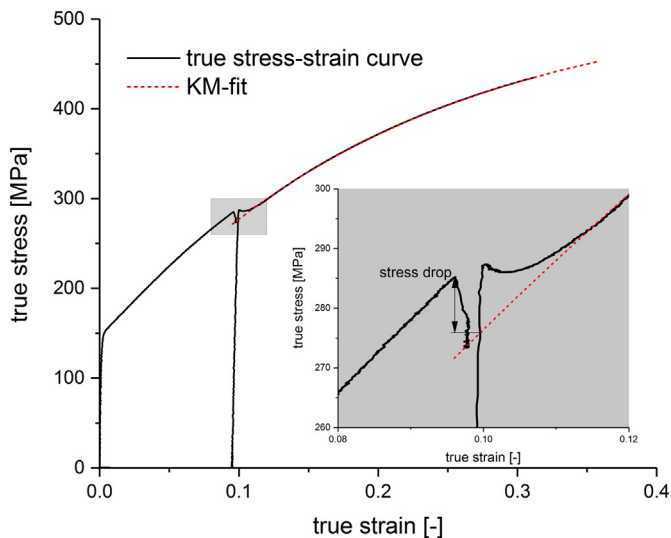
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## Data availability

The raw/processed data required to reproduce these findings cannot be shared at this time due to technical or time limitations.

## Appendix A

As discussed in Section 3.2 it is difficult to evaluate the actual stress drop from Fig. 3 and Fig. 4 due to stress plateaus and yield points followed by a stress decrease evolution. To bypass the flow onset phenomenon, the tensile testing curves were converted into true stress/strain curves. The flow curves after the RT recovery were fitted with a Kocks-Mecking type function. By back-extrapolation of the KM-fit to the flow onset of the recovered flow curve the difference between the stress before and after recovery (stress drop) could be evaluated. A graphical outline to illustrate the process is displayed in Fig. 14.



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