On the impact of post weld heat treatment on the microstructure and mechanical properties of creep resistant 2.25Cr–1Mo–0.25V weld metal

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ABSTRACT
Creep resistant low-alloyed 2.25Cr-1Mo-0.25V steel is typically applied in hydrogen bearing heavy wall pressure vessels in the chemical and petrochemical industry. For this purpose, the steel is often joined via submerged-arc welding. In order to increase the reactors efficiency via higher operating temperatures and pressures, the industry demands for improved strength and toughness of the steel plates and weldments at elevated temperatures. This study investigates the influence of the post weld heat treatment (PWHT) on the microstructure and mechanical properties of 2.25Cr-1Mo-0.25V multi-layer weld metal aiming to describe the underlying microstructure-property relationships. Apart from tensile, Charpy impact and stress rupture testing, micro-hardness mappings were performed and changes in the dislocation structure as well as alterations of the MX carbonitrides were analysed by means of high resolution methods. A longer PWHT-time was found to decrease the stress rupture time of the weld metal and increase the impact energy at the same time. In addition, a longer duration of PWHT causes a reduction of strength and an increase of the weld metals ductility. Though the overall hardness of the weld metal is decreased with longer duration of PWHT, PWHT-times of more than 12 h lead to an enhanced temper resistance of the heat-affected zones (HAZs) in-between the weld beads of the multi-layer weld metal. This is linked to several influencing factors such as reaustenitization and stress relief in the course of multi-layer welding, a higher fraction of larger carbides and a smaller grain size in the HAZs within the multi-layer weld metal.

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Introduction

The creep resistant steel 2.25Cr–1Mo–0.25V was introduced as a further development of the conventional alloy 2.25Cr–1Mo in the early 1990s. Compared to its predecessor, the V-modified version possesses several advantages such as increased strength at elevated temperatures, improved resistance to hydrogen attack and temper embrittlement as well as sufficient toughness [1–5]. On account of these benefits, 2.25Cr–1Mo–0.25V is often used for heavy wall pressure vessels for high temperature hydrogen service in power stations as well as in the petroleum and chemical industry, e.g. for hydrocracking reactors [2, 6, 7]. For this purpose, the 2.25Cr–1Mo–0.25V steel plates are usually joined via multi-layer submerged-arc welding (SAW) [6, 8]. For the application in hydrogen bearing reactors, the 2.25Cr–1Mo–0.25V steel and in particular the numerous weld seams have to withstand combinations of high pressures and temperatures of more than 400 °C. Consequently, high temperature strength and toughness are required and have to be maintained over several years [8].

In steels exposed to creep environment, solid solution and precipitation hardening are the only strengthening mechanisms which are able to effectively hinder dislocation movement [9–12]. Therefore, in ferritic and martensitic creep resistant steels, especially the distribution, size and stability of fine precipitates are of utmost importance for well-designed mechanical properties at high temperatures [12–15]. These fine precipitates are usually carbonitrides formed by micro-alloying elements such as V, Nb and Ti and consist of a cubic crystal structure of type MX [16–20]. As long as they are small in size and homogeneously distributed within the matrix, they act as obstacles for dislocations. On account of their small size, they cannot be cut or bypassed by dislocations and therefore increase the creep strength by decelerating dislocation creep [10, 11]. However, the fine carbonitrides are only effective as dislocation barriers if they are stable at high temperatures [20–22].

As 2.25Cr–1Mo–0.25 V weld metal tends to strongly harden during welding and in general exhibits low as-welded toughness, it is usually subjected to a post weld heat treatment (PWHT) [2, 6, 23–25]. Common PWHT-temperatures are 705 ± 14 °C and the advised minimum PWHT-time is 8 h, as described in API recommended practice 934-A [23]. In general, the PWHT has two main functions which are the softening of the weld metal and the relief of stresses resulting from welding [26, 27]. In addition, the PWHT stabilizes the weld metals microstructure by carbide and carbonitride precipitation and improves its properties for the subsequent application under creep loading conditions [27, 28]. Notwithstanding that many researchers [2, 4, 6, 7, 29–35] have addressed the mechanical properties of low-alloyed CrMoV-steel in various conditions, little attention has been paid on the weld metal and the combined impact of multi-layer welding and PWHT-time on its mechanical properties and the underlying correlation with its sub-microstructure. Furthermore, even though it was described for a CrMoV-alloy with a high Cr-content in the as-welded condition by Liu et al. [36], the inhomogeneous micro-hardness distribution over the multi-layer weld metals cross section and especially its alteration with longer duration of PWHT was not thematised before. However, it has been shown that this inhomogeneous microstructure of the multi-layer weld metal caused by locally different temperature profiles is vitally important for its creep resistance [34–36].

This study aims to comprehensively describe the microstructure-property relationships of 2.25Cr–1Mo–0.25V multi-layer weld metal by analysing and comparing the microstructure and mechanical properties after various times of PWHT. The weld metals tensile properties, impact energy and stress rupture time after various durations of PWHT are correlated with the observed changes of the dislocation structure and nanosized precipitates. With regard to the application in creep environment, particular focus is placed on a balanced ratio of toughness and creep resistance.

In addition, this study intends to explain the alteration of the micro-hardness over the multi-layer weld metals cross section and the local enhancement of temper resistance with increasing PWHT-time for the first time. By discussing influencing factors from multi-layer welding as well as changes in grain size and carbide area fraction, potential causes for the local differences in temper resistance are established.
Material and methods

2.25Cr–1Mo–0.25V multi-layer weld metal

The investigated 2.25Cr–1Mo–0.25V all-weld metal specimens were produced by single wire multi-layer SAW. Two 24 mm thick 2.25Cr–1Mo–0.25V steel plates were joined with a root gap of 22 mm and a bevel angle of 0° using a 2.25Cr–1Mo backing strip. The weld metal consisted of nine layers in total with two weld beads per layer. The backing strip and the diluted area were removed from the weld metal before further investigations were conducted.

Table 1 shows the chemical composition of the 2.25Cr–1Mo–0.25V all-weld metal and Table 2 provides the welding parameters in detail.

Heat treatment

For the HV1 hardness mappings and the microstructural investigation, the all-weld metal was cut into pieces of approximately 20 mm thickness. A PWHT at 705 °C for different times between 0.5 and 100 h was performed in a laboratory scale Nabertherm N 11/HR box furnace followed by air cooling. Previously to the PWHT, the whole weld seam was subjected to a dehydrogenation treatment at 250 °C for 10 h to remove diffusible hydrogen. Specimens which only experienced this dehydrogenation treatment are referred to as-welded.

Table 1 Chemical composition of the 2.25Cr–1Mo–0.25 weld metal in wt%  

| Element | Fe | Cr | Mn | Mo | V | C | Si | Nb |
|---------|----|----|----|----|---|---|----|----|
| Value bal | 2.20 | 1.00 | 1.00 | 0.30 | 0.08 | 0.08 | 0.01 |

Table 2 Overview of the single-wire SAW parameters  

| Polarity | Welding position | Welding sequence | Travel speed | Welding current | Welding voltage | Preheat temperature | Interpass temperature |
|----------|------------------|------------------|--------------|-----------------|-----------------|---------------------|-----------------------|
| Alternating current | PA | left ↔ right alternating | 50 cm/ min | 550 A | 30 V | 200 °C | 230 °C |

Overview of experimental conditions

Table 3 provides an overview of the experimental conditions of the specimens for mechanical testing and microstructural investigation.

Mechanical testing

Mechanical testing of the 2.25Cr–1Mo–0.25V multi-layer weld metal was conducted on samples which were post weld heat treated for 8, 16, 32 and 100 h at 705 °C. From each of the PWHT conditions one tensile testing sample according to DIN EN ISO 5178:2019 [37] and five Charpy V-notch impact testing samples according to DIN EN ISO 148–1:2016 [38] were produced. The tensile testing samples exhibited an initial gauge length of 50 mm and an inner diameter of 10 mm and the Charpy V-notch samples exhibited a quadratic cross section of 10 × 10 mm² and a total length of 55 mm. The Charpy V-notch specimens were cross weld specimens and the tensile sample were all-weld metal samples taken in welding direction. The V-notch of the Charpy specimens was located in the middle of the weld metals cross section, where the weld beads overlap each other, respectively, as described in DIN EN ISO 9016:2020 [39]. The Charpy impact testing was performed at a temperature of −30 °C and the tensile testing was performed at room temperature using a strain rate of 20 MPa/s. Furthermore, uniaxial stress rupture tests of the weld metal heat treated for 8, 16, 32 and 100 h at 705 °C were carried out at a testing temperature of 540 °C under an applied constant load of 210 MPa according to the ASME Boiler and Pressure Vessel Code Section VIII Division 2:2019 [40] and ASTM E139-11(2018) [41]. The stress rupture test samples were taken in welding direction from the middle of the weld metals cross section and exhibited a gauge length of 62 mm and an inner sample diameter of 13 mm. In addition, Vickers hardness mappings with a load of 9.807 N (HV1) were performed on half of the weld metals cross section, respectively, utilizing
an automatic hardness tester Qness Q 60 A + . The HV1 indents were placed with a measurement grid with a distance of 500 μm in horizontal and vertical direction.

**Scanning electron microscopy**

Scanning electron microscopic (SEM) images were taken with a FEI Versa 3D DualBeam workstation at two positions within the cross section of the 2.25Cr–1Mo–0.25V weld metal after varying PWHT-times at 705 °C, respectively. More precisely, ten backscattered electron (BSE) SEM images at a magnification of 2500 were taken within the last deposited weld bead and in the heat-affected zone (HAZ) between the last and the subjacent weld bead, see Fig. 1.

The evaluation of the carbide fraction, relative to the total investigated area of the weld metal cross section, was conducted with an Olympus Stream Motion image processing software.

Furthermore, scanning transmission electron microscopic (STEM) images were taken on electrolytically prepared transmission electron microscopic (TEM) specimens from the upper layer of the weld metal in the as-welded condition and after a PWHT-time of 8, 16, 32 and 100 h. Therefore, lamellas with a thickness of 150 to 200 μm were cut in welding direction followed by mechanical thinning to a thickness of 90 μm. The lamellas were then cut to discs with a diameter of 3 mm which were further thinned utilizing a combination of electro- and ion-polishing. The electro-polishing was conducted with an electro polisher TenuPol-5 from Struers utilizing Struers A2-I electrolyte. The polishing temperature was around –5 °C, the voltage was 34–39 V and the flow rate was 14. The final ion polishing step was conducted in a GATAN PIPS II precision ion polishing system at a voltage of 4 keV with a thinning velocity of 2 rpm for approximately 10 min. The electro-polished TEM specimens were investigated in a FEI Versa 3D DualBeam workstation using a FEI STEM detector. The SEM was operated in standard mode and the voltage was set to 30 kV.

**Electron backscatter diffraction and transmission Kikuchi diffraction**

All electron backscatter diffraction (EBSD) and Transmission Kikuchi diffraction (TKD) measurements were taken on a FEI Versa 3D DualBeam

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**Table 3 Specimen conditions for mechanical testing and microstructural investigation**

| PWHT-time (h) | Hardness mapping | Mechanical testing | SEM | EBSD | STEM | TKD |
|---------------|------------------|-------------------|-----|------|------|-----|
| 0             | x                | x                 |     | x    |      |     |
| 0.5           | x                |                   |     |      |      |     |
| 2             | x                |                   |     |      |      |     |
| 4             | x                |                   |     |      |      |     |
| 6             | x                |                   |     |      |      |     |
| 8             | x                | x                 |     | x    |      | x   |
| 10            | x                |                   |     |      | x    |     |
| 12            | x                |                   |     |      |      |     |
| 16            | x                | x                 |     | x    | x    |     |
| 20            | x                |                   |     |      |      |     |
| 24            | x                |                   |     |      |      |     |
| 32            | x                | x                 |     | x    | x    |     |
| 100           | x                | x                 |     | x    | x    |     |

**Figure 1** Stereo-microscopic image of the macro-etched cross section of the 2.25Cr–1Mo–0.25V weld metal, displaying the positions where the SEM images were taken.
workstation equipped with an EDAX Hikari XP EBSD system. EBSD grain size analysis was performed on the specimen in the as-welded condition and the specimen with a PWHT of 10 h at 705 °C. One EBSD scan was taken in the last deposited weld bead and one in the HAZ between the last and the subjacent weld bead, respectively. The specimen tilt was 70° and the working distance was between 10 and 15 mm. A voltage of 30 kV was used to map an area of approximately 180 × 180 μm² with a step size of 0.3 μm and 6 × 6 binning. The EBSD maps were evaluated with OIM 7 data analysis software. A reconstruction of the prior austenite grain structure was performed with the program ARPGE by Cayron [42] using Nishiyama–Wasserman orientation relationship.

TKD measurements were conducted on carbon replicas which were prepared from the weld metals cross section according to the method described by Bhattacharya et al. [43]. A specimen tilt of 0°, a working distance of 10 mm and a voltage of 30 kV was chosen. For a more detailed description of TKD, the reader is referred to Sneddon et al. [44]. The Kikuchi pattern obtained from the TKD analysis were evaluated using EDAX TEAM software.

Results

Mechanical properties

The results from tensile testing are presented in Fig. 2. Both, ductility and strength of the multi-layer weld metal significantly depend on the applied PWHT-time. A longer PWHT-time at 705 °C leads to an increase of the reduction of area and elongation at fracture on the one hand and a decrease of the 0.2% yield strength and the ultimate tensile strength on the other hand.

Figure 3 displays the influence of the PWHT-time at 705 °C on the 2.25Cr–1Mo–0.25V weld metals Charpy impact energy and stress rupture time. The stress rupture time is used as estimation for the resistance against creep damage and is presented as normalized value, which is the total value divided by the minimal value of the data set, respectively. For 2.25Cr–1Mo–0.25V multi-layer weld metal, the Charpy impact energy at -30 °C and the stress rupture time at 540 °C behave contrarily. The Charpy impact energy increases with longer PWHT-time whereas the stress rupture time decreases. However, this increase in Charpy impact energy and decrease in stress rupture time is considerably strong for lower PWHT-times up to 32 h. For even longer PWHT-times up to 100 h, no significant change in impact energy occurs and the decrease in stress rupture time is less pronounced compared to the shorter PWHT-times.

To assess local differences in hardness over the multi-layer weld metals cross section, HV1 hardness mappings were performed on the specimens post weld heat treated for varying times at 705 °C. Figure 4 shows the mean hardness of the last deposited weld bead and the HAZ between the last and the subjacent weld bead determined from the HV1 hardness mappings, respectively, versus the applied
PWHT-time. Additionally, three selected HV1 hardness mappings of the weld metal samples with different PWHT-times are provided for a better visualization of the hardness evolution during the PWHT.

The overall hardness of the weld metal significantly decreases with longer PWHT-times, which is in accordance with the results from tensile testing presented in Fig. 2. However, Fig. 4 clearly shows that for shorter PWHT-times the hardness difference between the last deposited weld bead and the subjacent HAZ is more pronounced than for longer PWHT-times at 705 °C. In the as-welded condition (0 h) and after short PWHT-times of 0.5, 2 and 4 h, the hardness in the last deposited weld bead is higher than in the subjacent HAZ. After a PWHT-time of 8 and 10 h, the hardness of the weld metal cross section is nearly homogeneous. After these intermediate PWHT-times, no significant differences between the hardness of the HAZs in the multi-layer weld metal and the hardness within the weld beads are visible, see micro-hardness mapping after 10 h of PWHT. Interestingly, after 12 h of PWHT, the trend changes and the hardness mappings reveal that the HAZs between the weld beads are now harder than the weld beads itself. This higher hardness of the HAZs becomes more and more pronounced with increasing PWHT-times, see HV1 hardness mapping after 100 h of PWHT in Fig. 4. As the overall hardness of the multi-layer weld metal decreases with longer PWHT-time, the higher hardness of the HAZs compared to the weld beads is considered to be an enhanced temper resistance of the HAZs. In addition, the overall difference in hardness over the weld metal cross section decreases from 150 HV1 in the as-welded condition to 50 HV1 in the 100 h post weld heat treated condition.

**Carbide fractions and grain size analysis**

To clarify the reasons for the hardness differences over the multi-layer weld metal cross section with longer PWHT-time, carbide area fraction analysis of

![Figure 4](image-url)  
**Figure 4** Mean HV1 hardness of the last deposited weld bead and the HAZ between the last and the subjacent weld bead versus the PWHT-time. Additionally, HV1 hardness mappings of the weld metal cross section after a PWHT of 0 h (as-welded condition), 10 h and 100 h are presented.
SEM images was performed. Figure 5a shows a BSE SEM image taken in the last weld bead of the specimen post weld heat treated for 100 h. In BSE mode, the carbides appear lighter and the non-metallic inclusions appear darker than the surrounding bainitic matrix. Figure 5b displays the carbide area fractions in the last deposited weld bead and in the HAZ between the last and the subjacent weld bead after various PWHT-times at 705 °C. In addition, the corresponding HV1 hardness mappings are provided. The carbide area fractions in the last deposited weld bead and in the HAZ between the last and the subjacent weld bead both increase with longer PWHT-time. Furthermore, with longer duration of PWHT, the difference of the carbide area fractions in the two investigated areas increases. The increase in carbide area fraction with longer PWTH-time is more pronounced for the HAZ between the weld beads than for the last deposited weld bead. However, it has to be noted that with this analysis only carbides which are large enough to be visualized at a magnification of 2500 in the SEM could be taken into account.

Furthermore, EBSD measurements were conducted to assess the influence of multi-layer welding on the bainite and prior austenite grain size. Figure 6a gives

![Figure 5](image_url)

**Figure 5**  
(a) BSE SEM image of a region within the last deposited weld bead of the specimen with a PWHT-time of 100 h.  
(b) Carbide area fraction of the last deposited weld bead and the HAZ between the last and the subjacent weld bead for different PWHT-times and the corresponding HV1 hardness mappings.

![Figure 6](image_url)
an overview of the morphological changes of the prior austenite grain structure across the last deposited weld bead and the HAZ between the last and the subjacent weld bead of the specimen in the as-welded condition. Figure 6b, c displays more detailed EBSD scans taken at higher magnifications in both investigated regions of the same specimen. The reconstructed prior austenite grain boundaries are highlighted with black lines, respectively. The EBSD maps show that the prior austenite grains in the last deposited weld bead are large and elongated in the direction of solidification (Fig. 6b). On the contrary, the austenite grains within the subjacent HAZ are equiaxed and considerably smaller than those within the last deposited weld bead (Fig. 6c).

The weld metal specimen with a PWHT of 10 h at 705 °C showed a similar morphology of the prior austenite grain structure, not depicted here. Comparable to Fig. 6, the prior austenite grains are elongated and larger in the last deposited weld bead and equiaxed and smaller in the HAZ between the last and the subjacent weld bead.

Table 4 displays the mean bainite grain size in the last deposited weld bead and in the HAZ between the last and the subjacent weld bead of the specimen in the as-welded condition and after a PWHT-time of 10 h at 705 °C, also determined via EBSD. For both sample conditions the bainite grain size in the last deposited weld bead is slightly larger than in the HAZ between the last and the subjacent weld bead. Furthermore, the grain size in the specimen with a

![Figure 6](image-url)
PWHT-time of 10 h is higher than that in the as-welded condition for both investigated regions.

**High resolution investigation of the microstructure**

The dislocation structure as well as the distribution, size and shape of the precipitates in the as-welded condition and after different PWHT-times at 705 °C were investigated via STEM. Figure 7a shows the microstructure in the as-welded condition. The bainitic matrix exhibits a high density of severely tangled dislocations. No ordered structures such as dislocation nets or subgrain boundaries are visible. Figure 7b–e displays the microstructure of the multi-layer weld metal heat treated for 8, 16, 32 and 100 h at 705 °C. After 8 h PWHT-time (Fig. 7b) the dislocation density is high, but reduced compared to the as-welded condition. Dislocation nets have already formed locally, though no subgrain boundaries are visible yet. The dislocations within the grains are still severely tangled and a high density of fine spherical shaped precipitates is present. These fine precipitates are mostly situated in the direct proximity of
After longer PWHT-times at 705 °C the dislocation density is further reduced and the dislocations form more and more ordered arrangements such as dislocation nets or subgrain boundaries. Furthermore, after 16 h of PWHT (Fig. 7c) small oval and also rod-like shaped precipitates of up to 50 nm in length decorate the grain and subgrain boundaries. These precipitates along the boundaries are small, but larger than the fine precipitates within the grains and increase in number and size with longer duration of PWHT. The comparison of the STEM images of the microstructure after 16 and 32 h PWHT-time (Fig. 7c, d) reveals that with the growth of the oval and rod-like precipitates at the boundaries, the amount of the much finer spherical precipitates within the grains decreases significantly. Moreover, simultaneously with the dissolution of the fine spherical precipitates within the grains, the dislocation density is further decreased. In addition, after 32 h PWHT-time nearly no tangled dislocations are visible anymore. After 100 h of PWHT (Fig. 7e), the smaller oval and rod-like precipitates at the grain and subgrain boundaries are severely coarsened and exhibit lengths of about 100 nm. Additionally, the fine precipitates within the grains have nearly disappeared completely. In comparison to the microstructure of the specimen with 32 h PWHT-time, the dislocation density after 100 h of PWHT is only slightly reduced. The reduction of the dislocation density is clearly more pronounced for the shorter PWHT-times. For longer PWHT-times than 32 h, the dislocation density is already very low and primarily coarsened at the expense of the smaller precipitates seems to take place. In all post weld heat treated specimens, the grain boundaries are decorated with larger precipitates of maximum 500 nm in length. They exhibit various shapes from oval to rectangular to blocky or even spherical. In many cases there is an overlap of precipitates which makes it difficult to determine their exact shape and size. However, no significant changes regarding the appearance of these larger precipitates was detected in the specimens with increasing PWHT-time.

TKD was performed on carbon replicas of the weld metal sample with a PWHT-time of 100 h. In this PWHT condition, the smaller precipitates at the grain and subgrain boundaries are severely coarsened and therefore easier to identify. Figure 8 shows the results of the TKD analysis of these coarsened oval and rod-like shaped precipitates. The extraction from the matrix via carbon replica method prevented overlapping of the Kikuchi patterns of the precipitates with the Kikuchi patterns of the matrix. Therefore, it was possible to unambiguously identify the crystal structure of the oval and rod-shaped precipitates as type MX. In Fig. 8 the TKD Kikuchi patterns of four MX precipitates and the patterns superimposed with the corresponding planes of the cubic MX crystal are shown. The carbides 2 to 4, exhibit relatively similar orientations to the incident electron beam.

**Discussion**

**Correlation between PWHT-time, mechanical properties and microstructure**

The mechanical testing of the 2.25Cr–1Mo–0.25V weld metal revealed that the impact energy and stress rupture time behave contrarily depending on the PWHT-time. To achieve a beneficial combination of toughness and creep resistance, the PWHT-time has to be long enough to ensure higher toughness on the one hand, but as short as possible to achieve a high resistance against creep deformation on the other hand. For a PWHT temperature of 705 °C, an annealing time of 16 h provides a good compromise of a well-balanced relation of impact energy and stress rupture time. In addition, strength and ductility are comparatively high after this duration of PWHT. Annealing times of 32 h and more lead to shorter stress rupture times but do not further enhance the weld metals impact toughness.

The changes in mechanical properties depending on the PWHT-time can be traced back to microstructural changes such as recovery processes of the dislocation structure and Ostwald ripening [20, 45] of fine MX carbonitrides. As soon as the equilibrium amount of MX carbonitrides is reached at a certain temperature, coarsening of the larger particles takes place at the expense of the smaller ones [46]. The STEM analysis revealed that an increasing PWHT-time at 705 °C causes a significant reduction of the dislocation density compared to the as-welded state. Furthermore, the dislocations rearrange to dislocation nets and in further consequence form subgrain boundaries. This is linked to the coarsening of fine MX carbonitrides with increasing time at 705 °C. Due to the larger carbonitride size after longer PWHT-times, their ability to stabilize the subgrain
microstructure is significantly reduced, as also observed in other CrMoV-steels [47–50]. While the MX carbonitrides effectively pin dislocations after 8 and 16 h, they seem to lose their pinning force after longer PWHT-times. After prolonged PWHT-times, severe coarsening of the MX precipitates occurs and the remaining precipitates are mostly situated along grain and subgrain boundaries. Simultaneously, the smaller spherical MX carbonitrides within the grains nearly completely dissolve. The larger MX carbonitrides at the boundaries grow at the expense of the smaller ones within the grains. As these larger MX precipitates are not able to hinder the recovery process and the accompanying dissolution of dislocations effectively, the dislocation density is strongly reduced with increasing PWHT-time. Furthermore, the coarsening leads to a higher inter-particle spacing of the MX carbonitrides. Thus, the Orowan stress is decreased and the dislocations can move more easily when load is applied, which manifests as a decrease in strength and stress rupture time as well as an increase in ductility.

**Factors influencing the weld metals hardness distribution**

The lower hardness of the HAZs in-between the weld beads in the as-welded condition and up to a PWHT-time of 8 h is expected to be caused by reheating to the austenitic phase field during multi-layer welding. This reaustenitization causes smaller equiaxed prior precipitates obtained by TKD and the superimposed solution for MX crystals are displayed below.

**Figure 8** SEM image of small oval and rod-like shaped precipitates on a carbon replica of the specimen with a PWHT-time of 100 h. The corresponding Kikuchi patterns of the four carbides 1, 2, 3, and 4 are shown.
austenite grains and also a smaller bainite grain size compared to the regions within the weld beads, where the temperature increase by reheating during multi-layer welding was less pronounced. Moreover, the local reaustenitization reduces residual stresses from welding which leads to a lower hardness of the HAZs in-between the weld beads. Similar inhomogeneities in terms of microstructure and hardness over the multi-layer weld metals cross section are also described by Liu et al. for high-Cr CrMoV-steel in the as-welded condition [36].

After 8 and 10 h of PWHT, the weld metals hardness is nearly homogeneous. This is assumed to result from recovery processes of the microstructure. All regions within the multi-layer weld metal soften and the hardness of the weld beads assimilates with the hardness of the HAZs in-between. As the overall dislocation density is reduced with longer duration of PWHT, the local differences in hardness are minimized leading to a relatively homogeneous hardness.

In contrast, longer PWHT-times of 12 h and more again lead to a non-homogenous hardness of the weld metals cross section. The higher hardness of the HAZs in-between the weld beads correlates with a higher amount of larger carbides in these regions of
the multi-layer weld metal. As only the larger carbide types which were visible at a magnification of 2500 in the SEM could be analysed, they were most likely M$_{23}$C$_6$ and M$_7$C$_3$ carbides [31, 51–54]. The carbide area fraction analysis revealed that the amount of the larger carbides is higher in the HAZ between the last and the subjacent weld bead than within the last deposited weld bead and that this difference in carbide area fraction increases with longer PWHT-time. This increasing difference in carbide area fraction is in consistence with the higher hardness of the HAZs in the multi-layer weld metal compared to the weld beads. Like the difference in carbide area fraction, also the difference in hardness becomes more pronounced with increasing PWHT-time. However, the higher hardness of the HAZs within the multi-layer weld metal is not a secondary hardening effect indeed. The HAZs exhibit an enhanced temper resistance and soften less than the regions within the weld beads. The overall hardness of the weld metal still decreases with longer PWHT-time due to the recovery of the microstructure and consequently due to a reduction in dislocation density. Figure 9 provides a schematic overview aiming to visualize the discussed changes of the 2.25Cr–1Mo–0.25V weld metal microstructure and the correlation with its micro-hardness distribution with increasing duration of PWHT.

Except from the enhanced temper resistance of the HAZs in-between the weld beads, also a gradient in hardness from the top to the bottom weld layer appears in all PWHT conditions. This hardness gradient is referred to the tempering treatment by the overlying weld beads in the course of multi-layer welding. The weld beads in the lower layers are reheated multiple more times than the weld beads in the upper layers of the weld metal.

Finally, it has to be emphasised that the enhanced temper resistance of the HAZs in-between the weld beads of the 2.25Cr–1Mo–0.25V multi-layer weld metal is most probably caused by multiple influencing factors. These include local variations of the temperature profile during multi-layer welding, a varying prior austenite and bainite grain size, segregations of C and carbide forming elements as well as local differences in carbide area fraction.

### Conclusions

The present study sheds light on the impact of PWHT-time at 705 °C on the microstructure and mechanical properties of 2.25Cr–1Mo–0.25V multi-layer weld metal fabricated by SAW. Based on the results from mechanical testing and the high resolution investigation of the dislocation structure and precipitates by means of STEM and TKD, the following conclusions can be drawn:

- Toughness and creep resistance of the 2.25Cr–1Mo–0.25V multi-layer weld metal behave contrarily: A longer PWHT-time at 705 °C leads to an increase of the impact energy and a reduction of the stress rupture time.
- The loss of stress rupture time with longer duration of PWHT is caused by recovery processes of the dislocation structure by enforced severe coarsening of fine MX carbonitrides.
- A PWHT-time of 16 h at 705 °C leads to a beneficial combination of Charpy impact energy and stress rupture time.
- A PWHT of 12 h and more causes an enhanced temper resistance of the HAZs in-between the weld beads, which becomes more pronounced with increasing PWHT-time.
- The enhanced temper resistance of the HAZs in-between the weld beads of the multi-layer weld metal is assumed to be linked to multiple influencing factors. These influencing factors include:
  - A smaller prior austenite and slightly smaller bainite grain size in the HAZs in-between the weld beads than in the regions within the weld beads caused by the reaustenitization in the course of multi-layer welding.
  - A higher amount of larger carbides in the HAZs in-between the weld beads compared to the regions within the weld beads.
  - An increasing difference of the amount of larger carbides between the HAZs in-between the weld beads and the regions within the weld beads with longer duration of PWHT.
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Declarations

Conflict of interest The authors declare that they have no conflict of interest.

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