Distortion and residual stress measurements of induction hardened AISI 4340 discs

J. Yi, M. Gharghouri, P. Bocher, M. Medraj

ABSTRACT

Induction hardened discs with two initial hardness levels were used for exploring the influences of the variation of initial hardness as well as induction hardening (IH) recipes on the heat treatment distortions and hardening depth. The results show that for the same initial hardness, the larger the energy input, the higher the distortion size as well as the hardening depth. For a given induction hardening recipe, the increase in initial hardness leads to a deeper hardening depth but a smaller distortion. One disc was selected for the residual stress investigation in three orthogonal directions by neutron diffraction (ND). The corresponding stress-free lattice spacing \(d_0\) was measured from the same material using both ND and X-ray diffraction (XRD) methods. The ND results show that the variation of \(d_0\) in the hardened layer is significant and should be taken into account for stress calculation. However, regarding the core region, the \(d_0\) value measured by XRD is more reliable. Accordingly, a combination of the ND-measured \(d_0\) profiles in the hardened layer and the XRD-measured \(d_0\) value in the core was adopted for the determination of residual stress distributions.

1. Introduction

High performance mechanical components used for aerospace applications require high strength, high hardness and satisfactory toughness. In order to meet these requirements, a variety of heat treatments, especially surface hardening techniques, are employed. In recent years, induction hardening (IH) has attracted considerable attention because it provides significant improvements in fatigue strength of the mechanical parts. Moreover, IH treatment can be performed very quickly and it is an environmentally friendly process.

In IH, the work-piece is first heated quickly using an oscillating electromagnetic field, and then quenched to room temperature [1]. Only material at the surface of the work-piece undergoes phase transformation, resulting in a hardened surface layer. The core material maintains a high toughness. The treatment results in compressive stresses at the surface, balanced by tensile stresses in the bulk of the component, which results in improved fatigue performance.

In the past ten years, many researchers have studied the correlation between IH process parameters and the resulting residual stress distributions [2,3]. The distortions associated with the IH treatment have also been studied [4].

X-ray diffraction (XRD) and neutron diffraction (ND) are non-destructive techniques widely used in residual stress examination. XRD is a powerful tool for near-surface stress measurement, but its low penetration depth makes it hard to determine the residual stress profile in the bulk. By alternately removing...
material and performing XRD measurements, it is possible to probe the residual stress to depths of several millimeters. However, this technique depends on a stress relaxation correction (usually based on the model given by Moore and Evan in 1958 [8]) which leads to certain uncertainties. These uncertainties have been found to increase significantly with increasing depth. For example, Coupard et al. [9] claimed that the Moore and Evan correction only worked well when the whole removed depths lower than one-tenth of the outside diameter of the disc-shaped sample. For greater depths, a more reliable experimental investigation is required. Accordingly, in the present study, ND was adopted to determine the bulk residual stress profile due to its strong penetration in most engineering materials (maximum 25 mm penetration depth for steel [7,10,11]).

Although many studies of the residual stress and distortions associated with heat treatments have been performed in recent years, an exploration of the correlation between the IH process parameters and their effects on distortions and residual stresses is still needed. This is the purpose of the present study.

The difficulties associated with surface residual stress measurements after various surface treatments have attracted considerable attention. For example, Hutchings et al. [11] and Prevey [12] pointed out that a significant stress gradient in the ND or XRD sampling volume will affect the stress results. Besides, in measuring residual strain by neutron or by X-ray diffraction, it is critical to obtain accurate measurements of the stress-free lattice parameter, \( d_0 \). This is because even a small change in \( d_0 \) can result in significant changes in the corresponding stress value.

Generally, IH parts exhibit severe hardness variations from the surface to the bulk, due to the changes in microstructure which resulted from the phase transformation. This difference in microstructure can in turn lead to considerable variations in \( d_0 \) values at corresponding positions. Therefore, the investigation of \( d_0 \) from sample’s surface to the core in IH parts is very important. By virtue of specific formula developed, Prevey and Mason [13] calculated the \( d_0 \) distribution in the normal direction as a function of depth from the axial and hoop residual stress measured from an IH sample by XRD. Their \( d_0 \) profile showed smaller \( d_0 \) values at the surface (suggesting some decarburization of the part) and some fluctuations in \( d_0 \) in the core material. Ezeilo and Webster [14] measured \( d_0 \) at the surface and in the base of a laser surface remelted specimen by neutron diffraction. For the intermediate material, a linear variation of \( d_0 \) was used. They calculated the residual stress based on this neutron-measured \( d_0 \) and compared it with that obtained by XRD and by thermal-mechanical model. They indicated the importance of applying appropriate values of the stress-free lattice spacing in neutron diffraction stress calculations, if the material under investigation has considerable variation in microstructure.

### 2. Experimental procedure

#### 2.1. Material and specimen geometry

Samples used in this study were made of AISI 4340 steel. The chemical composition of the specific alloy used is provided in Table 1 [15]. The geometry of the samples is sketched in Fig. 1. It is a 7.0 mm thick hollow disc with outside diameter of 106.1 mm and inside diameter of 13.5 mm prior to induction hardening.

Two batches of five disc samples were prepared. Group I samples had an initial hardness of 43 HRC (423 HV) and group II samples had an initial hardness of 27 HRC (279 HV).

#### 2.2. Induction surface hardening

The aim of induction hardening is to produce a hard surface layer while maintaining the initial hardness (the hardness before induction hardening) in the core. Fig. 2 contains photographs of the induction hardening process used in this work.

The inductor was a coil with a coupling distance (the gap between the inside radius of the coil and the outside radius of the disc) of 1.5 mm. All samples were induction hardened using a frequency of 200 kHz. The rotation speeds of the part during heating and quenching were 360 rpm and 240 rpm respectively. The heating time and power were varied to generate a range of IH conditions. The disc was firstly quickly heated by the current generated on the surface and then quenched by the Aqua-Quench solution (water + 12% to 12.5% polymer) sprayed from the quenching rig (Fig. 2) to reach room temperature. No additional tempering treatment was applied. Table 2 shows the selected IH parameters for the two batches.

#### 2.3. Distortion measurement

Distortion measurements were conducted on each disc sample using a Mitutoyo Bright-STRAT0 7106 coordinate measuring machine (CMM) at room temperature. The probe used in the present study was a ruby ball with a diameter of 1.5 mm. The resolution of this equipment is close to \( \pm 0.1 \mu m \) and the experimental repeatability was found to be \( \pm 0.6 \mu m \).

Variations in the coordinates of points located on the three faces of the disc before and after the IH treatment were measured. As shown in Fig. 3, for the top and bottom faces, distortion measurements were carried out from the OD (outside diameter) to 10 mm towards the center of the disc. For the lateral face, measurements were performed along the thickness of the disc. For all three faces, measurements were performed at 48 equally-spaced circumferential locations as shown in Fig. 3b. Identical measurement locations were used on the top and bottom faces, resulting in 2880 points on each face. Each measuring line comprised 60 measuring points. The spacing in the radial direction between measuring points. The spacing in the radial direction between measuring points.

### Table 1

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<td>Fe</td>
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**Fig. 1. Specimen geometry (dimensions in mm).**
2.4. Residual stress investigation

Neutron diffraction (ND) investigations were carried out on the L3 spectrometer of the Canadian Neutron Beam Centre, National Research Council, Canada. A monochromatic neutron beam was obtained using the (115) reflection of a germanium mosaic single crystal at a take-off angle of 98.77°, (\(\lambda = 1.653\)A). The neutrons' wavelength was determined using four diffraction peaks from a nickel powder standard sample. With this wavelength, the a-Fe (211) reflection occurred at a scattering angle 2\(\theta\)(112) = 89.90° which, being close to 90° provided optimal spatial resolution and avoided peak asymmetry arising as a result of axial divergence. The (211) reflection was selected for the measurements as it has been shown to be insensitive to inter-granular strains for metals with a body-centered cubic unit cell. A 32-wire based multi-wire detector spanning a scattering angle of 2.5° was used and the specimen was continuously oscillated \(\pm 0.5°\) during data acquisition to ensure that enough grains were sampled.

2.4.1. Stress-free lattice spacing determination

Stress-free lattice spacing (\(d_0\)) was obtained from measurements on a notionally stress-free reference sample, a comb produced from an extra disc that had undergone the same IH process which was applied on disc #5 (the disc selected for residual stress investigation by ND). The comb is composed of 40 (0.3 mm\(^2\)) teeth Xmm distant apart (Fig. 4).

Measurements were performed on the first 10 teeth of the comb only, along the transverse direction as indicated in Fig. 5. This is because the lattice spacing should not vary significantly beyond this 10-tooth distance (around 5.0 mm), considering the desired thickness of the hardened layer is less than 1.5 mm. The center of the sampling gauge volume (SGV) was located at tooth mid-height and in the center of each investigated tooth.

2.4.2. Residual stress determination

The ND set-up for the targeted sample (disc #5) is shown in Fig. 6. Measurements were carried out at the disc's mid-thickness at a series of locations along the radius, from a depth of approximate 0.2 mm—5.0 mm. The measuring step was 0.1 mm within the depth of 2 mm and was 0.5 mm in the following 3 mm, giving 25 measuring points in total. ND data were acquired for the hoop (X), axial (Z) and radial (Y) directions.

Fig. 7(a)–(c) represents the diffracting volume used for the radial, hoop and axial residual stress measurement in present ND study. The diffracting volume (sampling gauge volume, SGV) was of a rectangular shape: Ymm in the radial direction, Xmm in the hoop direction and Zmm in the axial direction. As shown, the ND-measured lattice strain is the averaged lattice strain over the SGV of the sample, which is decided by the width of the two beam slits (incident and diffraction beam slits) and the gauge height applied. The SGV was 0.36 mm\(^3\) (0.3 × 0.3 × 4) for radial direction and were 2.7 mm\(^3\) (3 × 3 × 0.3) for both hoop and axial directions lattice strain measurements.

2.5. Evaluation of the transformed region

The approximate size of the transformed region of all samples (two batches) was measured by a visual method. Without cutting process, samples were mechanically polished by sandpaper and then etched immediately by Nital liquid on the edges of both top and bottom faces. By virtue of optical microscope, the approximate transformed depth could be seen and measured directly. Because this method is not accurate enough (only for giving an estimated length of hardened layer), the main purpose here is to make a series comparisons among samples. The real transformed region of the hardened sample can be obtained by measuring the hardness profile of the investigated sample.

2.5.1. Micro-hardness measurement

In this study, Vickers micro-hardness test was applied only on the specimen (disc #5) used for residual stress investigation. A small piece was cut from the disc and mounted. The hardness investigation was carried out at ETS by using FUTURE-TECH FM-1 machine with the constant load of 200 g. The measuring step was 0.1 mm in the hardened and transition zones, and 0.5 mm in the non-affected region of the sample. Hardness profile was established along the cross-section (radial direction) at three positions (top-, mid- and bottom-thickness) separately, as illustrated in Fig. 8.

2.5.2. Microstructure investigation

Microstructural analysis was carried out on the cross-section of disc #5. This was performed using a Hitachi S-3400N SEM (Scanning Electron Microscopy).

3. Results and discussions

3.1. Transformed depth results

3.1.1. Transformed depth result of group I

Photographs of the transformed regions after IH for the discs in group I (with initial hardness of 43HRC) are shown in Fig. 9. The images show disc #1 to #5 top and bottom surface edges after etching. The overall hardening depth order is as follow: disc
disc #4 > disc #5 > disc #2 > disc #3 which follows the level of energy input (Table 2) (i.e. longer heating time and/or larger input power). Moreover, it should be noted that all five discs in this group display a similar discrepancy in hardening depth between the top and bottom faces (see Fig. 9). The hardening depth on bottom face is slightly larger than the top one for all cases. Although such discrepancy is relatively small and therefore the hardening depth on the top and bottom faces can be seen as similar, the possible reasons for this discrepancy need to be discussed. Firstly, this hardening depth difference probably suggests that the disc sample was not perfectly centered in the coil before performing the IH process—the disc may be positioned a little higher which makes less intense of the magnetic field at the bottom face. On the other hand, this discrepancy may indicate a non-perfect-flat coil surface. Similar results were obtained for the other group (group II) of discs.

3.1.2. Transformed depth comparison between groups I and II
A comparison of the hardening depths for groups I and II is shown in Fig. 10. It is clearly shows that for the same IH recipe, discs with a higher initial hardness of 43 HRC (423 HV) display a greater hardening depth. The results suggest that in order to obtain the same induction hardening depth, steel samples with a lower initial hardness require more heat input than do steels with a higher initial hardness. This could be due to the fact that for a given composition, the lower hardness steel has a relatively higher austenite transformation temperature according to the relevant phase transformation diagram [16]. The graph in Fig. 10 also shows that the hardening depth increases linearly with increasing input energy.

3.2. Distortion results
3.2.1. Distortion results of group I
Only distortion results for group I (initial hardness of 43 HRC (423 HV)) will be discussed here. Similar trends were observed in group II. Fig. 11(a)–(e) illustrates the dimensional changes of discs #1 to #5 in the axial (left column) and radial (right column) directions. The average hardened depth (Havg) measured from the top and bottom faces of the corresponding disc are also included for reference. The origin of the coordinate axes was located at the bottom surface of the disc and therefore Z = 0 mm and Z = 7 mm (approximately) represent the bottom and the top surfaces, respectively, as illustrated in Fig. 8.

The ΔZ profiles on the top and bottom surfaces are approximately symmetric for all samples. The axial distortion patterns are almost the same on these two faces for every disc. For all of the discs, the region close to the disc center shows no significant axial distortion (ΔZ). In contrast, near the periphery of the disc, the distortion becomes significant, showing a convex pattern, as if the materials had undergone some expansion. This could be partly due to the phase transformation at the surface hardened layer due to the induction hardening. If comparing the size (maximum of absolute ΔZ) and range of axial distortions among these five discs, it can be found that the axial distortion sequence yields: disc #1 > disc #4 > disc #5 > disc #2 > disc #3. It is not surprising to note that disc #1 has the largest while disc #3 has the smallest distortion, since they correspond to the strongest (longest heating time with highest input power) and weakest (shortest heating time with smallest input power) IH recipes, respectively.

For the radial distortion (ΔR) profiles, disc #1 has a different radial distortion profile from the other discs and distinctly large...
radial distortion, with $\Delta R$ in the range from approximately $-0.06$ mm to $0.10$ mm. It displays two noticeable protuberance patterns at each side. This is consistent with its extraordinary large axial distortions measured on top and bottom faces. In fact, during the induction heating step of disc #1, obvious sparks were observed on the edge of the disc, which suggests the input energy is large enough to melt the sample at the edge. Therefore, the particular shape of the axial and radial distortions of disc #1 can be attributed to the strongest IH recipe applied. As for the other four discs, the radial distortion profiles demonstrate a similar convex pattern over the whole disc thickness ($Z$), with the maximum radius nearby the mid-thickness of the disc and the minimum radius at the upper or lower part of the lateral face display as shown in Fig. 11(b)–(e).

3.2.2. Distortion comparison between group I and II

Distortion as a function of input energy is plotted for both groups of samples in Fig. 12. Samples from both groups show a similar trend but distortions for group I are systematically higher than for group II. Therefore, it can be concluded that, under the same IH treatment, samples with lower initial hardness undergo more distortion than those with higher hardness. This result is in contrast to the hardening depth observation between these two groups, where the discs with 27 HRC (279 HV) initial hardness demonstrated smaller hardening depth than those with 43 HRC (423 HV). If distortions were only due to the transformation process, one would expect that the more regions transformed, the larger the resulting distortions. However, the comparison between the results from group I and group II show a different tendency. This means the phase transformation during the IH treatment is not the only factor that can affect the distortion results.

3.3. Residual stress results

3.3.1. Stress-free lattice spacing $d_0$ results

Fig. 13 shows the stress-free lattice spacing $d_0$ profiles in the hoop and radial directions measured by ND on the reference (comb) sample. It is compared with the two $d_0$ values in radial direction measured from two thin (0.07 mm) layers of a similar sample by XRD.

The ND-measured $d_0$ in the hoop direction firstly demonstrates an increase near the surface, reaching a maximum of $1.1729 \pm 0.00016$ followed by a diminishment with increasing depth through the hardened case and finally approaching a constant value of approximately $1.1715 \pm 0.00009$ at a depth of around 2.0 mm. For the ND-measured $d_0$ in the radial direction, there is only a small variation in the stress-free lattice spacing near the surface followed by a smooth decrease towards the center and...
reaching a constant value of approximately 1.1718 Å ± 0.00013 at 2.2 mm.

Clearly, the ND-measured $d_0$ values in both directions (the hoop and radial directions) present a relatively higher value close to the surface than towards the core. This can be explained as the presence of carbon in solid solution in the martensite lattice leading to the increase of atomic spacing in the hardened layer. In the core region, since the material does not undergo any phase transformation during IH treatment, it keeps the original microstructure, the unstable tempered martensite. Therefore, the $d_0$ value in the core is relatively smaller than that in the hardened zone and nearly keeps a constant. Regarding the transition area, which has a mixed microstructure of fresh and over-tempered martensite, the corresponding $d_0$ value is consequently in between the one in the hardened and in the core region.

In order to make comparison, two XRD-determined $d_0$ values, the $d_0$-surface of 1.17195Å ± 0.0001 and the $d_0$-core of 1.17079Å ± 0.0001, are indicated by two dashed lines in Fig. 13. These two $d_0$ values were measured from two thin (0.07 mm) sheets located at a depth of 0.6 mm and 20 mm below the free surface of a backup sample of disc #5. Obviously, the XRD results are in agreement with the ND outcome. The $d_0$ has a larger value in the hardened layer than in the core. A comparison was made between the two ND-determined $d_0$ profiles and two $d_0$ values measured by XRD. It can be seen that the XRD-determined $d_0$-surface is close to the ND-determined $d_0$ in the hardened layer. However, a clear discrepancy was found in the core material, where the XRD-determined $d_0$-core is significantly smaller than that measured by ND. The possible reasons for this discrepancy can be viewed from two aspects. Firstly, the reference comb sample used for ND $d_0$ determination may not be completely stress relieved, which results in inaccurate measurement. Considering that the tooth of the comb is very tiny, it is hard to perfectly machine each tooth in spite of using the wire-EDM cutting. Few teeth located in the core region were even bent to some extent, which implies a certain amount of residual stress may exist in those teeth. Secondly, there are experimental uncertainties in both ND and XRD measurements. For the $d_0$ investigation by ND, the uncertainty is mainly due to the fact that the number of counts (neutrons) detected for the diffraction peak formation is restricted by the small dimensions of the tiny teeth. This means, for each tooth, the amount of counts received by the detector may not be sufficient enough to give a very representative peak shape. Consequently, the center of the diffraction peak (used to calculate $d_0$) can vary within a small range and therefore the $d_0$ values may not be very representative at certain locations, especially the ones located in the core region where the bent teeth were observed. For the $d_0$ investigation by XRD, however, the main uncertainty is considered from the measuring system itself.

Since the $d_0$ values determined by ND display a certain amount of uncertainties, the $d_0$ investigating along the axial direction is not very necessary. The average value of $d_0$ in radial and hoop direction was therefore used as the $d_0$ value in axial direction to calculate the residual stress in corresponding direction.

### 3.3.2 Residual stress results

Fig. 14 demonstrates the residual stress distribution of the targeted disc in the hoop, axial and radial directions calculated based...
Fig. 11. IH distortion results of group I: (a) to (e) represent the axial distortion $\Delta Z$ (the left column) on the top/bottom face along radius of disc #1 to #5, and the radial distortion $\Delta R$ (the right column) on the lateral face along thickness of disc #1 to #5.
on the ND-measured \(d_0\) profiles. Hoop and radial stresses were obtained from \(d_0\) profiles in hoop and radial directions, respectively.

As can be seen from Fig. 14, a compressive residual stress field in the hoop and axial directions exists in the area close to the surface. This is similar to the work of Coupard et al. [9] and Hornbach et al. [17]. A reversal into tensile stress happens at the radial depth of 1.2 mm from the surface in hoop direction and 1.5 mm in axial direction. The maximum compressive stress in the hoop and axial directions are of the order of \(-1120\) MPa, and they are observed at the same radial position (\(\sim 0.65\) mm) from the surface. A maximum tensile stress is found in hoop direction with an approximate value of 930 MPa. On the other hand, the axial stress is much smaller only 250 MPa. It implies that the hoop direction of the disc encountered the most critical stress condition (largest tensile stress), which is consistent with the most heat-treated cylindrical samples reported in the literature [18,19].

In addition, the radial stress is generally expected to be zero at the surface and the remaining radial (pseudo-normal) stresses are expected to be low due to the force balance requirement [20] considering the geometry of the sample. It is supported by literature which the heat-treated residual stress profiles show an almost constant value close to zero or only showing a slight variation over the whole radial direction [21]. In the present ND study, the radial stress is found to be compressive at the surface and demonstrates considerable variations along the radial positions within 2 mm from the surface. This finding is somewhat puzzling and is in contradiction with the recent results obtained by Coupard et al. [9] and Prevey et al. [13]. Only Hornbach et al. [17] reported large variation in radial residual stress for induction hardened gear using XRD. Nevertheless, this obvious variation in radial direction can be somewhat related to the noticeable experimental errors and uncertainties observed in \(d_0\) measurements (Fig. 13). Since the residual stress calculation is based on the reference \(d_0\) value, the larger the possible error and uncertainty, the higher the effects on the residual stress value. Moreover, the ND experiment itself has certain uncertainties [11], especially at the region where phase transformation occurred (i.e. the near-surface area in the present study).

It is interesting to see that after 2.5 mm from the surface, the three stress profiles gradually merge and nearly coincide at the radial depth of around 5.0 mm at \(-250\) MPa. It probably insinuates the beginning of the non-affected core region of the sample. However, the three component residual stresses should be all approaching to zero in the core material according to the principle of surface hardening stress distribution [22], which has been proved by numerous experimental works (e.g. the radial residual stress depth profile measured by ND [23] and XRD [24]). Therefore, the present constant stress value of about \(-250\) MPa in the core should also be considered as an error which could be attributed to the corresponding uncertainty and errors existed in stress-free \(d_0\) measurement such as being not 100% stress-free specimens.

If one uses the \(d_{0\text{-surface}}\) and \(d_{0\text{-core}}\) measured by XRD to recalculate the residual stress in these three directions, the overall profiles have shifted up and the results become more balanced in the core material (Fig. 15).

This observation is easy to be understood from Fig. 13 because the two XRD-determined \(d_0\) values are relatively smaller than that obtained by ND, therefore the corresponding stress profiles in Fig. 15 move upwards. However, comparing Fig. 15 with Fig. 14 in terms of core region, it seems that at the depth relatively far from the surface the \(d_{0\text{-core}}\) measured by XRD may be closer to the real stress-free lattice spacing of the sample. The most reasonable explanation for this could be that the \(d_{0\text{-core}}\) measured by XRD on the thin layer is more reliable than the \(d_0\) values obtained by ND (considering the significant uncertainty and error of \(d_0\) measurements in the core region). Nevertheless, in terms of surface and phase transition areas, although the uncertainty of ND is relatively large, the \(d_0\) profiles obtained by ND are more reliable. Since \(d_0\) values should vary with the distance due to the phase
transformation occurred during the IH surface treatment, it is much better to use the ND \( d_0 \) profile to analyze the case rather than using only one value \( d_{0\text{-surface}} \).

Consequently, a 3-dimensional residual stress distribution by a combination of applying the ND-measured \( d_0 \) profiles in the surface region and the XRD-measured \( d_{0\text{-core}} \) in the core area is given in Fig. 16. Comparing Fig. 16 with Figs. 14 and 15, it is noted that the three stress profiles shown in Fig. 16 is a combination of that in Fig. 14 (in the surface and transition area) and in Fig. 15 (in the core region). Stress profiles demonstrate a near constant value (approaching to zero) in the core material while display a large compressive residual stress value in the surface hardened layer. In present study, Fig. 16 is more reasonable and may reflect the real case.

3.4. Micro-hardness results

Micro-hardness profiles for the targeted disc after induction hardening through three positions of top-, mid- and bottom-thickness as specified earlier are shown in Fig. 17(a). The corresponding vertical cross-section view of the hardening depth contour across the thickness of the disc is shown in Fig. 17(b).

It has been observed from Fig. 17(a) that the micro-hardness distributions along radial directions measured at mid-thickness, top-thickness and bottom-thickness have similar trends. As shown, all three curves demonstrate a typically high hardness value varying from 625 HV0.2 to 700 HV0.2 near the surface. This region of high hardness is called the hardened zone and is quantified by the depth \( z_1 \) as displayed in Fig. 17. This region is followed by a sharp decrease until reaching a minimum around 320 HV0.2, indicating a transition zone (which size is \( z_2 \)). From that minimum value, the hardness gradually goes up progressively to reach the core hardness (470 HV0.2), forming the over-tempered zone (\( z_3 \)). Afterwards, the hardness keeps a nearly constant value around 470 HV0.2 in the core material, which is the same value as the initial hardness before induction hardening and therefore reveals the non-affected zone (\( z_4 \)).

The slight increasing trend in the hardness of the surface hardened layer with moving towards the center was also observed by Grum [25]. This could be explained by the formation of a very fine martensitic microstructure in the inner region of the hardened layer due to the non-uniform cooling rates in the disc surface during the IH process. The “groove” in the hardness profile, in the over-tempered zone (\( z_3 \)) is a fairly common feature when tempered martensite is subjected to IH. This is due to the tempering effect of the thermal flow happening during the IH treatment. More detailed study on the entire hardness profile especially the over-tempered zone of the induction hardened specimen was reported by Ducassy [26].

Analyzing the hardening depth at the mid-, top- and bottom-thickness, a distinct difference has been noted. In the present study, the hardening depth is defined as the radial position in region \( z_2 \) where the hardness reaches the core hardness. The value of hardening depth measured at top-thickness and bottom-thickness are very close to each other (around 1.6–1.7 mm) and they are also similar to the estimated hardening depth value obtained by image...
analysis (as described in Section 2.5, using optical microscopy). In Fig. 17(b), the red arrows indicate the hardening depth at the top-, mid- and bottom-thickness of the disc. However, they are obviously larger than the mid-thickness hardening depth, which is only 1.2 mm, leading to a clear and characteristic hardened case profile has been exposed with a convex shape towards the lateral face (Fig. 17(a)). This variation in the hardening depth at different positions has been viewed as a result of the edge effect caused by the concentration of the magnetic field at the disc edges [27].

If one compares the residual stress profile with the hardness profile (Fig. 17(a)), a sharp transition area can be observed in both profiles at a similar location, suggesting that the tendency of compressive residual stress profile is in accord with the trend of the corresponding hardness profile. This finding is further supported by the work of Grum [25] stating that a steeper reduction in hardness may contribute a sharp change of residual stress. Moreover, with respect to the radial positions after 2.5 mm, both the hardness distribution and the three residual stress profiles show a consistent trend. They actually approach to a nearly constant value towards the core around the same depth value (3 mm).

3.5. Full width half maximum (FWHM) results

The FWHM profile of the Fe{112} Kα1 diffraction peak is illustrated in Fig. 18. The relevant hardness curve is also given for better explanation. As seen, the overall FWHM profile in the hoop, axial and radial directions are quite similar. The only noticeable difference in peak width among them appears within the hardened layer. Regarding the transition zone and core material region, the three FWHM curves are very close to each other even overlapping at many radial positions. Besides, all the three FWHM profiles display a visible variation in the hardened region up to a radial depth around 1.0 mm below the surface, and such variations could be related to the likewise hardness fluctuation in surface hardened region, suggesting the inhomogeneous microstructure generated in the surface layer, possibly caused by the dislocations in the hardened layer.

Moreover, after the hardened region, the peak width declines quickly towards the center until reaching a minimum value around 0.6° at a radial depth of 1.5 mm, approximately. This is closely followed by a gradual increase to approach a nearly constant value of 1.0°, revealing the core material zone. The remarkable trough appeared in the FWHM curve is consistent with the similar groove

![Image](https://example.com/image1.png)

**Fig. 18.** Fe{112} diffraction peak FWHM distributions along radius.

![Image](https://example.com/image2.png)

**Fig. 19.** Microstructures over the cross-section of disc #5: (a) hardened region; (b) transition region; (c) over-tempered region; (d) non-affected region.

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pattern found in the hardness profile (Fig. 17(a)), which could be seen as the material over-tempering nature taking place at the end of the hardened region.

3.6. Microstructure results

The corresponding SEM microstructural analysis of the cross-section of disc #5 is given in Fig. 19. Fig. 19(a)–(d) represents the microstructure of hardened zone (z1), transition zone or otherwise called hardness loss zone (z2), over-tempered zone (z3) and non-affected core zone (z4).

It can be seen from Fig. 19(a) that the microstructure in the hardened zone is completely hard martensite. This is because the heating temperature in this area is above Ac3 and based on the fast cooling, the microstructure can be completely transformed from austenite to martensite. A mixed microstructure of hard martensite and over-tempered martensite is observed in Fig. 19(b), since the temperature in the transition zone is between Ac1 and Ac3. Fig. 19(c) displays a completely over-tempered martensite microstructure. In this region, the corresponding temperature is still high (due to the heat flow effect), but without reaching the critical point Ac1, the material is only tempered without being austenitized. It finally transforms to a more stable state (i.e., ferrite with dispersive carbides) which is associated with a loss of hardness, as can be seen from the groove in the hardness profile in Fig. 17(a). Fig. 19(d) displays the microstructure of the core of the disc. It is composed of the initial microstructure (the quenched and tempered martensite) of the sample without being affected by the IH treatment. Also, one may observe directly from Fig. 19(c) that the over-tempered martensite microstructure seems to be more disordered compared with that in the hardened and core region.

Moreover, since the tempered martensite has larger grain size than the quenched one, it can be understood that the grain size in the over-tempered region is larger than that in the core area and followed by the hardened zone. Therefore, the FWHM plot demonstrates a drop from surface to core and there is a groove in the over-tempered zone, considering the Scherrer formula (FWHM ∼ 1/d) [11].

4. Conclusions

The present study shows that both the IH parameters (heating time and input power) and the initial hardness affect the distortion and hardening depth of the samples. Under the same initial hardness, increasing IH parameters (heating time or input power) increase both the hardening depth and distortions of the disc. However, under the same IH parameters, increasing the initial hardness of the disc increases the hardening depth as it decreases the distortions of the disc.

Due to the severe variations of the hardness at the surface of an IH part, the knowledge of d0 depth distribution is necessary and it can significantly affect the residual stress results. A variation of 0.001A in d0 in hardened layer can result a difference in residual stress around 200 MPa in the same region. Both ND- and XRD-measured d0 results display a visible difference in d0 values between hardened layer and core material. Therefore, it is more reasonable to apply ND-measured d0 profile to calculate the residual stress near the surface than using a constant d0-surface value measured by XRD. However, for the core material, the XRD-measured d0-core is more reliable. The final residual stress distributions are constructed based on the combination of ND and XRD d0 results.

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References