Internal Friction Behavior Associated with Martensitic Decomposition in Low-carbon Dual-phase Steel

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(Received on November 1, 2018; accepted on February 5, 2019; J-STAGE Advance published date: May 18, 2019)

In this study, the martensitic decomposition behaviors of a low-carbon dual-phase steel were investigated by the low-frequency internal friction method, combined with various structural analysis techniques including X-ray diffraction, scanning electron microscopy, and transmission electron microscopy. Two internal friction peaks were observed at 418.4°C and 448.1°C, and were attributed to desolvation of supersaturated carbon atoms from martensite and formation of Cr3C7 precipitates, respectively. The results indicated that the two-step process during the martensitic decomposition, involving carbon atom diffusion and carbide precipitation, could be well explained by the internal friction technique. The microstructural mechanisms associated with the generation of the two internal friction peaks during the martensitic decomposition are discussed.

KEY WORDS: DP steel; internal friction; martensitic decomposition.

1. Introduction

Advanced high-strength steels, such as dual-phase (DP), transformation-induced plasticity (TRIP), and martensitic steels, have been developed and widely used in the automobile industry.1–3) Phenomena such as carbon segregation to structural defects, carbide precipitation in martensite or bainite, carbon partitioning from martensite to austenite during quenching, partitioning, and tempering involve carbon diffusion in a supersaturated matrix.4,5) However, the measurement of carbon diffusivity in steels is still very challenging.

Internal friction (IF) technique has been proved as an effective approach to study the microstructural evolution, defects characteristics, or phase transformations inside the metal materials.6–9) For steels, an IF measurement with a high sensitivity could be utilized to characterize the interstitial or dissolved carbon atoms in body-centered cubic (BCC) structural phases such as ferrite and martensite phases. Some typical internal friction peaks observed in steels include the Snoek peak, the Snoek-Köster peak10) and the spurious peak induced by reduction of background internal friction, and so on. Such internal friction characteristics have been verified related with various occupation or movement behavior of the carbon atoms. Therefore, this suggests that the IF technique can be also used to investigate other structural evolutions associated with movements of carbon atoms in steels.

Extensive studies are ongoing to understand the martensitic decomposition behaviors and associated mechanisms, which could be potentially utilized to advance the methods of precise microstructure control of steels. In this study, we investigated the IF behaviors accompanying the martensitic decomposition and carbide precipitation in DP steels, and discuss the underlying mechanisms.

2. Experimental

A low-carbon martensitic DP automobile steel was used in this study. Its chemical composition expressed by mass fractions is: 0.081% C, 0.0419% Si, 1.88% Mn, 0.0124% S, 0.03% P, and 0.278% Cr. The initial volume fraction of martensite is approximately 25%, while the other volume fraction corresponds to ferrite.

The mechanical spectrum was measured using an FTA-700 low-frequency IF tester (Kë torsion pendulum) in the free-attenuation mode. The measurements were performed at a heating rate of 2°C·min⁻¹, strain amplitude of 4.8 × 10⁻⁵, and atmosphere pressure of 6 × 10⁻² Pa. Standard specimens with thicknesses, widths, and lengths of 1, 3, and 55 mm, respectively, were cut by a slow-feeding wire-cut machine. The specimens were then abraded and polished with a metallographic abrasive paper, followed by application of 4% nitric acid/ethanol to corrode their surfaces. The attenuation degree of the specimen’s amplitude was...
used to calculate the IF using a Kê torsion pendulum. In order to improve the measurement accuracy, the system measures several free-attenuation vibration cycles and calculates the logarithmic mean value of the amplitude to determine the IF:

\[ Q^{-1} = \frac{\delta}{\pi} = 1/(\pi \cdot m) \ln(A_n / A_{n+m}), (\delta << 1) \]  \tag{1}

where \( m \) is the number of continuous vibration waves, \( A_n \) is the amplitude of the \( n^{th} \) vibration, \( A_{n+m} \) is the amplitude of the \((n+m)^{th}\) vibration, and \( \delta \) is the natural-logarithm ratio of \( A_n \) to \( A_{n+m} \).

The increase in temperature generally leads to larger distances between atoms and smaller binding forces between them, yielding a smaller modulus. The specimen’s lower end is held by the lower chuck of the torsion pendulum of the low-frequency inverted-torsional IF tester, whereas the other end of the specimen is connected to the rod of the inverted pendulum so that it can freely rotate. Therefore, the modulus recorded in the IF test reflects the specimen’s shear modulus, which is proportional to the square of the vibrational frequency of the torsion pendulum. \(^{14}\)

\[ G = \frac{8\pi I f^2}{R^2} \]  \tag{2}

Where \( I \) is the moment of inertia of the torsional pendulum, \( l \) is the specimen effective length, \( R \) is the specimen radius, approximately equal to \( \sqrt{a^2 + b^2} \) \( (a \) is the specimen’s width, \( b \) is the specimen’s thickness), and \( f \) is the resonance frequency of the torsional pendulum.

The evolutions of the thermophysical properties were measured at a heating rate of \( 2^\circ\text{C}\cdot\text{min}^{-1} \) by differential scanning calorimetry (DSC, PE DSC-8000) and thermal expansion analysis (NETZSCHDIL 402 Expedis Classic). The phase structure of the specimen was analyzed by means of X-ray diffraction (SHIMADZU XRD-7000) using Co K\( \alpha \) radiation. The scanning rate is \( 2^\circ\text{C} / \text{min} \). In addition, a Hitachi S2570 scanning electron microscope (SEM) was used to observe the metallography structure. A microstructural characterization was performed by transmission electron microscopy (TEM) with a JEOL JEM-2010F microscope operated at 200 kV. TEM samples were prepared by a twin-jet polisher operated at 30 \( ^\circ\text{C} \) and 15 V, using a solution of 5% perchloric acid and 95% ethanol (expressed in volume percentages). An MTS Exceed 64 tensile tester was used to evaluate the mechanical properties of the specimens.

3. Results and Discussion

In order to characterize the martensitic decomposition process, the thermal expansion, heat flow, and mechanical spectrum as a function of the temperature during heating of the studied low-carbon DP steel from room temperature to 500 \( ^\circ\text{C} \) at a heating rate of \( 2^\circ\text{C} / \text{min}^{-1} \) are analyzed, as shown in Fig. 1. The thermal expansion method is a typical method used to characterize phase transformations in steels. Figure 1(a) shows that the expansivity of the DP steel has a linear evolution during the heating until the temperature reached approximately 455 \( ^\circ\text{C} \), where a significant decrease is observed. The decrease in the expansivity usually indicates the occurrence of a phase change. In this case, it indicated the martensitic decomposition. DSC is also considered an effective method to study first-order phase transformations of materials by monitoring the latent heat accompanying the transformation. However, no obvious endothermic or exothermic peak can be distinguished from the DSC curve in Fig. 1(b). The mechanical spectra including the IF and shear modulus of the DP steel are shown in Fig. 1(c). Three IF peaks are observed, marked by the arrows. The peak at approximately 250 \( ^\circ\text{C} \) is believed to be the Snoek–Kê–Köster (SKK) peak, which has been well studied, attributed to movable dislocations interacting with interstitial atoms. \(^{15,16}\) The other two peaks, denoted as P1 and P2, are located in a narrow temperature range of 400 to 460 \( ^\circ\text{C} \).

Figure 2 shows the regional analysis of the temperature-dependent IF and modulus of the DP steel at the temperature range of 400 to 480 \( ^\circ\text{C} \).

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with a phase transition. The peak spectrum of \( P_2 \) is also irregular accompanied by modulus changes. The decrease in the temperature-dependent modulus deviates from the linearity; a small increase is observed during the heating beyond the critical temperature of approximately 448.1°C. The temperature is in agreement with that of the expansivity changes.

Figure 3 shows the results of the IF measurements at the vibration frequencies of 1.1 and 3.5 Hz at a heating rate of 2°C·min⁻¹. The resonant frequency was adjusted by changing the mass and length of the pendulum rod. The temperatures corresponding to the \( P_1 \) and \( P_2 \) peaks did not change upon the adjustment of the frequency, which indicates that the positions of both IF peaks are frequency-independent. Therefore, it can be concluded that the two IF peaks are not relaxation-type peaks, but associated with phase transformations and microstructural evolutions.

Figure 4 shows the IF spectra as a function of the temperature of the DP steel for two thermal cycles, in the range of room temperature to 500°C. The \( P_{SKK} \), \( P_1 \), and \( P_2 \) IF peaks observed in the first heating cycle disappeared, while the background IF spectra were significantly reduced. This indicated that the original structure of the DP steel was modified during the first cycle of heating to 500°C. The disappearance of \( P_{SKK} \) could be easily explained by the significant decrease in the dislocation density and interstitial carbon atom concentration, likely attributed to the martensitic decomposition or carbide precipitation during the thermal cycle. The reduction in the background IF may be attributed to the decrease in the density of movable dislocations.\(^{17,18}\) In addition, the disappearance of the \( P_1 \) and \( P_2 \) IF peaks indicated that the phase transformation and microstructural change during the heating in the first thermal cycle were irreversible.

Therefore, the observed temperature-dependent spectra of both IF and modulus reflect the martensitic decomposition of the DP steel. A further microstructural analysis is performed to reveal the underlying mechanism of the complex IF behaviors.

Figure 5(a) shows the XRD pattern for the DP steel specimen in the initial state and those of the specimens tempered for 20 min at 450 and 500°C. The patterns of the as-received DP steel and specimens tempered at 450 and 500°C showed the typical single BCC structure of martensite or ferrite. It is typically difficult to distinguish between martensite
and ferrite by XRD. A precise analysis of the evolution of the $\alpha$ (110) peak for the three XRD diffraction patterns is presented in Fig. 5(b). The results showed that the $\alpha$ (110) peak is observed at 44.58° for the original specimen, which decreased to 44.50° after the tempering at 450°C, followed by an increase to 44.64° upon the tempering at 500°C. According to the Bragg equation from the crystal diffraction theory, the diffraction angle is inversely proportional to the interplanar spacing. Therefore, in this study, the interplanar spacing increased upon the tempering at 450°C, and then decreased upon the tempering at 500°C.

The martensitic decomposition in steels is generally described as ordering of the carbon atoms as well as some carbide precipitation under thermal activation. For the DP steel composed of 75% ferrite and 25% martensite, the martensitic decomposition involved diffusion of carbon atoms from the carbon-rich martensite to the carbon-lean ferrite and carbide precipitation. The IF peak $P_1$ accompanied with the modulus decrease below 450°C is believed to originate from the diffusion of carbon atoms from martensite to ferrite. The non-relaxation-type IF peak is generally associated with the emergence or evolution of the interface during the phase transformation, such as the thermoelastic martensitic transformation in shape memory alloys. In this study, it may be attributed to the interface migration upon the martensitic decomposition. Meanwhile, upon the migration of the carbon atoms into ferrite, the macroscopic lattice constant increased, as shown in the XRD pattern. This led to decreases in the density and modulus.

When the specimen was heated to the higher temperature, the carbon atoms in the matrix started to desolventize and form the carbide precipitates; in addition, the number of dissolved carbon atoms in the matrix rapidly decreased. This well explains the significant decrease in the lattice constant of the specimen tempered at 500°C. With this process, we observed an irregular so-called IF peak $P_2$ accompanied with a slight increase in the modulus. However, the mechanism of the mechanical spectroscopy associated with precipitation transformation in steels or alloys is unclear yet. For carbon steels, a reasonable explanation is the presence of an internal friction athermal background that generated by the interaction of dislocations with point defects situated away from the dislocations glide plane. This background is proportional to carbon content in solid solution, and decreases due to the precipitation of carbides, producing a curve with a peak shape. In this study, the IF spectrum of the DP steel during cooling showed that the background of the IF spectrum significantly decreased after the thermal cycle. Additionally, a slight increase in the modulus accompanied this process. These results strongly support our conclusions that $P_2$ is not a peak, but a local maximum or shoulder caused by the reduction in the background of the mechanical spectrum.

A microstructural observation and mechanical analysis were carried out to provide more intuitive evidences for these phenomena.

**Figure 6** shows the morphologies of the original specimen and two tempered DP steel specimens measured by SEM. A typical metallographic structure of the martensite/ferrite DP steel was observed in the as-received specimen, in which martensite exhibits an island-like distribution in the ferrite matrix. The size of the martensite bulk is in the range of 1 to 15 μm. The clear edges and smooth surface of the martensite/ferrite interface are shown in Fig. 6(a). After the tempering at 450°C for 20 min, the morphology of the DP steel specimen was considerably changed. Although the original martensite/ferrite interface can be still resolved, the interface is not smooth and the martensite bulk is not clearly distinguishable. This indicates that the interface migration occurred during the martensitic decomposition in the DP steel tempered at 450°C, according to the observed structural evolution behaviors. As mentioned above, a phase-transformation IF peak is generally associated with interface generation or movement. This well explains that the $P_1$, IF peak emerges by the migration of the original martensite/ferrite interfaces.

**Figure 7** shows representative TEM micrographs of the DP steel specimens that tempered at 450°C (a) and 500°C (b–c) for 20 min, respectively. The area with a higher density of dislocations corresponds to martensite, while the other area corresponds to ferrite. After the tempering, a considerable number of dislocations are still observed in the martensite phase. A part of the martensite/ferrite interface was modified and became unclear and rough for both specimens, as shown in the bright-field images in Figs. 7(a) and 7(b). This further confirms the previous results about the interface migration. The diffraction pattern on the martensite phase of the specimen tempered at 450°C is marked as a single BCC phase. However, the diffraction pattern of
the specimen tempered at 500°C is different that a second phase was clearly observed. The analysis of the diffraction spots showed the presence of a second phase around the martensite/ferrite interface. A few granular features were observed in the corresponding dark-field image in Fig. 7(c). The composition of the second phase can not be intuitively characterized by EDS technique due to the ferromagnetism of the specimen. So the precipitate phase were inferred from two criteria: one is the orientation relationship between the precipitate and the martensite matrix as well as the lattice constants obtained from the diffraction spot. After calibration, the results reveal that the new phase most likely corresponds to the Cr7C3 precipitates. The second one is from the reference 23–25) in which it was mentioned that the formation temperature of the microalloyed Cr7C3 carbides in steel is around 550°C. It is close to the experimental results in our case. According to the alloy composition of the DP steel in this study, we inferred that the precipitates formed at 500°C is the Cr7C3 phase. As we checked that no precipitates could be distinguished even after aging at 450°C, it indicated that the Cr7C3 carbides precipitated during the tempering at 500°C, which is characterized by the appearance of the so-called P2 IF peak and relative increase in the modulus.

Figure 8 shows the mechanical properties of the original DP steel specimen and tempered specimens. The as-received DP steel exhibits a lower yield stress, continuous yielding, and increased strain hardening rate, which is consistent with the typical mechanical stress–strain curve of the martensite/ferrite DP steel. The DP steel usually consists of hard martensite islands embedded in a soft ferrite matrix, leading to a relatively high ultimate tensile strength (UTS), low yield–to–tensile-strength ratio, absence of yield point elongation, and good balance of strength and formability. After the specimens were tempered, the strain-dependent stress significantly changed so that a yield platform was observed followed by the strengthening segment of reinforcement. For the specimen tempered at 450°C, the yield strength was approximately 457.5 MPa, similar to that of the as-received DP steel. Upon the tempering at 500°C, the yield strength increased to approximately 543.1 MPa, which is an increase of approximately 18.7%. The yield strength of may lie on the ferrite structure, which acts as the soft phase in the DP steels. This indicates that the ferrite structure did not significantly change, although the structural characteristics of
the martensite/ferrite DP structure were destroyed and the number of solid-solution carbon atoms in the ferrite phase increased after the tempering at 450°C. After the tempering at 500°C, the carbides were precipitated in the ferrite phase and led to a significant increase in the yield strength.

In general, the evolutions of the mechanical properties of the original DP steel and tempered specimens further confirm the two processes in the martensitic decomposition: carbon atom diffusion from martensite to ferrite and carbide precipitation. These two processes are reflected by the two peaks, P1 and P2, observed in the IF spectrum of the DP steel during the heating.

4. Conclusions

This study showed that the IF technique combined with the microstructural analysis has a very unique advantage in the characterization of structural evolution or phase transitions related to the movement of carbon atoms. Compared to the traditional testing technologies, such as thermal analysis (DSC) and thermal expansion, the IF measurement could provide more information for the characterization of the martensitic decomposition of the DP steel. It will be of significance to further develop the spectrum analysis technique and analyze the IF physical mechanisms to understand the phase transitions or structural evolutions associated with carbon atom movements in steels.

In this study, the martensitic decomposition behaviors of the low-carbon DP steel were studied by the low-frequency IF method, combined with XRD, SEM, and TEM analyses. Two IF peaks associated with the martensitic decomposition were observed at 430.4 and 448.1°C. The lower-temperature peak was attributed to the migration of the martensite/ferrite interface as the supersaturated carbon atoms moved from martensite to ferrite, while the higher-temperature peak was attributed to the formation of the Cr3C7 precipitates. This indicates that both carbon atom diffusion and carbide precipitation, i.e., the two-step process of the martensitic decomposition, can be well described by the IF technique. The related mechanisms of the phase transition and microstructural evolution during the martensitic decomposition were also explained.

Acknowledgement

This work was supported by the National Key Research and Development Program of China (Program Number: 2016YFB0300400) and the Suzhou Vocational University Foundation for Research (Grant No. SVU2016YY03). The author would like to thank Prof. Shen Zhongcheng and Mr. Lan Jinshan (Suzhou Siyuan material Measurement Technology Institute), and the Institute of Solid State Physics, Chinese Academy of Sciences for the supports in measuring the internal friction spectra and appreciate the valuable advice from Prof. L. B. Magalas (AGH University of Science, Poland).

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