Compositional Change of Refractory Elements in Solution during Aging in High Cr Heat Resistant Ferritic Steels

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1. Introduction

The improvement in the creep strength has been achieved by the addition of boron as well as refractory elements, Mo and W1–3) in high Cr heat resistant ferritic steels. It has been accepted that the improvement by the refractory elements originates mainly in the solid-solution hardening of steels. In those steels containing a comparatively large amount of the refractory elements, on the other hand, the creep strength is improved by the presence of fine coherent precipitates of intermetallic compounds due to the precipitation hardening in them.4) However, in the steels containing an excess amount of refractory elements, in particular, W, it is known that the creep strength in a long time becomes lower than the values extrapolated from the creep strength in a short-time.

Recently, it has been found that Re is an effective element for improving a long-time creep-strength of high Cr ferritic steels.5–7) This is in contrast to the effect of W on the ferritic steels. As mentioned above, it is important to investigate the solid solubility limit of refractory elements in the matrix when considering the creep strength of high Cr ferritic steels. This is usually obtained indirectly by the thermodynamic calculation based on the chemical composition as well as the weight of the extracted residue from the steels.8,9) so that some errors are included inevitably in the calculated values. Then instead of such an indirect method, a direct analyzing method is employed in this study. In this method, the amount of refractory elements dissolved in the matrix is analyzed directly by means of the ICP emission spectrochemical analysis of the electrolyte which is prepared by electrolyzing the matrix phase in the steel, as explained later.

2. Experimental Procedure

2.1. Steel Preparation

Seven ferritic steels have been used in this study. The chemical compositions are listed in mass% units in Table 1. Of these steels, both A and B steels contain 4% W, and their chemical compositions are similar to each other except for Re. B steel contains 0.2% Re, but A steel contains no Re. 1% W of A steel is replaced by 1% Re in C steel, so this C steel contains 3% W and 1% Re. Also, Mo and V compositions are modified slightly in C steel.

Table 1. Chemical compositions of steels used in this study, mass%.

|   | C   | Cr | Ni | Mn | Mo | V | Cu | Al | Fe | Si | P  | S  | Cr% | Nb |
|---|-----|----|----|----|----|---|----|----|----|----|----|----|-----|----|
| A | 0.08 | 10.1 | 0.01 | 0.00 | 0.00 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |
| B | 0.08 | 10.2 | 0.01 | 0.00 | 0.00 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |
| C | 0.18 | 10.3 | 0.0 | 0.00 | 0.00 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |

D, E, F and G steels listed in Table 1(b) contain about 2% W, and the Re compositions are changed systematically in these four steels. Here, G steel contains the largest amount of Re among the four steels, and its Mo composition is lowered slightly in order to suppress the ferrite formation in it. The chemical compositions of these experimental steels are controlled using the d electrons parameter, Md, in order to suppress the formation of the ferrite phase in them. The critical Md, value for suppressing the ferrite formation is about 0.862 for steels containing 3% Co.4) The Md, parameters listed in Table 1 are much lower than this critical value. In accordance with this prediction, no ferrite is formed in any steel.

The 50kg ingots of these seven steels were prepared by vacuum induction melting, and then processed by hot forging in a conventional way. The slab size was 35mm thick, 130mm wide and 1000mm long. In order to control the grain size, each slab was normalized at 1373 K for 18ks (5h), followed by annealing at 993 K for 72ks (20h). Subsequently, they were austenitized at 1343 K for 18ks (5h) followed by quenching and then tempered first at 843 K for 72ks (20h) and secondly at 953 K for 72ks (20h) for the quality heat treatment. The heating rate was controlled to be 5.56×10⁻² K s⁻¹ (20K/h), whereas the cooling rates were controlled to be 2.78×10⁻² K s⁻¹ (100K/h) for quenching and 1.39×10⁻² K s⁻¹ (50K/h) for tempering, respectively. These rates were set for a simulation of the thermal hysteresis which appears in the central part of a large-scale rotor. After tempering, every steel has been supplied for the creep rupture test at 923 K.

2.2. ICP Emission Spectrochemical Analysis

Specimens taken from the grip part of test pieces crept at 923 K have been electrolyzed using a 10% hydrochloric acid–methanol solution under a constant-potential.10) It is known that MC carbide,11) M₂₃C₆ carbide10) and the Laves phase10) are extracted effectively by using this solution. In fact, MX carbonitride, M₂₃C₆ carbide and the Laves phase have been observed in an extraction replica made by using this solution.

In order to remove the extracted residues, the electrolyte has been first passed through a membrane filter having 0.2μm pores, and then a membrane filter having 0.025μm pores. In general, a filtering using a membrane filter having 0.2μm pores is enough to remove fine precipitates such as MX, BN, etc. from the electrolyte.12) In fact, the amount of the extracted residues after the first-step filtering does not change after the second-step filtering in this study. To make sure, however, the two-steps filtering has been adopted to remove the precipitates thoroughly from the electrolyte.

Since organic solvents are not suitable for the ICP emission spectrochemical analysis, the filtered electrolyte has been diluted by distilled water so that the total content of organic solvents becomes less than 5%. Then this diluted electrolyte has been supplied for the ICP emission spectrochemical analysis. In order to obtain calibration curves for
the ICP analysis, two kinds of aqueous solutions have been used as the standard solutions. One is a 1 ppm solution of Mo, W or Re, and the other is a 5 ppm solution. The calibration curve for each element has been obtained by the least square fit using three data, that is, 0 ppm (distilled water), 1 ppm and 5 ppm standard values.

3. Results and Discussion

3.1. Contents of Refractory Elements in the Matrix of High W Containing Steels

The results of the ICP emission spectrochemical analysis are shown in Fig. 1 for A and B steels and in Fig. 2 for C steel. In these figures, the vertical axis shows the analyzed value in mass%, and the abscissa shows the aging time at 923 K. Here, “zero” of the aging time corresponds to the as-tempered state. In the bar charts, gray, white and black show the respective contents to W, Re and Mo, respectively. Also, line graphs exhibit the amount of extracted residues. Here, the analyzed data is represented in the mass% units in these figures, because it is obtained in the mass% units directly from the ICP emission spectrochemical analysis.

It is seen from Fig. 1 that the total amount of the refractory elements in the matrix of A and B steels is about 1.2 and 1.4%, respectively, in the as-tempered state. It decreases with increasing aging time in both steels. For example, it is 0.95% in A steel aged for 26.11 Ms (7 253 h) and 0.85% in B steel aged for 30.33 Ms (8 425 h). The matrix of B steel contains about 0.2% Re after aging for 30.33 Ms (8425 h). Interestingly, this corresponds to the Re composition in the steel and is similar to the content of Re dissolving in the matrix in the as-tempered state. This result indicates that almost all the alloyed Re remains in the matrix phase of B steel even after aging. Therefore, the observed decrease in the total amount of the refractory elements is attributable mainly to the decrease in the W content in B steel. For example, the W content in the matrix decreases from 1.2% in the as-tempered state to 0.65% in the aged state for 30.33 Ms (8425 h).

As shown in Fig. 1, the W content in the matrix of A steel decreases monotonously from 1.25% in the as-tempered state to 0.8% in the aged state for 39.72 Ms (11 033 h). For advanced high Cr ferritic steels the contents of Mo and W in the matrix after a long time creep test at 923 K have been investigated by analyzing the chemical compositions of the extracted residues. It has been shown that the solubility of Mo+(1/2)W in those ferritic steels is 0.5–0.8 mass%. Also, by using Atom-Probe-Field-Ion-Microscopy (APFIM), the contents of Mo and W in the matrix have been measured to be 0.26 mass% and 0.82 mass%, respectively. As explained before, the result of A steel obtained from this study is about 0.8%, which is consistent with the previous reports.

In contrast to such a decrease in the matrix content of the refractory elements by aging, the amount of extracted residues increases in both A and B steels in the beginning of aging, and then it is saturated to be about 8%.

Figure 2 shows the results on C steel, in which 1% W in A steel is replaced by Re. The total amount of refractory elements in the matrix is 2.7% in the as-tempered state of this steel, and it decreases with aging time, resulting in 1.5% after aging for 25.77 Ms (7 159 h). This value of 1.5% is still much higher than that in A or B steel after aging (Fig. 1). This is caused simply by a high Re content in the matrix, i.e., 10% in the as-tempered state, 0.8% after aging for 16.02 Ms (4449 h) and 0.65% after aging for 25.77 Ms (7 159 h), as shown in Fig. 2. In contrast to the Re content, the W content in the matrix of C steel decreases considerably. For example, it changes from 1.6% in the as-tempered state to 0.75% after aging for 25.77 Ms (7 159 h). As a result, the dissolved W content becomes comparable to the Re content in the aged steel, although the W composition is three times higher than the Re composition in C steel (see Table 1). Thus, it can be said that Re has a larger solubility in steel than W.

The amount of the extracted residues in C steel also increases with increasing aging time and it is saturated to be about 6.5%. This value is slightly lower than the saturated values, about 0.8%, observed in both A and B steels (see Fig. 1). This is consistent with the result that the total amount of the refractory elements dissolved in the matrix of C steel is larger than that of A or B steel.

3.2. Change in the Dissolved Re Content in the Matrix with Re Composition in Steels

Figures 3 and 4 show the results of ICP emission spectrochemical analysis on the (b) group steels, D, E, F and G.
This is because those four steels contain a smaller amount of Mo, W, Re, and Mo, dissolved in the matrix of high Cr and Re-containing heat resistant steels, and its change with aging time at 923 K has been investigated systematically. The results obtained are as follows:

1. The total amount of the refractory elements in matrix in the steel with 4% W is about 2% in the as-tempered state when the steel contains no Re. This amount keeps decreasing with increasing aging time and finally reaches to about 0.95%.

2. When the steel contains Re, the total amount of refractory elements in the matrix increases with increasing Re composition in the steel in both the as-tempered state and the aged state. For example, it is about 1.5% even after aging for 25.77 Ms (7.159 h) when the steel contains 1% Re substituting 1% W.

3. Such a large amount of the refractory elements in the Re containing steels is attributable to the large solid solubility of Re in high Cr ferritic steels.

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