Production and Properties of Nano-scale Oxide Dispersion Strengthened (ODS) 9Cr Martensitic Steel Claddings

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The 9Cr–ODS martensitic steel claddings were developed by cold-rolling and subsequent heat-treatment. The standard chemical composition is Fe–0.13C–9Cr–2W–0.2Ti–0.35Y2O3. The substantially elongated grains formed by cold-rolling turned out to be into equi-axied grains by ferrite to austenite phase transformation at the final heat-treatment. The produced claddings have the tempered martensitic structure and excess oxygen of 0.060 mass%. The superior tensile and creep rupture strength were shown in the produced cladding, compared with conventional ferritic (PNC–FMS) and even austenitic (PNC316) claddings at higher temperature and extended time. The strength improvement is attributed to finely distributed nano-scale complex oxide. The coarser ferrite grains produced by slow cooling make further improvement in the tensile and creep rupture strength beyond those of tempered martensite at high temperature and longer testing time. The higher excess oxygen content of 0.137 mass% prevents fine distribution of the oxide particles that lead to inferior high temperature tensile and creep strength.

KEY WORDS: ODS; yttria; phase transformation; martensite; ferrite; excess oxygen; cladding; tensile strength; creep rupture strength; cold rolling; grain size.

1. Introduction
To improve thermal efficiency of plants operating at high temperature, efforts have been made to develop high temperature strength of ferritic–martensitic steels (FMS). The effects of alloying elements and heat-treatment to stabilize carbide precipitates have been studied in the field of the power-generation industry.1) The oxide dispersion strengthened (ODS) martensitic steel is expected to be a good candidate of the materials having a potential of superior high temperatures strength over 923 K. Since FMS is well known to have far superior radiation resistance compared with austenitic steels, the ODS martensitic steel is expected to be a promising cladding material in the long-life core of fast reactor fuel pins2–4) as well as a low activated fusion reactor materials.7) The development of ODS martensitic steel has being conducted in Japan Nuclear Cycle Development Institute (JNC). The chemical composition of the material for fuel cladding application to fast reactors was proposed to be Fe–0.13C–9Cr–2W–0.2Ti–0.35Y2O3, and substantially elongated grains formed by cold-rolling were made into equi-axied grains by means of α to γ phase transformation.5,6)

In this study, claddings with a standard composition of Fe–0.13C–9Cr–2W–0.2Ti–0.35Y2O3 are developed using the cold-rolling process, and the high temperature mechanical properties of the produced claddings are investigated. The effects of difference of the final heat-treatment and oxygen content on the mechanical properties of the 9Cr–ODS martensitic steels are extensively studied.

2. Experimental Procedure
2.1. Cladding Production and Heat Treatment
The preparation process of the ODS martensitic claddings is shown in Fig. 1. As the raw material powders, two kinds of powders: the Fe–0.13C–9Cr–2W–0.2Ti powders atomized in argon gas with a particle size below 150 μm and mixture of each element powder (Fe, C, Cr, W, Ti) measured in the same nominal composition were used for making M11/MS11 claddings and Mm13 claddings, respectively. The development of ODS martensitic steel has been conducted in Japan Nuclear Cycle Development Institute (JNC). The chemical composition of the material for fuel cladding application to fast reactors was proposed to be Fe–0.13C–9Cr–2W–0.2Ti–0.35Y2O3, and substantially elongated grains formed by cold-rolling were made into equi-axied grains by means of α to γ phase transformation.5,6)

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The raw bars were machined and drilled to form the mother tubes with 18 mm outer diameter, 3 mm wall-thickness and 350 mm length. M11 and MS11 claddings in the dimension of 8.5 mm outer diameter, 0.5 mm thickness and near 3 m length were produced by four-pass cold rolling using pilger mill. While, the Mm13 cladding was produced into the dimension of 6.9 mm outer diameter, 0.4 mm thickness and near 3 m length by the same process of four-pass cold rolling. The cold-rolling of these claddings was conducted at a state of the ferritic structure obtained by low cooling rate of 150 K/h in the furnace after normalizing at 1323 K.

Figure 2 represents the results of hardness change in the course of cold-rolling process of M11 cladding. A reduction (Rd) of the cold-rolling attains around 45% at each pass. The employed normalizing (1323 K for 30 min) and furnace cooling (cooling rate of 150 K/h) after each cold-rolling successfully softened the claddings.

After final cold-rolling, normalizing at 1323 K for 1 h in vacuum and subsequent cooling in argon gas atmosphere at cooling rate of 5000 K/h and tempering at 1073 K for 1 h were applied for M11 and Mm13 claddings. These heat treatments were applied to change elongated grain structure generated in the cold-rolling process to equi-axied one, by means of α to γ and γ to martensite phase transformations. On the other hand, MS11 cladding was normalized at 1323 K for 1h followed by the furnace cooling at cooling rate of 5 K/h to make the ferrite phase at the final stage.

2.2. Test of Mechanical Properties and Microstructural Observation

The produced claddings are tested to determine their high temperature mechanical properties by means of ring tensile and internally pressurized creep rupture tests. These tests were selected to prove their resistance against the dominant direction of loading in an actual fuel pin, i.e. hoop direction. In order to characterize the mechanical properties of the manufactured claddings, the optical micrographs were taken. The transmission electron microscopy (TEM) observations of the thin foil were made using a Hitachi H-800 to determine the dispersion parameters, especially the size and number density of oxide parti-
cles. The electron diffraction and EDX analyses for precipitates in the extracted replicas were conducted to characterize the carbide and oxide precipitates.

3. Result

3.1. Chemical Composition and Microstructure

The measured chemical composition of the manufactured claddings are listed in Table 1. The target chemical composition is Fe–0.13C–9Cr–2W–0.2Ti–0.35Y₂O₃, where the amount of 2.0 mass% W for solution hardening and 0.20 mass% Ti were added. The value of excess oxygen (Ex.O) was estimated by subtracting oxygen contained in Y₂O₃ from the total oxygen determined by inert gas melt method. The Y content was analyzed by the inductively coupled plasma method, from which the Ex.O and Y₂O₃ contents were estimated. The Mm13 cladding has higher content of excess oxygen of 0.137 mass% compared with M11 and MS11 claddings. This could be due to the introduction of higher energy and use of elemental powders during mechanically alloying.

The optical micrographs of M11, Mm13 and MS11 claddings are shown in Fig. 3. The tempered martensitic structure is formed in the whole area of M11 and Mm13 claddings. The finely elongated phases of about 30 to 100 μm in length are also observed in strips by the white color in M11 cladding. The white color is considered not to be α-ferrite, but could be non-transformed α phase remained without transformation during normalizing and cooling, since this material was produced by mechanically alloying at room temperature without melting. In M11 cladding with 0.060 Ex.O, it is supposed that the carbon content in the matrix is likely reduced due to Ti–C formation rather than Ti–O formation. On the other hand, in Mm13 cladding with

### Table 1. Chemical composition on produced claddings (mass%).

| Cladding | Chemical composition (mass%) |
|----------|-------------------------------|
| M11, MS11 | 0.13 9.00 1.95 0.021 0.20 0.29 0.013 0.0025 0.37 0.064 |
| Mm13     | 0.14 8.85 1.95 0.010 0.20 0.27 0.010 0.0054 0.34 0.137 |

*Si : 0.05 (M11, MS11), <0.005(Mm13)
Mn : 0.04 (M11, MS11), <0.01(Mm13)
P : 0.002(M11, MS11), 0.001(Mm13)
S : 0.002(M11, MS11), 0.003(Mm13)

*1 : Estimated from yttrium content assuming it to be contained in the form of Y₂O₃ (yttrium mass % x 1.27)
*2 : Excessive oxygen estimated as total oxygen content minus oxygen content in Y₂O₃ (total oxygen content - yttrium mass % x 0.27)
0.137 Ex.O, C content could be maintained, and α to γ phase transformation is perfectly taken place. The MS11 cladding has the coarser ferrite grains with about 20 μm, which were formed by γ to α phase transformation and grain growth occurred concomitantly during furnace cooling at cooling rate of 5 K/h after normalizing at 1 323 K for 1 h.

### 3.2. Tensile Properties

The 0.2% yield strength and ultimate tensile strength of M11, MS11 and Mm13 claddings in the hoop direction measured in the temperature range from room temperature to 1 073 K are shown in Figs. 4 and 5, together with the corresponding data of ferritic–martensitic stainless steel (PNC–FMS) used conventionally as fast reactor fuel cladding. All the three claddings present superior strength compared with conventional PNC–FMS. Among the claddings, the Mm13 showed the lowest strength, and this reason will be discussed later. Comparing M11 and MS11, the strength of MS11 is higher at above 900 K. This is probably due to the coarser ferritic grains in MS11 formed by low cooling rate (5 K/h) from the austenite.

The uniform and total elongations from room temperature to 1 073 K are shown in Figs. 6 and 7, together with

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**Fig. 3.** Optical micrographs of M11, Mm13 and MS11 claddings.

**Fig. 4.** 0.2% yield strength in the hoop direction using the ring specimens of M11, MS11 and Mm13 claddings at different temperature, comparing with conventional PNC–FMS.

**Fig. 5.** Ultimate tensile strength in the hoop direction using the ring specimens, together with PNC–FMS.

**Fig. 6.** Uniform elongation in the hoop direction using the ring specimens of M11, MS11 and Mm13 claddings, comparing with PNC–FMS.

**Fig. 7.** Total elongation in the hoop direction using the ring specimens of M11, MS11 and Mm13 claddings, comparing with PNC–FMS.
those of conventional PNC–FMS. In the temperature range from 673 to 973 K, where the fast reactor is commonly operated, the measured uniform elongation satisfies the value of 0.3% required for design. Particularly, all the produced claddings present superior uniform elongation to PNC–FMS at above 800 K, although their total elongations are smaller than PNC–FMS over the entire temperature range. This advantage of superior elongation in the produced claddings is probably ascribed to the pining of dislocations by Y2O3 particles, which retard recovery and sustain work-hardening. Smaller total elongation of the produced claddings could be caused by the limited amount of necking after uniform elongation.

3.3. Creep Rupture Properties

The measured data on creep rupture strength in hoop direction under internal pressure (hoop stress) at 973 K are indicated in Fig. 8. Mm13 exhibits significantly low creep rupture strength, similar to the strength level of PNC–FMS. The M11 and MS11 have almost same creep rupture strength, where the regression line for M11 is the result of analysis using the creep rupture data at temperature of 923 K, 973 K and 1023 K, that is expressed in detail in the next session. The rupture time for MS11 seems to extend to the longer time, because MS11 specimen did not rupture up to 10 784 h at 130 MPa, as indicated by arrow mark.

4. Discussion

4.1. Internal Creep Rupture Properties of M11 Cladding

The internal creep rupture data of M11 cladding were acquired at 923 K, 973 K and 1023 K, and were analyzed applying Larson–Miller-Parameter (LMP) that is expressed in terms of temperature and rupture time. The regression analysis of the least square method was conducted for the hoop stress vs. LMP to minimize the standard deviation. The best-fit curve with the optimized values is represented in Fig. 9. The largest value of LMP tested is 22 180 at 1023 K, where the cladding has not yet ruptured, as indicated by the arrow mark in Fig. 9. The value of LMP=22 180 is converted to the equivalent rupture time at the typically operational temperature of 973 K in fast reactor as 106 900 h at 973 K. This period is adequately longer than 5 years service of fuel elements.

The lifetime of the fast reactor fuel pin is dominated by the internal creep rupture strength of the cladding induced by the fission gas internal pressure at temperature around 973 K. In Fig. 10, internal creep rupture data are shown at 923 K, 973 K and 1023 K for M11 cladding. The best-fit curves for hoop stress vs. rupture time derived above are also shown by solid curves at each temperature. The corresponding creep rupture curves for PNC–FMS and austenitic PNC316 are also presented for comparison in this figure. PNC316 is the typical austenitic cladding developed by JNC in the fast reactor program.12) Notably superior performance in rupture time is shown in M11 cladding compared

Fig. 8. Internal creep rupture curve of M11, MS11 and Mm13 claddings at 973 K, comparing with conventional PNC–FMS.

Fig. 9. Best-fit curve of M11 rupture strength at 923 K, 973 K and 1023 K drawn by regression analysis against Larson–Miller parameter.
with PNC–FMS. The slope of PNC316 is steeper than M11, crossing over that before 1000 h at 1023 K and before 10000 h at 973 K. The stress condition of the fast reactor fuel pin gradually increases due to accumulation of fission gases up to around 100 MPa during service duration of 44000 h at 973 K. Therefore, it is to be noted that M11 cladding is of advantage compared with austenitic PNC316 cladding under the real fast reactor operation condition.

Figure 11 is a transmission electron micrograph of M11, showing nearly spherical shape of Y₂O₃ particles in the martensite matrix. The oxide particle size is around 3 nm, and inter-particles spacing is significantly shortened. Considering that high temperature strength of ODS steels is owing to an existence of oxide particles that act as a resistance for dislocation gliding,¹³ the creep rupture strength measured in M11 is dominated by those oxide particles distributed finely. The creep rupture strength of austenitic PNC316 is attributed to the formation of Ti–Nb carbide precipitates. Those precipitates have a tendency to become coarser, and thus creep rupture strength concomitantly falls at longer time and at higher temperature.¹⁴ On the other hand, Y₂O₃ particles are thermally stable at the extended duration, which is owing to the sustainable creep rupture strength of M11 even at temperature over 973 K and longer time.

4.2. Effect of Final Heat-treatment on the High Temperature Strength (M11 vs. MS11)

When compared MS11 with M11 using the data shown in Figs. 4 to 7, MS11 cladding exhibited superior tensile strength at above 873 K and adequate ductility over entire temperature range. The creep rupture strength also showed the extended lifetime in MS11 cladding at 973 K. These results indicate that high temperature mechanical properties of 9Cr–ODS martensitic steel cladding are improved by morphological change induced by the final heat-treatment.

The continuous cooling transformation (CCT) diagram⁹ of M11 cladding in the cooling rates ranging from 50 to 18000 K/h is shown in Fig. 12. In this diagram, martensite start (Ms) and finish (Mf) lines and ferrite start (Fs) and finish (Ff) lines are shown. M11 and Mm13 was cooled at rate of approximately 5000 K/h from austenite and tempered at 1073 K for 1 h. Thus M11 and Mm13 claddings have tempered martensite, as shown in Fig. 3. On the other hand, the microstructure of MS11 cladding is coarse ferrite grains of around 10 μm size and carbide, since its cooling rate is 5 K/h that intersect with only Fs and Ff lines, as shown in Fig. 12.

Since Y₂O₃ particles distribution in M11 and MS11 is almost same, the superior strength of MS11 cladding at high temperature could be attributed to the grain structure: a coarser ferrite grain in MS11 vs. extremely fine martensite grains of less than 1 μm size in M11. In the previous study,⁹ the existence of grain-boundary sliding was pointed out for the creep deformation of the martensitic 9Cr–ODS cladding with extremely fine and homogeneous grains at 973 K. Therefore, it is considered that the suppressed grain
boundary sliding contribution due to coarser grains in MS11 cladding is attributed to the superior tensile and creep rupture strength at higher temperature and longer time.

4.3. Effect of Excess Oxygen on the High Temperature Strength (M11 vs. Mm13)

Mm13 exhibited the inferior tensile and creep rupture strength compared with M11. The transmission electron micrograph of the oxide particles in Mm13 specimen is shown in Fig. 13. The size of the observed oxide particles is more than 5 nm, which is slightly coarser than that of M11 specimen shown in Fig. 11. In particular, the number density of the oxide particles is low in Mm13, and relatively longer inter-particles spacing in Mm13 than M11 leads to the lower strength in Mm13 from the inverse relation of a strength level to the inter-particles spacing.13)

The co-authors found out by means of X-ray diffraction analysis that Y2O3 particles are dissolved in the matrix during mechanically alloying process, and then fine complex oxide with Y and Ti precipitates during the hot-extrusion process at 1423 K.15) Hence, the size of precipitates which appear during the hot-extrusion at 1423 K would depend on the amount of Y, Ti and excess oxygen contents in the matrix. Among these elements, major difference between M11 and Mm13 is excess oxygen content as shown in Table 1. Therefore, the effect of the excess oxygen on the size of precipitates is investigated by means of TEM observation and thermodynamic analysis, focusing on partition of titanium element between Y2O3 particles and others.

TEM observation of extracted replicas was carried out for the precipitates of M11 and Mm13 normalized at 1323 K, and results are summarized in Table 2. In M11 with lower excess oxygen of 0.06 mass%, three types of precipitates are identified: M23C6 containing Ti (several hundred nm), Ti oxide (Ti–O) (a hundred nm) and Y–Ti oxide (3 nm size). The M23C6 was identified from the lattice constant measured by electron diffraction analysis. Ti oxide and Y–Ti oxide were estimated from the EDX analysis of each particle. Their chemical composition can not be exactly distinguished by electron diffraction analysis. In Mm13 specimen with higher excess oxygen of 0.137 mass%, however, several hundred nm size of Cr2O3 containing titanium was identified, together with a hundred nm size of YCrO3 oxide containing titanium and finely distributed Y–Ti oxide particles in more than 5 nm size. The Cr2O3 and YCrO3 were identified from the electron diffraction analysis. The carbide containing titanium was not be observed.

The co-authors investigated the effect of excess oxygen on the oxide formation thermodynamically. Using Thermo-Calc code,16) equilibrium phases are calculated at 1423 K that corresponds to the oxide precipitation temperature during hot-extrusion. In the lack of thermodynamic data of Y–Ti complex oxide, however, complex oxide formation can not be taken into account; Y2O3 and Ti oxide are separately calculated in this calculation. The titanium partitioning between oxide and carbide is paid attention in this calculation. In the case of M11 with lower excess oxygen content, Ti exists in three phases as shown in Table 2: 0.05 mass% in TiC, 0.10 mass% in Ti2O3, and 0.05 mass% in matrix g phase. On the other hand, in Mm13 containing higher excess oxygen content, all of Ti precipitate as Ti4O7. With increasing excess oxygen content from 0.06 mass% (M11) to 0.137 mass% (Mm13), the higher order of titanium oxide, Ti4O7, instead of Ti2O3 is predicted. Furthermore, chromi-
um oxide as a type of Cr₂O₃ is predicted for the higher excessive oxygen specimen of Mm13. These thermodynamic predictions well coincide with the results of TEM observation. The improved tensile and creep rupture strength in M11 cladding compared with that of Mm13 cladding is owing to the formation of finer nano-meter size Y–Ti complex oxide. This finer complex oxide formation could be associated with the formation of lower order of titanium oxide, Ti₂O₃, with lower excess oxygen. It is supposed that nano-meter size Y–Ti complex oxide would be formed by the precipitation as Y₂O₃–Ti₂O₃ during hot-extrusion process under lower excess oxygen content.

5. Conclusion

In this study, 9Cr–ODS steel claddings were originally developed and produced by cold-rolling in ferrite condition with parameters of final heat-treatment and excess oxygen content. The results of study are summarized as follows.

(1) The produced claddings in tempered martensitic structure (M11) attained the noticeable improvement in tensile and internal creep rupture strength due to fine distribution of the oxide particles in martensite matrix.

(2) A coarser ferrite grains of about 20 μm size were produced by slow cooling rate of 5 K/s in 9Cr–ODS steel (MS11). Those coarser ferritic grains promoted to the further improvement beyond that of tempered martensitic steels in the tensile and creep rupture strength at high temperature and longer time.

(3) The excess oxygen has significant influence on the high temperature mechanical properties; higher excess oxygen content (Mm13) prevents formation of finely distributed oxide particles, which induces inferior high temperature tensile and creep strength.

REFERENCES

1) Proc. Conf. Advanced Heat Resistant Steels for Power Generation, ed. by R. Viswanathan and J. Nutting. IOM Communication Ltd., London, (1999).
2) J. J. Huet: Nucl. Technol., 70, (1985), 215.
3) A. Alamo, H. Regle, G. Pons and J. L. Bechade: Mater. Sci. Forum, 88–90, (1992), 183.
4) P. Dubuisson, R. Schill, M.-P. Hugon, I. Grisdin and J.-L. Seran: 18th Int. Symp., Effects of Radiation on Materials: ASTM STP 1325, ed. by R. K. Nanstad, M. L. Hamilton, F. A. Garner and A. S. Kumar, American Society for Testing and Materials, West Conshohocken, (1999), 882.
5) S. Ukai, M. Harada, H. Okada, M. Inoue, S. Nomura, S. Shikakura, K. Asabe, T. Nishida and M. Fujiwara: J. Nucl. Mater., 204 (1993), 65.
6) S. Ukai, M. Harada, H. Okada, M. Inoue, S. Nomura, S. Shikakura, T. Nishida, M. Fujiwara and K. Asabe: J. Nucl. Mater., 204 (1993), 74.
7) R. L. Klueh and D. R. Harries: High-Chromium Ferritic and Martensitic Steels for Nuclear Applications, ASTM Stock Number: MONO 3, (2001).
8) S. Ukai, T. Nishida and T. Yoshitake: J. Nucl. Sci. Technol., 35 (1998), No. 4, 294.
9) S. Ukai, S. Mizuta, M. Fujwara, T. Okuda and T. Kobayashi: J. Nucl. Sci. Technol., 39 (2002), No. 7, 778.
10) J. L. Seran, V. Levy, P. Dubuisson, D. Gilbon, A. Maillard, A. Fissolo, H. Touron, R. Cauvin, A. Chalony and E. Le Boulbin: 15th Int. Symp., Effects of Radiation on Materials: ASTM STP 1125, ed. by R. E. Stoller, A. S. Kumar and D. S. Gelles, American Society for Testing and Materials, Philadelphia, (1992), 1209.
11) A. Uehira, S. Ukai, T. Mizuno, T. Asaga and E. Yoshida: J. Nucl. Sci. Technol., 37 (2000), No. 9, 780.
12) I. Shibahara, S. Ukai, S. Onose and S. Shikakura: J. Nucl. Mater., 204 (1993), 131.
13) S. Ukai, T. Okuda, M. Fujwara, T. Kobayashi, S. Mizuta and H. Nakashima: J. Nucl. Sci. Technol., 39 (2002), No. 8, 872.
14) Y. Tateishi, S. Yuhara, I. Shibahara, M. Ito, S. Nomura, Y. Sato, E. Yoshida and S. Shikakura: J. Nucl. Sci. Technol., 26 (1989), No. 1, 132.
15) T. Okuda and M. Fujwara: J. Mater. Sci. Lett., 14 (1995), 1600.
16) B. Sundman: Thermo-Calc User’s guide, Division of Computational Thermodynamics, Dept. Materials Science and Engineering, Royal Institute of Technology, Stockholm, Sweden, (1993).