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Zirconia matrix composite dispersed with stainless steel particles: processing and oxidation behavior

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Abstract

Materials with non uniform properties are being developed to optimize several functions of industrial components in severe atmospheres at high temperature. These composites called M(p)-CMC(s) : "ceramic matrix composites dispersed with metal particles" are candidates for high-temperature structure materials as functionally graded materials (FGMs) such as intermediate components between electrolyte and interconnecting components in SOFC.

Preparation of a model composite M(p)-CMC(s) is described. Powder metallurgy process is used to obtain a dense composite (98 % of the theoretical density) based on yttrium stabilized zirconia for the ceramic part and on 304L stainless steel for the metallic part. The characteristics of this material as well as it behavior under oxidation at high temperature are reported.

Keywords: ZrO₂, powder metallurgy process, composite, corrosion, solid fuel.

1. Introduction

New challenges of modern high technology require the development of technical materials with combined properties or functions [1, 2].

Ceramic matrix composites dispersed with metallic particles, called M(p)-CMC(s) [3], are one of the promising materials for high performance applications under severe environment such as high temperature. This new material design offers the possibility to combine heat resistance, corrosion resistance and wear resistance due to the ceramic phase, with mechanical strength and electric conductivity provided by the metal phase [4].

M(p)-CMC(s) are nowadays the candidates for functionally graded materials (FGMs) developed for their multiple functions and reduced cost [5, 6]. Recent studies highlight the use of graded M(p)-CMC(s) materials as intermediate components between interconnectors and electrolyte in the oxide fuel cell SOFC [4, 7].

The model M(p)-CMC(s) described in this study is based on yttrium stabilized ZrO₂ and 304L stainless steel materials.

Zirconia is an interesting ionic ceramic material with high oxygen diffusion at temperature around 800 $^{\circ}$ C. No interaction between ZrO₂ and stainless steel has been reported [8, 9]. The thermal expansion coefficients of zirconia $(11.10^{-6} \text{ °K}^{-1})$ and of the metal phase $(18.10^{-6} \text{ °K}^{-1})$ for temperature close to 1000 °C are close together which should lower thermal stresses during the sintering of the composite.

The present work is dedicated to the 304L(p)-ZrO₂(s) composite manufacturing based on powder metallurgy process : blending of powders, granulation, pressing and sintering at 1350 °C under dry hydrogen and argon atmosphere. This process allows controlling the composition and the microstructure of the material [10, 11]. The characterizations of the sintered samples are reported. Finally, the oxidation behavior of the model 304L(p)-ZrO₂(s) composite, studied by means of thermogravimetry (TGA) under 20 % O₂ in helium at 700 °C and 800 °C, is presented. Based on Nanko's study [3], the oxidation will normally only be due to stainless steel particles inside the zirconia matrix.

2. Experimental procedure

2.1 Basic materials

The starting ceramic powder is yttrium oxide doped ZrO_2 with a specific surface area of 14.6 m²/g. This value corresponds to a crystallite size, considering a spherical particle, around 67 nm. The atomized TZ-3Y-E zirconia powder doped with 3 mol % of yttrium oxide was supplied by the Japanese company Tosoh. Its composition is given in Table I.

Chemical composition (mass %) ZrO_2 Y_2O_3 SiO₂ Al_2O_3 Zirconia TZ-3Y-E base 5.2 0.25 < 0.002

Table I. Chemichal composition of ZrO₂ powder.

Laser granulometry analysis (Malvern instruments, Mastersizer 2000) of TZ-3Y-E zirconia powder shows a multimodal size distribution in the range 0.1 to 100 μ m (figure 1). The mode close to 100 μ m corresponds to the primary atomized particles. The other modes correspond to broken particles due to the sample preparation (powder dilution in water).



Fig.1. Laser granulometry of TZ-3Y-E ZrO₂ powder

The starting metal powder is austenitic 304L stainless steel atomized in water as supplied by Hoganas Belgium. The powder density measured with picnometry method is closed to 99.65 % of the theoretical density. Its specific surface area measured due to BET Krypton adsorption method is 0.06 m²/g. This value corresponds to a crystallite size, considering a spherical particle, around 13 μ m which is ~200 times bigger than the zirconia crystallites.

The chemical composition of the 304L stainless steel powder is given in Table II.

Chemical composition	Fe	Cr	Ni	Si	С	S	Mn	Р	0
mass (%)	base	19.1	11	0.8	0.013	0.004	0.1	0.011	0.27

Table II. Chemical composition of 304L stainless steel powder.

Laser granulometry (figure 2) shows the particle size distribution with one main mode of particle diameter in the range 10 to 200 μ m with a maximum at ~80 μ m.



Fig.2. Laser granulometry of 304L stainless steel powder

Figure 3-a, shows individual particles of 304L stainless steel powder. In figure 3-b, at a higher magnification, a native oxide layer can be observed around the particles. XRD analysis (D 5000 Siemens) of the metallic powder does not show any oxide diffraction peaks (figure 4). However, it can be supposed that the amount of oxide is too low to be detected (<1vol %). The oxide layer is probably Cr_2O_3 as expected for this kind of alloy.



Fig.3. SEM observations of 304L stainless steel powder



Fig.4. XRD analysis of 304L stainless steel powder

2.2 Preparation of the model 304L(p)-ZrO₂(s) composite

304L(p)-ZrO₂(s) composite was obtained due to a powder metallurgy process. It is composed of 60 vol% TZ-3Y-E zirconia with 40 vol% 304L stainless steel. To reduce the porosity in the sample, organic binder was avoided during the powder preparation. The ceramic and metal powders were mixed and homogenized during 30 min in a planetary blender TURBULA. To prepare a homogeneous blend at the particles scale, a milling step was added to the process. The mixed powder was milled during 5 min at a low speed in an annular carbide ring mill.

After the powder preparation, a two steps compressive process was used to optimize the flowability of the powder in order to improve the feeding of the matrix press.

2.2.1 Granulate step

The granulate powder was prepared by light compaction of blended powder at an optimal value of 250 MPa. Samples (8 mm diameter discs) were compacted with an INSTRON press. These compacts were manually gently broken and sieved to produce 1mm agglomerates. The resulting granules were mixed and rounded in a TURBULA for 10 min just before the compression step.

2.2.2 Compression step

During a second step, granules were compacted at a pressure higher than for the granulation step with the same INSTRON press. Cylindrical pellets (8 mm diameter, 6 mm length) are prepared. Magnesium stearate was used as an external die lubricant.

2.2.3 Sintering step

Differential Thermal Analysis (Setaram TG DTA 92-16/18) measurement on stainless steel powder indicated a melting temperature of 1451 °C whereas the melting temperature of the ZrO_2 powder is around 2000 °C. The supplier's recommendations for the sintering temperatures are respectively 1250 °C for the metal powder and 1350 °C for the oxide powder.

The densifications of the metal and the ceramic powders were individually studied with a SETARAM TMA 92 dilatometer at 1400 °C under Ar-10 % H_2 atmosphere. This gaseous mixture was chosen to guaranty a reducing environment and to avoid oxidation of the metal phase.

2.3 Characterization of the composites after sintering and after oxidation

The microstructure of the sintered 304L(p)-ZrO₂(s) composite samples were examined with a JEOL 6400 model Scanning Electron Microscope (SEM). Samples were coated with a hot mounting conductive resin with carbon fillers. They were abraded with 100 grains/cm² to 1200 grains/cm² SiC paper using water and 9 µm to 1 µm diamond polishing solution. The samples are finally cleaned with water. Before observations, a 3 nm gold layer was sputtered on the sample with a BALZERS SCD 050 coater.

The internal microstructure was observed with a Transmission Electron Microscope (MET Philips CM 2000). The specimens were small discs with a thickness less than 1 mm. A dimpling procedure was used to thinning the disc in the central region while leaving a thick supporting rim to protect the sample from damage. After dimpling, the samples were ion beam polished to generate a large electron transparent area due to a Gatan Model Precision Ion Polishing System (PIPS) instrument. The voltage of the ion beam was kept at 5.0 kV until sample perforation. Once the perforation was detected, the voltage of the ion beam was reduced to 2.5 kV and maintained during 20 min. Then, the voltage was lowered to 1.5 kV again and kept to this value during 20 min.

X-ray diffraction (XRD) with a D 5000 Siemens diffractometer at ambient temperature and with a D8 advance Bruker diffractometer + HTK Anton-Paar chamber at high temperature was used to characterize the initial and oxidized samples.

Auger analysis AES was carried out on a radial cross section sample surface with a spectrometer Riber-Cameca ultra vacuum chamber equipped with a non monochromatic X-ray beam and associated with a Mac3 analyzer. Such an equipment provides high spatial resolution (~200nm corresponding to the beam size). It offers the possibility to detect elements at low concentrations (0.5-1% of the atomic composition of the sample).

2.4 Corrosion of 304L(p)-ZrO₂(s) composite – Thermogravimetric analysis (TGA)

The M(p)-CMC(s) samples used for TGA analysis were 1 mm thin disks cut in the center of the sintered pellets. This design allows considering that the mass change is mainly due to the contribution of the two based faces of the disc. Oxidation experiments are performed in a symmetric thermogravimetric analyzer (SETARAM TGA16 with Pt-Rh 6 %/ Pt-Rh 30 % thermocouple). Open alumina crucible was used to maintain the sample in the isothermal area of the furnace. The temperature was raised up to 700 °C or 800 °C (30 °C/min) under free flowing helium atmosphere at atmospheric pressure (2 L/h). The oxygen impurity in helium was analyzed at the furnace outlet with an oxygen probe (Systech Instruments 900, accuracy: 1 ppm at 20 °C).

After 7 min at 700 °C or 800 °C, helium was switched to a mixture of 20 % oxygen in helium (2 L/h) for six hours. At the end of the plateau, the temperature was decreased as fast as possible (approximatively 10 °C/min) to avoid microstructure change during the cool down.

Irreversible transformations such as cracks can occur during the oxidation of the composite. These events are emissive and cause acoustic waves. Some of the thermogravimetric tests were done with an *in situ* coupling device (figure 5) to follow the emitted acoustic waves. Specific care should be brought to the device in order to not disturb the thermogravimetric analysis [12].



Fig.5. Acoustic emission associated to TGA measurement

The sample was deposited on an alumina waveguide in order to transmit acoustic signals from the sample to the acoustic emission transducers. The sensors (Pico type, 4 mm diameter) were linked to an acquisition chain which amplified (acquisition threshold is about 30 dB) and recorded all acoustic signals. The parameters of the acquisition chain are given in Table III:

Instrumentation	Data	Filter of the	Frequency	Model of the					
	acquisition card	preamplifier	(KHz)	amplifier					
Characteristic	PIC 2	10 - 1200	300	2/4/6 gain :					
	2 chains	KHz		$40 \ dB_{AE}$					

Table III. Parameters of the acoustic acquisition chain.

3. Results and Discussion

3.1 Preparation of 304L(p)-ZrO₂(s) composite

3.1.1 Compaction

Several pressures from 250 to 500 MPa were tested for the compression step in order to obtain the best initial green density and samples free of cracks (end capping could easily occurs during the ejection of the pellet as there was not counter pressure available on our press [13]). The optimal green density value was obtained with a pressure of 400 MPa. The geometrical density of the pellet is equal to 65 % of the theoretical density after the second step compaction pressure at 400 MPa.

3.1.2 Sintering of 304L(p)-ZrO₂(s) composite

The stainless steel and ZrO_2 shrinkage curves at 1400 °C are shown in figure 6-a and 6-b respectively. During the temperature rise, ZrO_2 exhibits a fast and important shrinkage compared to stainless steel which principally presents a thermal expansion up to 2 hours after the beginning of the thermal treatment. The metal particles embedded in the zirconia matrix which percolates are not responsible for the sintering. The ZrO_2 particles sinter around the metal ones.





Fig.6. Shrinkage curves at 1400 °C, a) 304L, b) ZrO₂.

A dilatometric curve of a composite pellet is presented in figure 7. During the temperature rise, the density of the sample increased to more than 95 % of the theoretical density. The maximum of the densification rate occurs around 1250 $^{\circ}$ C, 2 hours after the beginning of the thermal treatment, in good agreement with the zirconia shrinkage. Then, the rate of densification decreased to a stable value during the plateau. The final geometrical density reaches 96 % of the theoretical density.



Fig.7. Dilatometric curve of 304L (p)-ZrO₂(s) composite versus time of thermal treatment.

The sintering temperature of the composite was then fixed to 1350 °C which is 100 °C below the stainless steel melting point. This prevents from a liquid phase formation and any exudation phenomenon. The pellets were thus sintered at 1350 °C for 4 hours under Ar-10 % H₂ atmosphere.

To optimize the sintered density of the 304L(p)-ZrO₂(p) composite samples, a parametric study has been done considering the second step compaction pressure. Figure 8 shows the influence of the compaction pressure on the density of the composite samples sintered at 1350 °C. An optimal pressure was found at 400 MPa leading to 97 % of the theoretical density which ensures a closure of the porosity.



Fig.8. Density of 304L(p)-ZrO₂(p) composite after sintering at 1350 °C versus second step compaction pressure.

3.2 Characterization of the sintered 304L(p)-ZrO₂(s) composite

SEM observation of 304L(p)-ZrO₂(p) sintered composite is presented in figure 9. The final density of this sample, measured with a pycnometry method, is 97.6 ± 0.4 % of the theoretical density. The figure shows uniformly dispersed metal particles in the ZrO₂ matrix. The size of the metal particles is coherent with the initial stainless steel particle size. This dense sample was free of internal cracks and only small pores can be distinguished.



Fig.9. SEM observation of 304 L(p)-ZrO₂(s) sintered composite (dark grey : metal particles dispersed in ZrO₂ matrix; back dots : pores)

At the nanoscale level, TEM observations exhibited a well defined metal/zirconia interface (figure10). It is interesting to notice that the native Cr_2O_3 scale is not present at this interface. It indicates that the pre-existing chromia layer may has been dissolved in zirconia during the sintering [14]. However, the corresponding solid solution is not observed by TEM at the vicinity of the $ZrO_2/304L$ interface, probably due to a long range diffusion of Cr in ZrO_2 and/or a low concentration.



Fig.10. TEM observations of sintered 304 L(p)-ZrO₂(s) composite

The XRD diagram (figure11) of the sintered 304L(p)-ZrO₂(p) composite presents mainly peaks of the austenitic stainless steel phase and of the high temperature tetragonal ZrO₂ phase (with small amount of monoclinic ZrO₂); those peaks are not well centered to the reference peaks. This shift can be attributed to under stoechiometric zirconia. The presence of an iron phase is due to the decomposition of the alloy during the cooling at the end of the sintering cycle, the cooling speed being too low to freeze the high temperature structure. In fact we could observe using *in situ* XRD that iron dissolves in the alloy if samples are sumitted again to temperature as high as 800 °C. Finally, stainless steel is the only metallic phase to be taken into account during the oxidation treatment.



Fig.11. XRD diagram of sintered 304 L(p)-ZrO₂(s) composite.

3.3 Oxidation of the 304L(p)-ZrO₂(s) composite

Figure 12 presents the kinetic curves of 304L(p)-ZrO₂(p) composite during the oxidation at 700 °C and 800 °C. Fractional conversion (α) represents the ratio between the experimental mass obtained from TGA and the theoretical mass gain necessary to totally oxidize the metal phase. The first section of both curves (a) corresponds to the mass gain due to the presence of 50 ppm of oxygen in helium before introduction of oxygen at the plateau. In section (b), the rapid mass gain corresponds to the introduction

of oxygen and thus to the increase in the oxygen partial pressure to the desired value. Finally, section (c) presents the isothermal and isobaric part of the experiment. The fast initial increase in mass gain shows that the zirconia matrix allows the diffusion of oxygen species. Diffusion of oxygen in ZrO_2 is in fact very fast as expected with the high ionic conductivity of yttrium doped zirconia. The composite oxidation behavior is not the usual stainless steel parabolic ones nor in section (b) neither in section (c).



Fig.12. Fractional conversion (α) versus time for the oxidation of 304L(p)-ZrO₂(s) composite at 700 °C and 800 °C .

A SEM microview at low magnification of the oxidized composite oxidized at 800 °C is presented in figure 13. The surface of metallic particles directly in contact with the oxidizing atmosphere is covered with a thin oxide scale. On the other hand, in the composite core, all the metallic particles which are not directly in contact with the oxidizing atmosphere are surrounded by an oxide layer whose the thickness appears to be irregular. Many cracks in the zirconia matrix can be observed; most of them are connected to the oxidized stainless steel particles.



Fig.13. SEM observation of 304L(p)-ZrO₂(s) composite oxidized at 800 °C during 6 hours



Fig.14. a) SEM observation of oxidized 304L(p)-ZrO₂(s) composite at 800°C, b) oxygen, c) chromium and d) iron cartographies.

Figures 14-a shows a microview of the same composite at higher magnification than in figure 13. Figur14-b-c-d show O, Cr and Fe cartographies of the oxidized particles indicating that Cr is present all around the metallic particles whereas Fe is visible in the outermost part of the layer oxide.

At much higher magnification, figure 15 shows a stainless steel particle embedded in zirconia matrix for a sample oxidized with a fractional conversion α equal to 0.04. Small nodules of Cr₂O₃ are observed all around the internal metallic interface. Those nodules are characteristic of an internal isotropic growth due to an anionic diffusion mechanism.



Fig.15. SEM of oxidized at 800°C stainless steel particles in ZrO₂ for small value of fractional conversion α (white: ZrO₂, dark: Cr₂O₃ nodules grey: 304L stainless steel)

Acoustic emission (AE) was used to follow the oxidation behavior of a sample placed in the thermobalance. The mass gain was monitored by TGA simultaneously with the acoustic waves created during the oxidation. The AE signals together with the mass gain curve are shown in figure 16. It can be seen that the signal due to acoustic waves begins to increase as soon as the mass gain increases. Intense acoustic emission signals with a large number of counts occurred during all the oxidation treatment (figures 17-18) with a sharp increase in the part (b). The first initial increase in AE signal associated to the initial oxidation steps can be correlated to the oxidation process occurring at the $ZrO_2/304L$ interface (nucleation and growth of Cr_2O_3 nodules). In the last part of the oxidation plateau, the increase in the AE signal was most probably due to cracks initiation and propagation in the composite. If we consider the Pilling and Bedworth ratios for Fe/Fe₂O₃ (2.10) and Cr/Cr₂O₃ (2.07), the

volume expansion of the metal particles due to oxidation induces stresses in the ceramic matrix which produces cracks. Similarly, a high AE signal is observed during the cooling of the sample, due to the difference in thermal expansion coefficients between ZrO₂, the oxide layer and the metal phase.



Fig 16. Thermogravimetry associated with acoustic emission of 304L(p)-ZrO₂(s) composite during oxidation under 20% O₂ at 800°C.



Fig 17. Acoustic emission (AE) amplitude of counts versus time.



Fig 18. Parameters and wave form of a 38dB count.

Auger electron spectrometry analysis (AES) (figure 19-a-b) of the ZrO₂/304L interface region in the oxidized composite gave additional information on the oxidation layer; figure 19-a is a microview of the interface directly obtain with the secondary electron detector connected to the Auger spectrometer. The AES profile (figure 19-b) describes the elements present along a straight line starting from the stainless steel particle, passing through the oxide layer and ending into the zirconia matrix. Iron, chromium and nickel were the major elements found in the metallic phase, as expected. At the metal-oxide interface, the amount of chromium increases, whereas those of iron and nickel decrease. Deeper in the oxide layer, it can be observed a decrease in the amount of chromium and an increase in the iron amount and, at a lesser extent, in the amount of nickel. Then near the zirconia/oxide interface, the oxide layer is rich in iron. While passing the zirconia phase, metallic elements are no more detected thus no diffusion of the oxide cations in the zirconia matrix can be suspected.



a) SEM picture



b) AES profile



To ascertain the composition of the oxide layer, XRD analysis (figure 20) has been performed on a composite sample oxidized at 800 °C. The diffraction peaks correspond to various oxide phases: Fe_2O_3 , $(Fe_{0.6} Cr_{0.4})_2O_3$ and tetragonal ZrO₂. Tetragonal ZrO₂ peaks are well centered; this shows that, as expected, ZrO₂ is stoechiometric after oxidation.



Fig.20. XRD diagram of oxidized 304L(p)-ZrO₂(s) composite at 800 °C.

This assumption was verified using thermogravimetric analysis (figure 21). A pure yttrium stabilized ZrO_2 pellet, sintered as the composite in the H₂ reductive atmosphere, was summit to an oxygen exposure (20 % in helium) at 800 °C. After oxidation, we observed a mass gain and the black color of the sintered pellet disappears for a white one. The sintered ZrO_2 passes from the under stoechiometric state to the stoechiometric one. Base on the TGA result and on equation E1, we can calculate the value of the non-stoechiometry x : x = 0.03.

$$Y_{0.03}Zr_{0.97}O_{2-x} + xO_{2} = Y_{0.03}Zr_{0.97}O_{2}$$
(E1)



Fig.21. Thermogravimetric analysis of TZ-3Y-E ZrO₂ sintered pellet during an oxidation at 800 °C.

4. Discussion

From TGA, SEM, AES and XRD characterization of the oxidized 304L(p)-ZrO₂(s) composite, an oxidation scenario can be proposed. During the initial oxidation period (section (b) in figure 12: high kinetic rate), at the sample surface, a protective chromia scale appears on the metal particles. In the core of the composite, nodules of Cr₂O₃ start to nucleate and grow around the stainless steel particles to form an inward layer. Such a non protective scale is produced both because the metal particles are not covered with a thin native and protective oxide layer after the sintering process and because the oxygen diffusion through ZrO₂ is fast even in the absence of cracks at the beginning of the oxidation process. Then, after the appearance of cracks due to the volume expansion of the oxide layer, a complex

multiphase layer, mainly based on iron and chromium, develops outside the initial discontinuous layer $(Cr_2O_3 nodules)$ all around the metal particles.

We can calculate the critical thickness X* of the oxide layer around the metal particle (radius r) which can induce a critical stress σ^* and the fracture of the zirconia matrix using equation 2 [15] and the mechanical parameters of stabilized ZrO₂:

$$\sigma^* = \frac{E}{(1-\nu)} \left(\frac{r}{L/2}\right)^3 \frac{(1-1/R_{pb})}{r} X *$$
(E2)

where R_{pb} is the Pilling and Bedworth ratios for Cr/Cr₂O₃ (2.07); zirconia Young's modulus E and Poisson's ratio v are respectively 180 GPa and 0.34, zirconia fractured stress σ_f is 500 MPa and L is the average distance between the metal particles (50 µm). In the very beginning of the composite oxidation process, if the oxide layer thickness exceeds 0.05 µm, the stress exceeds also σ_f and cracks occur in the ZrO₂ matrix.

The oxidation of pure 304L stainless steel plate [16] in oxygen at high temperature usually shows parabolic behavior; the oxide grows outwards with an initial external Cr_2O_3 layer followed by growth of $(Cr, Fe)_2O_3$ layers with variable Cr and Fe proportions. The oxidation of the 304L(p)-ZrO₂(s) model composite begins with an internal growth of Cr_2O_3 nodules. There is an inward diffusion of oxygen anions in the Cr_2O_3 layer instead of a chromium cations outward diffusion. Due to well defined interfaces between ZrO_2 and the metal particles in the sintered composite, the stabilized zirconia matrix blocks the outward growth of chromium oxide until the appearance of cracks. This phenomenon was also reported in literature [15] for the oxidation of ZrO_2 composites dispersed with Ni particles.

5. Conclusion

Such a comprehensive study has put in evidence an unusual behavior of a stainless steel alloy inserted in a dense Y_2O_3 partially stabilized ZrO₂ matrix with respect to oxidation. An optimized powder metallurgy process was used to produce composites dispersed with metal particles with 97% of the theoretical density. The oxidation under 20% O_2 in He of the 304L(p)-ZrO₂(s) composite at 700°C and 800°C exhibits a different corrosion behavior than pure 304L stainless steel, beginning with an internal growth of Cr₂O₃ nodules instead of a continuous external chromium oxide layer. The zirconia matrix plays a major role on the corrosion behavior of the composite. Some additional works are in progress to deeper study the kinetics and the mechanisms of oxidation of the model 304L(p)-ZrO₂(s) composite. It would be interesting to test the composite resistance to oxidation by changing the stabilized ZrO₂ matrix with MgO matrix. Lower oxygen diffusion coefficient would lead to a better corrosion resistance. Moreover oxidation behavior may change with the amount of open porosity in the composite.

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