High temperature chromium volatilization from Cr2O3 powder and Cr2O3-doped UO2 pellets in reducing atmospheres

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Abstract
Chromium volatilization from Cr₂O₃ powder and Cr₂O₃-doped UO₂ pellets during sintering in reducing atmospheres has been studied by thermogravimetry (TG) coupled with differential thermal analysis (DTA) up to 1700°C. The sintering of Cr₂O₃-doped UO₂ pellets was also followed by dilatometry. Oxygen partial pressures in the range 10⁻²⁰-10⁻¹¹ atm (10⁻¹⁵-10⁻⁶ Pa) have been fixed in all the experiments thanks to mixtures of hydrogen and carbon dioxide. A linear heating rate of 20°C min⁻¹ was applied to all the experiments. The dopant amount was in the range 0.18-0.9 mol% Cr in UO₂. For all the oxygen potentials, the mass loss of Cr₂O₃ powder was found to start at temperatures as high as 1470°C due to Cr₂O₃ dissociation, the lower the oxygen potential, the lower the starting temperature and the higher the volatilized amount. For intermediate oxygen potentials, an exothermic DTA peak observed during cooling, from 1700°C to room temperature, attested for the crystallization of a liquid phase which was attributed to CrO₃ according to thermodynamic predictions. Then, the dilatometry and TG studies allowed following the behavior of Cr₂O₃-doped UO₂ pellets. The mass loss at temperatures higher than 1470°C was attributed to chromium volatilization for all the doped samples. During the sintering of doped UO₂ pellets, the liquid phase CrO₃ seemed to appear at a lower oxygen potential than in Cr₂O₃ powder, which probably contributed to enhance the densification rate. For the highest dopant amount, 0.9 mol% Cr, the volatilization process was found to be rather similar to that of Cr₂O₃ powder, due to the part of chromia not solubilized in the UO₂ crystal. Moreover, as the initial pellets were not dense, as long as the pellet porosity remained open, the chromia particles were able to dissociate as in the Cr₂O₃ powder. Thus the volatilization of chromium from doped UO₂ pellets under sintering in reducing atmospheres must be understood as the result of several phenomena whose contribution depends on temperature, oxygen potential and heating rate: before the porosity closure, both dissociation of chromia into Cr₉O₉ and oxygen from excess Cr₂O₃ particles, and Cr volatilization from doped UO₂ particles; then chromium volatilization from the doped UO₂ ceramic during further densification process.

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I. Introduction
Upgrading the economics of light-water reactor plants leads to increased demands on reactor maneuverability with continued enhancement of fuel performance and reliability. Research on fuel pellets focuses on the ability of large grain and visco-plastic UO₂ microstructure to increase fission products retention and pellet mechanical compliance [1]. High expectations are placed on the optimized chromia-doped UO₂ fuel developed by Areva NP which exhibits significantly higher performance compared to the reference UO₂ fuel [2]. In the first development step started in the 1990’s, the fundamental mechanisms governing UO₂ doping with Cr₂O₃ were clarified. The sintering conditions have been optimized to enhance chromia dissolution in the UO₂ matrix and to increase the solubility limit of chromium into UO₂ [3] and [4]. To master the doped UO₂ fuel fabrication technology, a large industrialization program has been launched to study the evolution of the pellet characteristics as a function of various parameters: U₃O₈ recycling, pellet sintering conditions, etc. Optimized manufacturing conditions are now defined to ensure the reproducibility of the targeted microstructure changes with a reduced scattering inside a pellet lot or between lots.

In parallel, fundamental researches are in progress such as the present study which aims to understand in depth the behavior of Cr₂O₃ at high temperature. Toker et al. [5] studied the thermodynamic properties and phase relations of the Cr-O system at temperatures from 1500 up to 1825°C. The outcome thermodynamic diagram, shown in Figure 1, defines domains of existence for several phases Cr₁(s), Cr₂O₃⁶(h), Cr₃O₄(d) and CrO₂(l) versus temperature and oxygen potential (oxygen partial pressures in the range of 10⁻²⁰ - 10⁻¹⁴ atm/10⁻¹⁵ - 10⁻⁶ Pa). Johnson and Muan [6] also studied a binary phase diagram from mixtures of Cr₂O₃ and Cr powders in a purified argon atmosphere and showed the existence of an eutectic at 1665°C. This value is closed to the one found previously by Ol’Shanskii and Shlepov [7] who reported an eutectic temperature at 1660°C. In such high temperature and low oxygen pressure conditions, chromium is suspected to "evaporate" from the solid phase; however at our knowledge, neither mechanism nor quantitative data on this phenomenon in reduced atmospheres are available. This article presents first a thermogravimetry study coupled with differential thermal analysis (TG-DTA) of a chromia powder in the range of temperatures 1100-1700°C in reducing atmospheres (oxygen pressure in the range of 10⁻²⁰ - 5 x 10⁻¹¹ atm/10⁻¹⁵ - 10⁻⁶ Pa). Then the behavior of chromia-doped UO₂ pellets followed by TG and dilatometry in the same thermodynamic conditions is discussed with the help of the results obtained with the Cr₂O₃ powder.

II. Experimental
II.1. Materials
The Cr₂O₃ powder (Elementis) used in this study, has a purity grade of 99.794%, the main impurities are calcium (410 ppm) and silicium (1100 ppm). Its specific surface area has been determined by the BET method with the Kr adsorption isotherm at a temperature of 75 K. A value of 6.0 m² g⁻¹ was found, to which corresponds an average diameter of 0.19µm for primary particles according to the assumption that Cr₂O₃ particles are spherical. The chromia-doped UO₂ pellets (noted Cr-doped UO₂ in the following) used for the TG analyses are small disks with a reduced thickness of ~2 mm and a diameter of ~9 mm. Such geometry is required to avoid edge effects and consider a planar symmetry. The samples have been prepared by Areva NP (manufacturing plant of FBFC Dessel, Belgium) from depleted UO₂ powder converted by the dry route process and without any other additives. This UO₂ powder is slightly overstoichiometric (UO₂.₀₆₉) and its purity grade is 99.9954%. The specific surface area of this powder is equal to 2.4 m² g⁻¹ with an equivalent particle diameter of 0.23µm. The chromia powder is directly added to UO₂ at the blending step. After pre-compaction and granulation operations, the pellets have been compacted to a green density level of ~6 g cm⁻³.

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The doped UO$_2$ samples studied by dilatometry were manufactured according to the same process and same materials. However, sample pellets with a cylinder form (–7 mm thickness and –10 mm diameter) were required to obtain a measurable signal during densification. For the present work, several Cr$_2$O$_3$ contents have been studied: undoped UO$_2$, UO$_2$ + 0.18 mol% Cr (589 ppm Cr$_2$O$_3$), UO$_2$ + 0.44 mol% Cr (1419 ppm Cr$_2$O$_3$) and UO$_2$ + 0.9 mol% Cr (2878 ppm Cr$_2$O$_3$).

II.2. Experimental methods

II.2.1. TG and DTA experiments

The thermogravimetry (TG) experiments have been performed with a symmetrical balance (SETARAM TAG 24) equipped with a differential thermal analysis (DTA) device (tungsten crucibles and W-5%Re/W-26%Re thermocouples) appropriated to high temperature conditions (maximum 1700°C) and to reducing atmospheres. The device is connected to a gas panel equipped with H$_2$ and CO$_2$ mass flowmeters to fix the desired amount of CO$_2$ in hydrogen: 0.6% CO$_2$, 1.11% CO$_2$, 1.47% CO$_2$, 2.06% CO$_2$ and 3.2% CO$_2$. Before introduction of the gas mixture, the device is evacuated to 10$^{-4}$ atm (10 Pa) to eliminate residual gases. The CO$_2$ content in the exit gas was ascertained by infrared spectrometry at room temperature. The measured error was ±0.02% CO$_2$ in the mixture. The oxygen potentials equivalent to the H$_2$/CO$_2$ mixtures were obtained according to Deines et al.’s data [8]; they have been reported as a function of temperature in Figure 1a; it can be seen that the lines corresponding to the experimental conditions cross interesting domains of the thermodynamic diagram. The estimated error on the oxygen potential, based on the CO$_2$ content measurement, corresponds to ±250 J mol$^{-1}$. The experiments have been conducted up to 1700°C with a heating rate of 20°C min$^{-1}$ followed by a soaking time of five minutes to compensate the thermal inertia of the furnaces and a cooling rate of 30°C min$^{-1}$. As a consequence, the very last part of the DTG and DTA curves will not be commented.

To reach uniform thermal and gaseous composition conditions within the powdered samples, the TG-DTA experiments were achieved with ~3 mg of Cr$_2$O$_3$ which corresponds to a powder bed as thin as of ~0.5 mm in the crucible. TG-DTA blank tests were performed without any sample in the crucible to monitor the baseline of the mass and DTA signals. The mass signal exhibited a continuous drift which did not exceed 35µg from 1400°C to 1700°C. The mass loss curves were corrected by subtraction of this baseline. The reproducibility of the mass loss curves versus temperature was good: ±0.3 wt%. Concerning the DTA curves, the onset and maximum peak temperatures were reproducible with an accuracy of ±1.5°C. However some problems of reproducibility were encountered in the case of DTA since for two identical experiments, the shape of the signals was not rigorously reproducible, probably due to the small amount of sample in the crucible. To follow the chromium volatilization from Cr-doped UO$_2$ pellets, TG experiments were done placing a pellet of ~0.8 g in a tungsten crucible suspended to a tungsten suspension (it was not possible to use the TG-DTA device due to the limited amount of sample in the DTA crucible).

II.2.2. Dilatometry tests

All the dilatometry tests were performed at the Areva NP Ceramic Laboratory in Erlangen (Germany) by means of a Netzch device (DIL 402 C) allowing to work up to 2000°C under H$_2$/CO$_2$ mixtures. The shrinkage measurement resolution, gave by the provider, is 0.125 nm/1.25 nm but the operational shrinkage precision was not better than 1µm. The tests were conducted up to 1700°C in the same atmospheres than those used in the thermobalance, with a heating and a cooling rate of 20°C min$^{-1}$. The results were analyzed by reference to the theoretical density which is 10.96 g cm$^{-3}$ for UO$_2$, and respectively 10.95, 10.94 and 10.93 for the 0.18%, 0.44% and 0.9 mol% Cr doped samples.
III. Results and discussion

III.1. TGA–DTA study of Cr₂O₃ powder

Figures 2a-e shows the experimental curves of mass loss (TG), derivative of mass loss (DTG) and heat flow (DTA) obtained for the Cr₂O₃ powder under the various H₂/CO₂ atmospheres. Neither mass loss nor DTA signals were observed below 1400 °C, so the results are presented between 1400 °C and 1700 °C. The mass loss due to the chromium volatilization was observed to begin in the temperature range 1430-1660 °C, depending on the oxygen potential, the higher the oxygen potential, the higher the onset temperature. All the experiments except for the highest oxygen potentials (3.2% CO₂) exhibited an endothermic signal during the heating treatment. An exothermic signal was observed during cooling for the experiments done with 1.47% CO₂ and 2.06% CO₂ mixtures (cf. Figures 2c and d). The results of the TG and DTA tests (onset and peak temperatures, final mass loss) are summarized in Table 1. The onset temperature for TG and DTA signals during heating are quite similar for 0.6% and 1.11% CO₂, whereas the DTA endothermic peak starts before the TG mass loss for 1.47% and 2.06% CO₂. It can be noticed that the higher the oxygen potential, the lower the total mass loss (cf. 4th line of Table 1). In order to explain the TG-DTA results, we have listed in Table 2 the most probable reactions that could occur considering the various condensed phases appearing in the thermodynamic diagram (Cr₂O₃(s), Cr(s), Cr₃O₄(s) and CrO₃(l)):

- dissociation of Cr₂O₃ into gaseous or solid metallic chromium and oxygen (reactions R1 or R2, respectively),
- chromium evaporation from solid metallic chromium (reaction R3)
- decomposition of Cr₂O₃ into CrO₃ (reaction R4),
- decomposition of Cr₂O₃ into gaseous or solid metallic chromium and oxygen (reactions R5 or R6, respectively),
- decomposition of Cr₂O₃ into liquid CrO and oxygen (reaction R7),
- decomposition of CrO₃ into gaseous chromium and oxygen (reaction R8),
- decomposition of CrO₃ into CrO₃l and oxygen (reaction R9),
- vaporization of CrO₃ into gaseous CrO (R10),
- reaction of Cr(l) with Cr₂O₃ to form liquid CrO (R11).

For each reaction the values of the corresponding standard enthalpy variations and Gibbs energy variations calculated at 1400°C and 1700°C [9] and [10] have been reported in Table 2 as well as the theoretical mass losses at total conversion. In Figure 1a have been reported the temperatures observed for the DTA peak onsets during heating (dots) and the mass loss onsets (triangles). It can be seen that for all atmospheres, the mass loss and the endothermic effect begins well before the limits of the phase diagram. This means the volatilization process begins to be detected without appearance of a new solid or liquid phase. Moreover, the onset temperature increases when the oxygen potential increases.

Figure 3 represents the equilibrium pressures of the gaseous species Cr(g), CrO(g), CrO₂(g), CrO₃(g) as a function of temperature, calculated considering the change in oxygen potential with temperature in the atmosphere 1.11% CO₂ (cf. line 2 in Figure 1a) and for the various condensed phases of Toker’s diagram. It can be seen that gaseous chromium is the most probable species that can be released whatever the condensed phase would be; considering the other atmospheres, the calculations (results not shown) lead to the same observation. Moreover Figure 4, which represents for reaction R1 the gaseous chromium equilibrium pressure vs. temperature for the various atmospheres, indicates that the higher the content in CO₂, the lower the Cr(g) partial pressure. It can thus be inferred from these thermodynamic considerations that the beginning of TG and DTA curves of Figure 2 corresponds to the dissociation of chromia according to reaction R1, in agreement with a high standard enthalpy value (1891 kJ mol⁻¹ at 1400°C). When the temperature increases, the equilibrium pressure in
Cr(s) increases, and so does the mass loss. The \( \text{Cr}_2\text{O}_3 \) dissociation process will continue provided the surface of the chromia particles remains in contact with the gas phase.

In the case of the most reducing atmosphere (0.6% CO\(_2\)), the thermodynamic diagram predicts that chromia would start to dissociate into solid metallic chromium \( \text{Cr}(s) \) and oxygen according to reaction R2 at 1470°C; since the final mass loss was found to exceed 31%, such a transformation must take place in parallel with R1 until chromia particles become totally covered with metallic chromium. Obviously mass loss due to reaction R3 has to be considered as soon as \( \text{Cr}(s) \) appear at the surface of the oxide particles (cf. Figure 2a). The large difference between the enthalpies of reactions R2 and R3 probably explains the difference in the peak temperatures of both DTG and DTA signals.

For the 1.11% CO\(_2\) atmosphere, in addition to reaction R1 which is detected from 1499°C from TG and 1487°C from DTA (cf. Figure 2a), the formation of \( \text{Cr}_2\text{O}_4(s) \) phase is expected at a temperature around 1650°C according to Toker et al. thermodynamic diagram. Above 1650°C four different reactions can occur: the formation of \( \text{Cr}_2\text{O}_4(s) \) (R4), the dissociation of \( \text{Cr}_2\text{O}_4(s) \) in gaseous chromium (R5), the dissociation of \( \text{Cr}_2\text{O}_4(s) \) in solid chromium (R6) and the vaporization of solid chromium (R3). Reactions R4 and R5 could start at 1650°C against 1665°C for reactions R3 and R6. In this temperature range, the corresponding mass losses should be low because during the experiment, the \( \text{Cr}_2\text{O}_4(s) \) domain is crossed during 10-20 s only, due to the high heating rate (20°C min\(^{-1}\)). In consequence the formation of \( \text{Cr}_2\text{O}_4(s) \) should be rather limited; that could explain why the shape of the DTA and mass losses signals observed in Figure 2b are very similar to those of Figure 2a. When the temperature reaches 1665°C, reaction R2 may occur and subsequently reaction R3, both responsible for the mass loss. The final mass loss is lower than in the most reducing atmosphere since the reactions R2 and R3 are shifted at higher temperatures and the \( \text{Cr}(s) \) equilibrium pressure is lower.

With the 1.47% CO\(_2\) atmosphere (cf. Figure 2c), similarly to the previous atmospheres, the phenomenon observed to begin at 1526°C for the mass loss and at 1470°C for the endothermic DTA signal should correspond to the reaction R1. Then, according to the thermodynamic diagram, two new phases can be considered: \( \text{Cr}_2\text{O}_4(s) \) that can be formed by reaction R4 (chromia decomposition) and \( \text{CrO}_3(l) \), that can be formed by reactions R7 (\( \text{Cr}_2\text{O}_4 \) decomposition) and R9 (\( \text{Cr}_4\text{O}_7 \) decomposition). These three reactions might all contribute to the mass loss and DTA signals, and in addition, the dissociation (R8) of the liquid phase \( \text{CrO}_4(l) \) and its vaporization (R10) may also be involved. However, no particular feature could be noticed on the curves of Figure 2c. Thus as previously proposed, the period in the \( \text{Cr}_2\text{O}_4 \) domain is too brief to consider a significant contribution of reaction R4 to the mass loss. During the cooling, an exothermic peak without mass change has been identified which starts at 1663°C. This indicates a crystallization phenomenon from a liquid phase, i.e. the presence of \( \text{CrO}_3(l) \) seems to be confirmed. Due to their low standard enthalpies, reactions R7 and R9 (cf. Table 2) do not produce any characteristic peak on the endothermic DTA signal during heating.

In the 2.06% CO\(_2\) atmosphere, according to Figure 1, only the dissociation of \( \text{Cr}_2\text{O}_3 \) can be envisaged. However, an exothermic signal has been observed during the cooling, at 1662°C, indicating again the crystallization from a liquid phase. This means that \( \text{CrO}_3(l) \) can be produced at lower temperature than predicted by the thermodynamic diagram (Figure 1) since according to Toker’s data the \( \text{CrO}_3(l) \) phase should form at 1716°C. First, experimental errors on temperature or oxygen potential values have been considered to explain such difference from thermodynamic prediction. Considering the tungsten thermocouple precision (1.5°C), a temperature error can be excluded. As illustrated by Figure 1b, the error on the oxygen potential cannot explain the difference between the experimental oxygen potential and the \( \text{Cr}_2\text{O}_4(s)/\text{CrO}_3(l) \) equilibrium value at 1700°C. Another possible explanation could be the effect of impurities, as reported by Li et al. [11] who studied the sintering behavior of \( \text{Cr}_2\text{O}_3 \) in presence of \( \text{ZrO}_2 \). The thermodynamic diagram for the Cr-O system calculated for a \( \text{CrO}_3(l) \) activity of 0.8 reveals a larger domain of existence of \( \text{CrO}_4(l) \) than that proposed by Toker. So since the chromia powder contains silicium and calcium, their effect on the temperature of appearance of \( \text{CrO}_3(l) \) cannot be excluded.

In the case of the less reducing atmosphere 3.2% CO\(_2\), the mass loss has been observed to start at 1659°C and there is no visible DTA signal. The mass loss seems to correspond to the
dissociation of \( \text{Cr}_2\text{O}_3 \) (reaction R1), the equilibrium metallic chromium partial pressure being
\(5.41 \times 10^{-5} \text{ atm} \) at 1659°C. The mass loss is very low even at 1700°C, which is in agreement with
a lower equilibrium partial pressure for higher oxygen potentials, as previously discussed and
illustrated by Figure 4. For this atmosphere, the conditions at 1700°C are quite far from the
thermodynamic predictions for the \( \text{CrO}_6 \) domain (1750°C), which is consistent with the
absence of a DTA exothermic signal during the cooling, probably due to an insufficient
impurities effect.

III.2. Cr-doped UO\(_2\) pellets

III.2.1. Dilatometry results

Table 3 reports some characteristic temperatures and density values obtained in all the
dilatometry experiments with the undoped and 0.18%, 0.44% and 0.9% mol Cr-doped UO\(_2\)
pellets. Figure 5a represents the dilatometry curves vs. temperature for a sintering cycle in
2.06% CO\(_2\) atmosphere, and the corresponding derivative curves showing the rate of
densification are shown in Figure 5b. All the doped samples show a delay in the densification
compared to UO\(_2\). It is worthwhile to notice for doped samples the presence of a slight
acceleration of the densification rate at temperatures around 1250°C. The temperature of the
corresponding maximum has been reported in the 4\(^{th}\) line of Table 3. This phenomenon, which
is absent of the curves with the undoped UO\(_2\), will be discussed in the next sections. The effect
of the dopant on the final density, reported in the 6\(^{th}\) line of Table 3, appears to depend both on
the amount of dopant and on the atmosphere composition. The temperature corresponding to
94-95% of the theoretical density was seen to depend on the gaseous atmosphere, as illustrated by
Figure 6 which represents the derivative of the density vs. density obtained with undoped and
0.9 mol% Cr-doped UO\(_2\) for 0.6%, 1.47%, 2.06% and 3.2% of CO\(_2\). On each graph of Figure 6,
the vertical line corresponds to 95% of the theoretical density, which is considered as the
theoretical limit of existence of open pores in ceramics; beyond that value, it can be considered
that there is no longer open porosity in the pellets. For 0.6% CO\(_2\) the porosity remains open
even at 1700°C whereas for 1.47%, 2.06% and 3.2% CO\(_2\) the closure of the porosity is found
near 1700°C. A maximum in the densification rate appears in the case of the doped samples
near 1660°C for 1.47% CO\(_2\) and 2.06% CO\(_2\) and near 1680°C for 3.2% CO\(_2\). These temperatures
correspond rather well to those of the porosity closure. This will be discussed later in Section
III.3.

III.2.2. TG analysis at H\(_2\) + 1.47% CO\(_2\)

The curves of mass loss from room temperature to 1700°C in the 1.47% CO\(_2\) atmosphere are
shown in Figure 7a for 0.9 mol% Cr-doped and undoped samples. Figure 7b represents the rate
of mass loss deduced from the curves of Figure 7a. During the heating up to around 1400°C,
after a first mass loss (peak at 155°C and 270°C) due to water desorption, a second mass loss
(peak at 420-440°C) is observed as the result of the UO\(_{2+x}\) reduction. The reduction process is
achieved at about 1480°C for both samples. This can easier be observed on the curves of
Figure 8 which represents an enlargement of the TG and DTG signals in the region 800-
1700°C for undoped and doped samples (0.18, 0.44 and 0.9 mol% Cr) in 1.47% CO\(_2\)
atmosphere. In addition, in the DTG curves of Figure 8b, it appears a small shoulder close to
1200°C on the high temperature side of the large peak of the end of the UO\(_{2+x}\) reduction. Such a
feature must be compared to the acceleration in the densification previously mentioned in
Section III.2.1. A significant increase in the mass loss rate of the doped sample begins when the
temperature exceeds 1480°C. This temperature is nearly the same as that observed for the
beginning of chromia volatilization in the TG/DTA experiments of Cr\(_2\)O\(_3\) powder sample at
1.47% CO\(_2\). By analogy to the chromia powder results, this phenomenon thus can be mainly
attributed to the dissociation of chromium oxide Cr\(_2\)O\(_3\) (reaction R1).
III.3. Discussion

III.3.1. Volatilization from Cr₂O₃ powder

The TG–DTA experiments with chromia powder have shown that chromia may dissociate releasing metallic gaseous Cr (and oxygen) as the major volatile species whatever the atmosphere composition and the temperature could be. According to the oxygen potential and the temperature, other reactions should contribute to the volatilization process, as for example the vaporization of solid metallic chromium or liquid CrO₃. The presence of CrO₃ in intermediate atmosphere compositions (1.47% and 2.06%) has been attested by an exothermic peak in the DTA signal during cooling. The presence of chemical impurities such as Ca and Si may explain the appearance of the liquid phase CrO₃ at temperatures lower than that expected by the thermodynamic predictions. The amount of Cr volatilization depends of temperature and oxygen potential in the same way as does the CrO₃ equilibrium partial pressure of the considered reactions. Such conclusions will help to understand the behavior of Cr-doped UO₂ pellets during similar conditions of temperature and oxygen potential.

III.3.2. Solubilization of Cr₂O₃ in UO₂

The doped samples show a delay in densification compared to undoped UO₂ for each studied atmosphere. Bourgeois [12] observed this phenomenon during the sintering of UO₂ doped pellets which contain 0.31 mol% of Cr and using the following experimental conditions: 1%H₂O in hydrogen, heating rate of 2.5°C/min, use of spray-dried UO₂ powder. He explained this delay in densification by the development of a Cr₂O₃ film around the UO₂ particles. Delafoy [13] described the same effect, which increases with chromium content. So Cr₂O₃ fine particles dispersed in the UO₂ pellet probably inhibit the intergranular diffusion of ions during the first stage of sintering. Another mechanism can be considered, the surface diffusion of chromium cations on the UO₂ grains surface can also delay the intergranular diffusion.

As we previously noticed in the Section III.2.2 dedicated to the thermogravimetry experiments with doped pellets (for the 1.47% CO₂ atmosphere), it is interesting to compare the densification rate with the rate of mass loss. The maximum of the mass loss rate at 1200°C is close to that of the densification peak at 1157°C. These peaks could be due to the chromia solubilization in UO₂ accompanying the final step of the UO₂₂⁺ reduction. The mass loss rate difference when compared to the undoped UO₂ is probably due to the modification of the point defects nature and concentration in the doped UO₂ crystal structure. Similarly these defects contribute to the increase in the sintering rate. The slight differences between the peak temperatures of the DTG and densification rate curves probably come from the difference in the shape of the samples used in both experimental devices. The limit of solubility of chromia into UO₂ has been studied by several authors [4], [13], [14] and [15]. According to the Riglet-Martials data [15], it is possible to calculate the solubility for every condition of temperature and oxygen potential. The maximal fraction of dopant in solution into UO₂ in the case of 0.44 mol% Cr, and only 30% in the case of 0.9 mol% Cr. At 1640°C, these values move to 80% for 0.44 mol% Cr, and only 40% in the case of 0.9 mol% Cr. Thus, in reality, only a part of chromia enters in solution into the UO₂ crystal, the other part remaining in the form of Cr₂O₃ particles dispersed in the pellet. Since during the temperature increase, the porosity remains open during a given period, chromia is expected to dissociate according to the same process as that observed in the case of Cr₂O₃ powder. This certainly explains the similarity between the mass loss curves observed in Cr₂O₃ powder and Cr-doped UO₂ pellets. This is especially clear in the case of the 0.9 mol% Cr doped sample for which the mass loss rate continuously decreases after 1500°C up to 1700°C (Figure 8b). For a fixed oxygen potential, the higher the temperature, the higher the solubility. Considering the rate of mass loss and densification curves, it can be inferred that for kinetic reasons the solubilization process of Cr₂O₃ occurs majoritarily near 1200°C, then the "excess" chromia may be progressively incorporated in UO₂ as far as the solubility limit increases with temperature. However, the fraction of chromia which remains as Cr₂O₃ particles cannot be neglected with respect to their contribution to the volatilization process.
III.3.3. Sintering behavior of undoped and Cr-doped UO.

For all the atmosphere compositions, a second acceleration of the densification occurs giving a maximum rate in the range 1500-1600°C, the higher the dopant amount, the higher the densification rate. Such an acceleration has to be compared to Chevrel's study [16] showing a similar effect during the sintering of overstoichiometric UO$_2$ containing numerous Willis type defects which are known to enhance the ions diffusion and thus accelerate the densification. In the present study, this phenomenon can be related to an increase in the Willis defects concentration as the result of chromium insertion at uranium sites of UO$_2$ crystals and distortion of the oxygen lattice around chromium ions [17]. Chromium ion in solution into UO$_2$ takes III + valency and disturbs a lot the chemical defects in the UO$_2$ matrix, producing mainly Willis type defects. For the 0.6% CO$_2$ atmosphere, in agreement with the thermodynamic diagram proposed by Toker et al. [5], the dissociation of Cr$_2$O$_3(s)$ in Cr$_{(g)}$ occurs at 1473°C (R2). Consequently, the presence of metallic chromium precipitates can pin the grain boundaries [12] explaining the low final density of the sample compared to the less reducing atmospheres. In the doped pellets the Cr volatilization process is expected to depend of the competition between the sintering and solubilization rates since the "excess" Cr$_2$O$_3$ particles will behave differently before and after the closure of the porosity. Before this point, the surface of the Cr$_2$O$_3$ particles is in contact with the surrounding atmosphere allowing Cr$_{(g)}$ transport in the gas phase, and beyond it, the volatilization of chromium necessitates bulk or/and grain boundary diffusion inside the UO$_2$ ceramic matrix.

III.3.4. Liquid phase

For the 0.9 mol% doped sample and the 1.47% and 2.06% CO$_2$ atmospheres, the densification rate curves exhibit a second peak close to 1660°C. Based on microstructure examinations, Bourgeois [12] explained such a phenomenon by the hypothesis of a liquid phase for the high Cr$_2$O$_3$ contents. TG-DTA results on chromia powder, for 1.47% CO$_2$ and 2.06% CO$_2$, validate this interpretation due to the exothermic peak associated to the crystallization of a liquid phase detected during the cooling period. For the atmosphere 3.2% CO$_2$, and 0.9 mol% Cr doped sample, the dilatometric curve of the sintered doped pellet also shows a second acceleration of the densification, but at a higher temperature, near 1680°C. However in this case any exothermic DTA peak was observed during the cooling of the Cr$_2$O$_3$ powder. Open porosity can disappear when the ceramic density exceeds 95% $d_{th}$, in the doped pellet the closure can begin around 1680°C. At this temperature, the decomposition of Cr$_2$O$_3(s)$ in Cr$_{(g)}$ can take place with a chromium partial pressure close to $1 \times 10^{-3}$ atm. Thus Cr$_{(g)}$ could be trapped in the closed porosity and react with Cr$_2$O$_3(s)$ to form liquid CrO$_{(l)}$ according to reaction R11. Moreover, considering the experiments done in the dilatometer furnace, the thermodynamic conditions may not be strictly the same outside and inside the sample just after the closure of the porosity. This may also contribute to the formation of the liquid phase. The appearance of the CrO$_{(l)}$ liquid phase thus favors the sintering process and, as previously explained with the porosity closure effect, it may play indirectly a significant role in the chromium volatilization process.

III.3.5. Volatilization process from Cr-doped UO$_2$ pellets

The volatilization process from Cr-doped UO$_2$ pellets may be explained due to the comparison of the results obtained with the Cr$_2$O$_3$ powder. During the high temperature treatment from 1000 to 1700°C, the following phenomena have to be considered:

- A part of the added chromia solubilizes with a maximum rate around 1200°C, then the solubilization process may continue as far as the temperature increases, in relation to the corresponding solubility limit.

- At temperatures higher than 1480°C, and in all atmospheres, as far as the porosity remains open, the volatilization process consists of two contributions: one is the dissociation of excess chromia (not yet solubilized chromia) with a behavior similar to that of Cr$_2$O$_3$ powder (reaction R1); a second smaller contribution may come from Cr$_{(g)}$ volatilization from effectively Cr-doped UO$_2$ particles (soluble Cr) with a kinetic
behavior expected to be different from that of the chromia particles; indeed, in that case, the volatilization process needs chromium to diffuse into the UO$_2$ particles.

- After the porosity closure, which has been shown to be influenced by the appearance of the CrO$_6$ liquid phase, only one process should be considered: the volatilization needs the chromium to diffuse into the UO$_2$ matrix.

IV. Conclusions

The behavior of a chromium oxide powder at high temperature and under reducing atmospheres has been studied in details by combining DTA and thermogravimetry experiments. Experiments with a linear temperature increase up to 1700°C under various oxygen potentials where done to follow the chromia volatilization process. For all conditions, the results showed mass losses associated to endothermic signals. Based on thermodynamic considerations, the mass loss was principally attributed to the dissociation of Cr$_2$O$_3$ into gaseous metallic chromium and oxygen, and depending on the atmosphere composition, the formation of a liquid phase CrO$_6$ could be evidenced.

Dilatometry tests performed with undoped and Cr-doped UO$_2$ pellets allowed to determine the temperature domain where chromium is solubilized in UO$_2$, i.e. around 1100-1200°C, as well as the temperature at which the open porosity vanishes in the pellet. The presence of a liquid phase for the highest content of chromium was suggested from dilatometry measurements in good agreement with the DTA results with chromia powder.

The volatilization process during the sintering of doped UO$_2$ pellets has been shown to result from different phenomena: both decomposition of Cr$_2$O$_3$ particles (not solubilized in UO$_2$) and volatilization from Cr solubilized in UO$_2$ particles before the porosity closure, then Cr volatilization from the doped UO$_2$ ceramic matrix after the porosity closure. In the last case, both grain boundary and lattice diffusion processes have to be considered before volatilization, whereas only lattice diffusion is involved in the first case.

Concerning the application to the industrial nuclear fuel processing, further developments based on these conclusions could be achieved. In order to lower as far as possible the amount of chromium volatilization. Detailed kinetic study is in progress in order to quantify the kinetic rates of these various contributions and to obtain a predictive rate law able to find the optimal processing conditions as regards to both the chromia volatilization and the final density of the ceramic.

References


Figures

Figure 1: (a) Thermodynamic diagram for the Cr–O system according to Toker et al. [5] with the DTA signal start (●) and mass loss start (▲) for all atmospheres studied in this paper. H₂ + 0.6% CO₂ (1), H₂ + 1.11% CO₂ (2), H₂ + 1.47% CO₂ (3), H₂ + 2.06% CO₂ (4), and H₂ + 3.2% CO₂ (5). (b) Magnification of the diagram centered on the Cr₂O₄ domain and representation of the oxygen potential error (dashed line) for H₂ + 2.06% CO₂ atmosphere.

Figure 2: Mass loss Δm in% (–), rate of the mass loss d(Δm)/dt in% min⁻¹ (––) and DTA signal (⋯) in arbitrary units for Cr₂O₃ powder under various H₂/CO₂ atmospheres with %CO₂ equal to 0.6 (a), 1.11 (b), 1.47 (c), 2.06 (d) and 3.2 (e).
Figure 3: Calculated pressures of gaseous species Cr\(_2\)O, CrO, CrO\(_2\) and CrO\(_3\) in equilibrium with solids or liquid Cr\(_2\)O\(_3\) (a), Cr\(_2\)O\(_3\) (b), Cr\(_2\)O\(_4\) (c) and CrO\(_3\) (d) versus temperature for H\(_2\) + 1.11% CO\(_2\) atmosphere.

Figure 4: Calculated Cr(g) pressure in equilibrium with Cr\(_2\)O\(_3\) versus temperature for the various atmospheres used in this study.
Figure 5: (a) Densification curves in %dth and (b) rate of densification in %dth min\(^{-1}\) for UO\(_2\), UO\(_2\) + 0.18 mol% Cr, UO\(_2\) + 0.44 mol% Cr and UO\(_2\) + 0.9 mol% Cr in H\(_2\) + 2.06% CO atmosphere.

Figure 6: Rate of densification in %dth min\(^{-1}\) for UO\(_2\) (---) and UO\(_2\) + 0.9 mol% Cr (--) under various H\(_2\)/CO atmospheres with %CO, equal to 0.6 (a), 1.47 (b), 2.06 (c) and 3.2 (d). The vertical lines correspond to 95% dth.
Figure 7: (a) Mass loss $\Delta m$ in% and (b) rate of mass loss $d(\Delta m)/dt$ in% min$^{-1}$ for $\text{UO}_2$ (--) and $\text{UO}_2 + 0.9\text{ mol\% Cr}$ (--) in $H_2 + 1.47\% \text{CO}_2$ atmosphere.

Figure 8: Mass loss $\Delta m$ in% and (b) rate of mass loss $d(\Delta m)/dt$ in% min$^{-1}$ for $\text{UO}_2$, $\text{UO}_2 + 0.18\text{ mol\% Cr}$, $\text{UO}_2 + 0.44\text{ mol\% Cr}$ and $\text{UO}_2 + 0.9\text{ mol\% Cr}$ in $H_2 + 1.47\% \text{CO}_2$ atmosphere.

Tables

Table 1: TG and DTA results for Cr$_2$O$_3$ powder.

<table>
<thead>
<tr>
<th>% CO$_2$ in hydrogen</th>
<th>0.6%</th>
<th>1.11%</th>
<th>1.47%</th>
<th>2.06%</th>
<th>3.2%</th>
</tr>
</thead>
<tbody>
<tr>
<td>TG: onset temperature (°C)</td>
<td>1430</td>
<td>1499</td>
<td>1526</td>
<td>1521</td>
<td>1659</td>
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<tr>
<td>dTG: peak temperature (°C)</td>
<td>1663</td>
<td>1686</td>
<td>1697</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>TG: Mass loss (%) at 1700°C</td>
<td>35.9</td>
<td>23.8</td>
<td>15.9</td>
<td>7.4</td>
<td>1.9</td>
</tr>
<tr>
<td>DTA: onset temperature (°C) (heating)</td>
<td>1435</td>
<td>1487</td>
<td>1470</td>
<td>1508</td>
<td>-</td>
</tr>
<tr>
<td>DTA: peak temperature (°C) (heating)</td>
<td>1635</td>
<td>1671</td>
<td>1689</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>DTA: onset temperature (°C) (cooling)</td>
<td>-</td>
<td>-</td>
<td>1663</td>
<td>1662</td>
<td>-</td>
</tr>
<tr>
<td>DTA: peak temperature (°C) (cooling)</td>
<td>-</td>
<td>-</td>
<td>1624</td>
<td>1634</td>
<td>-</td>
</tr>
</tbody>
</table>
Table 2: Standard (P = 1013 hPa) molar enthalpy variations and Gibbs energy variations of reactions at 1400 °C and 1700 °C [8,9] and theoretical mass loss for each reaction R1.

<table>
<thead>
<tr>
<th>Reactions</th>
<th>ΔH° (kJ.mol⁻¹)</th>
<th>ΔG° (kJ.mol⁻¹)</th>
<th>Theoretical mass loss (% of the reactive compound)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1400°C</td>
<td>1700°C</td>
<td>1400°C</td>
</tr>
<tr>
<td>R1: Cr₂O₃(s) → 2Cr(s) + 3/2 O₂(g)</td>
<td>1891</td>
<td>1881</td>
<td>1028</td>
</tr>
<tr>
<td>R2: Cr₂O₃(s) → 2Cr(s) + 3/2 O₂(g)</td>
<td>1129</td>
<td>1134</td>
<td>702</td>
</tr>
<tr>
<td>R3: Cr(s) → Cr(g)</td>
<td>381</td>
<td>373</td>
<td>156</td>
</tr>
<tr>
<td>R4: Cr₂O₃(s) → 2/3 Cr₃O₄(s) + 1/6 O₂(g)</td>
<td>-</td>
<td>191</td>
<td>-</td>
</tr>
<tr>
<td>R5: Cr₂O₃(s) → 3Cr(g) + 2 O₂(g)</td>
<td>-</td>
<td>2533</td>
<td>-</td>
</tr>
<tr>
<td>R6: Cr₃O₄(s) → 3Cr(s) + 2 O₂(g)</td>
<td>-</td>
<td>1414</td>
<td>-</td>
</tr>
<tr>
<td>R7: Cr₂O₃(s) → 3CrO₂(l) + 1/2 O₂(g)</td>
<td>-</td>
<td>583</td>
<td>-</td>
</tr>
<tr>
<td>R8: CrO₂(l) → Cr(g) + 1/2 O₂(g)</td>
<td>656</td>
<td>650</td>
<td>376</td>
</tr>
<tr>
<td>R9: Cr₂O₃(s) → 2 CrO₂(l) + 1/2 O₂(g)</td>
<td>580</td>
<td>579</td>
<td>267</td>
</tr>
<tr>
<td>R10: CrO₂(l) → CrO(g)</td>
<td>444</td>
<td>437</td>
<td>233</td>
</tr>
<tr>
<td>R11: Cr₂O₃(s) + Cr(g) → 3CrO₂(l)</td>
<td>-76</td>
<td>-71</td>
<td>-109</td>
</tr>
</tbody>
</table>

Table 3: Dilatometry results for undoped and Cr₂O₃-doped UO₂ pellets.

<table>
<thead>
<tr>
<th>%CO₂ in hydrogen</th>
<th>0.6%</th>
<th>1.47%</th>
<th>2.06%</th>
<th>3.2%</th>
</tr>
</thead>
<tbody>
<tr>
<td>mol% Cr</td>
<td>0%</td>
<td>0.9%</td>
<td>0%</td>
<td>0.18%</td>
</tr>
<tr>
<td>Onset T(°C)</td>
<td>1198</td>
<td>1175</td>
<td>-</td>
<td>1157</td>
</tr>
<tr>
<td>Peak T(°C)</td>
<td>1585</td>
<td>1608</td>
<td>1669</td>
<td>1597</td>
</tr>
<tr>
<td>Density (% d/th) at 1700°C</td>
<td>90</td>
<td>87</td>
<td>93</td>
<td>96</td>
</tr>
</tbody>
</table>