Calcined Resin Microsphere Pelletization (CRMP) a novel process for sintered metallic oxide pellets synthesis.

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Calcined resin microsphere pelletization (CRMP): A novel process for sintered metallic oxide pellets

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Abstract

This study deals with the preliminary development of a powder-free process called calcined resin microsphere pelletization (CRMP) used for the fabrication of metallic oxide pellets. This dustless process could be used for the fabrication of mixed U1−xAm2xO2 pellets dedicated to the transmutation of Am in fast neutron reactors. In this study, porous CeO2 microspheres, used as a surrogate of AmO2, were obtained after mineralization of cerium loaded ion exchange resin beads. These millimetric oxide microspheres were die-pressed into pellets which were then sintered under air to form ceramic pellets. Their densities approached 95% of theoretical density of CeO2 and a homogeneous microstructure was obtained by using optimized microspheres. The influence of calcination parameters on the characteristics of microspheres and on the properties of sintered pellets is discussed.

Keywords: Microsphere; Calcination; Pressing; Sintering; Precursors-organic

1. Introduction

One of the most promising options under study to reduce the radiotoxicity and the heat load of nuclear waste storage is the transmutation of the long-lived radionuclides such as minor actinides (MA), americium (Am) for instance, in a fast neutron reactor. In this recycling process, MA are transformed by transmutation, consisting of diluting MA in the whole core of the reactor and heterogeneous transmutation, where MA are concentrated in uranium dioxide fuels introduced in the periphery of the core. These specific fuels are oxide materials called minor actinide bearing blanket (MABB). Their fabrication requires special attention due to the high radiotoxicity of MA. Currently, the fabrication of these mixed oxide pellets follows a powder metallurgy route which consists in numerous steps of grinding and milling of UO2 and AmO2 powders, pressing and sintering. Preparation of granular material before compaction and especially granulation, which is required to produce free-flowing feed material for automatic pellet presses, generates fine and highly contaminant particles. Dustless processes which avoid the presence of highly contaminating powder in the fabrication line, using oxide in the form of free flowing spherules with millimetric size, are recommended.

The first process developed in the 70s, based on the compaction of millimetric spheres to form pellets, was called the sphere-cal process. Authors studied the compaction of UO2 and ThO2 beads, obtained by sol–gel process, into pellets and
the subsequent sintering to form simple or mixed oxide fuel pellets for advanced pressurized heavy water reactors (PHWRs). Their work focused especially on the influence of the gelation and calcination processes on bead compaction into pellets and on the subsequent sintering. Exhaustive investigations were carried out to optimize heat treatment parameters in order to obtain UO₂ microspheres suitable for pelletization. More recent studies have been published on this process renamed sol–gel microsphere pelletization (SGMP) to produce ThO₂–UO₂ pellets.⁹–¹² Pelletization of soft porous microspheres with a low crushing strength leads to the synthesis of homogeneous and dense sintered pellets (≥94% TD). The SGMP process was then studied for the elaboration of UO₂ pellets,¹³,¹⁴ and mixed uranium, cerium or plutonium oxide pellets for liquid-metal cooled fast-breeder reactors (LMFBR). In these studies, cerium was used as a surrogate of plutonium.¹⁵,¹⁶ Pellets with high density (≥95% TD) and a tailored microstructure containing uniformly distributed closed pores were produced by using porous microspheres, obtained by the addition of carbon black as a pore former or by adjusting gelation parameters.¹⁴,¹⁷ The viability of the SGMP route for fabrication of (U, Pu)C and (U, Pu)N has also been established.¹⁸,¹⁹ Afterwards, uranium–plutonium nitrides as low-density pellets have been prepared and characterized for transmutation application.²⁰

As with the sphere-cal or the SGMP programme, our study deals with the development of a spherule route which is called the calcined resin microsphere pelletization (CRMP) process. It consists of the synthesis of mixed oxide precursors of spherical form and millimetric size obtained by an adaptation of the weak acid resin (WAR) process²¹–²⁴ and their compaction into pellets before sintering. The microsphere synthesis is based on the fixation of cerium cations into beads of ion exchange resin (IER) and their mineralization in air to form the oxide.

This present study constitutes a first step in the elaboration of mixed oxide pellets by the CRMP process with the elaboration of cerium (as a surrogate of americium) oxide pellets. It describes the conversion method, the optimization of the calcination step and the results of compacting and sintering by examination of the pellets’ microstructure and density.

2. Materials and methods

2.1. Starting materials

The acrylic resin was obtained from Dow Chemicals Company (Rohm and Haas, Chauny, France) and consisted of an IMAC HP335 gel type acrylic exchanger in the form of beads of 3 different size distributions (batches 1, 2 and 3). Their characteristics are listed in Table 1.

Concentrated ammonia solution (25%, Merck, Pro Analysis) and concentrated nitric acid solution (70%, Fisher Chemical, Certified ACS Plus) were used to prepare washing solutions. Hexahydrate cerium(III) nitrate (Ce(NO₃)₃·6H₂O, 99.99% pure, Prolabo) was used to prepare 0.25 M Ce(III) stock solution.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Characteristics of the 3 batches of IMAC HP335 ionic exchange resin (nc, not communicated).</th>
</tr>
</thead>
<tbody>
<tr>
<td>IMAC HP335 resin</td>
<td>Commercial form: batch 1</td>
</tr>
<tr>
<td>Structure</td>
<td>Gel —COOH</td>
</tr>
<tr>
<td>Functional group</td>
<td>Exchange capacity (meq/mL RH)</td>
</tr>
<tr>
<td>Physical form</td>
<td>Moisture holding capacity</td>
</tr>
<tr>
<td>Matrix</td>
<td>Density (H⁺ form)</td>
</tr>
<tr>
<td>Exchange capacity (meq/mL RH)</td>
<td>Harmonic mean size (μm)</td>
</tr>
</tbody>
</table>

2.2. Solution analysis

The cerium content of the different samples was determined by atomic emission spectrometry (ICP-AES) using an Activa HORIBA LAB equipment. A calibration was performed using diluted Ce certified standard (Merck, Certipur, 1000 mg/L in 2–3% HNO₃) in 0.5 M HNO₃. Scandium was also used as an internal standard (Merck, Certipur, 1000 mg/L in 2–3% HNO₃) at a fixed concentration of 25 mg/L. Two wavelengths were chosen for cerium analyses (394.275 and 413.380 nm) and one for scandium measurement (358.963 nm).

The ammonium concentration was measured by acid–base titration carried out with a Titripro 809 Metrohm automatic titrator in the presence of potassium oxalate used to shift cerium hydrolysis. The evolution of pH was followed continuously using a Consort C861 analyser and a glass electrode (Blueline 14 pH, Schott Instruments).

2.3. Synthesis of the solid oxide microsphere precursor

2.3.1. Preparation of resin beads

The wet resin was first mechanically screened with an AS200 Basic Retsch apparatus under deionized water through successively finer standard Prolabo sieves (100 μm, 250 μm, 400 μm, 630 μm, 800 μm, 1000 μm and 1250 μm). Then a manual second screening was performed on each size range in order to limit clogging of the sieves.

The selected resin was washed in a column with deionized water under successive finer standard Prolabo sieves (100 μm, 250 μm, 400 μm, 630 μm, 800 μm, 1000 μm and 1250 μm). Then a manual second screening was performed on each size range in order to limit clogging of the sieves.

The selected resin was washed in a column in two cycles consisting of percolating successively a 1 M aqueous nitric acid solution, deionized water, 1 M aqueous ammonia solution and deionized water. Two cycles were necessary to remove contaminants such as sodium and synthesis impurities. The resin in its final ammonium form was drained and recovered semi-wet.

2.3.2. Resin metal loading

A batch method was used to perform the cation exchange on the washed resin in its ammonium form. Those resin beads were fully rehydrated on a column 2 h before exchange in deionized water and their packed bed volume was measured (15–45 mL). Different size distributions of resin were contacted with the 0.25 M cerium nitrate stock solution (100–300 mL) for 4 h in...
a vessel at room temperature (22 ± 2 °C) under gentle agitation with a magnetic stirring bar. The resin was then transferred to a column and washed with deionized water. After draining under vacuum, the loaded resin was dried at 110 °C overnight. During exchange, the bulk solution was sampled to follow cerium and ammonium concentrations.

2.3.3. Calcination
Calcination of the loaded resin was performed in a tubular furnace (Nabertherm, HTRH 100-300; alumina tube 1200 cm in length and 10 cm in internal diameter) under synthetic air flow (Air Liquide, 20% O2–80% N2). Temperatures in the range of 500–1400 °C were tested. The microspheres were placed as a single layer in an alumina crucible in order to obtain a homogeneous calcination as mentioned by Ferreira et al.25

2.3.4. Pressing and sintering
Cerium oxide microspheres obtained after calcination were cold compacted into green pellets in a 10 T pneumatic Enerpac press at 500 MPa. Tungsten carbide die-punch sets of 5.165 mm diameter were used and were lubricated by zinc stearate prior to each compaction. The green pellets were sintered at 1400 °C for 6 h in the same furnace and same air flow as previously described. The heating and cooling rates were fixed at 10 °C/min.

2.4. Solid characterization

2.4.1. Loaded resin
An elemental analysis was performed on a sample of the 630–800 μm loaded resin. A mass of 150 mg of this sample was placed in a column and contacted by percolating 25 mL of 1 M HNO3 during 100 min. The percolate was recovered in a 50 mL volumetric flask and completed by washing water from the resin. ICP-AES analyses were then conducted on this solution to determine the concentration of cerium released during dissolving corresponding to the resin metal content.

The metal content was also estimated by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) performed with a STA 449 C Netzsch equipment. Microspheres were placed in an alumina crucible fitted with a cover and were calcined up to 1100 °C in air flow (3 NL/h) at a heating rate of 5 °C/min with an isothermal treatment of 1 h at 1100 °C. The residue was pure cerium oxide which was confirmed by X-ray diffraction (XRD) and carbon analysis.

2.4.2. Oxide microspheres
The effective density26 of oxide microspheres was measured by helium pycnometry (Accupyc 1330 Micromeritics pycnometer). The pycnometer was equipped with a 10 cm³ volume cell and a sample module of 1 cm³. It worked with helium (Alphagaz; Air Liquide, 99.995% pure). The 1 cm³ volume module was calibrated using a 0.718502 cm³ standard ball. Each sample was degassed before measurement overnight at 140 °C under vacuum using a VacPrep 061 Micromeritics apparatus.

Apparent density26 of microspheres was determined by volume and mass measurements of a sample of a few hundred beads. Number of spheres and the sample volume were estimated from image analysis of a layer of microspheres (considered as perfect spheres). Images were taken by an optical video microscope (Fort SVM 01) and treated by Ellix pattern recognition software (Microvision). Unitary weight equivalent to the weight of one oxide microsphere was also determined by this method.

Specific surface area was obtained from N2 adsorption and desorption at 77 K using the Brunauer–Emmett–Teller (BET) method. A Tristar II 3020 Micromeretics apparatus was used for measurements. Samples were pretreated in a vacuum oven to remove volatiles.

The residual carbon content in oxide microspheres was measured with a CS 230 LECO carbon–sulphur analyser calibrated with carbon standards obtained from LECO (501-024, 501-676 and 502-809). The crystallographic structure of cerium oxide microspheres was characterized by powder XRD (D8 advanced BRUCKER diffractometer, CuKα radiation). Gold (NBS Standard) was added to all samples as an internal standard to calibrate the angular positions of the observed XRD lines. XRD diagrams were treated by Diffracplus Eva software. The lattice parameters of the oxide were refined by pattern matching using FULLPROF Suite.27

2.4.3. Green and sintered pellets
Effective density of sintered pellets was measured by helium pycnometry. Bulk density of pellets was obtained by volume measurement and gravimetric analysis. A dilatometric test was carried out on a green pellet with a Setsys evolution Setaram dilatometer which allowed continuous monitoring of the shrinkage kinetics up to 1400 °C with a heating rate of 10 °C/min under air flow (synthetic air, Alphagaz, Air Liquide, 20% O2–80% N2).

2.4.4. Microstructure analysis
Scanning electron microscopy (SEM) micrographs of pellets and microspheres were carried out with a Supra 55 ZEISS instrument. Microspheres were cold mounted with epoxy resin within a ring mould and then grinded with SiC sandpaper (1200, 2400 and 4000 grit sizes) to access internal microstructure. A diamond suspension (10 µm) was used to polish samples and a final polishing with colloidal SiO2 suspension was performed. Internal microstructure of microspheres was observed on the polished surface.

Sintered pellets were cut and sections were cold mounted and polished following the same process. In situ high temperature environmental scanning electron microscopy (HT-ESEM) experiments were performed using a field emission gun environmental scanning electron microscope (model FEI QUANTA 200 ESEM FEG) equipped with a 1500 °C hot stage.28 During this work, observations were performed under water vapour at an operating pressure of 233 Pa and a specific detector was used for in situ gaseous secondary electron imaging at high temperature. An individual bead of cerium loaded resin was fixed
on refractory cement and was heated to 1000 °C. Micrographs were recorded at regular intervals during the heat treatment of the microsphere.

3. Results and discussion

3.1. Exchange capacity

The resin was sorted first to work on homogeneous batches for exchange kinetics. The size distributions of the three batches of IMAC HP335 resin were obtained by sieving (see Table 2). Four size ranges were qualified in this study: 100–250, 400–630, 630–800 and 1000–1250 μm.

Exchange capacity of the resin was determined by measuring the amount of ammonium and proton exchanged during the last washing cycle. Average values of scientific weight capacity of two size distributions (Q_{weight}) in milliequivalent per gram of dried resin in proton form (meq/g dry H^+) and technical volume capacity (Q_{eq}) in milliequivalent per milliliter of packed bed (meq/mL packed bed) are shown in Table 3. The swelling factor which corresponds to volume expansion of the packed bed after H^+/NH_4^+ conversion is also mentioned. Swelling is important because the action of ammoniac causes complete ion dissociation in the resin but also introduces a large solvated ammonium counter ion.\(^{29,30}\)

The scientific weight capacity does not vary significantly with size distribution and is equal to 12.6 meq/g dry H^+ form, which is not far from the theoretical value of 13.3 meq/g dry H^+ form (knowing the weight of 1 mol of carboxylic moiety to be 75 g, which represents 1 equivalent, the theoretical weight capacity corresponds to \(1/75 = 13.3\) meq/g dry H^+).

3.2. Kinetics of fixation

Evolutions of pH, ammonium and cerium concentrations during conversion step of the 630–800 μm size distribution are plotted in Fig. 1.

The amount of cerium in solution rapidly decreased during the first hour of conversion corresponding to the loading of cerium in the resin and the subsequent release of ammonium. Accordingly, the quantity of NH_4^+ in solution increased. After 1 h, the pH stabilized at a value between 5 and 6 corresponding to achievement of equilibrium.

3.3. Cerium content in the loaded resin

3.3.1. Balance from solution analyses

Experimental conditions and results of conversion of each size distribution, such as final mass of the loaded resin, quantity of cerium fixed in the resin, quantity of ammonium released in solution, effective exchange volume capacity and weight percentage of cerium in the dried loaded resin are listed in Table 4. The quantity of cerium fixed in the resin is calculated on the basis of the difference between the initial quantity of cerium and the amount of Ce found in the solution at equilibrium, in sampling and washing. The weight percentages of Ce (wt% Ce) for every size distributions are quite homogeneous values and their mean value equals 35 ± 3%.

The quantity of ammonium released is calculated in the same way by adding the amount of ammonium in solution at equilibrium, in sampling and washing.

The stoichiometry of the exchange reaction (ratio of the quantity of ammonium released in solution over the quantity of cerium fixed in the resin) is close to 3 which confirms that the lanthanide cation takes up 3 exchanging sites in the resin and therefore releases 3 ammonium cations in solution. This ratio of 3 has already been reported in the literature\(^{31}\) for La^3+, Ce^3+, Pr^3+ in the case of a Na-form resin at pH 6–7. The cation exchange reaction can be written as Eq. (1):

\[
3RNH_4^+ + Ce^{3+} \rightarrow R_3Ce + 3NH_4^+ \tag{1}
\]

where R represents an exchanging site of the resin.

3.3.2. Balance from solid characterization

Elemental analysis also indicates that the weight percentage of cerium equals 33 ± 2% for the 630–800 μm loaded beads. Metal content in the fully loaded resin can also be measured after calcination under air at 800 °C for each size distribution considering that the residue is pure cerium oxide free of carbon (confirmed by carbon analyses presented later in Table 6), the residual mass is representative of the metal oxide materials left after calcination and allows assessment of the initial metal content in the resin. The weight percentage of Ce is similar for each size distribution and the mean value is 34.4 ± 0.2%.

---

Table 2
Size distributions of the IMAC HP335 resin.

<table>
<thead>
<tr>
<th>Batch</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sieve aperture (μm)</td>
<td>Non cumulative total (between sieves) (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>100–250</td>
<td>–</td>
<td>100</td>
<td>–</td>
</tr>
<tr>
<td>250–400</td>
<td>5.4</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>400–630</td>
<td>47.2</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>630–800</td>
<td>40.9</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>800–1000</td>
<td>6.4</td>
<td>–</td>
<td>19.7</td>
</tr>
<tr>
<td>1000–1250</td>
<td>–</td>
<td>–</td>
<td>79.0</td>
</tr>
<tr>
<td>1250–1400</td>
<td>–</td>
<td>–</td>
<td>1.3</td>
</tr>
</tbody>
</table>

Fig. 1. Evolution of Ce^{3+} and NH_4^+ concentrations in solution during the conversion step.

---

Table 3
Technical volume and scientific weight capacities of different size distributions of IMAC HP335.

<table>
<thead>
<tr>
<th>Size distributions (µm)</th>
<th>100–250</th>
<th>250–400</th>
<th>400–630</th>
<th>630–800</th>
<th>800–1000</th>
<th>1000–1250</th>
<th>Mean values</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Q_{\text{vol}} ) H+ form (±0.2) (meq/mL)</td>
<td>4.1</td>
<td>4.1</td>
<td>3.8</td>
<td>3.7</td>
<td>3.6</td>
<td>3.9</td>
<td>3.9 ± 0.1</td>
</tr>
<tr>
<td>( Q_{\text{vol}} ) NH4+ form (±0.1) (meq/mL)</td>
<td>2.66</td>
<td>2.56</td>
<td>2.52</td>
<td>2.67</td>
<td>2.63</td>
<td>2.53</td>
<td>2.60 ± 0.04</td>
</tr>
<tr>
<td>( Q_{\text{weight}} ) H+ form (±0.1) (meq/g)</td>
<td>12.5</td>
<td>–</td>
<td>12.6</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>12.6 ± 0.1</td>
</tr>
<tr>
<td>Swelling factor (±0.1)</td>
<td>1.4</td>
<td>1.5</td>
<td>1.4</td>
<td>1.4</td>
<td>1.4</td>
<td>1.4</td>
<td>1.4</td>
</tr>
</tbody>
</table>

Table 4
Results of Ce conversion for different size distributions.

<table>
<thead>
<tr>
<th>Size distribution (µm)</th>
<th>100–250</th>
<th>400–630</th>
<th>630–800</th>
<th>1000–1250</th>
</tr>
</thead>
<tbody>
<tr>
<td>( V_{\text{resin}} ) NH4+ (mL)</td>
<td>15.0 ± 0.3</td>
<td>15.0 ± 0.3</td>
<td>42.0 ± 0.6</td>
<td>15.0 ± 0.3</td>
</tr>
<tr>
<td>( V_{\text{Ce(NO3)3}} ) (mL)</td>
<td>100.0 ± 0.6</td>
<td>100.0 ± 0.6</td>
<td>300.0 ± 0.6</td>
<td>100.0 ± 0.6</td>
</tr>
<tr>
<td>Final pH</td>
<td>4.38 ± 0.05</td>
<td>4.68 ± 0.05</td>
<td>5.28 ± 0.05</td>
<td>5.22 ± 0.05</td>
</tr>
<tr>
<td>( n ) (Ce3+) loaded (mmol)</td>
<td>14.0 ± 0.8</td>
<td>12.6 ± 0.8</td>
<td>38.9 ± 2.0</td>
<td>13.0 ± 0.8</td>
</tr>
<tr>
<td>( n ) (NH4+) released (mmol)</td>
<td>43 ± 2</td>
<td>40 ± 2</td>
<td>121 ± 1</td>
<td>39 ± 2</td>
</tr>
<tr>
<td>Effect. Capa. (meq/mL)</td>
<td>2.8 ± 0.1</td>
<td>2.5 ± 0.1</td>
<td>2.8 ± 0.2</td>
<td>2.6 ± 0.1</td>
</tr>
<tr>
<td>Equivalent NH4+/Ce3+</td>
<td>3.1 ± 0.3</td>
<td>3.2 ± 0.3</td>
<td>3.1 ± 0.4</td>
<td>3.0 ± 0.3</td>
</tr>
<tr>
<td>mresin Ce (g)</td>
<td>5.88 ± 0.01</td>
<td>5.43 ± 0.01</td>
<td>14.25 ± 0.01</td>
<td>4.92 ± 0.01</td>
</tr>
<tr>
<td>wt% Ce loaded resin</td>
<td>33 ± 2</td>
<td>33 ± 2</td>
<td>38 ± 5</td>
<td>37 ± 2</td>
</tr>
</tbody>
</table>

The size distribution has no influence on the weight percentage of Ce in the dried resin which is about 34% according to all characterizations. Those results are in agreement with results concerning the fixation of lanthanide(III) cations studied by Mokhtari. The theoretical value of weight percentage of Ce in the dried resin corresponding to a full exchange is 38.7%, calculated from the following equation:

\[
\begin{align*}
\text{wt% Ce}_{\text{th}} &= \frac{\text{MW(Ce)}}{\text{MW(Ce)} + \text{MW(R)}} \\
&= \frac{140.12 + 3 \times 74}{140.12 + 3 \times 74} = 38.7
\end{align*}
\]

with MW being the molecular weight and R the carboxylic inorganic group in its deprotonated form.

The resin exchange capacity is not fully used owing to the establishment of equilibrium generated by the accumulation of NH4+ in solution during the batch exchange. The effective capacity reaches 90% of theoretical capacity. This process can be divided into three exothermic mechanisms, with a first degradation step between 290 °C and 420 °C causing a diameter drop of 12% followed by a second step between 420 °C and 490 °C concomitant with a 3% diameter decrease and a final degradation until 690 °C corresponding to a diameter reduction of 4%. The microsphere continues to shrink above 690 °C. This phenomenon is attributed to the beginning of sintering. Up to 800 °C, a decrease of 24% in microsphere diameter is observed.

3.4.2. Microsphere characteristics

The diametral and volumetric shrinkages of the loaded resin (R3Ce) for calcination up to 800 °C were determined by image analysis for each size distribution. Mean values are 22 ± 3% and 53 ± 8% respectively. The former value of diametral shrinkage is in agreement with the data of 24% obtained from HT-ESEM images (Fig. 2).

3.4. Solid analyses

3.4.1. TGA

The thermal conversion of a sample of 630–800 µm loaded resin was investigated using TGA at a heating rate of 5 °C/min from room temperature to 1100 °C in air (Fig. 2). The thermal behaviour of one individual microsphere was also characterized by HT-ESEM which enables to follow the evolution of the microsphere diameter with temperature (see Fig. 2 and Appendix B). Series of images is given in Fig. 3. The first significant weight loss (5%) between room temperature and 290 °C can be assigned to dehydration and loss of weakly adsorbed water coupled with reduction of 3% in microsphere diameter. The elimination of carbon matter in the loaded resin beads occurred in the temperature range of 290–690 °C with a loss of 52% of original mass.

3.4.3. Crystalline evolution

X-ray diffraction patterns were obtained on oxide microspheres calcined at 600 °C, 800 °C and 1000 °C. For all calcination temperatures, the diffraction patterns are characteristic of the face-centred cubic CeO$_2$. The pattern corresponding to the temperature of 800 °C is reported in Fig. 4. The refined lattice parameter (Table 5), $a = 5.408$ Å is not significantly different from the value of 5.411 Å published in the literature. The oxide is already formed at 600 °C. The variation of the crystallization state reported in Fig. 4 is estimated through the average full width at half maximum (fwhm) from the more intense XRD lines. The crystallite size increases with temperature and the evolution is in good agreement with published data.

### Table 5

<table>
<thead>
<tr>
<th>Temperature of calcination (°C)</th>
<th>Lattice parameter ($\pm$0.001) (Å)</th>
<th>Average crystallite size ($\pm$3) (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>600</td>
<td>5.407</td>
<td>19</td>
</tr>
<tr>
<td>800</td>
<td>5.407</td>
<td>32</td>
</tr>
<tr>
<td>1000</td>
<td>5.408</td>
<td>48</td>
</tr>
</tbody>
</table>

3.4.4. Microstructural evolution

ESEM micrographs realized on the calcination of a unique loaded resin microsphere enabled identification in situ the microstructural changes of the microsphere during the heating process (Fig. 5).

After dehydration and at the beginning of mass loss, cracks appear at the surface of the beads corresponding to the beginning
of the removal of carbon in the form of carbon dioxide. Open porosity is created at this moment and seems to disappear when temperature increases and the diameter of the sphere decreases.

SEM micrographs performed on as-polished sections of CeO$_2$ beads reveal the homogeneity of their internal microstructure and micrometre-sized grains (Fig. 6). The porosity evolution versus temperature was investigated and results will be presented hereafter.

3.5. Calcination optimization

Different thermal cycles were applied to 630–800 µm beads to compare the influence of each parameter of the cycle such as: heating rate (HR) and final temperature (FT). A dwell time of 4 h was applied at final temperature to all batches of microspheres except for microspheres calcined at 500 °C.

3.5.1. Heating rate (HR)

Different heating rates were applied during the calcination of the loaded resin beads: 0.1, 0.2, 0.5, 1, 2.5, 5 °C/min (with a FT = 800 °C).

Heating rate has an important influence on the microsphere fracture as shown in Fig. 7.

An excessive heating rate of 5 °C/min causes violent outgassing of CO$_2$, leading to important mechanical stress on the beads and their disintegration. The percentage of broken beads calcined at 5 °C/min is about 20%. Moreover, an increase of heating rate results in greater residual carbon content into the oxide bead (2000 ppm at 5 °C/min compared to 570 ppm at 1 °C/min). Removal of carbon is not complete for excessive rate.

A slow heating rate ($\leq$ 1 °C/min) was then selected during the mineralization to form homogeneous oxide microspheres with enough mechanical strength to avoid emission of small particles during the handling step.

3.5.2. Final temperature (FT)

Final temperatures of 500 °C (with no dwell time) and 600 °C, 800 °C, 1400 °C were tested for optimizing the calcination of loaded resin. Characteristics of calcined beads are summarized in Table 6.

A first observation is that the effective density of oxide beads increases with temperature which is linked mostly with the decrease of closed porosity from 4% at 500 °C to a value inferior at 1% at 1400 °C. Open porosity also decreases from 800 °C but still remains elevated for beads calcined at high temperature (68% at 1400 °C) as we can observe in Figs. 8 and 9 which explains that bulk density of beads is limited to 31% of TD in those conditions.

Those results are in agreement with those of TGA and the evolution of mean diameter during mineralization (Fig. 2). They reveal the densification of microsphere above 690 °C in relation with a grain growth observable in Fig. 9 and a decrease of the specific surface of oxide beads with temperature (from 45 m$^2$/g at 500 °C to 0.7 m$^2$/g at 1400 °C).

A second observation concerns the evolution of carbon content into the oxide which is about 0.15% below 600 °C independently of dwell time and which is not significant above this
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Fig. 6. SEM micrographs performed on as-polished section of CeO₂ beads heat treated at 1400 °C.

Fig. 7. CeO₂ microspheres obtained by calcination in air with different TR: (a) 0.2 °C/min; (b) 1 °C/min and (c) 5 °C/min.

Fig. 8. Evolution of effective density, open and closed porosity of CeO₂ beads calcined at different temperatures.

Fig. 9. SEM micrographs of the surface of CeO₂ beads calcined at (a) (600, 1) and (b) (1400, 1).


Final temperature is a critical parameter and enables synthesis of microspheres with different final amounts of carbon, specific surface, density and porosity. These parameters will have important consequences for the microstructure of sintered pellets, as will be discussed in the following section.

Unitary weight corresponding to the weight of one oxide microsphere can be also calculated and is about 75 µg/microsphere equivalent to approximately 61 µg of Ce per microsphere. Those data have a great interest in the determination of capacity requirements as mentioned by Haas et al. For instance, the volume of initial resin in its ammonium form, necessary for the production of 1 g of CeO₂ (mCeO₂) which represents approximately \( \frac{1}{75} \times 10^{-6} = 13,300 \) beads of oxide (\( n_{\text{microspheres}} \)), is about 10 mL of resin bed. Detailed calculations are given in Appendix A.
3.6. Compacting and sintering

Microspheres calcined above 800 °C were uniaxially pressed at 500 MPa in a 5.165 mm matrix. The microspheres obtained at lower temperatures were discarded because their carbon content was too high. The green pellets were subsequently sintered at 1400 °C for 6 h in air. A dilatometric analysis performed on a green pellet following this thermal cycle is shown in Fig. 10. Sintering begins at 800 °C with a maximum linear shrinkage rate at 1090 °C. The sintering is ended at 1400 °C. The final linear shrinkage after cooling is about 14%. The shrinkage curve is classical and in accordance with the sintering behaviour of green pellets obtained from the compaction of cerium oxide powder prepared from oxalate precursors even if the final shrinkage is slightly inferior: a mean value of 18% is obtained with these pellets.37

The green pellet dimensions were measured before sintering to determine bulk density. Effective and bulk densities of ceramic pellets are shown in Table 7. The bulk density of green pellets is found to be between 62 and 65% of TD which is greater than the value of 52% obtained for CeO₂ pellets fabricated by powder compaction.37 A slight influence of calcination parameters on the green density is observed, which increases with heating rate.

Green pellets obtained from the compaction of microspheres calcined at 1400 °C were too brittle and not suitable for handling. This can be explained by the hardness of beads which is too high and not appropriate for pressing. This property has been widely described in the literature as a consequence of the use of “non-porous” and hard microspheres which do not disintegrate during pelletization.9,25

Sintering of green pellets at 1400 °C for 6 h resulted in effective density between 85 and 94% of TD depending on the heating rate. In particular, a heating rate of 5 °C/min during calcination leads to sintered pellets with an important closed porosity approaching 15% probably due to a heterogeneous population of beads before compaction as mentioned previously. SEM micrographs of as-polished diameter sections of sintered pellets are given in Fig. 11.

Pellets obtained from compaction of beads calcined at a low rate of 1 °C/min up to 800 °C are dense and present a...
homogeneous microstructure with 6% of closed porosity and no open porosity. On the contrary, a high heating rate causes a heterogeneous and porous microstructure. A final calcination temperature of 800°C produces beads with specific surface of 8.2 ± 0.1 m²/g and bulk density of 1.18 ± 0.15 g/cm³ and seems to be an optimized treatment to obtain dense and homogeneous pellets. Actually, those two characteristics have been described in the literature²⁵ as fundamental properties in the case of the optimization of the pelletization process of microsphere precursors obtained from sol–gel technology. The authors’ conclusion is to use low densified kernel (~2 g/cm³) with low specific surface (~5 m²/g) obtained at 850°C.

4. Conclusion

This work constitutes a first approach concerning the elaboration of dense cerium oxide pellets by the calcined resin microspheres pelletization (CRMP) process based on the compaction of oxide microspheres into green pellets and their subsequent sintering to obtain dense ceramic pellets. Those homogeneous and porous oxide beads of 150–750 μm in diameter were obtained from mineralization of resin beads loaded in cerium following different thermal treatments. Different calcination temperatures and heating rates have been investigated to produce oxide beads suitable for pressing. In particular, compaction of porous oxide beads calcined at slow mineralization rate (~1 °C/min) and a temperature of 800 °C results in dense and homogeneous metal oxide pellets (95% of TD).

In contrast, an overly high temperature of calcination (1400 °C) resulted in hard beads not suitable for compaction.

The mechanical behaviour under compression of those oxide beads produced in different calcination conditions is under study and should improve the control of the ceramic microstructure. Eventually the optimized CRMP process should be applied to the elaboration of mixed (U,Ce)O₂ pellets and (U,Am)O₂ ceramics.

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Appendix A. Determination of the volume of initial resin in its ammonium form, necessary to the production of 1 g of CeO₂ (mCeO₂)

\[
m_{\text{RCe}} = \frac{m_{\text{CeO}_2}}{1 - \text{WL}} = \frac{1}{1 - 0.57} = 2.32 \quad (A.1)
\]

where \(m_{\text{RCe}}\) represents the mass of the cerium loaded resin and WL is the weight loss in percentage during calcination taken from TGA equal to 57%.

\[
m_{\text{Ce}} = \text{wt}% \quad \text{Ce} \times m_{\text{RCe}} = 0.344 \times 2.32 = 0.8 \quad (A.2)
\]

with wt% Ce representing the weight percentage of Ce in the loaded resin which is equal to 34.4%.

So

\[
V_{RH}(\text{mL}) = \frac{m_{\text{Ce}} \times 3 \times 1000}{\text{MW}_{\text{Ce}} \times \frac{Q_{\text{volH}^+}}{3 \times 1000}} = \frac{0.8}{140.12 \times \frac{2.6}{2}} = 6.6 \quad (A.3)
\]

where MW_Ce is the Ce molecular weight and \(Q_{\text{volH}^+}\) is the technical volume capacity of the proton resin expressed in meq/mL (Table 3).

The estimation of the resin bed volume in its ammonium form (\(V_{RNH_4}\)) is obtained by taking into account the swelling factor described in Table 3: in this case, \(V_{RNH_4}\) is equal to 6.6 × 1.4–10 mL.

Appendix B. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jeurceramsoc.2012.04.011.

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