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# Ultrafast high-temperature sintering (UHS) of cerium oxide-based compound

Ahsanul Kabir<sup>a,\*</sup>, Bartłomiej Lemieszek<sup>b</sup>, Jakub Karczewski<sup>c</sup>, Emanuele De Bona<sup>d</sup>, Maxim Varenik<sup>e</sup>, Sebastian Molin<sup>b</sup>, Mattia Biesuz<sup>d</sup>

<sup>a</sup> Institute for Manufacturing Technology of Ceramic Components and Composites, University of Stuttgart, 70569, Stuttgart, Germany

<sup>b</sup> Advanced Materials Center, Faculty of Electronics, Telecommunications, and Informatics, Gdańsk University of Technology, Ul. G. Narutowicza 11/12, 80-233,

Gdańsk, Poland

<sup>c</sup> Institute of Nanotechnology and Materials Engineering, Faculty of Applied Physics and Mathematics, Gdańsk University of Technology, 80-233, Gdańsk, Poland

<sup>d</sup> Department of Industrial Engineering, Via Sommarive 9, 38123, Trento, Italy

e Department of Molecular Chemistry and Materials Science, Weizmann Institute of Science, Rehovot, 7610001, Israel

 A R T I C L E I N F O
 A B S T R A C T

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 Ultrafast high-temperature sintering (UHS) is an innovative sintering technique that can densify ceramics in a few seconds, dramatically reducing the carbon footprint and firing costs. In this work, the feasibility of applying UHS in Gd-doped ceria (GDC) and GDC-Er-stabilized bismuth oxide (ESB) composite powders was investigated. At high UHS currents (22–24 A), fully dense GDC samples with a large grain size were obtained. Nonetheless, most of the GDC pellets exhibited micro/macro cracks, which were reduced by lowering the sample thickness. Interestingly, the GDC-ESB composite samples exhibit no cracks or fragmentation at all, thanks to ESB as a

#### 1. Introduction

Electrostriction

The sintering of ceramics, i.e., consolidation of loose powders into a bulk solid component requires high sintering temperatures ( $\approx 2/3 T_m$ ) usually between 1000 °C and 2000 °C, and long processing times, which account for more than 80% of the energy consumed for their production. According to recent studies, ceramics manufacturing yearly accounts for the emission of 1.3 trillion kg of CO<sub>2</sub> [1], giving a market value of over \$100 billion per year. From a materials standpoint, such lengthy high-temperature exposures may have a detrimental impact on the microstructures of ceramics such as undesirable chemical interactions, the loss of volatile components, and abnormal grain growth [2]. Therefore, modern ceramic processing methods are pushing towards the reduction [3] of firing energy either by lowering the sintering temperature or shortening processing time. In this context, advanced sintering techniques with enhanced energy efficiency have emerged over the last few years such as fast firing [4,5], laser sintering [6], microwave sintering [7], spark plasma sintering [8], flash sintering [9,10], cold sintering [11, 12], etc. To date, they have been partially established at the industrial scale. However, sintering in these methods relies on specific material characteristics such as laser and microwave absorption, solubility, and electric conductivity which limits their widespread applicability. Very recently, a novel sintering technology called ultra-fast high-temperature sintering (UHS) has been developed by Wang et al. [13]. UHS is a pressure-less sintering technique that can densify ceramic material more than 1000 times faster than conventional sintering. In this method, a green body is sandwiched between two carbon felts in an inert atmosphere. The process employs an electric current that rapidly heats the carbon felt by Joule heating and provides sufficient heat up to  $10^3$ – $10^4$  °C/min, bringing the material rapidly to the sintering temperature (up to 2000+  $^{\circ}$ C) through radiation and thermal conduction [14, 15]. As a result, near-full densification of the powder compacts occurs in a few seconds. Such high kinetics also provide additional benefits such as limited grain coarsening, reduced loss of volatile elements (Li/Na), and hindered inter-diffusion in multi-layered/composite ceramics [16, 17]. Moreover, UHS can bring the materials to a non-equilibrium state, which may have unanticipated advantages like lowering the activation energy for densification [13], enhancing or impairing specific phase

sintering aid. This samples were further characterized from an electrochemical and electromechanical point of

view. The GDC-ESB material displays an ionic conductivity value of  $\sim$ 1.5  $\times$  10<sup>-2</sup> S/cm at 600 °C and frequency-

stable (0.1–350 Hz) room temperature electrostriction strain coefficient of  $\sim 10^{-18}$  (m/V)<sup>2</sup>.

\* Corresponding author. *E-mail address:* ahsanul.kabir@ifkb.uni-stuttgart.de (A. Kabir).

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Fig. 1. (a) X-ray diffraction (XRD) patterns and (b) Raman spectra of GDC pellets, sintered via UHS. (c) SEM image of the cold-fractured surfaces of the GDC samples. GDC-0.1–24A and GDC-30-20A reveal intergranular cracks.

changes, and formation of grain boundary segregation [18,19].

So far, UHS has been explored in various ceramics compounds such as  $Al_2O_3$  [20],  $Si_3N_4$  [21], YSZ [13], ultra-high-temperature ceramics [19,22], high-entropy compounds [23–25] and solid-state electrolytes of LLZO/LATP [13,26] as well as samples with complex shapes [27] and large thicknesses [14,28]. However, to the best of our knowledge, there was little interest in studying ceria-based compounds (just one work in the literature [29]), a model material that is extensively used in electro-ceramics as a solid electrolyte and electrode material for solid oxide cells (SOCs) [4], catalysts [30], gas sensors [31], memristors [32], gas separation membranes [33], etc. Furthermore, unusual electrome-chanical activity was recently demonstrated in both thin films and bulk ceramics of ceria-derivates [34–36].

The main aim of the current work is to examine the feasibility of applying the UHS process in consolidating ceria-based ceramics with or

without the addition of bismuth oxide and shedding light on essential properties such as densification behavior and microstructure evolution, electrical and electromechanical properties.

#### 2. Experimental Procedures

#### 2.1. Powder synthesis

Nanosized gadolinium doped ceria (GDC) powder with different compositions (Ce<sub>1-x</sub>Gd<sub>x</sub>O<sub>2- $\delta$ </sub> where x = 0.01, 0.05, and 0.3) were synthesized by the co-precipitation method in an aqueous solution, as described in details elsewhere [37]. GDC and Er-stabilized Bismuth Oxide (ESB) composites were obtained using commercial 20 mol% Gd-doped ceria (GDC) powders (DC-20 K, Daiichi Kigenso Kagaku Kogyo Co., Japan) and 20 mol% Er- stabilized bismuth oxide (ESB),

synthesized by solid-state reaction [38]. For the composite, three different compositions were selected (1, 2, and 4 wt% of ESB in GDC). At first, the appropriate amount of powder was dispersed into ethanol and the dispersion was ball-milled overnight in a plastic bottle. The milling balls were then separated from the slurry and then oven-dried at 100 °C and calcined at 300 °C for 1 h in a hot plate. The powder was then gently crushed using a mortar and pestle and sieved (150  $\mu$ m).

#### 2.2. Pellet preparation

The green pellets were produced by uniaxial pressing at 200 MPa for 30 s with a diameter of 8 mm. The relative density of the green samples was ~45–50%. UHS was carried out in an Ar-filled borosilicate glass flask using graphite felt (70 × 24 × 5 mm<sup>3</sup>). About 20 mm of the felt were clamped on both sides by two steel electrodes so the heating zone was  $30 \times 24 \times 5$  mm<sup>3</sup>. The green sample was put into the middle of the felt which was cut by a razor blade. A DC electric current was applied stepwise (1A every 10 s, starting from 15A), to maximum currents between 18 and 24 A. After reaching the maximum current, it was held for 15 s. After that, the power was turned off which led to the fast cooling off of the sample. The total time of the sintering process was less than 5 min. The samples were mechanically polished using SIC grinding papers up to grit 2500.

#### 2.3. Materials characterization

The bulk density of the sintered ceramics pellets was measured by Archimedes' method in a water medium at room temperature. The crystallographic phase purity was analyzed by X-ray diffraction (XRD) technique, Philips PANalytical X'Pert) with CuK<sub>a</sub> radiation, and the Raman spectroscopy using a LabRAM Aramis equipped with an excitation laser 532 nm. The microstructure was examined by a scanning electron microscope (SEM), model FEI Quanta FEG 250 coupled with an energy dispersive x-ray spectroscopy (EDS), EDAX Genesis APEX 2i with ApolloX SDD spectrometer. The grain sizes were calculated by the linear intercept method and multiplied with a correction factor of 1.57 [39]. The electrical properties were investigated by electrochemical impedance spectroscopy (EIS) using a Solarton (1260) at a temperature between 300 and 600  $^\circ C$  in the frequency range of 0.05 Hz–3 MHz with an AC signal of 30 mV. The platinum paste (electrode) was applied on opposites of the pellet and fired at 700 °C for 1 h in air to ensure good contact with the platinum mesh of the setup. The EIS data were fitted employing an equivalent circuit model consisting of a resistor connected in parallel to a constant phase element and analyzed by the ZView software. The electromechanical properties were measured under ambient conditions (23 °C, relative humidity 20-55%) using a proximity sensor (Capacitance, Lion). The signal from the sensor was read by a lock-in amplifier, as explained in detail elsewhere [40,41]. The electrostrictive strain was calculated as a ratio between the field-induced displacement and the original thickness of the pellets.

#### 3. Results and discussions

#### 3.1. Gd doped ceria (GDC)

The synthesized powders are nanometric in size (15–30 nm), loosely agglomerated, and have no impurities (Fig. S1). The cold-pressed green GDC samples were consolidated by UHS at different currents ranging from 20A to 24A, for 15 s holding. At 20A, the predicted temperature at the felt is around 1150 °C, and with a 2A of current escalation a 100 °C increment is also anticipated [27]. At currents  $\geq$  22A, most of the samples show an inhomogeneous dark color (the usual color is yellowish) suggesting a partial chemical reduction of the sample (Ce<sup>4+</sup>  $\rightarrow$  Ce<sup>3+</sup>) [42]. Discoloration has been observed previously for ceria compounds sintered in UHS or similar field-assisted sintering technologies using graphite tooling and at low oxygen partial pressure [42,43].

Thus, all the samples were post-annealed at 800 °C for 1 h in an air atmosphere. The crystallographic phase composition and microstructure of the sintered GDC samples are shown in Fig. 1. As expected, the diffraction pattern (Fig. 1a) shows a single-phase cubic fluorite structure like pure ceria (Fm-3m). Within the resolution limit of the XRD technique, no secondary phases (double fluorite) were noticed, even for the GDC-30 sample. The crystallographic phase is further confirmed by Raman spectra (Fig. 1b) and displays a sharp peak at about 460 cm<sup>-1</sup> which corresponds to the F<sub>2g</sub> mode of the CeO<sub>2</sub> fluorite structure and is ascribed to symmetrical stretching of the Ce-O bond in eightfold coordination [44,45]. With increasing dopant concentration, the peak shifts and broadens, highlighting the dissolution of the solute/dopant into the  ${\rm GeO}_2$  lattice. The extra shoulder peak at 561  ${\rm cm}^{-1}$  for the GDC-30 sample is assigned to dopant-oxygen vacancy defect clusters [46]. As shown in SEM images (Fig. 1c and Fig. S2), the samples sintered at 20A are still largely porous exhibiting the typical microstructure of the intermediate sintering stages, the grain size is submicrometric. Moreover, with increasing current to 22A/24A, densification as well as grain growth increases. Nevertheless, small pores are visible in the microstructure. The EDS elemental mapping and average grain size of the samples are listed in Figs. S3 and S4, respectively. The EDS map confirms a homogenous solid solution of GDC without any impurity phase. For GDC-0.1, the grain size increased from  $\sim$  300 nm at 20 A to  $\sim$ 7  $\mu$ m at 24 A, by nearly a factor of 20. The extremely high grain growth in such a short time strongly suggested extremely fast diffusion through the grain boundaries at 24A, which can be attributed to a reduction of the cation. It has already been reported that in fluorite-structured ceramics the energy barrier for ion migration in reducing conditions is heavily reduced [47] resulting in extraordinary grain growth [48]. The grains have a fully relaxed shape, grain boundaries are flat and reveal intergranular micro-cracks. Such an effect is attributed to the chemo-mechanical stress that is generated due to the reduction (chemical expansion) during sintering and following room temperature oxidation (chemical contraction). For ceria, micro-crack formation was typical for sintering under a reducing atmosphere and has already been reported in the works of literature [49-51]. Moreover, restricted grain growth is observed for highly doped ceria samples (GDC-5 and GDC-30) because mass diffusion is strongly limited by the dopant solute drag effect. These outcomes are exactly opposite to D. Ni et al. who showed restricted and enhanced grain growth in low and highly-doped ceria, respectively sintered under reducing conditions [52]. It is worth mentioning that oxygen partial pressure (1400 °C, for 4 h, 9% H<sub>2</sub>-N<sub>2</sub>,  $10^{-31} < pO_2$ ) in their study was much lower than this work. The solute drag effect is also experiential for the sample, fabricated via conventional free sintering in the air (Fig. S5). Point to note that most of the UHS sintered samples demonstrated micro and/or macro cracks and were highly sensitive to fragmentation (Fig. S6), thus they were not subjected to further property characterization. Moreover, by decreasing the sample thickness, fragmentation was avoided and the number of microcracks was reduced to a minimum.

## 3.2. Gd doped ceria (GDC) and Er-stabilized bismuth oxide (ESB) composite

An alternative way to mitigate cracking problems was to add a highly functional bismuth oxide phase in GDC. The bismuth oxide additives melt during sintering. Thus, the volume (chemical) expansion from Ce<sup>4+</sup> to Ce<sup>3+</sup> is accommodated by the liquid phase and hinders cracking. Recent studies also demonstrate that the introduction of bismuth oxide in ceria-based compounds significantly increases electrostrictive, ionic conductivity, and mechanical properties [53,54]. Among all stabilized bismuth oxide, 20 mol% Er-stabilized bismuth oxide (ESB) exhibits the highest conductivity [55] and stabilized bismuth oxide is also a non-classical electrostrictor [40]. The sintering protocol of the ESB-GDC composite was the same as the pure GDC as mentioned above. These samples were also post-annealed at 800  $^{\circ}$ C for 1 h in air. The as-sintered



Fig. 2. (a) X-ray diffraction (XRD) patterns and (b) Raman spectra of GDC-ESB composite pellets, sintered via UHS method employing 13V-22/24A.



Fig. 3. Scanning electron microscope (SEM) image of the cold-fractured surfaces of the GDC-ESB composite (4 wt% ESB in GDC) samples, sintered at 20-24A.

GDC-ESB composite pellet was crack-free and mechanically stable (Fig. S7). Visual inspection exhibited that compared with pure GDC, the ESB-GDC compound has a relatively high shrinkage. It was reported for

conventional sintering that  $Bi_2O_3$  additives with concentration  ${\geq}0.5$  wt % have a factor of 2–5 higher shrinkage rate compared to pure-GDC one, significantly impacting the sintering mechanism [56]. The bulk density



**Fig. 4.** (a–b) Representation of the impedance spectra i.e. Nyquist plot (Re Z vs Im Z) at temperatures of 300 and 600 °C of ESB1-24A pellet, measured in synthetic air with Pt electrode. Impedance was also performed on additional samples to confirm the reproducibility of the results. (c) Arrhenius plot for the estimation of the temperature-dependent total electrical conductivity of the GDC-ESB composite samples.

of the sample is listed in Table S1. The crystallographic phase purity and micrographs of the fracture surfaces are reported in Fig. 2. As expected, no crystal planes other than cubic fluorite structures like pure CeO<sub>2</sub> are observed in the XRD diffraction pattern (Fig. 2a). The peak intensity at (200) and (400) crystalline planes are unexpectedly higher as compared to the UHS sintered GDC compound, albeit the reason for this is unclear. A possible explanation might be the development of some texture which is not uncommon in liquid phase sintering, occurs due to grain re-arrangement and dissolve precipitation stages. Raman spectra (Fig. 2b) show dominant peaks around the 495, 515, and 553 bands, attributed to  $\gamma$ -Bi<sub>2</sub>O<sub>3</sub> phase (space group, I23) [57]. The shoulder peak near ~445 cm<sup>-1</sup> Raman band resembles the F<sub>2g</sub> mode of the GDC.

The exemplary SEM images of the sintered samples are demonstrated in Fig. 3 and Fig. S8, further confirming a good level of densification behavior. ESB has a melting point of around ~850 °C, therefore it melts during sintering, enabling rapid densification by liquid phase sintering. Since the solubility limit of cubic-structured  $Bi_2O_3$  in GDC was found to be about 0.8 wt % [58], ESB is expected to be present at the grain boundaries or triple points. As observed, the grain structures are well-homogenized having equiaxed grains whose size is between 1.8 and 2.0  $\mu$ m and could be attributed to the intermediate-stage sintering, i.e., in which the samples are >95% dense and closed porosity [56].

Gil et al. claimed that the retarded or not promoting grain growth in ESB-GDC composite can be assumed to be due to poor wetting properties of the formed liquid phase [56]. Zhao and coworkers also disclosed similar results showing that the addition of  $Bi_2O_3$  did not considerably affect the grain growth kinetics of the doped ceria compounds [59]. Since UHS is occurring at extremely high temperatures but for a short time, a few  $Bi_2O_3$  contents can evaporate (sublimation temperature is around 1000 °C [60]). At 22A–24A, the predicted temperature of the felt is around 1250–1350 °C as mentioned above [27]. The elemental analysis obtained by EDS shown in Fig. S9, confirms a loss of Bi from the samples and local segregation at the triple point.

The electrical properties of the samples are measured using impedance spectroscopy in the air at the temperature of 300-600 °C. Fig. 4a and b exemplifies a typical EIS spectrum i.e., a Nyquist plot of ESB1-24A sample at two different temperatures. At 300 °C, the sample exhibits a slightly deformed single semicircle. Typically, ceramics oxide exhibits two well-resolved semi-circles: at high and low frequencies, corresponding to the transport phenomena in the bulk/grain interior and grain boundaries, respectively [4,43]. A single semicircle is often observed for nanocrystalline ceramics and can be attributed to overlapped bulk and grain boundary contribution [61]. Moreover, its characteristic frequency value (~1 MHz) suggests that the semicircle is probably dominated by bulk contribution (Bode plots Fig. S10). Its capacitance value (4.5  $\times$  10<sup>-9</sup> F) is roughly two orders of magnitude higher than the expected bulk value. With increasing temperature, the superimposed contribution of bulk and grain boundary decreases significantly and a single semicircle due to electrolyte-electrode interfaces is observed because the relaxation frequency is shifted towards high frequencies at a higher temperature [62,63]. This advocates that ionic conductivity is a thermally activated process and it increases as the relaxation time response of the oxygen ion decreases [64,65]. The temperature-dependent total electrical (ionic) conductivity ( $\sigma$ ) of the samples is plotted in an Arrhenius relationship in Fig. 4c. The Arrhenius equation follows,  $\sigma T = \sigma_0 \exp(\frac{-E_a}{kT})$  where  $\sigma_0$ , k,  $E_a$  denotes the pre-exponential factor, Boltzmann constant, and activation energy of long-range oxygen ion migration respectively. The conductivity value conductivity is similar among the investigated samples and conventionally sintered reference GDC-20. The conductivity value at 600 °C is  $\sim 1.5 \times 10^{-2}$  S/cm and is in line with previous reports [58,66]. The activation energy is obtained from the slope of the straight line. All samples have a similar activation energy of approx.  $\sim 0.85$  eV and in good agreement with the literature data [37]. In summary, UHS allowed to get in a few minutes defect-free and well-densified ESB-GDC



**Fig. 5.** Electrostriction strain coefficient ( $M_{33}$ ) as a function of applied AC frequency. The data is compared with 20 mol% Gd-doped ceria [67].

composites with electrochemical properties substantially analogous to those of electrolytes obtained by slow conventional firing.

The electromechanical properties of the sample were measured at room temperature and illustrated in Fig. 5. The investigated compound responds at the second harmonic of the applied electric field and contracts in the direction of the applied electric field, confirming its electrostriction behavior [68,69]. They exhibit longitudinal strain that agrees with previous reports of the GDC and other doped ceria compounds [70]. The electrostrictive strain coefficient M<sub>33</sub> value was high in the ESB1-24A sample and interestingly, it was found that M<sub>33</sub> decreases with ESB concentration. Although the magnitude of M<sub>33</sub> is one order of value lower than GDC-20 [67], these compounds do not display Debve-type frequency-dependent strain relaxation. Frequency-associated strain relaxation is typical for Gd-doped ceria and is described in various previous reports [71,72]. The M<sub>33</sub> value at 100 Hz in the ESB1-24A sample is  $10^{-18}$  (m/V)<sup>2</sup> is still one order of magnitude larger than the Newnham scaling law prediction and gigantic in nature.

#### 4. Conclusion

In this work, UHS was applied to sinter pure GDC and GDC-ESB composite ceramics. The entire sintering process took less than 5 min, including heating, dwelling, and cooling steps which are at least 100 times shorter than conventional firing cycles of ceria-based ceramics. The pure GDC ceramics showed macrocracks and fragmentation due to chemical reduction during sintering. The addition of ESB reduces the cracking problem. In both types of samples, densification and grain growth increase with increasing the applied UHS current. The addition of ESB did not change the electrical properties, however, the electromechanical response generally decreased with ESB addition. Despite this reduction, the  $M_{33}$  coefficient was found to be gigantic for the GDC sample doped with 1 wt% of ESB and UHSed under 24 A. Interestingly, the electromechanical properties are extremely robust and frequency-stable in the 0.1–400 Hz frequency range.

#### Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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#### Appendix A. Supplementary data

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