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ATMOSPHERE- AND VOLTAGE- DEPENDENT ELECTRONIC CONDUCTIVITY OF OXIDE ION CONDUCTING $Zr_{1-x}Y_xO_{2-x/2}$ CERAMICS

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Abstract

Cubic, fluorite-structured solid solutions Zr_{1-x}Y_xO_{2-x/2}, YSZ, x=0.4-0.7, were prepared by solgel synthesis. Impedance measurements on pellets of approximate density 85% sintered at 1300 °C for 24 h, showed strong evidence of oxide ion conduction with an inclined Warburg spike at low frequencies and capacitance values $\sim 10^{-6}$ Fcm⁻¹ at 40 Hz. Arrhenius plots of total pellet conductivities were linear with activation energies 1.4 to 1.56 eV. Conductivity decreased with x and was 2 to 4 orders of magnitude lower than with optimised YSZ, x=0.08. On changing the atmosphere from N₂ to O₂ during impedance measurements, two reversible effects were seen: the Warburg spike contracted greatly and the sample resistance decreased. These effects were more noticeable at higher x and are attributed to the introduction of p-type electronic conduction, in parallel with pre-existing oxide ion conduction. A similar reversible result was observed on application of a dc bias during impedance measurements. On either increasing pO_2 or application of a dc bias, hole creation is believed to arise by ionisation of underbonded oxide ions situated near the Y^{3+} dopant ions. The ionised electrons are trapped at surface oxygen species and the holes that are left on oxygen are responsible for p-type conduction. The electrolytic domain of x=0.4-0.7 extends up to approximately 10^{-2} atm O₂. before p-type conduction is observed. The upper pO_2 limit of the electrolytic domain of x=0.08 is not known but is likely to be close to or slightly above 1 atm O_2 .

Keywords: Y₂Zr₂O₇; YSZ; electronic conduction; atmosphere/voltage dependence

Introduction

Yttria-stabilised zirconia is the most widely-used oxide ion conductor in solid oxide fuel cells, SOFCs, because of its high conductivity, resistance to reduction, good mechanical strength and stability at high temperatures [1]. The key to this application is the high concentration of mobile anion vacancies created to charge-compensate for introduction of Y^{3+} dopant onto the Zr^{4+} sites. YSZ is a solid solution series of general formula $Zr_{1-x}Y_xO_{2-x/2}$ which has the fluorite structure in which the concentration of oxygen vacancies depends on the value of x. According to the ZrO₂-Y₂O₃ phase diagram [2], Fig. 1(a), the cubic YSZ solid solution extends over the composition range 0 < x < 0.7 at high temperatures. At lower x values the solid solutions transform rapidly to tetragonal and/or monoclinic structures on cooling. At higher x values the cubic solid solutions should transform or precipitate secondary phases on cooling but transformation rates may be very slow. The coordination numbers of Zr and Y, which are disordered, depend on x and are less than the ideal value of 8 in the fluorite structure [3]. The cation distribution may not be completely random, however; in composition x = 0.5, corresponding to formula $Y_{0.5}Zr_{0.5}O_{1.75}$, there is a preference for a higher coordination number for Y of 7.2 compared to 6.8 for Zr [3]. Composition x = 0.5 is sometimes referred to as a pyrochlore [4], but diffraction data and crystal structure modelling [5] show little evidence of a pyrochlore structure apart from this indication of a non-random cation distribution.

The composition most widely used in SOFC applications with highest oxide ion conductivity is x=0.08, and is labelled as YSZ08 in this work [1]. At higher x, oxygen vacancy

concentration increases, but defect interactions are held responsible for mobile ion trapping leading to reduction in conductivity and increase in activation energy [6]. For instance, x=0.5 has oxide ion conductivity, $3.75 \times 10^{-3} \text{ Scm}^{-1}$ at 800 °C [7] compared to $2.50 \times 10^{-2} \text{ Scm}^{-1}$ [1], for YSZ08 used in SOFCs.

For all oxide ion conducting solid electrolytes, the electrolytic domain is limited at low oxygen partial pressures, pO_2 , by the onset of n-type electronic conduction and at high pO_2 by the onset of p-type conduction, as shown schematically in Fig 1(b) [8]. YSZ08 is generally regarded as stable down to a pO_2 of 10^{-20} atm which leads to its use as electrolyte in contact with the anode and reducing atmospheres. Little information is available on the upper pO_2 limit of the electrolytic domain for YSZ08, probably because of the difficulty in achieving a wide range of pO_2 values greater than atmospheric pressure. Nevertheless, in the p-type region, and using Kroger-Vink notation, it is assumed that uptake of oxygen may occur by a mechanism summarised as:

$$\mathbf{V}_{\mathbf{o}}^{\prime\prime} + 1/2\mathbf{O}_2 \rightarrow \mathbf{O}_{\mathbf{o}}^{\prime \mathbf{x}} + 2\mathbf{h}^{\prime} \tag{1}$$

leading to a gradient of +1/4 in plots of log σ vs log pO₂. Usually, holes are assumed to be located on transition metal ions, present either as controlled dopants or unavoidable impurities, but direct evidence for the site of hole location seems not to be available.

Recently, Masó and West [9] demonstrated that electronic conduction is introduced into YSZ08 under the action of a small dc bias at high temperature in air. Similar conductivity changes were seen with acceptor-doped titanate perovskites including Ca-, Zn- and Mg-doped BaTiO₃ [10-12] and Ca-doped BiFeO₃ [13], which all show enhanced conductivity on either increasing pO_2 in the measuring atmosphere or application of a small dc bias. The enhanced conductivity was attributed to ionization of underbonded O²⁻ ions in the vicinity of

acceptor dopants leading to hole creation on oxygen which conceptually, is the same as the creation of O⁻ ions:

$$O_0^x \rightarrow O_0^+ + e^{\prime}$$
 (2)

The O⁻ ions are responsible for the enhanced electronic conductivity, which is p-type, since the ionized electrons have been immobilized at sample surfaces by participating in one, or more, of the equilibria shown in:

$$O_2(g) \leftrightarrow O_2(ads) \stackrel{e}{\leftrightarrow} O_2^- \stackrel{e}{\leftrightarrow} O_2^{2-} / 20^- \stackrel{e}{\leftrightarrow} 20^{2-} (surface) \leftrightarrow 20^{2-} (bulk)$$
 (3)

The p-type conduction in the acceptor-doped titanate perovskites is thermally activated which means that the holes are localised rather than delocalised in the valence band. It is a logical step to refer to the holes localized on oxygen as O^- ions, therefore.

Although the conductivity of YSZ with x = 0.08 is not sensitive to pO_2 in atmospheres such as O_2 , N_2 or air at atmospheric pressure, preliminary tests on compositions with higher x showed that their conductivity is indeed, sensitive to pO_2 . The present work reports a systematic study of the conductivity of YSZ with 0.4<x<0.7, to determine the effects of pO_2 and dc bias on the electrical properties.

Experimental

Four compositions based on the general formula $Zr_{1-x}Y_xO_{2-x/2}$, with x = 0.4, 0.5, 0.6 and 0.7 and labelled YSZ4, YSZ5, YSZ6 and YSZ7 respectively, were prepared by a polymeric solgel procedure using Y(OOCCH₃)₃·H₂O (99.9%, Strem Chemicals) and Zr(OCH₂CH₂CH₃)₄ (70% Sigma-Aldrich) as precursors. All reagents were of analytical grade and used without further purification. Absolute ethanol (Scharlab, 99.9%) was used as solvent with the necessary amount of acetylacetone, acacH (99% Alfa Aesar) as a stabilizing agent for the Zr alkoxide precursor. The Zr alkoxide was dissolved in an ethanol-acetylacetone mixture (acacH:Zr molar ratio 4:1), the Y(OOCCH₃)₃·H₂O added, the mixture stirred for 10 min and then transferred to a balloon flask and heated at 70 °C for 72 h. A transparent gel formed and was dried in air under an IR lamp at room temperature for approximately 1 day. Finally, the dry gel was ground with an agate mortar and pestle, decomposed by heating at 5 °C min⁻¹ to 500 °C, left at 500 °C for 1 h and then heated to 1300 °C at the rate of 5 °C min⁻¹ with a final hold at 1300 °C for 12 h. Powder samples were cooled, crushed and pressed into pellets of 5 mm diameter at 1 ton by uniaxial pressing, reheated at 1300 °C for 24 h and cooled slowly inside the furnace. The four compositions chosen lie in the fluorite-structure phase field at high temperatures, as shown in Fig. 1(a). Pellet densities for all compositions were ~85%.

The phase(s) present were analyzed by X-ray powder diffraction, XRD, using a Bruker D4 Endeavor diffractometer, Karlsruhe, Germany CuK α radiation. Data were collected by stepscanning from $2\theta = 20$ to 70 ° with a step size of 0.03 ° and 6 s counting time at each step. The cubic lattice parameter was determined by least-squares refinement for reflections in the range 15 < 2 θ < 70, using WinXPow version 1.06 (Darmstadt, Germany). Scanning electron micrographs of the pellets were taken on a field emission scanning electron microscope (FE-SEM) JEOL 7001F, equipped with a spectrometer for energy dispersion analysis of X-rays (EDX) from Oxford instruments, using acceleration voltage 15 kV. The samples were deposited on an Al holder and coated with Pt.

For electrical property measurements, electrodes were fabricated on opposite pellet faces from Pt paste (Pt Ink 6082, Metalor, Barcelona, Spain), which was dried and decomposed by gradually heating to 900 °C. Samples with electrodes attached were placed in a conductivity jig and measured using an Agilent 4294A (Agilent, Madrid, Spain) analyzer over the

frequency range 40 Hz to 13 MHz and temperature range, room temperature to 900 °C. Impedance data were corrected for overall pellet geometry and are reported in units of specific resistance: ohm cm and capacitance: Fcm^{-1} ; data were not corrected for the geometry of regions such as grain boundaries since this was not known accurately. Data were not corrected for sample porosity. In order to avoid any effect of water and therefore, any possible proton conduction, impedance data were recorded in dry atmospheres. Measurements in atmospheres of different pO₂ were carried out in a specially-designed cell fitted with a built-in zirconia probe (MicroPoas by SETNAG) next to the sample, to measure the pO₂ value.

Results

Pellets fired at 1300 °C for 24 h were ground and analysed by XRD. All samples were single phase and were indexed on a cubic unit cell, space group Fm-3m, fully consistent with literature data. XRD data for x = 0.5 are shown in Fig. 2(a). There was no evidence of a possible pyrochlore structure, whose XRD pattern is expected to contain extra peaks, arrowed, at $2\theta = 36.3$ and 43.6 compared to the defect fluorite pattern [14]. Lattice parameters as a function of x, inset of Fig 2(a), showed linear increase with x.

The structure of the YSZ solid solutions therefore contains a disordered mixture of Y^{3+} and Zr^{4+} distributed over the 8-coordinate sites of the fluorite structure, with some oxide ion sites vacant, depending on composition x. From Fig 1(a), cubic solid solutions at 1300 °C are expected to form over the range $0.15 \le x \le 0.52$ [2,15,16]. Our results showed single phase cubic solid solutions over a more extensive range of compositions including x = 0.6 and 0.7. We do not know at present whether single phase cubic solid solutions at these compositions

are thermodynamically metastable although clearly, they are kinetically stable under the conditions of synthesis used here. These results may be consistent with the possibility to form metastable phases by sol-gel synthesis [17-20]. A typical microstructure of the pellet surface for YSZ5 is shown in Fig 2(b). Similar microstructures and grain sizes in the range 0.2 - 0.8 µm were observed for all compositions. An SEM of a fracture surface through a pellet with the Pt electrode attached is shown in Fig 2(c) for sample YSZ7; the inset of this figure corresponds to a magnification of the YSZ7 microstructure and confirms the expected porosity. A homogenous layer of the Pt electrode, in contact with the sample surface, is observed in this figure, showing that there is no penetration of the electrode into the interior of the sample. EDX mapping showed a homogeneous distribution of Y and Zr with no evidence of impurities for any of the compositions.

A typical set of impedance data, recorded in dry N_2 at 650 °C and presented in different formats is shown in Fig. 3 for a pellet of YSZ5 sintered at 1300 °C for 24 h with density 85%. Similar response was observed for all compositions. The impedance complex plane plots (a) show a broad, depressed arc at high frequencies and an inclined spike with some curvature at low frequencies. The total resistance obtained from the intercept of either the arc or the spike on the Z' axis had a value of ~20 k Ω cm. The Z["]/M["] spectroscopic plots (b) show peaks at high frequencies which are much broader than expected for an ideal Debye peak, indicating some degree of probable electrical inhomogeneity in the sample but without clear separation of bulk and grain boundary impedances.

The plot of log C[']/log f (c, d) for the same data at two temperatures, 650 and 823 °C, shows evidence for a high frequency plateau of value ~10 pF which corresponds to the limiting high frequency permittivity of ~ 112. The C['] values increase with decreasing frequency, with some

evidence for a poorly-resolved intermediate plateau at ~ 30 pF. This value is too small for it to represent a conventional grain boundary since it corresponds to a significant volume fraction of the sample. Instead, we attribute it to either a combination of a constriction grain boundary associated with sample porosity and possible inhomogeneity in distribution of the Y^{3+} dopant or to a dipole orientation-related impedance such as was reported for YSZ08 [21]. Further studies are in progress to understand the nature of this impedance.

With decreasing frequency, C' increases rapidly to reach a value of approximately 1µF at 40 Hz and 823°C (d); this value is typical of ion blocking at a sample-electrode interface and formation of a double layer capacitance. The inclined spike shown in (a) at low frequencies is attributed to diffusion of oxygen molecules towards and away from sample-electrode interfaces, which therefore places a limit on the rate of oxygen exchange reactions that occur during low frequency measurements. Although this spike, attributed to a Warburg impedance, is not directly associated with oxide ion conduction, it is nevertheless an intrinsic component of redox reactions involving oxygen and is an indirect indication of oxide ion conduction. We therefore interpret the low frequency impedance data to indicate that conduction is primarily by oxide ions but with the possibility of a small amount of electronic conduction (see later).

From data such as these, total sample resistances were obtained from intercepts of either the arc or the Warburg spike on the Z' axis and are shown as conductivity Arrhenius plots as a function of reciprocal temperature in Fig. 4; literature data for YSZ, x=0.08 [22], are shown for comparison. The total conductivity of YSZ solid solutions decreases with x and is 2 to 4 orders of magnitude lower than that of YSZ with x=0.08. In addition, the activation energy increases with x. It is possible that the bulk conductivities are somewhat higher than the total conductivities shown in Fig 4 since either grain boundary constriction impedances or dipole

reorientation impedances are included in the total resistance values; also, conductivities were not corrected for sample porosity. The impedance measurements were obtained on both cooling and heating and the data were reproducible, showing linear behaviour.

The data shown in Figs. 3 and 4 were recorded in an atmosphere of dry N_2 with a pO₂ value of approximately 10⁻⁴ atm. Data are shown in Fig. 5 for pellets of each composition measured in different atmospheres. Data were measured first in dry N_2 (pO₂ ~ 10⁻⁴ atm: Hector check); the atmosphere was then changed to dry O₂ (pO₂ ~1 atm) and the system allowed to stabilise for a few minutes before impedance data were collected in O₂. Two effects are seen in the impedance complex plane plots, (a-d), which were reversible on switching between O₂ and N₂. First, the Warburg spike contracts greatly and becomes a small arc in O₂; second, the total sample resistance decreases. The effects were more noticeable with higher x compositions. It is most unlikely that these two phenomena could be attributed to changes in oxide ion conductivity in the samples simply as a consequence of changing pO₂ in the measuring atmosphere.

Changes in conductivity with pO_2 are widely used as an indication of extrinsic electronic conduction in materials since reactions such as eq'n 1 lead to changes in electronic carrier concentration, of either electrons or holes. We therefore interpret the data shown in Fig 5 to indicate the presence of p-type conduction when samples are measured in O_2 since chemisorption of oxygen with increasing pO_2 leads to hole creation, eq'n 1. This means that a certain amount of p-type electronic conduction [23, 24] occurs in parallel with the preexisting oxide ion conduction. The electronic conduction must arise from reactions involving O_2 in the atmosphere. Eq'n 1 shows the creation of holes but gives no indication as to their location. Traditionally, holes are associated with transition metal impurities but we believe that a more likely location, at least in our materials, is underbonded oxide ions.

The loss of the Warburg spike is also attributed to creation of a parallel electronic conduction pathway [9]. It is further emphasised in the capacitance data, insets Fig 5(a-d), in which the large, high-valued low frequency capacitance decreases because it is no longer associated specifically with charge transfer at the thin, double layer interface.

Impedance data were sensitive to pO_2 over the complete temperature range studied, 500-850 °C and are summarized in Arrhenius plots of total conductivity for one composition, YSZ7 in Fig 6. The data recorded in O_2 are higher by a factor of 2 to 3 with a smaller activation energy and clearly must contain a contribution of both electronic and ionic conductivity. We do not know if the data in N_2 also contain a small electronic component or whether they represent exclusively ionic conduction. These results show that, with reference to Fig 1(b), samples YSZ4 - YSZ7 in air at atmospheric pressure are in the crossover region between the electrolytic domain and p-type conduction. At present, we have not performed a detailed determination of the electronic transport number in our compositions, which would allow the contribution of each conduction mechanism to the electrical response to be determined as a function of temperature, pO_2 and x. Different methods are reported in the literature to determine the electronic and ionic components of the conductivity [9, 25].

The effect on impedance data of application of a small dc bias, in the range 0.5-2V, was found to be very similar to that of increasing pO₂. A selection of results is shown in Fig 7 for YSZ7 and YSZ4. In (a), data are shown in N₂ and O₂ without an applied bias and in N₂ with 2V bias. The data in O₂ and in N₂ with 2V bias are similar and both show reduced total resistance and a much reduced Warburg spike. In (b), data are shown for dc bias in the range 1-10V; a systematic decrease in both sample resistance and Warburg impedance is observed.

The changes in conductivity with changing pO_2 are not instantaneous but take a few minutes to reach the steady state, as shown in Fig 8 for YSZ5 at 750 °C. The sample was first equilibrated in N₂ for 24 h and the resistance measured as a function of time on switching to O₂. Once a steady state had been reached, the atmosphere was switched back to N₂; several minutes were needed to reach a steady-state in both directions. Fig 8 therefore corresponds to a continuous time plot with a switch in atmosphere at intermediate time.

Discussion

The electrical properties of YSZ, x=0.4-0.7, without application of a dc bias, are consistent with oxide ion conduction, as is, of course, widely-recognised for YSZ08 [26]. The conductivity is lower than that of YSZ08, which is usually used as a ceramic electrolyte in SOFC systems. Although the compositions prepared with higher x have a higher concentration of oxygen vacancies, the reduction in ionic conductivity with increasing x is attributed to trapping of oxygen vacancies in defect complexes associated with the Y^{3+} dopant; this is also consistent with the observed increase in activation energy for ionic conduction in higher x compositions [6].

YSZ08 is used as the ceramic electrolyte in SOFCs in part because it is very stable to reduction. However, it has recently been shown that electronic conduction can be induced in YSZ08 by application of a small bias voltage [9]. This causes YSZ08 to become a mixed

conductor with, for instance, an electronic transport number, t_e , of 0.5 with 10 V bias at ~ 550 °C. The electronic conductivity is lost on removal of the dc bias.

In the present study, we report a similar dependence of conductivity on dc bias, but also show that the conductivity is sensitive to pO_2 . The onset of electronic conductivity is shown by two changes to the impedance data: first, the Warburg spike is greatly reduced and second, the total sample resistance is reduced. Reversible changes of a similar nature are seen with both an increase in pO_2 and a dc bias. There is no evidence from the literature that YSZ08 is sensitive to an increase in pO_2 , although it is sensitive to dc bias [9]. From the results presented here, the introduction of electronic conductivity becomes easier with increasing x and it seems highly probable, therefore, that YSZ08 itself will be sensitive to pO_2 at pressures just above 1 atm O_2 . This result has clear significance for the use of YSZ-based materials either in an oxygen-rich atmosphere or in the presence of a bias voltage.

The sensitivity of the total sample conductivity to changes in pO₂, Fig 5, is evidence that the electronic conduction mechanism is p-type. This was presumed, but not established, in the earlier YSZ results [9] and follows previous studies of acceptor-doped titanate perovskites which showed enhanced p-type conductivity with a dc bias. In those studies [9-12], the mechanism of hole creation was attributed to ionisation of underbonded oxide ions associated with lower-valence acceptor dopants. The holes that are created are localised on oxygen, as O⁻ ions, and are responsible for the thermally-activated p-type conduction. The Y^{3+} component of the YSZ solid solutions may be regarded also as an acceptor dopant in which Y^{3+} substitutes for Zr⁴⁺; oxide ions in the immediate vicinity of Y^{3+} are effectively underbonded and can be readily ionised. The effect of increasing x is to increase the Y^{3+}

concentration and therefore, the concentration of underbonded oxide ions; this may account for the increased sensitivity to pO_2 that is seen with increasing x.

The chemisorption on sample surfaces that takes place with increasing pO_2 is a commonlyused diagnostic for extrinsic conductivity and to distinguish p-type from n-type conduction. Using eq'n 1 to represent the chemisorption, electrons are removed from the sample, presumably close to the absorption sites, to form reduced oxygen species. In reality, several equilibria at sample surfaces may be involved, as indicated schematically in eq'n 3, giving rise to species such as superoxide O_2^- , peroxide, $O_2^{2^-}$, O^- and finally O^{2^-} ions, whose net negative charges are balanced by the holes that are created as lattice O^- ions. Several of these equilibria require electrons and hence, with increasing pO_2 , lead an overall increase in p-type conductivity, as shown in Fig. 8. These processes commence at any part of sample surface exposed to the atmosphere.

Similar behaviour to that seen with increasing pO_2 is observed on application of a small dc bias which, for voltages of 1-2 V, is below the probable decomposition voltage of the samples. This behaviour may also be interpreted in terms of the equilibria shown in eq'n 3 as these equilibria are driven to the right hand side by a positive bias, leading to removal of electrons from the sample and an increase in p-type conduction. The positive electrode is therefore responsible for the observed conductivity changes and there is no evidence of any opposite effect occurring at the negative electrode, which would lead to negative charge injection. Thus, if charge injection were to occur, this would lead to the onset of n-type conduction, as represented by the cross-over region at low pO_2 shown in Fig 1(b), in contrast to the observed p-type behaviour. There are reported instances of blackening and electronic conduction in YSZ ceramics which can be attributed to charge injection from the negative electrode under highly reducing conditions. If changes to the sample do occur at the negative electrode under dc bias, they are not directly responsible for the effects reported here. Thus, we also exclude the possibility of Schottky barrier formation and charge injection resulting from a reduction in barrier height on application of the dc bias.

Conclusions

There is huge interest in yttria-stabilised zirconia, YSZ as the ceramic electrolyte for solid oxide fuel cell applications because, as well as its high oxide ion conductivity, it is very resistant to reduction; consequently, its 'electrolytic window' extends down to very low partial pressures of oxygen before it shows the onset of n-type conduction. This work concerns the upper limit of the electrolytic window before the onset of p-type conduction.

Conductivity measurements in different atmospheres show that with increasing Y content, YSZ compositions x=0.4-0.7 are already at the cross-over between electrolytic and p-type conduction on switching between N₂ and O₂ atmospheres; a similar conclusion is reached on conducting impedance measurements in the presence of a small dc bias, and is attributed to the creation of holes on underbonded O^{2-} ions associated with the Y³⁺ acceptor dopant. We do not know the value of the activation energy for the electronic component in the p-type region, but it is probably non-zero which means that the holes are localised on oxygen and not delocalised in the valence band structure. The activation energy of YSZ7 in O₂, 1.48 eV, is somewhat less than that in N₂, 1.56 eV, Fig. 6 and therefore, the activation energy for p-type conduction is presumed to be less than that for oxide ion conduction.

These results are significant for several reasons:

First, they demonstrate that, depending on Y content, YSZ samples may lose their exclusive oxide ion conductivity either in the presence of a small applied voltage or with increasing oxygen partial pressure. For samples of YSZ08 used in SOFC applications, the onset of the p-type conduction is likely to occur at oxygen partial pressures close to, or slightly above 1 atm.

Second, they demonstrate an alternative method by which electronic conductivity can be introduced into YSZ. Most focus in the literature is on the onset of possible n-type conductivity with either reducing atmospheres or decreasing pO_2 , whereas we demonstrate the very easy introduction of p-type conductivity under near-ambient conditions and in the presence of small dc voltages.

Third, they highlight that the mechanism of hole creation must involve redox activity of underbonded oxide ions, which challenges the widely-held belief that oxygen in oxides is present solely as the double negative oxidation state. In high purity chemicals such as used here, there seems to be no other possibility for location of the holes.

Fourth, the ready introduction of p-type conductivity into YSZ-based materials may have significant consequences for their application as electronically-insulating ceramics in either a high pO_2 environment or in the presence of extraneous voltages.

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Figure captions

- Fig 1. (a) Phase diagram of the ZrO₂-YO_{1.5} system [2]; the compositions prepared in this work are indicated. (b) Schematic ionic and electronic conductivity domains as a function of oxygen partial pressure, pO₂, adapted from reference [8].
- **Fig 2**. (a) XRD pattern and (b) SEM of the pellet surface of YSZ5 sintered at 1300°C. (c) SEM of the fracture surface of YSZ7 with the Pt electrode attached. Inset in (a): variation in lattice parameter with x. Inset in (c): magnification of the YSZ7 microstructure of the fracture pellet.
- Fig 3. (a) Impedance complex plane plot, Z*, (b) Z''/M'' and (c,d) C' spectroscopic plots at different temperatures for YSZ5. Measurements in dry N₂; $\omega = 2\pi f$.
- **Fig 4**. Arrhenius plots of total conductivities for samples sintered at 1300°C and measured in dry N₂. Error bars are within the size of data points. Activation energies in eV, with errors in the range 0.02-0.05 eV, are shown beside each data set. YSZ08 ceramic sintered at 1600°C for 6h is included for comparison [22].
- **Fig 5**. Impedance complex plane plots, Z*, with C' spectroscopic plots as insets for all compositions measured in dry N₂ and O₂ at 750°C.
- Fig 6. Arrhenius plots of total conductivity for YSZ7 measured in dry N_2 and O_2 . Activation energies in eV are shown beside each data set.
- Fig 7. (a) Impedance complex plane plots for YSZ7 measured in dry N₂ and O₂ at 750°C and with 2V bias voltage in dry N₂. (b) Impedance complex plane plots for YSZ4 measured in dry N₂ after applying different bias voltages at 800°C.
- Fig. 8. Time-dependent total resistivity for YSZ5 at 750°C in dry (a) O_2 and (b) N_2 .



Fig 1.



Fig 2.



Fig 3.



Fig 4.



Fig 5.



Fig 6.



Fig 7.



Fig. 8.

TOC graphic and synopsis

Cubic solid solutions based on $Zr_{1-x}Y_xO_{2-x/2}$, x = 0.4-0.7 were prepared by sol-gel synthesis. p-type electronic conduction was introduced by increasing pO₂ or dc bias voltage. The electrolytic domain extends up to ~10⁻² atm O₂ before p-type conduction is observed, which is different to the known YSZ, x=0.08.

The introduction of p-type conductivity into YSZ-based materials may have significant consequences for their application as electronically-insulating ceramics in either a high pO_2 environment or in the presence of extraneous voltages.

