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## Virtual cathode induced in  $Rb_2Ti_2O_5$  solid electrolyte

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#### **Abstract**

Rb2Ti2O<sup>5</sup> (RTO) has recently been demonstrated to be a solid electrolyte, producing colossal capacitance when interfaced with metals. In order to understand the mechanisms leading to such colossal equivalent permittivity (up to four orders of magnitude above state-of-the-art values), the charge distribution in RTO is a key feature to be investigated. In the present article, this charge distribution is probed using the pressure-wave-propagation method, in devices made of RTO single crystals or polycrystals sandwiched between two metallic electrodes. Remarkably enough, in both types of samples, negative charges are found to accumulate inside RTO, near the anode, while the electric field near the cathode remains zero. This proves that the ionic carriers are majoritarily negatively charged and provides an explanation for the colossal capacitance. The latter takes place only at the anode while the cathode is virtually shifted into the solid electrolyte.

*Keywords:* Ion conductor, Solid electrolyte, Rubidium titanate, Virtual cathode, supercapacitor material *2010 MSC:* 00-01, 99-00

#### **1. Introduction**

Solid electrolytes have attracted much attention during the last decade in particular in the search for solid oxide fuel cells [1], and high capacitance devices [2, 3]. When a voltage is applied to these high capacitance structures, a thin double layer of charges is formed under voltage at the interface between

- the electrodes, which are electron conductors, and the solid electrolyte, which is a purely ionic conductor. Taking advantage of the ultra-thin character of the double layer, and of the porosity of the materials which allows very large electrode equivalent area, it is possible to produce super-capacitors of EDLC (Electric double-layer capacitors) type. When an additional "blocking" mechanism is present such as an electrochemical reaction or an adsorption or intercalation effect, these devices are termed
- <sup>10</sup> pseudocapacitors and take advantage of cumulated EDLC and pseudocapacitive effects [4]. The use of such components in high power electrical applications, such as for instance electric vehicles, is growing rapidly [5].

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Figure 1: Photographs of a polycrystalline RTO pellet before electrode deposition. The diameter is 15mm. The differences in color are presumably due to different hydration degrees of RTO.

The perovskite-derived material  $Rb_2Ti_2O_5$  (RTO), belonging to the  $M_2Ti_2O_5$  family, which structure was identified in the 1960' [6, 7], has been recently found to display very interesting dielectric properties,

- $15$  showing an equivalent relative permittivity up to  $10^9$  at room temperature [8]. This is three to four orders of magnitude above competitive materials and was attributed to ionic motion. In addition, this material has been demonstrated to exhibit memory effects also related to ionic motion [9].For obtaining such high equivalent permittivity, it is necessary that the charges accumulate either (1) at both electrode interfaces or (2) at one interface only, the other interface exhibiting an Ohmic contact. In the first case,
- <sup>20</sup> the structure is equivalent to two capacitors in series and at least two mobile ionic species of opposite polarity should coexist in the electrolyte, the overall equivalent capacitance being controlled by the smaller of the two capacitances. In the second case, the structure is equivalent to a single capacitor and may involve only one type of mobile ionic species in the electrolyte. Previous work has been performed to calculate the charge distribution in the system under various hypotheses [9]. However, if these cal-
- <sup>25</sup> culations enabled to simulate the observed I-V curves, they failed at reproducing them in a quantitative way, with reasonable physical parameters, which was interpreted as pointing to a missing ingredient in the problem [9].

The present paper aims at identifying this missing ingredient, by investigating the charge distribution inside the samples. As a matter of fact, an extremely high density of charges is required at the interface

<sup>30</sup> to account for such a colossal equivalent capacitance (over  $10^5$  F g<sup>-1</sup>). Therefore we have used a space charge distribution measurement method, namely the pressure-wave-propagation method developed in our Laboratory [10], to follow the charge buildup under a continuous voltage bias [11].

#### **2. Materials and methods**

#### *2.1. Sample preparation*

<sup>35</sup> RTO crystals were synthesized [8, 12], and then annealed at 150 deg C for 24h under 5 mbar of nitrogen gas. Then, to fabricate samples, one single crystal per sample was selected and embedded in Epoxy resin (Araldite D). After polymerization at 50deg C, resin and crystal were cut in 1-mm thick slices in such a way that the crystalline *ab*-planes lay perpendicular to the surface of the sample. Finally two



Figure 2: (a) Schematics of the single crystal sample structure. (b) Schematics of the polycrystalline sample structure.

5-mm-thick aluminum electrodes were glued on both sides with carbon paste. This ensured a good <sup>40</sup> mechanical coupling as well as a stable sample structure for carrying out all the measurements. The full sample was 15-mm in diameter but the RTO single crystal cross section was typically 1 mm $\times$ 1.5 mm. The structure of a typical single-crystal-type sample is described in Figure 2a.

Alternately, as-grown crystallites were ground into powder and then pressed under  $5$  ton  $cm^{-2}$  during <sup>45</sup> 1 min in order to obtain ceramic pellets of 13 mm diameter and typically 1.8 mm thick, in order to fabricate polycrystalline samples. All this was performed under controlled water-vapor-free atmosphere. Then gold was evaporated on both sides of the pellets, and finally a gold foil was pressed against both surfaces in order to ensure optimal electrical and acoustical contact. Silicone oil was used to optimize the latter. The structure of the polycrystalline samples is depicted in Figure 2b. Since RTO is known to

## <sup>50</sup> be sensitive to humidity, the pellets were kept inside anhydrous silicone oil between the measurements.

#### *2.2. Charge distribution measurements*

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In the pressure-wave-propagation method, a short-duration pressure pulse is transmitted to the studied structure [13]. The pressure wave is created by a high power piezogenerator [ref]. A 20 mm waveguide is used to decouple the signal from the pressure pulse generator excitation. The principle of the <sup>55</sup> measurement method is illustrated in Figure 3. As the pressure wave propagates inside the sample, the charges encountered are slightly displaced which induces a current in the measurement circuit connecting the electrodes in short-circuit conditions, or a voltage in open-circuit conditions. The obtained signal represents an image of the charge distribution in short-circuit conditions, and of the internal electric field in open-circuit conditions, time and position inside the sample being connected by the velocity  $\int$ <sub>60</sub> of sound  $v_s$ .

All measurements were carried out at room temperature and hygrometry and atmospheric pressure. A typical signal is shown in Figure 4 for a single crystal sample. Piezoelectric transducers were coupled to the samples in preliminary measurements to determine the exact position of the electrodes in each



Figure 3: Principle of the pressure-wave-propagation method [13]. (a) A continuous voltage *Vapp* is applied across the sample. A pulsed pressure wave travelling from the left to the right provokes a displacement of the charges inside the sample, which generates in open-circuit conditions a voltage signal as function of time, the shape of which is an image of the electric field distribution inside the sample. (b) From top to bottom, charge distribution, electric field distribution, corresponding signal in open circuit conditions.

sample. It was determined that at  $t = 1970$  ns in Figure 4a, the pressure pulse enters the sample. Because <sup>65</sup> of the large sample capacitance, measurements were performed in open-circuit conditions. Therefore the first peak around  $t = 2 \mu s$  is proportional to the electric field at the interface. The rising edge is due to positive charges on the anode electrode and the falling edge is due to the presence of negative charges in the crystal. Such behavior is not visible on the other interface (cathode).

The presence of the epoxy and the complex geometry of the sample make it difficult to ensure a perfect  $\pi$ <sup>0</sup> pressure pulse propagation inside the single crystal sample. Thus a reliable analysis of the signal after the first peak can not be garanteed. Therefore the single-crystal sample was then mounted the opposite way in the sample holder and the voltage was reversed. As a consequence, anode and cathode remain the same with respect to the sample, but the pressure wave enters from the cathode instead of the anode (see Figure 4), in such a way that the integrity of the pressure pulse is now preserved at the cathode.

- $75$  Surprisingly enough, it is clear from Figure 4b that when the pressure wave enters the sample from the cathode, almost no interface peak is visible. On the contrary, the signal begins to rise later indicating that up to about the middle of the sample, there is no detectable electric field on the cathode side. The absence of electric field on the cathode side indicates that there is no charge buildup which implies in turn a substantial charge transfer across that interface, contrarily to what happens at the anode interface
- <sup>80</sup> where charges are accumulated. Moreover, since the zero electric field extends up to the middle of the sample, overall charge neutrality can be inferred in that region. On the anode side, a large amount of negative charges (negative slope in the signal) are clearly accumulated inside the RTO crystal in the vicinity of the anode, with positive charge (positive slope) on the surface of the anode.

In order to further confirm this surprising behavior by avoiding any parasitic signal due to the presence <sup>85</sup> of epoxy resin and to understand the previous observation of colossal capacitance also in polycrystalline samples, the same measurements were performed in the polycrystalline samples prepared as described



Figure 4: Electric field distribution inside a single crystal sample (See Figure 2a). The electrodes are materialized by the grey boxes.(a) Measurement when the pressure pulse enters through the anode. Electric field is clearly non-zero at the anode. (b) Measurement when the pressure pulse enters through the cathode. No electric field is detected at the cathode and well inside the material.

above. The results are presented in Figure 5 for three consecutive measurements on the same sample. The first observation, which comes as a surprise, is that the distribution of electric field is similar in a single crystal and a polycrystalline sample. In Figure 5a is plotted the output signal after applying

- <sup>90</sup> 20V to the sample during 5 h. Due to the absence of epoxy here, the positions of the input and output interfaces are easily determined by additional measurements during which first the sample and then the outcome contact are replaced by piezoelectrics. The electrodes are materialized by the gray boxes. Again here, negative charges are accumulated in the vicinity of the anode inside the material while the electric field is null at the cathode and well inside the sample on the cathode side. Figure 5b displays the
- 95 result of the same measurement taken 1 min after the voltage is reversed to  $-20$  V, which shows a rapid decrease of the electric field, and after 5h, where the signal is fully reversed with respect to Figure 5a. It is noteworthy that the expected dispersion of sound in the polycrystallline material is consistent with the decreasing signal resolution with depth.

#### **3. Results and discussion**

<sup>100</sup> The first outcome of the measurements is to unveil the sign of the ionic charge carriers without ambiguity. The ionic mobile species are negatively charged, and there are no detectable accumulated mobile cations. This comes at odd with previous assumptions [8, 12] where mobile carriers were hypothesized



Figure 5: Electric field distribution obtained in a polycrystalline sample (See Figure 2b). The electrodes are materialized by the grey boxes. (a) After applying a constant voltage of +20 V during 5 h. (b) 1 min (dotted line) and 5 h (solid line) after commuting the voltage from +20 V to −20 V. The electric field is rapidly decreased at the former anode and rapidly builds up in the vicinity of the new anode.

to be oxygen vacancies, and implies that the colossal capacitance takes place only at the anode. For this to occur, the charge transfer at the anode has to be almost zero. Possible candidates for mobile ions are  $105 \text{ }$  O<sup>2−</sup> ions or OH<sup>−</sup> ions coming from potential water contamination. The material is indeed found to be strongly hygroscopic [12] and bears similarities with vanadate structures [14] and in particular vanadium pentoxide gels [15, 16, 17, 18] or with hydrated ruthenium oxides [19]. This remains to be further investigated, as well as the full evolution of the distribution of electric field as function of time under various thermal, hygroscopic and electric conditions.

- <sup>110</sup> This accumulation of charge is a further confirmation of the predominant electrostatic nature of the giant capacitance. As a matter of fact, in the case of electrochemical capacitance, as described and modelled by Jamnik and Maier [20], the system would be locally electroneutral, and the pressure pulse propagation method would not reveal any charge accumulation. On the contrary, our measurements clearly demonstrate negative charges inside the RTO on the anode side and positive charges on the anode it-
- <sup>115</sup> self. The integral of the corresponding electric field obtained by calibrating the signal using a reference sample is indeed shown to correspond within 5% to the applied voltage, leaving very little margin for electrochemical capacitance.

The second consequence is that the cathode interface is purely Ohmic, i.e. there is a strong charge transfer at the interface and sufficient conduction (either electronic or ionic) in the material near the cathode

<sup>120</sup> to ensure a perfectly null electric field both on the cathode and inside the crystal over sizable length, about typically 0.5 mm. Therefore the system behaves as if the cathode had virtually shifted from the electrode to the middle of the sample. The mechanisms for conduction close to the cathode could be either that the departing ions have created electron or hole conduction, or that the ionic conductivity is sufficient to ensure metallic-like behavior in this region. Similar behavior has already been predicted

- in crystals including titanium [21]. A possible mechanism is electron hopping between  $Ti^{3+}$  and  $Ti^{4+}$ mixed valence cations leading to electron delocalization [22]. For reasons that are still to be understood, the cathode and the anode interfaces thus behave quite differently in this system, with substantial charge transfer at the cathode and immaterial charge transfer at the anode. This is true both for Al electrodes used for the single-crystals and Au electrodes used for the ceramics, thus pointing to an intrinsic phys-
- <sup>130</sup> ical property of the material. This could be related to a strong electron/hole asymmetry for transfer processes. Although these processes are still to be elucidated as well as the dominant mechanism for conduction near the cathode, the present observation can be an explanation for the colossal equivalent permittivity observed in RTO. Indeed, in contradiction to conventional electrolytes where both charge polarities coexist, our findings indicate a charge compensation in the bulk on the cathode side, which
- <sup>135</sup> enables much higher charge accumulation at the anode. In any case the mechanism at play has to be highly reversible since we did not observe degradation in the I/V curves after thousands of cycles.

Finally, the third remarkable observation is that single crystals and ceramics behave qualitatively the same way. This rules out the possibility of accumulation of ions inside the grains, since in this case, an accumulation of charges should be seen also on the cathode side. This therefore tends to indicate that

<sup>140</sup> ionic motion is not only intragrain but also intergrain. In addition it opens the way to easier and cheaper possibilities for application in the domain of supercapacitors for energy storage.

#### **4. Conclusion**

We have shown from pressure-wave-propagation measurements that negative ionic carriers accumulate in Rb<sub>2</sub>Ti<sub>2</sub>O<sub>5</sub> single crystals and polycrystalline samples. When anions (possibly O<sup>2−</sup> or OH<sup>−</sup>) accumu-<sup>145</sup> late at the anode, the cathode and the region nearby experience an Ohmic-like behavior resulting in a zero electric field in that region, thus leading to a "virtual cathode" extending deep inside the material. These findings potentially explain the colossal capacitance observed in  $Rb_2Ti_2O_5$  as it enables much higher accumulated charge density. Further work is required to understand the charge transfer processes at the cathode, the dynamics of the charge relaxation as well as the nature of the mobile ions.

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