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Site and Oxidation-State Specificity Yielding Dimensional Control in **Perovskite Ruthenates**

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In this communication we report the synthesis and structural characterization of the first ruthenium-based 2:1 ordered perovskite, Sr₃CaRu₂O₉. Ruthenium's unusual ability to readily adopt both IV(d⁴) and V(d³) oxidation states, when prepared in air, endows ruthenates with a host of different possible structures. The magnetic and electronic properties elicited by these many structural motifs have been equally diverse. For example, SrRuO₃ is a ferromagnetic metal¹ whereas CaRuO₃ is paramagnetic at all temperatures,² although ferromagnetism may be induced by transition-metal substitution.3 Sr₂RuO₄ is the only example of a layered noncuprate oxide superconductor⁴ and GdSr₂RuCu₂O₈ shows the microscopic coexistence of magnetic order and superconductivity. 5,6 These fascinating properties, which undoubtedly arise from the large degree of hybridization between oxygen p-orbitals and the extended ruthenium 4d-orbitals, once again focus attention on the synthesis of new materials and the interplay between stoichiometry and structure.

The structure of Sr₃CaRu₂O₉, or Sr(Ca_{1/3}Ru_{2/3})O₃, is characterized by a 2:1 ordering of Ru⁵⁺ and Ca²⁺ over the six-coordinate B-sites of the perovskite lattice (Figure 1). Similarly ordered materials based on Nb5+ or Ta5+ are known and serve as good dielectrics for microwave applications. 7 In the latter, the majority octahedral cation shifts slightly toward one face of its octahedron. Consequently, this type of ordering has only been observed when the majority site is occupied by a d⁰ cation (e.g., Nb⁵⁺, Ta⁵⁺, etc.). To our knowledge, Sr₃CaRu₂O₉ is the first reported example of this structure type based on a majority metal with d electrons.

Sr₃CaRu₂O₉ was synthesized by solid-state reaction of stoichiometric amounts of the alkali earth carbonates (99.99%) and dried RuO₂ (99.95%). The reactants were ground and calcined for 12 h at 850 °C in air. Additional cycles of grinding and heating to 1200 °C were carried out until phase purity was achieved as determined by powder X-ray diffraction (PXD).

Excluding several weak peaks ($I/I_{\text{max}} \leq 0.02$), the PXD pattern of Sr₃CaRu₂O₉ could be modeled by a 2:1 ordered perovskite framework with untilted octahedra8 using the trigonal space group P-3m1 with a = b = 5.69 Å and c = 7.02 Å. The presence of unindexed peaks suggested the existence of a supercell. A subsequent electron diffraction study confirmed the presence of a monoclinic unit cell with $a \approx 17$ Å, $b \approx 5.6$ Å, $c \approx 9.6$ Å and β $\approx 125^{\circ}$. The extinction conditions were compatible with space group $P2_1/c$. All peaks in the PXD pattern could be indexed using this unit cell. Energy-dispersive X-ray analysis performed on numerous crystallites confirmed the Sr:Ca:Ru ratios to be 3.01:1.02:1.97. Hydrogen reduction thermogravimetric analysis indicated a weight loss of 12.29% consistent with an oxygen stoichiometry of nine (calculated weight loss 12.33%). For the structure determination, neutron diffraction data were collected on

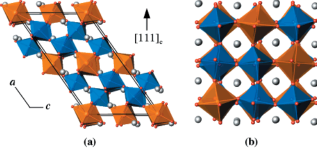


Figure 1. Structure of Sr₃CaRu₂O₉ viewed along the [010] direction (a) and along one of the pseudocubic axes (b). RuO₆ octahedra are blue, CaO₆ octahedra are orange, and Sr atoms are gray. The cubic [111] direction, perpendicular to the close-packed planes, is shown.

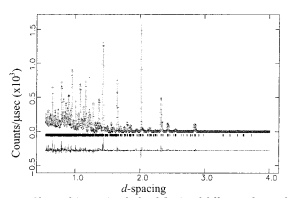


Figure 2. Observed (crosses), calculated (line) and difference (bottom line) neutron diffraction pattern of polycrystalline Sr₃CaRu₂O₉. Allowed Bragg reflections are marked (vertical bars).

a polycrystalline sample. The atomic positions used for the initial structural model were derived from the P-3m1 subcell. The model refined well in $P2_1/c$ with lattice parameters a = 17.1540(5) Å, b= 5.6891(2) Å, c = 9.8627(3) Å, $\beta = 125.0(1)^{\circ}$ and Z = 4 (Figure 2).9

Sr₃CaRu₂O₉ consists of corner-connected RuO₆ and CaO₆ octahedra sequenced {...-Ru-Ru-Ca-...} along each of the pseudocubic directions of perovskite (Figure 1b). Equivalently, owing to the ordering of the B-site cations, the structure can be viewed10 as layered parallel to the close-packed planes (i.e., the (111) planes of cubic perovskite) with one layer of CaO₆ octahedra alternating with two layers of RuO₆ octahedra (Figure 1a). The 2:1 cation ordering over the octahedral sites is due to the size and charge differences between ruthenium and calcium.

The sizable mismatch between Ru-O (1.97 Å) and Ca-O (2.4 Å) bond-lengths leads to significant tilting, twisting and deformation of the octahedra. The bond lengths reveal that while the two crystallographically distinct CaO₆ octahedra are under compression they remain relatively undistorted (standard deviation in bond lengths, $\sigma_{\rm BL} = 0.012$, 0.007). In contrast, the two distinct RuO₆

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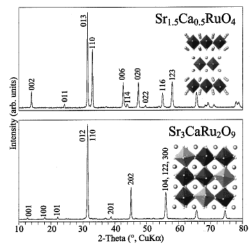


Figure 3. Top: X-ray diffractogram of polycrystalline Sr_{1.5}Ca_{0.5}RuO₄ synthesized by reduction of Sr₃CaRu₂O₉. Bottom: Diffraction pattern of polycrystalline Sr₃CaRu₂O₉ prepared by oxidation of Sr_{1.5}Ca_{0.5}RuO₄. For clarity, peaks in the Sr₃CaRu₂O₉ pattern are indexed based on the *P-3m*1

octahedra are under slight tension and more distorted ($\sigma_{BL} = 0.113$, 0.042). In the absence of cation ordering, the tilting may be described as $a^+b^-b^-$ using Glazer's notation. This tilt system, when combined with the 2:1 cation ordering, is consistent with the selection of P2₁/c as the space group. 12 Although Ca²⁺ and Ru⁵⁺ are expected to have close to regular octahedral coordination, such distortions are also observed in Ca₃CaNb₂O₉.¹³

Sr₃CaRu₂O₉ and Ca₃CaNb₂O₉ are strikingly similar. In each case, the atom connectivity and the B-site ordering vector, perpendicular to the AO₃ close-packed planes, are the same. Both share the same tilt pattern, space group and lattice parameters, however the a and c axes are switched. The altered orientation produces a subtle structural change owing to the different orientation of the c-glide. As represented in Figure 1a, the glide in $Sr_3CaRu_2O_9$ lies in the cdirection, whereas in Ca₃CaNb₂O₉ it is along the a direction. Levin et al. have anticipated this possibility in their description of Ca₃CaNb₂O₉, although it was not observed in that system.¹³

The formation of three-dimensional (3D) Sr₃CaRu₂O₉ is remarkable considering the existence of the two-dimensional (2D) Sr_{1.5}-Ca_{0.5}RuO₄ (i.e., Sr₃CaRu₂O₈)¹⁴ with the same ratio of cations and the pseudo-one-dimensional Sr₄Ru₂O₉. ¹⁵ Perhaps it is not surprising, then, that both were often encountered as secondary phases during intermediate reaction steps. Sr_{1.5}Ca_{0.5}RuO₄ is metallic and adopts a layered n = 1 Ruddlesden-Popper structure, wherein the Ca²⁺ and Sr2+ occupy the nine-coordinate A-sites between isolated RuO_{4/2}O₂ slabs.¹⁴ Although structurally very different, Sr_{1.5}Ca_{0.5}-RuO₄ and Sr₃CaRu₂O₉ have stoichiometries that differ only in their oxygen content. To elucidate the effect of the oxygen content and the ruthenium valence, experiments were undertaken in argon, air, and oxygen. Sr₃CaRu₂O₉, initially synthesized in air, was quantitatively converted to Sr_{1.5}Ca_{0.5}RuO₄ by treatment in 1 atm flowing argon for 24 h at 1200 °C. Reoxidation returns the original 3D structure after 24 h at 1000 °C in 1 atm flowing oxygen (Figure 3). This transformation can be repeated multiple times with the same result. Temperature-dependent resistivity measurements indicated Sr₃CaRu₂O₉ is semiconducting ($\rho_{100K}/\rho_{300K} = 17.8$). Thus, by controlling the synthesis conditions, structures of different dimensionalities and properties result.

This structural transformation is made possible by the unique combination of the oxidation state of ruthenium and the coordination of calcium. First, ruthenium's two available oxidation states allow two oxygen stoichiometries yielding two different structures. Second, calcium, which can occupy six-, nine- and twelvecoordinate sites, is essential for the formation of a 3D perovskite structure. Larger cations would not fit into the octahedral sites of the perovskite structure and would yield structures related to Sr₄Ru₂O₉ or multiphasic products.

The Sr-Ca-Ru-O system provides a significant opportunity with respect to thin-film fabrication of metal-insulator-metal (MIM) or metal-insulator-semiconductor (MIS) devices. Using a single precursor composition the properties of a deposited film could be regulated solely by the oxygen partial pressure.

Sr₃CaRu₂O₉ is the first example of a 2:1 ordered perovskite with a non-d⁰ majority metal. Previous examples had been limited to metals such as Nb5+, Ta5+, etc. In relation to other ruthenates, the 2:1 B-site ordering in Sr₃CaRu₂O₉ is intermediate between the allruthenium ferromagnetic SrRuO₃¹ and the 1:1 ordered antiferromagnetic Sr₂YRuO₆. ¹⁶ Work is in progress to demonstrate that, in general, we expect a number of homeotypic ruthenates with interesting magnetic and electronic properties to exist based on Aand B-site substitutions. 17,18

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Supporting Information Available: Tables listing detailed crystallographic data, atomic positions, bond lengths and selected bond angles based on the Rietveld refinement of the powder neutron diffraction data of Sr₃CaRu₂O₉; an indexed PXD profile and electron diffraction micrographs of Sr₃CaRu₂O₉ (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.

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