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Crossover from itinerant-electron to localized-electron behavior in 
Sr$_{1-x}$Ca$_x$CrO$_3$ perovskite solid solution

Youwen Long$^1$, Liuxiang Yang$^1$, Yuxi Lv$^1$, Qingqing Liu$^1$, Changqing Jin$^1$, Jianshi Zhou$^2$ and John B Goodenough$^2$

$^1$ Institute of Physics, Chinese Academy of Sciences, Beijing 100190, People’s Republic of China
$^2$ Texas Materials Institute, University of Texas, 1 University Station, C2200, Austin, TX 78712, USA

E-mail: ywlong@iphy.ac.cn and Jin@iphy.ac.cn

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Abstract
Polycrystalline samples of the perovskite family Sr$_{1-x}$Ca$_x$CrO$_3$ have been prepared at high pressure and temperature in steps of 1/6 over the range 0 ≤ x ≤ 1. Rietveld analysis shows a series of structural phase transitions from cubic to tetragonal to orthorhombic with increasing x. The cubic samples have no long-range magnetic order; the other samples become antiferromagnetically ordered below a $T_N$ that increases with x. At ambient pressure, the electric transport properties of the cubic and tetragonal phases are semiconducting with a small (meV range) activation energy that increases with x; the orthorhombic phase exhibits variable-range hopping rather than the small-polaron behavior typically found for mixed-valent, localized-electron configurations. Above a pressure $P = P_C$, a smooth insulator–metal transition is found at a $T_{IM}$ that decreases with increasing x for a fixed x; $P_C$ increases with x. These phenomena are rationalized qualitatively with a π* band model having a width $W_{π}$ that approaches crossover from itinerant-electron to localized-electron behavior as $W_{π}$ decreases with increasing x. The smaller size of the Ca$^{2+}$ ion induces the structural changes and the greater acidity of the Ca$^{2+}$ ion is primarily responsible for narrowing $W_{π}$ as x increases.

(Some figures in this article are in colour only in the electronic version)

1. Introduction

Since 1967 [1], it has been recognized that the Cr$^{4+}$:3d$^2$ electron configuration changes from itinerant-electron behavior in SrCrO$_3$ to strongly correlated electron behavior in CaCrO$_3$, but a systematic study of the evolution of the transition from itinerant to strongly correlated electronic behavior in Sr$_{1-x}$Ca$_x$CrO$_3$ solid solution was little known because these perovskites can only be synthesized under high pressure. Recently, SrCrO$_3$ and CaCrO$_3$ have received renewed interest due to the findings of an anomalous electronic state, microstrain-sensitive phase separation and orbital ordering [2–5]. Castillo-Martínez et al prepared Sr$_{1-x}$Ca$_x$CrO$_3$ solid solution and studied the structural properties in detail [6, 7]. Unfortunately, however, considerable impurity phases, which varied from 10% to 25% in weight for x ≥ 0.2, existed in their samples. It is therefore desirable to prepare higher quality samples and study the intrinsic physical properties.

It is also of interest to compare the properties of the Sr$_{1-x}$Ca$_x$CrO$_3$ system with those of the RVO$_3$ (R = rare-earth) perovskites in which the V$^{3+}$:3d$^2$ configuration is localized for all R$^{3+}$ ions. The Cr$^{4+}$ and V$^{3+}$ ions in these AMO$_3$ perovskites occupy corner-shared octahedral sites in which the strength of the (180° − $Φ$) M–O–M interactions (M = transition metal) determines whether the electrons are localized or itinerant. Moreover, the octahedral-site cubic-field splitting of a 3d$^2$ configuration into a t$^2_e^0$ configuration does not fully quench the orbital angular momentum of the threefold-degenerate π*-bonding t orbitals.
xy, yz ± izx. However, in the RVO\textsubscript{3} system, a cooperative rotation of the VO\textsubscript{6} octahedra lowers the crystal symmetry from its ideal cubic-perovskite structure with 180° M–O–M bonds to orthorhombic; and in the orthorhombic structure, local octahedral-site distortions split the threefold-degenerate t orbitals, suppressing the orbital angular momentum [8]. These distortions bias the ordering of the t orbitals in the RVO\textsubscript{3} perovskites [9]. CaCrO\textsubscript{3} has the same orthorhombic structure whereas SrCrO\textsubscript{3} has the ideal cubic structure [10, 11].

In this paper, we report on the synthesis of higher quality Sr\textsubscript{1−x}Ca\textsubscript{x}CrO\textsubscript{3} perovskite solid solution. By using a π∗-band model, the electronic state evolution from itinerant-electron to localized-electron behavior is studied on the basis of systematic changes in the crystal structure, electric transport, magnetism, and pressure-induced insulator–metal transitions.

2. Experimental details

Polycrystalline Sr\textsubscript{1−x}Ca\textsubscript{x}CrO\textsubscript{3} samples were prepared by using a high pressure and high temperature technique with highly pure SrO, CaO, and CrO\textsubscript{2} starting materials. The pressure we used was 6.0 GPa, and the temperature was changed from 900 to 1200°C with a step of 25°C to find the optimum synthesis condition for each sample. The appropriate reactants were thoroughly mixed and then pressed into pellets 6 mm in diameter and about 2 mm in height. After packaging in Pt foil, the pellets were inserted into pyrophyllite assemblies. Graphite tubes were used as heaters. Pressure was generated by a cubic-anvil-type high pressure apparatus. In order to obtain single-phase SrCrO\textsubscript{3} and CaCrO\textsubscript{3}, a 5% and 10%, respectively, excess SrO and CaO was used; the excess CaO and SrO after the reaction was washed out with dilute acid [10]. All the samples were black.

The sample quality and crystal structure were determined using powder x-ray diffraction (XRD) with a Rigaku diffractometer equipped with monochromatic Cu Kα radiation (50 kV, 200 mA). Diffraction data were collected in the 2θ range from 20 to 135° in steps of 0.02°. The counting time at each step was 5 s. Crystallographic parameters were obtained by a Rietveld full-profile refinement made with the GSAS program [12].

Four-probe ambient-pressure resistivity measurements were made in zero and 6 T magnetic fields from 5 to 300 K. The temperature and magnetic field dependence of the magnetization were measured with a commercial superconducting quantum interference device (Quantum Design, MPMS-5). A four-probe method was used to measure the resistance under pressure in a diamond-anvil cell; the culet of the diamond was 800 μm in diameter. T301 stainless steel was used as a gasket in which a 300 μm hole was drilled as the sample chamber. Gold wires with 12 μm diameter were used as electrical probes. To obtain a satisfactory hydrostatic environment, fine MgO powder (1 μm in diameter) was used for the pressure-transmitting medium. In addition, the gasket was covered with a thin layer of Al\textsubscript{2}O\textsubscript{3} powder to prevent any short circuit of the electrical probes. Pressures were calibrated by the ruby-fluorescence method [13].

3. Results

3.1. Crystal structures

Figures 1(a)–(c) display the results of the Rietveld analysis of three representative samples of the Sr\textsubscript{1−x}Ca\textsubscript{x}CrO\textsubscript{3} system. For x = 0 and 1/6, the XRD data fit well with the ideal simple-cubic structure, space group Pm\textsubscript{3}m (No. 221) as shown for x = 0 in figure 1(a). For x = 1/3 and 1/2, the room-temperature symmetry changes to tetragonal, space
CrO be a two-phase one; it refines in both the tetragonal 
Glazer notation [14]. The sample with 
$Pbnm$ (No. 62) as shown in figure 1(c), where the cooperative 
and 1.0 samples are refined in the orthorhombic space group 
increasing small amount of Cr
$Pbnm$ from 180 
not require a distortion of the local octahedra, but it bends 
and 440 as shown in figure 2. This cooperative rotation does 
peak into 206 and 422, and the cubic 400 peak into 008 
of CaCrO 
phases.
do signal the presence of cation vacancies in the perovskite 
perovskite phases does not appear to be significant, but they 
influence of the impurities on the intrinsic properties of the 
impurity amount reported in [6] (about 16 wt%). Fortunately, 
be 6.7 wt% (see figure 1(c)). This value is much less than the 
Table 1 summarizes the refined crystallographic data for 
the family of Sr$_{1-x}$Ca$_x$CrO$_3$. The crystal structural evolution 
observed in this study is consistent with previous reported 
results [6, 10, 11]. The $x$ dependence of the unit-cell volume 
per molecular $ACrO_3$ unit ($V_0/Z$) obeys Végard’s law in the 
region 0 $≤ x ≤ 2/3$ although a cubic-to-tetragonal phase transition occurs in this region; but as can be seen in figure 3, 
there is a discontinuity in the law, if not also in $V_0/Z$ itself, 
where there appears to be a first-order change from tetragonal 
to orthorhombic symmetry at $x = 2/3$. Note that the present 
$x − V_0/Z$ relationship is different from the reported results 
in [6]. This is probably related to the considerable amount of 
impurity in previous samples, which were prepared at 6.5 GPa and 
950 °C for all the intermediate $x$ values. In this study, to 
obtain higher quality samples, the synthesis pressure was set 
to 6.0 GPa but the temperature was gradually changed from 
900 to 1200 °C with a step of 25 °C. The data presented here 
were taken from the best sample for each composition.

3.2. Electric transport and magnetism at ambient pressure

Figures 4(a)–(c) show the temperature dependence of the resistivity $\rho(T)$ for the $x = 0$, 1/2, and 1.0 samples, which 
are representative of the three different structural phases. 
Above 80 K, the $\rho(T)$ of SrCrO$_3$ is only weakly temperature 
dependent, and the data can be fitted by using a thermal 
activation model with the function $\rho(T) \sim \exp(E_a/k_B T)$, 
as shown in the inset of figure 4(a). Here, $E_a$ and $k_B$ present 
the Boltzmann constant and activation energy, respectively. 
The fitted $E_a$ for the cubic SrCrO$_3$ is 5.6 $\times 10^{-3}$ eV. A similar $\rho(T)$ behavior above 150 K is found for the tetragonal 
sample with $x = 1/2$, but with an increased $E_a = 1.8 \times 10^{-2}$ eV (see the inset of figure 4(b)). However, the $\rho(T)$ of orthorhombic CaCrO$_3$ is higher by about four orders of magnitude and fits to $\rho(T) \sim \exp(T_0/T)^{1/4}$ as presented 
in the inset of figure 4(c), which is characteristic of Mott 
3D variable-range hopping [15]. When a magnetic field 
(6 T) is applied for the cubic SrCrO$_3$, the $\rho(T)$ curve is 
almost unchanged with respect to that for zero field. In the
Table 1. Crystal-structure parameters from the Rietveld refinements of Sr$_{1-x}$Ca$_x$CrO$_3$. (Note: in Pm3m symmetry, Sr/Ca, Cr, and O occupy the special sites 1a (0, 0, 0), 1b (0.5, 0.5, 0.5), and 3c (0.5, 0.5, 0.5), respectively. In the I4/mcm space group, Sr/Ca, Cr, O1, and O2 occupy the special sites 4b (0, 0.25, 0.25), 4c (0, 0, 0), 4a (0, 0, 0.25), and 8h (x, y, z), respectively. In Pbnm symmetry, Ca/Sr and O1 occupy the special sites 4c (x, y, 0.25) and Cr occupies the special site 4b (0.5, 0.5, 0).

Table 2. Antiferromagnetic transition temperatures $T_N$ (K) of the Sr$_{1-x}$Ca$_x$CrO$_3$ family. (Note: C = cubic, T = tetragonal, O = orthorhombic.)

3.3. The pressure-induced insulator–metal transition

Figure 5 shows the pressure-dependent resistance at 280 K for three representative phases: the cubic SrCrO$_3$, the tetragonal Sr$_{2/3}$Ca$_{1/3}$CrO$_3$, and the orthorhombic CaCrO$_3$. A sharp initial decrease in resistance with applied pressure is evident in these samples. The temperature dependence of the resistance of these three samples, plotted as ln $R$ vs. $T$, is shown for several pressures in figures 6(a)–(c). The insets show that the slopes d$R$/dT change sign at a critical temperature $T_M$(P) that decreases with increasing pressure. In SrCrO$_3$, $T_M$ falls below 300 K for pressures $P > 5$ GPa; in Sr$_{2/3}$Ca$_{1/3}$CrO$_3$ and CaCrO$_3$ this occurs at pressures $P > 15$ GPa.

4. Discussion

An itinerant-electron model of the CrO$_3$ array of the Sr$_{1-x}$Ca$_x$CrO$_3$ system would have an antibonding band of t-orbital parentage, 1/3 filled. However, on the approach to a transition from itinerant to localized electronic behavior, the on-site electrostatic energy $U_π$ may create an energy gap between the Cr$^{3+}$/Cr$^{4+}$ and Cr$^{4+}$/Cr$^{3+}$ redox couples that is larger than the $π^*$ bandwidth, i.e. $W_π < U_π$. Our previous measurements [2] of the thermoelectric power for SrCrO$_3$ and CaCrO$_3$ have indicated a small overlap of the Cr$^{4+}$/Cr$^{3+}$ redox couples ($W_π ≈ U_π$) in SrCrO$_3$ and a $U_π > W_π$ in CaCrO$_3$.

Actually, the existence of the Cr$_2$O$_3$ impurity signals that Cr vacancies are present; the presence of Cr vacancies means not only that the CrO$_3$ array is somewhat oxidized, which lowers the Fermi energy $E_F$ into the Cr$^{3+}$/Cr$^{4+}$ couple, but also that there is a perturbation of the periodic potential that would introduce Anderson-localized states [16] above a mobility edge μ in at the top of the Cr$^{3+}$/Cr$^{4+}$ $π^*$ redox band. An $E_F > μ$ would have localized states at $E_F$, and an $E_F < μ$ would have itinerant-electron states at $E_F$. As $E_F$ approaches $μ$ from above, thermal excitation of electrons from the itinerant-electron density of states to
Figure 4. Temperature dependence of the resistivity at ambient pressure and 0 T (open circles) and 6 T (solid circles) for (a) \( x = 0 \), (b) \( x = 1/2 \), and (c) \( x = 1 \) in the \( \text{Sr}_{1-x}\text{Ca}_x\text{CrO}_3 \) family. The solid lines in the insets show the fitting results as described in the text.

Figure 5. Pressure dependence of the resistance for \( x = 0, 1/3 \) (inset), and \( x = 1 \) samples in the \( \text{Sr}_{1-x}\text{Ca}_x\text{CrO}_3 \) family at 280 K.

Therefore, metallic conductivity. In order to interpret the observation of the insulator–metal transition at \( T_{\text{IM}}(P) \) and the other observations, we must consider how the structure and bandwidth are influenced by substitution of \( \text{Sr}^{2+} \) by \( \text{Ca}^{2+} \) ions.

The tight-binding bandwidth is \( W_{\pi} \approx 12b_{\pi} \), where \( b_{\pi} \approx \varepsilon_{\pi}(\psi_i,\psi_j) \) is the expectation for an electron hop between nearest-neighbor Cr atoms. The overlap integral \( \langle \psi_i,\psi_j \rangle \) of the ligand-field \( t \) orbitals on neighboring Cr atoms at \( \mathbf{R}_i \) and \( \mathbf{R}_j \) increases with the covalent admixture into the \( t \) orbitals of the 2p orbitals of the bridging oxygen that \( \pi \)-bond with the Cr \( t \) orbitals. Because the \( \text{Cr}^{3+}/\text{Cr}^{4+} \) empty redox couple lies close to the top of the O 2p bands, the covalent admixture of O 2p into the ligand-field \( t \) orbitals is large enough to give a significant \( \langle \psi_i,\psi_j \rangle \) overlap integral over the bridging oxygen. Moreover, there is no contribution from the cubic-field splitting of the Cr 3d electronic states to the intra-atomic electrostatic energy \( U_{\pi} \) that inhibits formation of itinerant-electron states. Consequently, we find a \( W_{\pi} \approx U_{\pi} \) in the \( \text{CrO}_3 \) array as is made evident by the antiferromagnetic order in \( \text{CaCrO}_3 \) and the lack of long-range magnetic order in \( \text{SrCrO}_3 \).

We note that the substitution of \( \text{Ca}^{2+} \) for \( \text{Sr}^{2+} \) in the present \( \text{Sr}_{1-x}\text{Ca}_x\text{CrO}_3 \) family has two effects: one is physical, which influences primarily the structure, and the other is chemical, which has the more important influence on \( W_{\pi} \). The smaller size of the \( \text{Ca}^{2+} \) ion decreases the geometric tolerance factor \( t \equiv (\text{A–O})/\sqrt{2}(\text{Cr–O}) \), where \( (\text{A–O}) \) is the mean equilibrium Sr/Ca–O bond length of A-site cations and \( (\text{Cr–O}) \) is the equilibrium Cr–O bond length. In general, an ideal cubic-perovskite structure is found for \( t \approx 1 \); the structure adjusts to \( t < 1 \) by introducing cooperative octahedral rotations. The orthorhombic \( \text{Pbnm} \) rotations are commonly found for \( t \approx 1 \); but the transition from orthorhombic to cubic symmetry as \( t < 1 \) increases to \( t \approx 1 \) is not second order. Rotations that give tetragonal or rhombohedral symmetry may be found in the tolerance-factor interval between that for the orthorhombic structure and the \( t \approx 1 \) for the cubic structure. The observation of a
The more important influence on $W_{\pi}$ is chemical in origin. Chemically, the Ca$^{2+}$ ion is more acidic than the Sr$^{2+}$ ion, which makes it compete more strongly for the O 2p orbitals. Since the O 2p orbitals that $\pi$-bond with the Cr t orbitals also $\sigma$-bond with the alkaline-earth atoms, this competition reduces the covalent O 2p admixture into the Cr t ligand-field orbitals and therefore the overlap integral of $h_{\pi t}$. Any influence of the A-site cation on the one-electron energy $\varepsilon_{p}$ of $b_{\pi}$ is not obvious, but it is universally observed that the substitution of smaller, more acidic A-site cations in the AMO$_3$ perovskites narrows $\pi$-bands and that the chemical influence is more important than the bending of the M–O–M bond [17–20].

From this analysis, we conclude that stoichiometric SrCrO$_3$ contains itinerant $\pi^*$ electrons in a Cr$^{4+}$/Cr$^{3+}$ redox couple having a bandwidth $W_{\pi} \approx U_{\pi}$. The instability of a $W_{\pi} \approx U_{\pi}$ may be the origin of the stabilization of some oxidation of the CrO$_3$ array by the introduction of cation vacancies [21]; these vacancies perturb the periodic potential sufficiently to introduce Anderson-localized states above a mobility edge $\mu_C$ at the top of the Cr$^{4+}$/Cr$^{3+}$ couple. For a fixed oxidation state of the CrO$_3$ array, the position of the Fermi energy $E_F$ with respect to $\mu_C$ is lowered by a broadening of $W_{\pi}$. The transport data for SrCrO$_3$, figure 4(a), suggest an $E_F$ only a few meV above $\mu_C$, which results in semiconducting behavior with a small activation energy below 300 K. The primary experimental results of synchrotron XRD under high pressure reveal no change in crystal structure for SrCrO$_3$ and CaCrO$_3$ [22], indicating that the pressure-induced metallization is due to broadening of $W_{\pi}$ by increasing its O 2p character. The application of hydrostatic pressure broadens $W_{\pi}$ to reduce $E_F - \mu_C$ until, above 5.4 GPa for example in SrCrO$_3$, $E_F$ falls below $\mu_C$ and the sample becomes metallic above $T_{IM} < 300$ K as shown in figure 6(a); $T_{IM}$ falls to lower temperatures as $W_{\pi}$ increases with pressure. Similar behaviors occur in the tetragonal and orthorhombic phases in the Sr$_{1-x}$Ca$_x$CrO$_3$ family with increasing pressure (figures 6(b) and (c)). Moreover, electrons in Anderson-localized states may introduce localized spins that introduce a temperature-dependent component to the paramagnetic susceptibility; but these spins are not expected to give long-range magnetic order at low temperatures. The paramagnetic susceptibility of SrCrO$_3$ reveals no evidence of long-range magnetic order above 5 K, but a small difference between the zero field cooled and field cooled curves below 50 K may reflect the onset of some short-range magnetic order [2]. The same type of spin order may also take place below 50 K in the $x = 1/6$ sample. The initial sharp drop in resistance with applied pressure in our polycrystalline samples (figure 5) shows the presence of a significant grain-boundary resistance that could be responsible for the small activation energies of the resistivity in the $x = 0$ and 1/6 samples. Recently, a poor metallic behavior was predicted in theory and also detected from optical measurements for CaCrO$_3$ [23, 24]. However, electric measurement on the same sample suggested a nonmetallic state with $\rho > 1000$ $\Omega$ cm at low temperature. It looks as if the grain boundaries play a role in the electric transport properties in the Sr$_{1-x}$Ca$_x$CrO$_3$ family. The grain-boundary effects may also be x dependent in this system, but they should not influence qualitatively the results that we concluded in this study.
The structural data of figure 3 signal that an important change in the electronic properties may occur at a first-order transition from tetragonal to orthorhombic symmetry near $x = 2/3$. It has been argued from the virial theorem [25] that more localized, i.e. more strongly correlated, electrons have a longer equilibrium (M–O) bond length, but figure 3 shows that $V_0/Z$ versus $x$ for $0 \leq x < 2/3$ falls below the extrapolation of Végard’s law for the orthorhombic phase in the range $2/3 < x \leq 1.0$. However, a reduction in the Cr–O–Cr bond angle in the orthorhombic phase can not only accommodate a larger Cr–O bond length, but also introduces a shorter competitive Ca–O bond. Therefore, we conclude that in the orthorhombic phase, the Cr 3d electrons are more strongly correlated, i.e. are more tightly bound, even though the superexchange interaction cannot be described by the localized-electron second-order perturbation theory. Moreover, the transport data of figure 4(c) are consistent with a large $E_F - \mu C$. However, variable-range hopping rather than small-polaron behavior of the holes in the Cr$^{5+}$/Cr$^{4+}$ couple suggests that the $\pi^*$ electrons give rise to a band antiferromagnetism rather than a localized-electron antiferromagnetism. This conclusion is reinforced by the observation [10] of $dT_N/dP < 0$ in CaCrO$_3$ and also by the increase in $T_N$ from 83 to 90 K as $x$ increases from 5/6 to 1.0 shown in table 2. Where the superexchange perturbation theory is valid [26], an increase in $b_\pi$ would increase $T_N \sim b_\pi^2/\Delta E$, and a reduction in $b_\pi$ would decrease $T_N$.

For the interval $0 < x < 1.0$, we comment qualitatively on the evolution with $x$ of the physical properties without consideration of the effect of the variance introduced by the coexistence of two different A-site cations. The smaller Ca$^{2+}$ ion lowers the tolerance factor $t_1$, inducing increasing cooperative rotations of the CrO$_{6/2}$ octahedral sites as $x$ increases. The cubic-to-tetragonal transition may be smooth; it does not introduce any local octahedral-site distortion that would bias orbital order. However, narrowing of $W_\pi$ and bending of the Cr–O–Cr bond angles in the $a$–$b$ plane induce long-range antiferromagnetic order of the $\pi^*$ electrons with a $T_N$ that increases with decreasing $W_\pi$. We can predict that the magnitude of the spin density of a Cr site may also increase with $x$. In the tetragonal phase, the spins are not canted, and we predict that they are oriented along the $c$ axis in the tetragonal structure. However, the tetragonal–orthorhombic transition is first order, and the larger bending of the Cr–O–Cr angle, to 156° in CaCrO$_3$, is accompanied by local site distortions that bias ordering of the $t$ orbitals as in YVO$_3$, which has a localized V$^{3+}$-$t_2^0$ configuration. A recent neutron-diffraction study [24] shows the same orbital and magnetic order setting in below $T_N$ in CaCrO$_3$ as is found in LaVO$_3$.

5. Conclusion

We have synthesized at 6.0 GPa and 900–1200°C the polycrystalline Sr$_{1-x}$Ca$_x$CrO$_3$ perovskite solid solution in steps of $x = 1/6$ over the range $0 \leq x \leq 1$. Three room-temperature structures were observed: cubic for $x = 0$ and 1/6, tetragonal for $x = 1/3$ and 1/2, and orthorhombic for $x = 5/6$ and 1, in agreement with previous reports [6, 10, 11]. The tetragonal and orthorhombic phases coexisted in the $x = 2/3$ sample, characteristic of a first-order change between these phases. No long-range magnetic order occurs in the cubic phase, whereas long-range antiferromagnetic order appears in the tetragonal and orthorhombic phases with $T_N$ increasing with $x$. In both CaCrO$_3$ and LaVO$_3$, orbital ordering below $T_N$ resolves the orbital–spin frustration of the superexchange interaction responsible for the magnetic order. It would be of interest to know whether a similar resolution occurs below $T_N$ in the more itinerant antiferromagnetism of the tetragonal phase.

The evolution of the structure is well understood from the decrease in the geometric tolerance factor; it follows the pattern observed for all AMO$_3$ perovskite families. Moreover, the evolution of electronic properties can be understood to be due to a narrowing of the $\pi^*$ band of itinerant electrons with Ca$^{2+}$ substitution for Sr$^{2+}$; this narrowing is characteristic of other perovskites as a result of a greater competition of the smaller, more acidic A cation for the O 2p orbitals that $\pi$-bond with the $\pi$-bonding d orbitals on the transition metal M cations. A slight oxidation of the CrO$_3$ array is characteristic of lattice instabilities found in perovskites near the transition from itinerant to localized electronic behavior. The introduction of cation vacancies and the variances associated with the coexistence of two different A-site cations complicates a quantitative analysis of the evolution of physical properties with $x$, but the model of a $\pi^*$ band with a width $W_\pi \approx U_\pi$ allows a straightforward rationalization of the observed data, indicating a crossover from itinerant-electron to localized-electron behavior in Sr$_{1-x}$Ca$_x$CrO$_3$ solid solution [2, 4, 24].

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