Temperature-dependent optical conductivity of layered LaSrFeO$_4$

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Compounds with intermediate-size transition metals such as Fe or Mn are close to the transition between charge-transfer systems and Mott-Hubbard systems. We study the optical conductivity $\sigma(\omega)$ of insulating layered LaSrFeO$_4$ in the energy range 0.5 – 5.5 eV from 15 K to 250 K by the use of spectroscopic ellipsometry in combination with transmittance measurements. A multiplet structure is observed in both $\sigma^+(\omega)$ and $\sigma^-(\omega)$. The layered structure gives rise to a pronounced anisotropy, thereby offering a means to disentangle Mott-Hubbard and charge-transfer absorption bands. We find strong evidence that the lowest dipole-allowed excitation in LaSrFeO$_4$ is of Mott-Hubbard type. This rather unexpected result can be attributed to Fe 3d - O 2p hybridization and in particular to the layered structure with the associated splitting of the $e_g$ level. In general, Mott-Hubbard absorption bands may show a strong dependence on temperature. This is not the case in LaSrFeO$_4$, in agreement with the fact that spin-spin and orbital-orbital correlations between nearest neighbors do not vary strongly below room temperature in this compound with a high-spin 3$d^9$ configuration and a Néel temperature of $T_N = 366$ K.

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I. INTRODUCTION

Many transition-metal compounds with a partially filled 3$d$ shell show charge localization and insulating behavior. Typically, this is driven by the large on-site Coulomb repulsion $U$, which splits the conduction band into a lower and an upper Hubbard band (LHB and UHB). According to the Zaanen-Sawatzky-Allen scheme, one distinguishes between two kinds of correlated insulators, depending on the relative size of $U$ and the charge-transfer (CT) energy $\Delta$ between the transition-metal 3$d$ band and the highest occupied ligand level, e.g. oxygen 2$p$. In a CT insulator with $U > \Delta$, the charge gap is formed between O 2$p$ and the UHB. In contrast, Mott-Hubbard (MH) insulators show $U < \Delta$, and the states closest to the Fermi level predominantly have transition-metal character (see Fig. 1). This character is decisive for a quantitative description of doped compounds. For instance in a hole-doped CT system such as the high-$T_c$ cuprates, the mobile carriers predominantly have oxygen character.

Early transition-metal compounds are typically classified into the group of MH insulators, whereas late ones are identified as being of CT type. With increasing atomic number, $U$ increases whereas $\Delta$ decreases which reflects the decreasing spatial extension of the 3$d$ orbitals and the increasing electronegativity of the transition-metal ions, respectively. However, the character of oxides with intermediate size transition-metal ions, in particular Mn$^{3+}$, has been discussed controversially. On the basis of optical data, both LaMnO$_3$ and LaSrMnO$_4$ have been interpreted either as of CT type or of MH type while recently a dual nature of the optical gap has been proposed. This controversy arises due to the strong hybridization between Mn 3$d$ and O 2$p$ states (see Fig. 1). Early on, Mizokawa and Fujimori pointed out that $U > \Delta$ in LaMnO$_3$, but that the highest occupied O 2$p$ band shows a large admixture of 3$d$ character. Goessling et al. emphasized that the symmetry of the highest occupied, strongly hybridized band is determined by the 3$d$ band, which is essential for the optical selection rules and thus for a quantitative analysis of the optical data. They suggested that the manganites can be viewed as effective Mott-Hubbard systems, where $U_{\text{eff}}$ is strongly renormalized by hybridization (see Fig. 1). This scenario is supported by recent measurements on transition-metal difluorides MF$_2$ using x-ray emission spectroscopy. Due to the element selectivity of this technique, the contribution of the transition-metal LHB to the highest occupied states can be revealed even for $U > \Delta$.

![FIG. 1: (Color online) Sketch of a Mott-Hubbard insulator (top row, $U < \Delta$) and of a charge-transfer insulator (bottom row, $U > \Delta$) for a single half-filled 3$d$ orbital and degenerate O 2$p$ orbitals. The black dashed line depicts the increase of $U$ from top to bottom, $E_F$ depicts the Fermi level (dashed-dotted). Due to hybridization, we have to distinguish bonding (B), non-bonding (NB), and anti-bonding (AB) bands. For $U > \Delta$, the highest occupied anti-bonding band may still be classified as the lower Hubbard band with symmetry properties derived from the 3$d$ character, yielding an effective value $U_{\text{eff}} < \Delta$ (cf. Fig. 1 in Ref. [18] and Fig. 2 in Ref. [2]).]
At first sight, the situation is more transparent in the case of the ferrites RFeO$_3$ with trivalent Fe ions. Compared to the 3$d^4$ manganites, the stability of the high-spin 3$d^5$ state of Fe$^{3+}$ gives rise to a comparably large energy of MH excitations of roughly $U + 4J_H$, where $J_H$ denotes the intra-atomic Hund exchange. Indeed these compounds commonly are identified as CT systems.

However, the case of layered LaSrFeO$_4$ is still under discussion. It shows the same Fe$^{3+}$ 3$d^5$ configuration and has also been interpreted as a CT insulator based on optical reflectivity data measured up to 36 eV at room temperature with in-plane polarization of the electric field. In contrast, Omata et al. conclude from their resonant photoemission data that the valence band in LaSrFeO$_4$ is formed by a mixture of Fe 3$d$ and O 2$p$ states. Thus they characterize LaSrFeO$_4$ as an intermediate type CT and MH insulator, but they also mention that the states at the valence band edge mainly show O 2$p$ character. Here, we address the optical conductivity $\sigma_1(\omega)$ of LaSrFeO$_4$. The anisotropy of $\sigma_1(\omega)$ of this layered structure provides the key to disentangle MH excitations and CT excitations. Neglecting hybridization, a MH excitation refers to an electron transfer between neighboring Fe sites $i$ and $j$, $|3d^5_{i}\rangle \rightarrow |3d^6_{j}\rangle$, whereas a CT excitation refers to a transfer from O 2$p$ to Fe 3$d$, $|3d^5_{i}2p^5_{j}\rangle \rightarrow |3d^{n+1}_{i}2p^4_{j}\rangle$. In the presence of hybridization, this distinction between MH and CT excitations is still a valid classification scheme since the hybridized states retain their original symmetry and thus follow the same selection rules. As discussed by Goessling et al. for the case of LaSrMnO$_4$, MH excitations only contribute to $\sigma_1^\omega$, i.e., for polarization of the electric field within the 2D layer, but not to $\sigma_1^\sigma$. This reflects that Fe-Fe hopping between adjacent FeO$_6$ layers is negligible. On the contrary, CT excitations are observed in both, $\sigma_1^\omega$ and $\sigma_1^\sigma$, as each Fe site is surrounded by an oxygen octahedron. We find a clear anisotropy of the lowest dipole-allowed excitations in LaSrFeO$_4$, which suggests that they are of Mott-Hubbard type. We propose that this is driven by Fe 3$d$ - O 2$p$ hybridization and by the layered structure, the associated crystal-field splitting of the $e_g$ level pulls the lowest MH absorption band below the onset of CT excitations.

Additionally, we address the temperature dependence of the spectral weight. In general, CT and MH excitations show different spin and orbital selection rules. The spectral weight of MH excitations is expected to be strongly affected by a change of the nearest-neighbor spin-spin and orbital-orbital correlation functions.

In agreement with theoretical expectations, the spectral weight of the lowest absorption band in LaMnO$_3$, LaSrMnO$_4$, and 3$d^2$ RVO$_3$ changes by a factor of 2 - 3 due to the ordering of spins (and orbitals). This clearly demonstrates the (effective) MH character of these systems. However, the behavior of the MH insulators YTiO$_3$ and SmTiO$_3$ is still puzzling in this context. In YTiO$_3$, the spectral weight of the lowest MH excitation is expected to change by 25% between the paramagnetic and the ferromagnetic state. However, the increase around the ordering temperature $T_C$ amounts to only 5%, while, at the same time, larger changes are observed up to 300 K. Due to the three-dimensional character of the magnetic order, these larger changes far above $T_C$ can not be explained by a change of the spin-spin correlations. In the G-type antiferromagnet SmTiO$_3$, spin ordering is expected to suppress the spectral weight of the lowest MH excitation by about 50% for all crystallographic directions, but the observed effects are again much smaller and show even the wrong sign along $b$ and $c$. This behavior has been attributed to small changes of the orbital occupation. It is important to quantify the possible strength of other effects such as excitonic contributions, the thermal expansion of the lattice or bandstructure effects. In this context, layered LaSrFeO$_4$ with its stable 3$d^5$ state is an interesting candidate for a reference system.

Long-range antiferromagnetic spin order sets in at $T_N = 366$ K thus one expects only very small changes of the spin-spin and orbital-orbital correlation functions below room temperature. This far, not much is known about the optical spectra of LaSrFeO$_4$. Room temperature data has been obtained by the means of diffuse reflectance on powdered samples and by reflectivity measurements on single crystals, but only for in-plane polarization of the electric field.

The paper is organized as follows. The experimental details are described in Sec. II followed in Sec. III by a short description of the crystal structure and the electronic structure. A discussion of the expected multiplet splitting of MH and CT absorption bands is given in Sec. IV. Finally, in Sec. V we present our experimental data (Sec. V A) together with a discussion of the peak assignment (Sec. V B) and an analysis of the temperature dependence of the MH excitations (Sec. V C).

II. EXPERIMENT

Single crystals of LaSrFeO$_4$ have been grown using the floating-zone method. The purity, stoichiometry, and single-phase structure of the crystals was checked by neutron diffraction and by x-ray diffraction. Typical dimensions of crystals used in this study are a few mm along all three crystallographic axes. Ellipsometric data in the energy range 0.75 – 5.5 eV was obtained using a rotating-analyzer ellipsometer equipped with a retarder between polarizer and sample. The angle of incidence was $\theta = 70^\circ$. The ellipsometric measurements have been performed from 15 K to 250 K in a UHV cryostat with $p < 10^{-9}$ mbar. Window effects of the cryostat have been corrected using a standard Si wafer. Ellipsometry is a self-normalizing technique and does not require reference measurements, furthermore it yields the complex dielectric function directly without a Kramers-Kronig transformation. These are two significant advantages over conventional reflection measure-
is a valid approximation below 2 eV. Using this result for the analysis of the low-energy ellipsometry data yields a surface roughness with a thickness of approx. 6 nm.

In the top panel of Fig. 2, we compare our result for \( \varepsilon_2(\omega) \) with the so-called pseudo-dielectric function \( \varepsilon_2^{c,ps} \). Aspnes \[22\] proposed that the latter may serve as a reasonable approximation for the former under certain conditions. The main advantage of the pseudo-dielectric function is that it can be determined directly from the data measured for a single orientation. Figure 2 shows that the overall features are well reproduced, but large discrepancies are observed below the onset of strong absorption at about 3.5 eV, where \( \varepsilon_2^{c,ps} \) shows a spurious background. As far as Fe\(^{3+}\) compounds are concerned, a similar feature has been discussed in BiFeO\(_3\). In the case of LaSrFeO\(_4\), the transmittance data prove that this background is an artefact present only in \( \varepsilon_2^{c,ps} \) but not in \( \varepsilon_2 \).

III. STRUCTURE, SPINS, AND ORBITALS

The compound LaSrFeO\(_4\) crystalizes in the single-layered structure of K\(_2\)NiF\(_4\) with tetragonal symmetry \( I\bar{4}/mm\). The Fe\(^{3+}\) ions are octahedrally coordinated by oxygen ions, building perfect FeO\(_2\) square planes with 180° Fe-O-Fe bonds. The lattice constants at room temperature are \( a = 3.8744(1) \text{ Å} \) and \( c = 12.7134(3) \text{ Å} \).\[22\]

Nominally, there are five electrons in the 3d shell per Fe site. In the high-spin ground state, these five electrons yield a total spin of 5/2. Antiferromagnetic order has been observed below \( T_N = 366 \text{ K} \).\[22\] Our crystals do not exhibit any evidence for an additional magnetic phase transition.\[22\] In cubic approximation, the 3d level is split into the lower-lying \( t_{2g} \) and the higher-lying \( e_g \) levels. The magnitude of the \( t_{2g} - e_g \) splitting \( \Delta_{t_{2g}-e_g} = 10 \text{ Dq} \) is mainly determined by the Fe-O bond lengths, and it can be estimated to be roughly 10 Dq = 1.0-1.5 eV in LaSrFeO\(_4\).\[16\]\[18\]

The OFeO octahedra show a sizeable tetragonal distortion with Fe-O bond lengths of 1.9354 Å in the plane and 2.1486 Å perpendicular to it at 10 K.\[22\] Therefore, the \( t_{2g} \) manifold is split into the lower-lying doublet \( e_g' \) (yz and zx) and the higher-lying \( b_{2g} \) (or \( xy \)) level. At the same time, the \( e_g \) level splits into \( a_{1g} \) (3z\(^2\) - r\(^2\)) and \( b_{1g} \) (x\(^2\) - y\(^2\)), where the energy of the former is significantly reduced compared to the energy of the latter. In LaSrMnO\(_4\), these splittings have been determined from optical data, yielding \( \Delta_{t_{2g}}(d^4) = 0.2 \text{ eV} \) and \( \Delta_{e_g}(d^4) = 1.4 \text{ eV} \).\[15\]

For LaSrFeO\(_4\), we expect a similar value of \( \Delta_{t_{2g}} \) but a smaller value of \( \Delta_{e_g} \), since \( \Delta_{e_g} \) is enhanced in LaSrMnO\(_4\) due to the additional Jahn-Teller splitting of the singly occupied \( e_g \) level, which is evident from the larger value of \( c/a \). We emphasize that the pronounced elongation of the octahedra in LaSrFeO\(_4\) gives rise to a sizeable value of \( \Delta_{e_g} \), even in the absence of a Jahn-Teller contribution. Our results below yield \( \Delta_{e_g} \approx 0.8 \text{ eV} \).
IV. CHARGE-TRANSFER AND MOTT-HUBBARD EXCITATIONS

First, we focus on the physics of CT excitations. These result from the transfer of an electron from a ligand O 2p orbital into a Fe 3d orbital, [3d^5 2p^6] → [3d^4 2p^5]. Pisarev et al. present a detailed theoretical analysis of the CT excitations for undistorted FeO₆ octahedra. The highest occupied O states are of non-bonding character with symmetry t₁g(π), t₂u(π), t₁u(π), and t₁u(σ). Their relative energies are determined by, e.g., the different Madelung energies of 2p(π) and 2p(σ) orbitals and by the 2p(π) - 2p(σ) overlap. The t₁g(π) level is expected to be the highest in energy. Quantum-chemistry calculation for LaFeO₃ predict that t₂u(π), t₁u(π), and t₁u(σ) are lower by 0.8 eV, 1.8 eV, and 3 eV, respectively, and these are split by Δt₂u-Δt₁u = 10 Dq.

According to the parity selection rule, the even-even (from g type to g type) transitions from the t₁g(π) level at the top of the O band to the unoccupied t₂g(π) and e_g(σ) orbitals are forbidden. Additionally, the matrix elements for transitions from π to σ levels vanish for a single octahedron, thus only π-π and σ-σ transitions give rise to strong absorption. In summary, the onset of CT excitations is governed by the dipole-forbidden transition t₁g(π) → t₂g(π), followed by the strong dipole-allowed transitions t₂u(π) → t₂g(π) and t₁u(π) → t₂g(π). The next transition t₁g(σ) → e_g(σ) is roughly 2 eV higher in energy due to the splitting between t₂g(π) and e_g(σ) on the one hand and between 2p(π) and 2p(σ) states on the other hand. Thus in cubic approximation there are only two strong excitations in the energy range relevant to us. For the layered structure of LaSrFeO₄, we have to expect additional splittings. However, the large splitting Δe_g of the e_g level is not important for the lowest CT excitations, which correspond to an electron transfer into the t₂g(π) level.

We now address the MH excitations, which result from the transfer of an electron between neighboring Fe sites i and j via the σ or π bonding of the ligand O 2p orbital, [3d^5 3d^5] → [3d^4 3d^6]. Starting from the cubic approximation, the initial 3d^5 state has A_1 symmetry, corresponding to the (2S + 1) = 6-fold degenerate t^2g^2e^g high-spin state (see Fig. 3 (a)). According to the orbital selection rule, there is no overlap between t^2g and e_g orbitals on neighboring sites due to the undistorted 180° bonds in LaSrFeO₄. From the [6 A_1g(d^5)] initial state one can reach the final states [5 T^2g(t^2g, e^g)], [5 T^2g(t^2g, e^g)] (corresponding to an electron transfer between neighboring orbitals, see Fig. 3 (b)) and [5 E_g(t^2g, e^g)], [5 E_g(t^2g, e^g)] (transfer between e_g, orbitals, see Fig. 3 (c)). All reachable 3d^4 and 3d^6 states have total spin S = 2. The spectral weight of these transitions therefore strongly depends on the nearest-neighbor spin-spin correlations favoring antiparallel alignment of spins on neighboring sites (see Fig. 3), as given in the antiferromagnetically ordered state below T_N = 366 K. Since the orbital quantum number is preserved in the transition (cf. Figs. 3 (b) and (d)), both excitations have approximately the same energy. As a consequence, only one MH peak is expected in the optical spectra in the cubic approximation.

However, deviations from cubic symmetry play an important role in the tetragonal structure of LaSrFeO₄. First of all, the matrix elements for MH excitations between different FeO₂ layers can be neglected, thus MH excitations do not contribute to σ_1(ω). Second, lifting the degeneracy in particular of the e_g level enhances the number of observable absorption bands in σ_1(ω). An e_g splitting Δe_g = Δ[(x^2-y^2)-(3z^2-r^2)] on the order of 1 eV is expected due to the elongation of the FeO₆ octahedra, as discussed in Sec. 111. In the following, we neglect the much smaller splitting Δt₂g within the t₂g levels. We will show that Δe_g is crucial in order to pull the lowest MH absorption band below the onset of CT excitations.
The matrix elements for nearest-neighbor Fe - Fe transitions are summarized in Table I. Due to the undistorted 180° bonds of LaSrFeO$_4$, these hopping process conserve the orbital character. The single exception is the finite overlap between $3z^2-r^2$ on site $i$ and $x^2-y^2$ on a neighboring site. One thus expects three different MH peaks (referred to as MH1, MH2, and MH3 in the following) in $\sigma_1^a(\omega)$ with energy separation $\Delta_{eg}$: (1) The electron transfer from $x^2-y^2$ to $3z^2-r^2$ (MH1, see Fig. 3(d)). (2) The excitation from any orbital on site $i$ to an orbital of the same type on the neighboring site (MH2, see Fig. 3(e)). This excitation is expected at an energy of $\Delta_{eg}$ above MH1. The individual contributions have approximately the same energy because the orbital quantum number is preserved. (3) The excitation from $3z^2-r^2$ to $x^2-y^2$ (MH3, see Fig. 3(f)). This excitation is expected at an energy of $2\Delta_{eg}$ above MH1.

We use the matrix elements of Table I to calculate the relative spectral weight of the MH peaks. Summing up the individual contributions we find that the spectral weight of MH1 and MH3 is identical, whereas the spectral weight of MH2 is expected to be 3.8 times larger.

V. RESULTS

A. Experimental data

Figure 4 displays the dielectric function $\varepsilon(\omega) = \varepsilon_1^l + i\varepsilon_2^l$ ($l = a, c$) from 1.5 eV to 5.5 eV as obtained from the ellipsometric measurements. The real part of the optical conductivity $\sigma_1^a(\omega) = (\omega/4\pi) \cdot \varepsilon_1^a(\omega)$ is shown in Fig. 5. Overall, our data agree with the room-temperature data of $\sigma_1^a(\omega)$ reported in Refs. 8,31. Thus far, data for $\sigma_1^a(\omega)$ has not been reported. We find a striking anisotropy. Both $\sigma_1^a(\omega)$ and $\sigma_1^c(\omega)$ show a strong absorption band at 4 - 5.5 eV. However, in $\sigma_1^c(\omega)$ we find an additional peak at 3 eV and the shoulder at 3.8 eV is much more pronounced. In the following we argue that the latter two features correspond to MH excitations.

B. Peak assignment

The observed absorption bands can be assigned to MH and CT excitations. The role played by local crystal-field (i.e., valence-conserving $3d^{5} \rightarrow 3d^{5,*}$) excitations can be neglected in our analysis of the ellipsometry data. In LaSrFeO$_4$ they are forbidden both by parity and by the spin selection rule. Accordingly, they show a very small spectral weight with typical values of $\sigma_1 < 10 \, \text{(\Omega cm)}^{-1}$. Their signatures are visible in the transmittance data below the charge gap, see lower panel of Fig. 2.

A first tool to distinguish CT and MH excitations is the spectral weight. In transition-metal oxides, typical values of $\sigma_1(\omega)$ are of a few $1000 \, \text{(\Omega cm)}^{-1}$ for CT excitations but only a few $100 \, \text{(\Omega cm)}^{-1}$ for MH excitations. The difference is due to the fact that the matrix elements are of first order in the Fe - O hopping amplitude for CT excitations and of second order for MH excitations. To disentangle CT excitations and MH excitations, we further make use of the observed anisotropy. In layered LaSrFeO$_4$, MH excitations do not contribute to $\sigma_1^a(\omega)$ since the interlayer Fe - Fe hopping is strongly suppressed.
The strong absorption band observed at 4 - 5 eV in $\sigma^0(\omega)$ clearly has to be attributed to CT excitations. Our analysis of the ellipsometric data uses three Gaussian oscillators to describe the line shape, see Fig. [3]. This does not imply the existence of three microscopically different excitations since the line shape of the absorption band is not necessarily Gaussian but reflects bandstructure effects. The analysis of the $a$-axis data requires three very similar oscillators, in particular with a similar spectral weight. The peak energies differ by up to 0.4 eV for the two crystallographic directions, which most probably reflects the different on-site energies of apical and in-plane O ions. We attribute this band at 4 - 5 eV to the two strongest dipole-allowed CT excitations $t_{2u}(\pi) \rightarrow t_{2g}(\pi)$ and $t_{1u}(\pi) \rightarrow t_{2g}(\pi)$ (in cubic approximation, see discussion in Sec. [IV]). The splitting between these two excitations is expected to be about 1 eV according to quantum-chemistry calculations for LaFeO$_3$[13]. In LaSrFeO$_4$, the next higher-lying peak is observed at about 7 eV in the in-plane data of Refs. [31]. This large energy difference to the peak at 5 eV supports our interpretation that both $t_{2u}(\pi) \rightarrow t_{2g}(\pi)$ and $t_{1u}(\pi) \rightarrow t_{2g}(\pi)$ contribute to the absorption band between 4 and 5 eV. Note that both excitations correspond to a transfer to a 3d $t_{2g}(\pi)$ state, and that the crystal-field splitting of the $t_{2g}(\pi)$ level is expected to be only small, about 0.2 eV (see Sec. [III]). Moreover, the matrix elements for transitions into the $t_{2g}$ manifold do not differ very strongly between $a$ and $c$, even for an elongated octahedron, in contrast to the matrix elements for transitions into the $x^2-y^2$ orbital. The similar spectral weights along $a$ and $c$ between 4 eV and 5 eV therefore support our assignment.

The $a$-axis data show two additional features at 3.0 eV and 3.8 eV, see Figs. [5] and [6]. For the feature at 3.0 eV, both its lower spectral weight with $\sigma^0(3 \text{ eV})$ peaking at about 250 ($\Omega$cm)$^{-1}$ and the observed anisotropy support an interpretation in terms of a MH excitation. Moreover, the splitting between the two lowest MH excitations is expected to be roughly $\Delta_{eg} \approx 1$ eV (see Fig. [3]), in very good agreement with the difference of 0.8 eV between the observed peak energies of 3.0 and 3.8 eV. The third MH excitation is expected roughly 2$\Delta_{eg}$ above the lowest one, i.e., within the strong CT band. As far as the relative spectral weight is concerned, we roughly expect a factor of 3.8 between the two lowest MH excitations, see Sec. [IV]. Experimentally, the spectral weight of MH1 and MH2 is rather similar. However, our simple estimate does not take into account hybridization effects and is based on a local approach.

At first sight, it is unexpected that the lowest absorption band is of MH type because the 3d$^5$ configuration is stabilized by the intra-atomic Hund exchange $J_H$. In cubic approximation, a first rough estimate of the MH excitation energy yields $U + 4J_H$ with $4J_H \approx 3$ eV. This is much larger than in the 3d$^4$ manganites, for which we expect $U - J_H$. However, MH1 in the layered structure of LaSrFeO$_4$ corresponds to a transfer from $x^2-y^2$ to $3z^2-r^2$, thus it is $\Delta_{eg} \approx 1$ lower in energy than in cubic approximation. In comparison, the lowest MH excitation in LaSrMnO$_4$ requires the opposite transfer from $3z^2-r^2$ to $x^2-y^2$, raising the excitation energy to $E_{\text{Mn}} = U - J_H + \Delta_{\text{Mn}}^0$. In LaSrMnO$_4$, this MH excitation is observed at $E_{\text{Mn}} \approx 2$ eV[13]. We thus expect MH1 in LaSrFeO$_4$ at about $E(\text{MH1}) \approx E_{\text{Mn}} + 5J_H - \Delta_{eg} - \Delta_{\text{Mn}}$, i.e., roughly at 3 - 4 eV. Here, we neglect the slight increase of $U$ from Mn to Fe, but we also neglect that the 3d$^4$ and 3d$^5$ states relevant for the MH excitation both are Jahn-Teller active (cf. Fig. [3]), which reduces $E(\text{MH1})$. Moreover, these estimates neglect the effect of hybridization depicted schematically in Fig. [1]. Therefore, the assignment of the peak at 3.0 eV to MH1 appears feasible.

However, we also have to discuss alternative scenarios. As discussed in Sec. [IV] the lowest CT excitation $t_{1g}(\pi) \rightarrow t_{2g}(\pi)$ is dipole-forbidden and expected at about 0.8 eV below the lowest dipole-allowed CT excitation. Firstly, the spectral weight of the peak at 3 eV is too large for a dipole-forbidden excitation, and secondly, we expect only a modest anisotropy of this excitation. Possibly, this dipole-forbidden excitation may explain the small but finite values of $\sigma^0(\omega)$ between 2 eV and 3.5 eV. In a further scenario, the peak at 3.0 eV may be interpreted as a CT exciton. Note that this peak is lying at about 0.5 eV below the CT absorption edge and that a truly bound state with such a large binding energy is very unlikely. Again, it is not obvious why such an exciton should show a pronounced anisotropy. Moreover, an exciton with such a large binding energy is expected
to show a larger spectral weight and a smaller line width.

Summarizing this section, we have found strong evidence that the lowest dipole-allowed absorption band in LaSrFeO$_4$ is of MH type, which is made possible by the strong splitting $\Delta_{eg}$ caused by the layered structure and by the Fe 3d - O 2p hybridization. We want to add that in LaSrFeO$_4$ the energy of the lowest dipole-forbidden CT excitation may be comparable to the energy of MH1. Furthermore, the MH excitation from $x^2-y^2$ at site $i$ to a $t_{2g}$ orbital on a neighboring site is lower in energy than MH1, but the matrix element for this excitation vanishes (cf. Table I). We emphasize that our results do not disagree with the common interpretation that non-layered ferrites belong to the class of CT insulators. The different character can be explained by the absence of a large $\Delta_{eg}$ in the non-layered compounds. Pisarev et al.$^{18}$ studied the optical properties of a series of different ferrites with trivalent Fe ions. Many of these compounds show a shoulder in the vicinity of the onset of strong CT absorption, which has been attributed to the parity-forbidden excitation $t_{1g}(\pi) \rightarrow t_{2g}(\pi)$. The peak observed at 3.0 eV in LaSrFeO$_4$ is much too strong for such a dipole-forbidden excitation. However, our results suggest that MH excitations may not be neglected for a quantitative analysis of the non-layered ferrites, they may for instance provide a better explanation for a shoulder close to the absorption edge than the dipole-forbidden excitation discussed above.

C. Temperature dependence

As discussed in the introduction, the temperature dependence of the spectral weight of MH excitations has attracted considerable interest in different transition-metal compounds because it allows to study the temperature dependence of nearest-neighbor spin-spin and orbital-orbital correlations.$^{13,15,23,29}$ In LaSrFeO$_4$ we expect that the temperature dependence of these correlations is only small below 300 K. Actually, the spin-spin correlations change only gradually even above the ordering temperature due to the two-dimensional character.$^{15}$ This ferrite thus may serve as a reference compound to study the role of other effects such as the thermal expansion of the lattice or bandstructure effects.

As expected, the MH excitations at 3.0 eV and 3.8 eV show only a modest temperature dependence, see top panel of Fig. 5. According to the fit using Gaussian oscillators (cf. Fig. 6), the spectral weight of MH2 at 3.8 eV changes only by about 10% between 5 K and 250 K, see inset of Fig. 7. At the same time, the peak width increases by about 15%, and the peak frequency decreases by about 1%. For the peak MH1 at 3.0 eV, a quantitative analysis is more challenging. Both the spectral weight and the width of the Gaussian oscillator depicted in Fig. 6 increase strongly with temperature, while the frequency of the oscillator increases by about 1% from 5 K to 250 K. As mentioned above, there is not necessar-
VI. SUMMARY AND CONCLUSIONS

We present a detailed analysis of the optical conductivity of layered LaSrFeO$_4$ for temperatures ranging from 15 K to 250 K in a broad frequency range from 0.5 eV to 5.5 eV. Both the anisotropy and the different spectral weight allow us to disentangle Mott-Hubbard and charge-transfer excitations. We arrive at a consistent assignment of all absorption bands and find strong evidence that the lowest dipole-allowed excitation is of Mott-Hubbard type. Remarkably, this result is in agreement with previous studies of non-layered ferrites which have been identified as charge-transfer insulators.

The difference can be explained by the large splitting of the $e_g$ orbital in the layered structure, which partially compensates the large intra-atomic exchange contribution and pulls the lowest Mott-Hubbard excitation below the onset of charge-transfer excitations.

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