Electronic Structure of Ladder Cuprates

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We study the electronic structure of the ladder compounds Sr$_{14-x}$Ca$_x$Cu$_2$O$_4$ and SrCu$_2$O$_3$. LDA calculations for both give similar Cu 3d-bands near the Fermi energy. The hopping parameters estimated by fitting LDA energy bands show a strong anisotropy between the $t_x$ and $t_y$ intra-ladder hopping and small inter-ladder hopping. A downfolding method shows that this anisotropy arises from the ladder structure. The conductivity perpendicular to the ladders is computed assuming incoherent tunneling giving a value close to experiment.

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Sr$_{14-x}$Ca$_x$Cu$_2$O$_4$ is the first material in which doped ladder (see Fig. 1) can be experimentally studied and compared to theoretical predictions of a Luther-Emery state with a spin-gap, hole-pairing and superconductivity. A spin-gap of $\Delta \approx 280$K (for $x=0$) has been measured and superconductivity under high pressure $P>3$GPa has been found in Ca-rich samples $x \approx 11$(Ca). Ladders having ladder layers doped with either 20% holes ($\delta \approx 0.2$) or Sr have been realized. The transport properties are dominated by holes in the ladder planes. The normal state of the Ca11 shows a strong anisotropy between the $c$-resistivity and the $\perp$ across the ladder direction with $\rho_{\perp}/\rho \parallel \approx 30$ at $T=100K$. For lower temperature, both resistivities increase exponentially due to localization effects. For higher temperatures, $\rho_\perp$ increases linearly in $T$ while $\rho_{\parallel}$ remains nearly constant, $\rho_{\parallel} = 12m\Omega$cm. The mean free path along the ladder is larger than the lattice constant while across the ladders it is smaller than the inter-ladder distance indicating incoherent transport in this direction.

Moreover, fits of the spin susceptibility have shown a large difference between the exchange coupling $J_{\perp}$ ($J_{\parallel}$) along the rungs (legs) of the ladder even if both involve similar $180^\circ$ Cu-O-Cu superexchange processes. Analysis of neutron scattering data gives $J_{\perp}=72$meV and $J_{\parallel}=130$meV. For these reasons a detailed examination of the electronic structure is desirable.

In this paper we present LDA calculations of the electronic structure which give estimates of effective hopping matrix elements between states on different copper ions. The LDA studies are performed for SrCu$_2$O$_4$, a compound which possesses the same kind of Cu$_2$O$_3$ ladder planes as Sr$_{14-x}$Ca$_x$Cu$_2$O$_4$(see Fig. 1). Recently, Arai and Tsunetsugu reported LDA calculations for M$_{14}$Cu$_2$O$_{41}$, (M=Sr or Ca) which give similar results.

The TB-LMTO ASA energy bands for SrCu$_2$O$_3$ are plotted in Fig. 2. The uppermost graph displays the bands on the path $I=(0,0,0)$ to $(2\pi/a,0,0)$. The two parallel bands near zero energy (Fermi energy of the half-filled band) are separated from the rest of the spectrum. They are due to hybridization through $\sigma$-bonds of the 2$p$ O- and 3$d_{x^2-y^2}$ Cu-orbitals. These bands hybridize with non-bonding O-bands near $\Gamma$. In the lower graph, the energy bands are shown up to the edge of the zone with $Z=(0,0,0)$ and $A=(2\pi/a,0,0)$. Due to destructive interference, the 2 D-bands do not display dispersion on the path $ZA$.

The low energy physics can be described by an effective model containing only these two bands with similar shape near the Fermi energy. Such a model includes only one state per Cu with effective hopping matrix elements. These 2 bands are the bonding (b) and anti-bonding (a) rung bands of the effective ladder model.

Note that the parallel nature of the bands at $k_z = \pi/2c$ (path $ZA'$) cannot be explained by effective interactions between nearest-neighbor (n.n.) Cu-sites only. In such a model, the dispersion along the $k_x$ direction is given by the inter-chain hopping matrix element between the second leg of one ladder and the first leg of the next ladder. Since the hopping matrix elements between two bonding (antibonding) states of the rung $r$ of two neighboring ladders $l$ and $l'$ $b_{\sigma,l,r}(a_{\sigma,l,r}) = \sqrt{2}(\phi_{\sigma,l,r} + (-)\phi_{\sigma,l,r})$ is given by

$$\langle b_{\sigma,l,r} H(t_{ll'}b_{\sigma,l',r}) = \langle a_{\sigma,l,r} H(t_{ll'}a_{\sigma,l',r}) = \phi_{\sigma,l,r} H(t_{ll'}\phi_{\sigma,l',r}) \phi_{\sigma,l,r} \phi_{\sigma,l',r} \phi_{\sigma,l',r} \rangle,$$

where the last index of $\phi_{\sigma,l',r}$ labels the ladder leg. Therefore b- and a-states should have an opposite dispersion in the $k_x$-direction. Thus, an effective atomic model must contain some longer range inter-ladder hopping to account for their parallel nature.

Arai and Tsunetsugu introduced a simpler rung parameterization of the band structure fitting the b- and a-bands separately, allowing n.n. and n.n.n. hopping leading to the form:

$$\epsilon(\mathbf{k}) = \epsilon_0 - 2h_{l_{l1}}\cos(k_z) - 2h_{l_{l2}}^2\cos(2k_z) - [4h_{l_{l1}} \cos(\frac{\pi}{2}k_z) + 4h_{l_{l2}}^2 \cos(\frac{3}{2}k_z)] \cos(k_x).$$

The values they obtained for Sr$_{14}$Cu$_2$O$_{41}$ are in good qualitative agreement with ours for SrCu$_2$O$_3$ (see Table 1). Note that the signs of the interladder hopping
parameter $h_{\perp,1}$ does not change between an a-band and b-band contrary to the expectations from Eq. 2. All hopping parameters apart from $h_{\parallel,1}$ are higher order in the Cu-O ($t_{pd}$) and O-O overlaps $t_{pp}$ and thus much smaller.

To gain more insight we introduce a single parameterization of both bands in terms of intersite hopping parameters shown in Fig. 3. The solution of this tight-binding model is

$$
\epsilon_{\pm}(k) = \epsilon_0 + \epsilon_{\parallel}(k_z) + \epsilon_{\perp,1}(k_z) \cos(k_x) 
$$

$$
\pm \sqrt{\epsilon_{\perp,3}(k_z)^2 + \epsilon_{\perp,4}(k_z)^2 + 2\epsilon_{\perp,3}(k_z)\epsilon_{\perp,4}(k_z) \cos(k_x)}
$$

(3)

where

$$
\epsilon_{\parallel}(k_z) = -2t_{\parallel,1}(k_z) - 2t_{\parallel,2}\cos(2k_z),
$$

$$
\epsilon_{\perp,1}(k_z) = -4t_{\perp,1}\cos\left(\frac{k_z}{2}\right) + 4t_{\perp,2}\cos\left(\frac{3k_z}{2}\right),
$$

$$
\epsilon_{\perp,3}(k_z) = 2t_{\perp,1}\cos\left(\frac{k_z}{2}\right) + 2t_{\perp,2}\cos\left(\frac{3k_z}{2}\right),
$$

$$
\epsilon_{\perp,4}(k_z) = 2t_{\perp,1}\cos\left(\frac{k_z}{2}\right) + 2t_{\perp,2}\cos\left(\frac{3k_z}{2}\right),
$$

(4)

The parallel nature of the bands at $k_z = \pi/2$ is recovered if the term in the square root of Eq. 3 is independent of $k_z$. This obtains if $\epsilon_{\perp,4}(\pi/2) \simeq 0$, leading to $t_{\parallel,2}, t_{\perp,5} \simeq 0$ or to $t_{\parallel,2} = t_{\perp,5}$. In these cases the dispersion at $k_z = \pi/2$ is due to the $\epsilon_{\perp,1}$ term being a function of $t_{\perp,3}$ and $t_{\perp,7}$. It clearly shows that an intersite model must contain at least the third order hopping term $t_{\perp,3}$.

The coupled ladder system has a glide symmetry given by the product of a reflection through the c-axis (see Fig. 3) and a translation of half a lattice constant along the ladder. When this operation is applied twice it is equivalent to a translation of one lattice constant along the ladder. This implies that the energy band at $k_x = 0$ and $k_z = \pi$ can be parameterized through one parameter. Actually $\epsilon_{\pm}(k_z = \pi, k_z) = \epsilon_{\parallel}(k_x = 0, 2\pi - k_x)$ as can be directly checked from Eq. 3. This allows one to use a single function containing the information about all hopping parameters. This symmetry also implies the lack of dispersion in the $k_z$ direction at $k_z = \pi$ as discussed above, as well as the square root form containing intra- and inter-ladder hopping terms.

Introducing $\sigma = 1(-1)$ for $k_x = 0(\pi)$ Eq. 3 reduces to the simpler form

$$
\epsilon_{\pm,\sigma}(k_z) = \epsilon_0 + \epsilon_{\parallel}(k_z) + \sqrt{\epsilon_{\perp,3}(k_z)^2 + \epsilon_{\perp,4}(k_z)^2 + 2\epsilon_{\perp,3}(k_z)\epsilon_{\perp,4}(k_z) \cos(k_x)}
$$

(5)

representing the 4 energy bands of the double-ladder system. Rewriting the bands from the rung form (3) to the intersite form (5) gives the values of Table II. They are consistent with each other and emphasize the dominance of the n.n. intra-ladder matrix elements $t_{\parallel,1}$ and $t_{\perp,1}$ w.r.t. the others. Moreover, they show surprisingly that these two parameters describing n.n. Cu-Cu processes differ from each other by $\sim 35\%$.

Recently, Andersen et al. introduced a systematic downfolding scheme to obtain an effective single (or few) band model capable of reproducing the details of the LDA bands close to the Fermi level. Effective hopping parameters are calculated by performing the Fourier transform of the downfolded Hamiltonian $H(k) \rightarrow H(R)$, for $|R|$ less than a cut-off radius $R_0$. This has the advantage among others that it allows the origin of the parameters in the effective single or few band model to be traced. We have applied this scheme to the LDA bands for SrCu$_2$O$_3$. The anisotropy $t_{\parallel,1} \neq t_{\perp,1}$ is best understood by starting with an effective model including the 3$d_{x^2-z^2}$, $4$s Cu- and $2p_x, 2p_y$ O-orbitals (dsp-model). The dominant parameters ($> 0.1$ eV) for in-plane hopping are given in the first rows of Table II. Here $r$ labels the O on the rung of one ladder while $l$ labels either a Cu or an O on the upper leg of the ladder. Vectors $\mathbf{e}_x = (a/6, 0, 0)$ and $\mathbf{e}_y = (0, 0, c/2)$ gives the translation vector from one O(Cu) to the neighboring Cu(O) in the respective direction. The notation is such that $t^R_{l(1)p_{z}(\pm e_x)}$ denotes the hopping between a 3$d$ Cu-orbital at I on the upper leg to the neighboring $2p_{z}$ O-orbital at $I \pm \mathbf{e}_x$. The on-site $t^R_{s(1)p_{d}(l)}$ hopping is non-zero as consequence of the downfolding of all other bands in the absence of local four-fold symmetry.

The on-site energies of the rung-oxygen $\epsilon(R)$ is slightly larger than that of the leg-oxygen $\epsilon(I)$ due to the local environments. The distance between Cu and O is $r_l = 1.98\AA \parallel \hat{z}$ and $r_r = 1.92\AA \parallel \hat{\mathbf{r}}$ implying a larger hopping along the $\mathbf{r}$-direction according to $t_{pd} \propto 1/r^4$ and explaining the anisotropy for the $t_{pd}$'s listed in first row of Table II. Moreover hopping processes involving s-orbitals are large with non-negligible $t_{sd}$ hoppings. They will contribute to the $t_{pd}$ hopping in second and higher orders through paths like $d-s-p, d-s-s-p$, etc. By downfolding the s-orbitals, these processes will strongly renormalize the $t_{pd}$ such that $t^R_{l(1)p_{z}(+e_x)} > t^R_{l(1)p_{z}(-e_x)}$. Lastly downfolding the p orbitals gives the effective (d)-model. Results are given in third column of Table II. They are consistent with our previous results. The anisotropy between $t_{\perp,1}$ and $t_{\parallel,1}$ can now be seen to be a consequence of different $t_{pd}$ hopping due to the downfolding of the s bands.

The exchange interaction $J$ between spins are usually difficult to estimate a priori. In perturbation theory, $J$ scales as $t_{pd}^4$ but given the large value of $t_{pd}$ relative to $(\epsilon_p - \epsilon_d)$ the results are not reliable. Our analysis shows that the energy difference $(\epsilon_p - \epsilon_d)$ does not differ much between rung and legs but $t_{pd}$ does. A ratio $J_{pd}/U_{pd} < 1$ is expected consistently with previous results. This will disfavor the hole coupling and the RVB liquid state and may explain why holes are unbound for $T > 100$K, however this point will not be discussed further.

In the following, two limiting estimates of the conductivity are investigated. First, the band structure model limit is considered ignoring magnetic interactions although they must be important. In a metallic ground state, the conductivity reduces to an integral over the Fermi surface.
\[ \sigma_{ij} = \frac{2e^2}{(2\pi)^2} \int d\mathbf{k} \tau(\epsilon(\mathbf{k})) v_i(\mathbf{k}) v_j(\mathbf{k}) \left( -\frac{\partial f}{\partial \epsilon} \right). \] (6)

Considering an electron filling of \( n = 0.8 \) the ratio between the conductivity perpendicular and parallel to the ladder can then be simply computed yielding very large values i.e. \( \sigma_\parallel / \sigma_\perp \simeq 75, 90, \) and 104, for the parameter in Table II. Thus roughly one has

\[ \sigma_\parallel / \sigma_\perp \simeq 100. \] (7)

This large anisotropy is a consequence of a warped Fermi-surface with very small Fermi velocity perpendicular to the ladders.

Second, as discussed previously, the resistivity data perpendicular to the ladder indicates a mean-free path smaller than the inter-ladder spacing. Holes seem to be confined to ladders and to hop incoherently between them. A detail structural refinement of Ca11Cu22O41 shows a complex distortion pattern of the CuO2-chains leading to substantial potential variations between rungs due to the proximity of apical O-ions at \( \sim 50\% \) of the rungs. An estimate of the interladder conductivity can be made by considering the limit where ladders form quasi one-dimensional metallic systems weakly coupled to each other. The conductivity is thus a consequence of the inter-ladder hopping term

\[ H = \sum_k T_k c_\uparrow^{\dagger}_k c_{2,k} + \text{h.c.}, \] (8)

where \( T_k = \sum_i h_{\perp,i}(k) \). The conductance \( \sigma_\perp \) is given by

\[ \sigma_\perp = 4\pi e^2 \hbar N(\epsilon_F)^2 |T(k_F)|^2, \] (9)

with \( N(\epsilon_F) \) denoting the density of state at \( E_F \) in the quasi-one dimensional metallic system. Results for the different parameter sets are \( \sigma_\perp \simeq 0.1, 0.09, \) and 0.074, respectively. They are thus all close the value of

\[ \sigma_\perp \simeq 0.08 \text{ m}\Omega^{-1}\text{cm}^{-1}, \] (10)

which corresponds well with the experimental result at \( T \geq 100\text{K} \).

In this paper, estimates of the hopping matrix elements based on LDA calculations gave three main results. First, the effective intra-ladder hopping between n.n. are not the same, \( t_\parallel \neq t_\perp \). This was explained as the consequence of anisotropic \( t_{pd} \) in the \((pd)\)-model due to effective hopping through paths involving Cu s-states. Second, inter-ladder hopping is much smaller than intra-ladder and longer range hopping must be included. Third, estimates of the conductivity in the model where holes are unbound and confined into the ladder give good agreement with the experiment at temperatures \( T \geq 100\text{K} \).

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FIG. 1. The Cu$_2$O$_3$ ladder plane.

FIG. 2. LDA band calculations on the path $\Gamma Z' A' X' \Gamma$ (uppermost graph) and the path $\Gamma Z A X \Gamma$ (lower graph).

FIG. 3. Coupled ladders illustrating effective hoppings between Cu sites (t) or Cu-Cu rungs (h).

TABLE I. The hopping parameters in eV for Sr$_{14}$Cu$_{24}$O$_{41}$ and SrCu$_2$O$_3$ in the rung scheme.

|                  | Sr$_{14}$Cu$_{24}$O$_{41}$ | SrCu$_2$O$_3$ |
|------------------|-----------------------------|---------------|
| $\epsilon_0$    | -0.31                       | 0.46          |
| $h_{||,1}$       | 0.41                        | 0.50          |
| $h_{||,2}$       | 0.08                        | 0.07          |
| $h_{\perp,1}$    | 0.07                        | 0.03          |
| $h_{\perp,2}$    | 0.00                        | -0.04         |

TABLE II. The parameters in eV from direct fitting and downfolding of the LDA bands.

|                  | Sr$_{14}$Cu$_{24}$O$_{41}$ | SrCu$_2$O$_3$ |
|------------------|-----------------------------|---------------|
| $\epsilon_0$    | 0.075                       | -0.045        |
| $t_{||,1}$       | 0.500                       | 0.565         |
| $t_{\perp,1}$    | 0.385                       | 0.395         |
| $t_{\perp,2}$    | 0.040                       | 0.040         |
| $t_{\perp,3}$    | 0.050                       | 0.050         |
| $t_{\perp,4}$    | -0.090                      | -0.115        |
| $t_{\perp,5}$    | 0.040                       | 0.040         |
| $t_{\perp,6}$    | 0.075                       | 0.075         |
| $t_{\perp,7}$    | 0.005                       | 0.005         |
| $t_{\perp,8}$    | -0.020                      | -0.020        |
|       | $\epsilon_d$ | $\epsilon_{p(r)}$ | $\epsilon_{p(l)}$ | $t_{d(l)p_z(+e_z)}$ | $t_{d(l)p_z(-e_z)}$ | $t_{d(l)p_z(+e_z+e_z)}$ | $t_{p_z(l)p_z(+e_z+e_z)}$ | $t_{p_z(l)p_z(-e_z+e_z)}$ | $t_{p_z(l)p_z(-e_z)}$ | $t_{p_z(l)p_z(+2e_z)}$ | $t_{p_z(l)p_z(+e_z)}$ | $t_{p_z(l)p_z(+2e_z)}$ | $\epsilon_s$ | $t_{s(l)d(l)}$ | $t_{s(l)d(+2e_z)}$ | $t_{s(l)s(+2e_z)}$ | $t_{s(l)s(-2e_z)}$ | $t_{s(l)p_z(+e_z)}$ | $t_{s(l)p_z(+e_z)}$ | $t_{s(l)p_z(+e_z)}$ |
|-------|--------------|-------------------|-------------------|---------------------|---------------------|-------------------------|---------------------------|-------------------------|---------------------|---------------------------|---------------------|---------------------------|-------------|-----------------|------------------|------------------|------------------|-----------------|------------------|
| (dsp) | -3.82        | -3.66             | -4.00             | 0.74                | 0.85                | -0.85                   | 0.30                      |                         |                    |                            |                     |                           |             |                 |                  |                  |                  |                 |                  |
| (dp)  | -3.83        | -4.59             | -4.53             | 0.65                | 0.51                | -0.63                   | 0.22                      |                         |                    |                            |                     |                           |             |                 |                  |                  |                  |                 |                  |
| (dsp) | 0.30         | 0.01              | -0.13             | 0.28                | -0.42               |                         |                           |                         |                    |                            |                     |                           |             |                 |                  |                  |                  |                 |                  |
| (dp)  | -0.17        | -0.62             | 2.43              | -0.24               | -0.14               |                         |                           |                         |                    |                            |                     |                           |             |                 |                  |                  |                  |                 |                  |
| (dsp) | -0.32        | 0.20              | -0.44             | 1.89                | $\pm2.10$           |                         |                           |                         |                    |                            |                     |                           |             |                 |                  |                  |                  |                 |                  |

**TABLE III.** Hopping parameters in eV from the downfolding method. Notation is explained in the text.