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Localisation of doped holes in edge-shared CuO₂ chain cuprates: consequences for dynamic spectral weight transfer


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We present a joint experimental and theoretical study of the electronic structure of the cuprate chain systems A₁₋ₓCuO₂ (A=Ca,Sr,Ba), as measured using O-K and Cu-L₃₂ spectra. The doping-dependent behaviour in these systems is radically different to that in conventional 2D cuprate networks formed from corner sharing CuO₄ plaquettes, and follows from the strongly suppressed inter-plaquette hybridisation resulting from the 90° Cu-O-Cu interaction pathway in the chain systems. Spectroscopically, this results in (a) a classical mixed-valent scenario whereby the different final states in the Cu-L₃ spectra can be used directly to 'read-off' the Cu valency and (b) a drastic reduction in the dynamic spectral weight transfer from the upper Hubbard band to the low energy scale in the O-K spectra. The final picture emerges of localisation of the doped holes, with the chain then comprised of a mixture of pure Cu(II)O₄ and Cu(III)O₄ plaquettes.

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Low dimensional cuprate systems, such as chains and ladders, gained interest originally in the context of their being model systems for the high temperature cuprate superconductors. It soon became clear that such quasi-1D systems offer a rich correlated electron physics of their own with the experimental observation of the separation of the spin and charge degrees of freedom being one of the highlights so far.

A second area to catch the imagination of researchers worldwide has been that of the spin ladders. Here, superconductivity has been predicted theoretically for a doped ladder but has not been observed so far in pure ladder compounds.

Consequently, the discovery of superconductivity under pressure in Sr₀.₄Ca₁₃.₆Cu₂₋ₓO₄₁₊₇ with no CuO₂ planes, but rather a mixture of linear CuO₂ chains (built up of edge-shared CuO₄ plaquettes) and two-leg Cu₂O₃ ladders has sparked still further interest in the properties of the ladders and the chains as individual elements.

In this paper, we present a joint experimental (O-K and Cu-L₃ XAS) and theoretical investigation of the electronic structure of hole-doped edge-sharing CuO₂ chains. The compounds under investigation are Ca₀.₈₃CuO₂, Sr₀.₇₃CuO₂ and Ba₀.₆₇CuO₂, all of which contain CuO₂ chains formed of edge-sharing CuO₄ plaquettes. The chains are doped with either 0.33, 0.52 or 0.67 holes per Cu site, respectively. Remarkably, the magnetic susceptibility of Ca₀.₈₃CuO₂ and Sr₀.₇₃CuO₂ - despite their high hole-doping levels - can be described within the dimerized alternating spin-1/2 Heisenberg-chain model with antiferromagnetic order setting in below 10 K and 12 K for Ca₀.₈₃CuO₂ and Sr₀.₇₃CuO₂, respectively. For the better known CuO₂ plane, neither Néel nor any other long-range magnetic order has been found for doping levels beyond 0.04 to 0.125 holes per Cu site (i.e. beyond the spin-glass and stripe region). Furthermore, there have been reports of superconductivity with Tₓ =13 K in the related compound Ba₂Cu₃₋ₓO₇₋ₓ which naturally would necessitate the delocalisation of doped holes in the A₁₋ₓCuO₂ systems.

The synthesis and the structural analysis of the polycrystalline samples of Ca₀.₈₃CuO₂, Sr₀.₇₃CuO₂ and Ba₀.₆₇CuO₂ have been described previously. The XAS measurements were performed at the SX700/II monochromator operated by the Freie Universität Berlin and the P25 beamline at the Berliner Elektronenspeicherring für Synchrotronstrahlung (BESSY). In all measurements the non-surface-sensitive fluorescence-yield mode was used. The energy resolution of the monochromator was set to 280 meV and 600 meV at the O 1s and the Cu 2p₃/₂ thresholds, respectively. The O 1s data were corrected for the energy-dependent incident flux and normalized 60 eV above the threshold. The self-absorption effects were taken into account as described in Refs. [12] and [13]. Prior to the measurements, the surface of the sintered tablets was scraped in-situ with a diamond file at a base pressure of 5x10⁻¹⁰ mbar.

The simulated Cu-L₃ and O-K spectra were calculated with the aid of an extended five-band pd Hubbard-model as described in Ref. [14]. The Cu-L₃ and O-K 1s-XAS spectral density was calculated by means of the exact...
diagonalisation of small periodic \((\text{CuO}_2)_n\) clusters (with \(n=3,4\)), following Refs. \[14\] and \[15\]. The calculated data were broadened with Lorentzian functions of width 0.4 eV, in order to ease their comparison with experiment.

Fig. 1 illustrates the systematic evolution of the Cu-L\(_3\) XAS spectra of Ca\(_{0.83}\)CuO\(_2\), Sr\(_{0.73}\)CuO\(_2\) and Ba\(_{0.63}\)CuO\(_2\). The left panel shows the experimental spectra, whereas the right panel contains the results of the extended five-band \(pd\) Hubbard model referred to above.\[19\] Also shown are data from Li\(_2\)CuO\(_2\) and NaCuO\(_2\), which serve as formally divalent and trivalent reference systems for the edge-shared CuO\(_2\) chain geometry. The spectral signatures of Li\(_2\)CuO\(_2\) and NaCuO\(_2\) are simple, single component white-lines at 930.7 and 932.4 eV, respectively.

Fig. 1 shows clearly that the edge-shared chain systems with non-integer formal valence exhibit a Cu-L\(_3\) XAS spectrum resembling a superposition of scaled versions of the divalent and trivalent reference systems. Based upon the excellent agreement observed between experiment and theory in Fig. 1, we can assign the features as follows. For Li\(_2\)CuO\(_2\) the single peak is described by the 2\(p^3d^{10}\) configuration (where 2\(p\) denotes the core hole). The dominant final state configuration representing the main peak at 932.4 eV in NaCuO\(_2\) is described by 2\(p^3d^{10}\)\(L\). Consequently, for Ca\(_{0.83}\)CuO\(_2\), Sr\(_{0.73}\)CuO\(_2\), and Ba\(_{0.63}\)CuO\(_2\) the low and higher energy features result from 2\(p^3d^{10}\) and 2\(p^3d^{10}\)\(L\) final states, respectively. Furthermore, the clear energetic alignment with the divalent and trivalent reference systems confirms that the features in the Cu-L\(_3\) XAS of the intermediate doped chains signal the existence of Cu(II)O\(_4\) and Cu(III)O\(_4\) plaquettes. In line with this picture, the spectral weight of the higher energy peak increases in line with the formal valence on going from Ca\(_{0.83}\)CuO\(_2\) to Sr\(_{0.73}\)CuO\(_2\) and further to Ba\(_{0.63}\)CuO\(_2\).

To put these results in context, we show in Fig. 2 a comparison of the Cu-L\(_3\) XAS spectra of two formally trivalent systems - NaCuO\(_2\) and LaCuO\(_3\) (the latter from Ref. \[17\]) - and two formally Cu 2.33+ valent systems - Ca\(_{0.83}\)CuO\(_2\) and La\(_{1.66}\)Sr\(_{0.34}\)CuO\(_4\) (the latter from Ref. \[18\]). The enormous differences between the spectra of compounds with the same formal valency is a signal of the crucial role played by the geometry of the Cu-O network, not only in the Cu 2\(p\) core level photoemission spectra\[14,15\] but also in Cu-L\(_3\) XAS\[16\]. This point is worth further explanation.

In the edge-sharing CuO\(_2\) chain systems the Cu-O-Cu interaction pathway is essentially 90°, which strongly suppresses the inter-plaquette hybridisation. For the undoped system Li\(_2\)CuO\(_2\), this has been observed to lead to a behaviour analogous to that of an isolated CuO\(_4\) plaquette.\[19\] This extreme localisation means that the spectroscopy of such cuprates is reminiscent of the situation in some mixed valent 4f compounds in which the intensity ratio of the features at the (Ln)-L\(_3\) thresholds can be used to determine the average valence.\[20\] In the case of the CuO\(_2\) chain systems, due to the geometrically mediated switching-off of the inter-plaquette hybridisation, one observes a clear feature at 932.4 in the Cu-L\(_3\) spectra whose weight scales with the formal copper valence.

In contrast, the systems containing strong 180° Cu-O-Cu interaction pathways (LaCuO\(_3\) and La\(_{1.66}\)Sr\(_{0.34}\)CuO\(_4\)), support significant inter-plaquette hopping, thus resulting in an increase in the effective covalency. This has two effects. Firstly, it reduces the contribution of the bare 3\(p^3d^{10}\) configuration in the ground state\[24\]. Secondly, the inter-plaquette hybridisation offers the system further opportunity to screen the copper core hole involving the transfer of electron density from ligand levels further away from the core ionised site. As a result of this, the Cu-L\(_3\) XAS spectrum of trivalent LaCuO\(_3\) exhibits a feature at the same energy as is usually associated with divalent copper, but which actually stems from a trivalent final state with additional (non-local) screening.\[22\] The same behaviour is seen for the formally 2.33+ valent cuprates: the edge shared chain exhibits two clear final state peaks, whereas the interplaque hopping prevalent in the 2D CuO\(_2\)-plane system results in a mere asymmetry at higher energy.\[24\]

The spectroscopic consequences of the geometry-induced localisation are not restricted to the Cu-L\(_3\) spectra, as is illustrated by the O-K spectra shown in Fig. 3. The pronounced peak directly above the absorption onset at 530.2 eV in the undoped Li\(_2\)CuO\(_2\) system is related to transitions into O 2\(s^\pi\) upper Hubbard band (UHB).\[1] The pre-edge peak in the O 1s XAS spectrum of the trivalent cuprate NaCuO\(_2\), which in the context of the HTSC would be assigned to the doped hole states, is down-shifted by 1.3 eV with respect to the Cu(II) UHB, since with increasing Cu valence the covalence increases.\[15,16,24\]

In conventional, corner-shared 2D-planar systems which have been hole-doped, two pre-edge peaks are observed on O-K XAS. With increasing doping level the UHB quickly loses its intensity, whereas the hole peak gains in intensity and shifts to lower energy as the doping progresses.\[23\] As can be seen from Fig. 3, the intermediate-doped edge-shared cuprate chains present a different picture. Firstly, here, as was the case for the Cu-L\(_3\) spectra, the two features remain at the same energies as their counterparts in the formally Cu(II) and Cu(III) reference systems. Secondly, the rate at which the UHB spectral weight is transferred to the low energy scale is drastically reduced in these compounds compared to their corner-sharing 2D cousins. This is much more than an XAS detail, as this spectral weight transfer is an important characteristic of doped effective Mott-Hubbard insulators and has been extensively studied in the corner-shared, layered cuprates both experimentally\[24,30,31\] and theoretically.\[32\] In the corner-shared systems such as La\(_{2-x}\)Sr\(_x\)CuO\(_4\)\[24,30,31\] and the two-leg ladder system La\(_{1-x}\)Sr\(_x\)CuO\(_{2.5}\)\[34\], this effect - expressed in the doping-mediated
destruction of the UHB intensity - has been found to be supralinear with $x$, and has thus been dubbed dynamical spectral weight transfer or DSWT \cite{Mizokawa}. For example, in the case of La$_{0.75}$Sr$_{0.25}$CuO$_4$ for $x=0.15$, the UHB is scarcely visible and for the analogous doping level in the two-leg ladder, the UHB peak has already lost more than 40% of the spectral weight it had for $x=0$ \cite{Boeske}.

However, as can be clearly seen in Fig. 3 for an even greater doping level of 0.34 in Ca$_{0.83}$CuO$_2$ we still find $\sim 67\%$ spectral intensity of the UHB compared with Li$_2$CuO$_2$, the latter acting as a reference point for the undoped case. Even Ba$_{0.63}$Cu$_{0.37}$O$_2$, which has a formal doping level of 0.67 holes per Cu, still shows a clear indication of an UHB feature in the O-K XAS. The suppression of the DSWT is a direct result of the extremely small inter-plaque hopping along the Cu$_2$O$_2$ chains, and is well reproduced by the extended five-band $pd$ Hubbard model,\cite{Kim} as can be seen from the good agreement between experiment (left panel) and theory (right panel) in Fig. 3.

In addition, this knowledge has played an important role in our understanding of recent polarisation-dependent XAS data from the (La,Y,Sr)$_{14}$ systems, which contain, apart from the 2-leg ladders, highly doped CuO$_2$ chains.

Fig. 4 summarises the Cu-L$_3$ and O-K core level excitation data of the edge-shared chain systems. In the upper panel, the mean Cu valencies of Li$_2$CuO$_2$ and the three doped chain compounds under investigation, derived from the intensity of the Cu(II) and Cu(III) signals in the Cu-L$_3$ XAS are compared to the formal valence derived from the stoichiometry, yielding an almost perfect agreement. This emphasises the unique nature of these cuprate chains, as, to the best of our knowledge, they represent the first examples of a mixed valent cuprate family, in which core level spectroscopy allows one to ‘read off’ the mean Cu valence merely by comparing the intensity ratio of the double-peaked Cu-L$_3$ XAS feature.

The lower panel of Fig. 4 shows an analysis of the doping dependent evolution of the spectral weight of the hole peak and the UHB in the O-K XAS spectra, whereby the doping dependence of the ZRS and UHB features in LS (from Ref. \cite{Shengelaya}) has been included for comparison. The strong suppression of the DSWT in the edge-shared chain systems is very clear, clearly indicating the supression of the dynamic spectral weight transfer in these systems.

To summarize, we have presented a joint experimental and theoretical study of the effects of hole doping on the electronic structure of CuO$_2$ chains built up of edge-shared CuO$_4$ plaquettes. Analysis of the data, both at the qualitative and quantitative level, illustrates the remarkable consequences of the edge-sharing geometry of these systems, which results in a 90° Cu-O-Cu interaction pathway, thereby essentially switching-off the inter-plaque hopping.

(1) The system is robbed of the possibility of using non-local processes to screen the core hole in Cu-L$_3$ XAS, thus resulting in a classical mixed valence behaviour in which the average Cu valence can be simply extracted from the relative intensity of the 2p$3d^{10}$ and 2p$3d^{10}L$ final state features. This is an unprecedently clear example of the impact of non-local effects in XAS.

(2) The dynamic transfer of spectral weight from the upper Hubbard band to the low energy scale is strongly suppressed, resulting in the observation of the UHB feature in the O-K spectra even up to for a doping level of 0.67 extra holes per Cu.

(3) Together with the observation of Neel order in the Sr$_{0.73}$CuO$_2$ and Ca$_{0.83}$CuO$_2$ systems,\cite{Kim} the spectroscopic data presented here provide clear evidence for the existence of individually identifiable Cu(II)O$_4$ and Cu(III)O$_4$ plaquettes in these materials.

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16. The following parameters were adopted. Undoped compounds: $U_d=8.8$ eV, $U_p=5$ eV, $t_{pd}=1.4$ eV, $V_{pd}=1$ eV, $V_{pp}=0.5$ eV, $U_{pp'}=0.8U_p$ ($pp'$ denotes the different O 2p-orbitals at the same oxygen site), and $U_{cd}=10$ eV. In addition $t_{pp}^{\perp} = 0.5 t_{pp,\perp}$ has been assumed, where $t_{pp}^{\perp} = (1/\sqrt{2})[(pp\sigma) - (pp\pi)]$. To correctly reproduce the spectra with increasing doping level, alteration of $\Delta_{pd}$ and $t_{pp}$ was necessary, which is ascribed to changes of the crystal potential and to a contraction of the oxygen Wannier functions, which are most extended in the undoped case. $t_{pp}^{\perp} \leq 0.3$ eV has been adopted for all CuO$_2$ chains (cf. a typical value of $\approx 0.65$ eV for layered cuprates).
The vast majority of doped CuO₂-plane systems studied with XAS exhibit similar Cu-L₃ spectra to that of the La-214 system [28], with the exception of the Hg-cuprates, in which the high energy feature is visible as a clear shoulder or sub-peak on the main white line. However, as the formal doping level in these systems is difficult to define, we use the La-214 system as a more clear-cut reference.

FIG. 1. Left panel: Cu-L₃ XAS spectra of edge-shared CuO₂ chain systems as a function of hole doping. From the bottom: Li₂CuO₂, Ca₀.₈₃CuO₂, Sr₀.₇₃CuO₂, Ba₀.₆₃CuO₂ and NaCuO₂. The right panel shows the same spectra simulated using an extended five band pd Hubbard model. The hole doping level appropriate for each system is indicated next to each theoretical curve. For details see text.

FIG. 2. Cu-L₃ XAS spectra of (from the top) the formally trivalent systems NaCuO₂ and LaCuO₃ (from Ref. [17]), together with the formally Cu 2.33+ valent systems Ca₀.₈₃CuO₂ and La₁₋ₓSrₓCuO₄ (the latter from Ref. [31]).

FIG. 3. Left panel: O-K XAS spectra of edge-shared CuO₂ chain systems as a function of hole doping. From the bottom: Li₂CuO₂, Ca₀.₈₃CuO₂, Sr₀.₇₃CuO₂, Ba₀.₆₃CuO₂ and NaCuO₂. The right panel shows the same spectra simulated using an extended five band pd Hubbard model. The hole doping level appropriate for each system is indicated next to each theoretical curve. For details see text.

FIG. 4. (a) The mean Cu valence (filled triangles) obtained from the Cu-L₃ XAS spectra of Li₂CuO₂, Ca₀.₈₃CuO₂, Sr₀.₇₃CuO₂, Ba₀.₆₃CuO₂ together with the formal valence (filled circles) from the stoichiometry plotted versus the hole doping level, x. (b) the relative spectral intensity of the doping-induced hole peak (ZRS, open squares) and the UHB (filled squares) in the O-K XAS spectra of the same systems, normalized to the intensity of the UHB for undoped Li₂CuO₂. The decay of the UHB intensity and the growth of the ZRS feature for the system La₁₋ₓSrₓCuO₄ (from Ref. [31]) is shown as filled and open grey circles for comparison.
Figure 1: Fluorescence yield as a function of photon energy for different doping levels of NaCuO$_2$, Ba$_{1-x}$CuO$_2$, Sr$_{1-x}$CuO$_2$, Ca$_{1-x}$CuO$_2$, and Li$_2$CuO$_2$. The doping levels are indicated by the solid lines: 1.0, 0.67, 0.5, 0.33, and 0.0. The data is compared to experimental results (open circles).
Fig. 2 Hu et al.
Fig. 3 Hu et al.
Fig. 4 Hu et al.