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Perspective on the phase diagram of cuprate high-temperature superconductors

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Universal scaling laws can guide the understanding of new phenomena, and for cuprate high-temperature superconductivity the influential Uemura relation showed, early on, that the maximum critical temperature of superconductivity correlates with the density of the superfluid measured at low temperatures. Here we show that the charge content of the bonding orbitals of copper and oxygen in the ubiquitous CuO₂ plane, measured with nuclear magnetic resonance, reproduces this scaling. The charge transfer of the nominal copper hole to planar oxygen sets the maximum critical temperature. A three-dimensional phase diagram in terms of the charge content at copper as well as oxygen is introduced, which has the different cuprate families sorted with respect to their maximum critical temperature. We suggest that the critical temperature could be raised substantially if one were able to synthesize materials that lead to an increased planar oxygen hole content at the expense of that of planar copper.

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he understanding of the complex properties of the cuprates, and what causes their high critical temperature of superconductivity (T_c) , is one of the greatest challenges in condensed matter physics. From it one expects clues that make the synthesis of cuprates with much higher T_c possible or of how to improve other fundamental properties. In particular, it is still not well understood what sets the very different, maximum T_c 's for different families of materials. An early experimental observation in this regard is the famous Uemura plot¹. It shows that the maximum T_c is correlated with the muon spin relaxation rate σ_0 (extrapolated to T=0 K) that is proportional to the superfluid density divided by the effective mass ($\sigma_0 \propto n_s/m^*$). This relation holds for the underdoped materials and orders different cuprate families with respect to their maximum T_{c} . The Uemura relation and subsequent scaling laws have remained stimulating, up to now, and some were shown to be valid for other superconductors as well²⁻⁹. While there are various attempts at a theoretical explanation of the Uemura relation^{4,10-12}, a connection to other experimentally probed properties of cuprates is still lacking, in particular to material chemistry parameters.

The various cuprate families have in common, see Fig. 1a, a CuO_2 plane and charge reservoir (CR) layers that separate the planes from each other. While the nearly square CuO_2 plane, defined by the Cu $3d_{x^2-y^2}$ orbital bonding to four O $2p_{\sigma}$ orbitals, is very similar for all systems, the CR chemistry can vary significantly. The nominal hole at Cu ($3d^9$ configuration) is responsible for strong magnetic correlations that make parent materials antiferromagnetic. Holes or electrons can be added to the CuO₂ plane by alteration of CR layers. As a result, static magnetism vanishes and new electronic phenomena emerge and the systems become conducting or superconducting. There are many similarities between different cuprate families, and one typically differentiates only between the hole and electron-doped phase diagrams, depicted in Fig. 1b, that appear to show a distinct

asymmetry. However, the extent in temperature and doping of the different phases and observed phenomena varies substantially between different families, also for the much more thoroughly investigated hole doped materials. In addition, the comparability between different families is somewhat obstructed with regard to doping. For example, while for $La_{2-x}Sr_xCuO_4$ the doping level can be varied over a large range quite reliably by stoichiometry, interstitial doping with oxygen (O_{δ}) as, for example, in HgBa₂CuO_{4+ δ} leaves uncertainties with regard to the actual doping level. In addition, ionic migration may cause phase separation or other ordering phenomena as in the YBa₂Cu₃O_{6+v} systems. Given the very different T_c for various cuprate families, it was questioned whether the average doping level of the CuO₂ plane is the appropriate chemical parameter for discussing all aspects of the complex physical properties of the cuprates, or whether other parameters should be considered as well, for example, distances within the plane, buckling, disorder, the role of the apical oxygen, or interlayer coupling. Nevertheless, from the understanding of the electronic properties we expect, in particular, clues as how to raise T_c .

Nuclear magnetic resonance (NMR), as a local probe of the magnetic spin susceptibility, focusses mostly on measurements of shift and relaxation caused by the interaction of the nuclear magnetic dipole moment with electronic magnetic moments. However, the nuclear electric quadrupole moment for nuclei with spin I > 1/2 (I = 3/2 for ^{63,65}Cu, I = 5/2 for ¹⁷O) interacts with the local electric field gradient (EFG) causing a quadrupole splitting (v_Q) of the NMR lines in high magnetic fields. The EFG at the nuclear site is very sensitive to the local charge symmetry, and has been useful for assigning NMR signals to the various lattice positions or detecting inhomogeneous charge distributions in the CuO₂ plane^{13,14}. Since the quadrupole splittings of planar Cu and O depend on doping, models have been put forward that attempted to understand these changes in terms of the hole content of certain orbitals, see for example, refs 15–18 and works

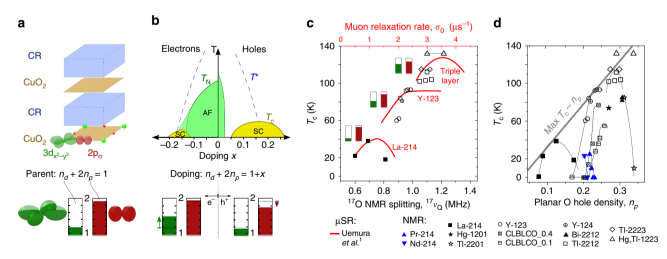


Figure 1 | General features of the cuprates. (a) The cuprates' layered structure consists of CR layers and CuO_2 planes with the bonding orbitals $Cu \, 3d_{x^2-y^2}$ and $O \, 2p_{\sigma}$, which share the nominal 3*d* hole of the Cu^{2+} ion. Columns indicate occupation of $Cu \, 3d_{x^2-y^2} \, (2-n_d)$ and $O \, 2p_{\sigma} \, (2-n_p)$, with hole contents n_d and n_p measurable with NMR. (b) Schematic representation of electronic phase diagram of the cuprates for electron and hole doping *x*: AF phase below Néel temperature (T_N), SC below critical temperature (T_c), and pseudogap regime below pseudogap temperature (T^*). Doped electrons (e⁻) go to the $Cu \, 3d_{x^2-y^2}$ orbital almost exclusively, while doped holes (h⁺) predominantly go to the $O \, 2p_{\sigma}$ orbital, arrows next to columns indicate changes of n_d and n_p caused by doping. (c) Solid red: Uemura plot¹, that is, T_c versus muon spin relaxation rate (σ_0 , upper abscissa); black symbols: T_c versus planar oxygen quadrupole splitting ${}^{17}v_Q$ (lower abscissa). For list of abbreviations see Table 1. For triple layer TI-2223 and Hg,TI-1223 the pairs connected with a dotted line belong to the same sample and correspond to planar O sites of inner and outer layer (smaller splitting corresponds to underdoped inner CuO_2 layer). (d) T_c versus planar O hole density n_p calculated from ${}^{17}v_Q$ for all available data (see text). Black dotted lines are guides to the eye and connect different doping levels for one family. Solid grey line indicates increase of the maximum T_c (that is, for the optimal doping level) as a function of n_p . AF, antiferromagnetic; SC, superconducting phase.

cited therein. While trends relating the local charge distribution with T_c could be established, the uncertainty with regard to the EFG contributions from a variable CR chemistry, limited quantitative agreement with charge contents expected from stoichiometry, as well as insufficient experimental data hampered advances of such analyses.

Here we show that if we plot T_c versus the planar ¹⁷O NMR quadrupole splitting, a functional dependence very similar to that of the Uemura plot emerges. This documents that the superfluid density is a function of the EFG at planar O. Based on very recent progress in the understanding of NMR quadrupole splittings in terms of the charge distribution in the CuO₂ plane¹⁹ we show that the maximum T_c increases with the hole content of the planar O $2p_{\sigma}$ orbital, at the expense of that at Cu $3d_{x^2-y^2}$. Thus, we identify material chemistry parameters, the hole contents at planar Cu and O, that are largely temperature independent, yet determine the superfluid density at low temperatures. This finding stimulates the use of these orbital hole contents, calculated from NMR literature data^{16,18-49}, to draw a threedimensional cuprate phase diagram that encompasses all cuprate families and has the superconducting domes ordered according to the maximum $T_{\rm c}$. We argue that such a phase diagram might be very useful in discussing the complex properties of the cuprates.

Results

Superfluid density and charge densities in the CuO₂ plane. We plot in Fig. 1c, together with the original Uemura plot (in red), T_c versus ${}^{17}\nu_{\rm Q}$ for similar materials and doping, and find a striking correspondence. This shows that the muon spin relaxation rate deep inside the superconducting state must be tied to the almost temperature independent EFG at the planar O nucleus, which determines the 17 O NMR splitting measured far above T_c , a rather unanticipated result.

It was confirmed recently, based on NMR data on the electron-doped and parent compounds, that NMR quadrupole splittings provide a quantitative measure of the charge distribution in the CuO₂ plane of apparently all cuprates¹⁹. A list of materials with abbreviations is given in Table 1. It was shown that the hole densities in the Cu $3d_{x^2-y^2}$ orbital (n_d) and the O $2p_{\sigma}$ orbital (n_p) in the CuO₂ plane are related to the experimentally measured splittings ⁶³ v_Q at ⁶³Cu and ¹⁷ v_Q at ¹⁷O as follows^{18,19}:

$$^{17}v_{\rm Q} = 2.45\,{\rm MHz} \cdot n_p + 0.39\,{\rm MHz},$$
 (1)

$$^{33}v_{\rm Q} = 94.3 \,\mathrm{MHz} \cdot n_d - 5.68 \,\mathrm{MHz} \cdot (8 - 4n_p).$$
 (2)

The planar oxygen splitting in equation (1) is only dependent on the hole content n_p of the onsite bonding orbital $2p_{\sigma}$ with the prefactor 2.45 MHz derived from the electric hyperfine interaction experimentally determined with atomic spectroscopy

Table 1 List of abbreviations.	
Abbreviation	Formula
Bi-2212	$Bi_2Sr_2CaCu_2O_{8+\delta}$
CLBLCO_x	(Ca _x La _{1-x})(Ba _{1.75-x} La _{0.25+x})Cu ₃ O _{6+y}
Hg-1201	$HgBa_2CuO_{4+\delta}$
Hg,TI-1223	$Hg_{0.5}TI_{0.5}Ba_2(Ca_{1-x}Sr_x)_2Cu_3O_{8+\delta}$
La-214	$La_{2-x}Sr_{x}CuO_{4}$
Nd-214	$Nd_{2-x}Ce_{x}CuO_{4}$
Pr-214	$Pr_{2-x}Ce_{x}CuO_{4}$
TI-2201	$TI_2Ba_2CuO_y$
TI-2212	$TI_2Ba_2CaCu_2O_{8-\delta}$
TI-2223	$TI_2Ba_2Ca_2Cu_3O_{10-\delta}$
Y-123	$YBa_2Cu_3O_{6+\gamma}$
Y-124	YBa ₂ Cu ₄ O ₈

of the O $2p^5$ state¹⁸. The term of 0.39 MHz is due to the charge symmetry at planar oxygen in the ubiquitous CuO₂ plane, and this term is found to be rather independent on doping and similar for all families¹⁹. Therefore, one can easily convert the experimentally measured ${}^{17}v_{\rm Q}$ into a reliable hole content n_p . The situation is somewhat more complicated for the Cu splitting in equation (2), which depends on the hole densities of both bonding orbitals. The first term is from the onsite hole content n_d of $3d_{x^2-y^2}$ with the prefactor 94.3 MHz, again derived from atomic spectroscopy of Cu $3d^9$ state¹⁸. The second term in equation (2) accounts for the EFG at the Cu nucleus caused by the charge in the bonding orbitals of the four surrounding planar O atoms, with the prefactor derived from the orbital overlap of O $2p_{\sigma}$ with the empty Cu 4p and the electric hyperfine interaction of the latter¹⁸.

First, we use equation (1) and convert all the planar oxygen splittings from the literature to n_p . The result is plotted in Fig. 1d, that is, we plot T_c versus n_p for the different materials. Of course, this plot is very similar to Fig. 1c, but it includes non-superconducting underdoped and parent materials with $T_c = 0$ K. We see that different cuprate families have rather different n_p , which results in the sorting of the families as in the Uemura plot. We also recognize that a large n_p is a prerequisite for a high maximum T_c , that is, at optimal doping. In Fig. 1d one can also notice a parabolic-like dependence of T_c on the oxygen charge n_p , which resembles the typical phase diagram that shows a dome-like dependence of T_c on the average doping level. The correlation between σ_0 and ${}^{17}v_Q$ is lost in the overdoped regime where σ_0 decreases with increasing doping^{50,51}, which was attributed to a decrease of n_s (ref. 52).

In Fig. 1d, we also included recent results for the electron-doped materials¹⁹. For Nd_{1.85}Ce_{0.15}CuO₄ the superfluid density was reported to be very similar to that of hole doped YBa₂Cu₃O_{6+y}, albeit measured optically and not by muon spin relaxation^{3,53,54}. We find that these results are also in agreement with ¹⁷v_Q splittings (see Supplementary Fig. 1) and corresponding hole contents for those two families, cf. Fig. 1d. Electron doping appears to be less efficient in providing a high T_c , but the rather high oxygen hole contents of the parent materials Pr₂CuO₄ and Nd₂CuO₄ suggest that hole doping should result in much higher T_c . The so-called infinite layer cuprate Sr_{1-x}La_xCuO₂, for which there are no reports of ¹⁷O splittings, has the highest T_c among electron-doped families and a very high muon spin relaxation rate ($\sigma_0 \approx 4.5 \,\mu s^{-1}$) (ref. 53), and we expect a high n_p . Indeed, a rather high T_c of more than 100 K was reported in the infinite layer system upon hole doping^{55,56}.

Clearly, a large n_p is a prerequisite for a high T_c , but is not sufficient, as expected for such a material chemistry parameter. If this empirical relation (max $T_c \propto n_p$) remains valid for higher oxygen hole content, the T_c of the cuprates might be raised substantially by the proper chemistry (we estimate 300-400 K per oxygen hole from the straight line in Fig. 1d).

The splittings of the ⁶³Cu NMR lines can only be converted into n_d if there are also ¹⁷O NMR data available, cf. equation (2). However, there are much less ¹⁷O splittings reported since the materials have to be enriched with ¹⁷O (the naturally abundant ¹⁶O nucleus has spin I=0) and therefore only part of ⁶³Cu splittings can be converted. (In Supplementary Fig. 1, we plot T_c versus experimentally measured splittings).

Note that the simple analysis using equations (1) and (2) gives hole densities that are in astonishingly good quantitative agreement with the total charge in the CuO_2 plane expected from the stoichiometry of the materials¹⁹, that is,

$$1 + x = n_d + 2n_p, \tag{3}$$

where the factor of 2 accounts for the two O atoms per CuO_2 .

This means that the sum of the hole contents n_d and n_p as determined with NMR (r.h.s.) equals the inherent Cu $3d^9$ hole content plus the hole content added by doping x (l.h.s.). This agreement was shown to apply for electron, as well as hole doping, and different parent materials differ only in terms of the charge transfer between Cu and O¹⁹. One can therefore also infer from Fig. 1d that compounds with the highest maximum T_c favour a smaller Cu hole content, and we conclude that it is the transfer of hole density to the O sites that is important for the highest T_c . With this result, one may ask if other properties of the cuprates should be discussed in terms of n_p and n_d , as well? This leads us to propose a cuprate phase diagram based on NMR.

Phase diagram of the cuprates based on NMR. In Fig. 2 we plot T_c as a function of n_d and $2n_p$ for all cuprates for which we could find both, Cu and O quadrupole splittings in the literature (see Supplementary Tables 1 and 2) with n_d and n_p calculated from equations (1) and (2). All materials appear in four separate groups, marked by colour: (1) $La_{2-x}Sr_xCuO_4$; (2) YBa₂Cu₃O_{6+ ν}, and other cuprates of that structure, for example, $(Ca_xLa_{1-x})(Ba_{1.75-x}La_{0.25+x})Cu_3O_{6+y}$ as well as YBa₂Cu₄O₈; (3) Bi, Tl and Hg based families; and finally, (4) the two electron-doped systems $Pr_{2-x}Ce_xCuO_4$ and $Nd_{2-x}Ce_xCuO_4$. The parent line, that is, the line that is given by $n_d + 2n_p = 1$ (bold dashed line) separates hole doped and electron-doped systems. Note that the lines parallel to the parent line are given by $n_d + 2n_p = 1 + x$, and represent constant hole (x = +0.1, +0.2) or electron (x = -0.1, -0.2) doping. While there may be material-specific uncertainties, for example, La₂CuO₄ is not located exactly on the parent line, our straightforward analysis uncovers simple systematic trends concerning all cuprates, and we discuss some salient features now.

While n_d and n_p change significantly between different parent compounds along the line $n_d + 2n_p = 1$, antiferromagnetism persists as long as there is one hole per CuO₂. Such a large range of variation in the charge transfer in different parent

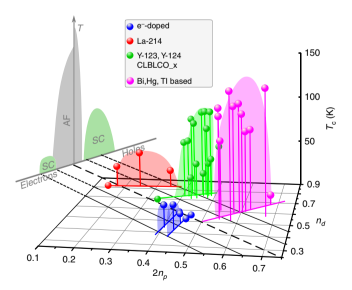


Figure 2 | Cuprate phase diagram from NMR. T_c as a function of oxygen $(2n_p)$ and copper (n_d) hole content for hole doped La-214, Y-123, Y-124, CLBLCO_x and Bi-, Hg-, Tl-based compounds, as well as electron (e⁻) doped Pr-214 and Nd-214. For list of abbreviations see Table 1. The parent line (dashed bold black) indicates expectation for the undoped case $(n_d + 2n_p = 1 \text{ from } x = 0)$, parallel lines (thin black) correspond to expectation for doping $x = n_d + 2n_p - 1$ changing with a step of 0.1. The commonly used phase diagram (*T* versus *x*) appears as a projection (upper left).

compounds, with $2n_p$ ranging from 0.15 to 0.45, is perhaps quite surprising, and its further increase, if possible, could raise T_c substantially.

Doping holes means entering the right upper half of the $(2n_p, n_d)$ -plane. While n_d and n_p increase with doping, the ratio of the respective changes $(\Delta n_d/2\Delta n_b)$ appears to be a family property, that is, parent materials with low n_p (for example, La_2CuO_4) add more holes to O than those with high n_p . With electron doping we enter the lower left half of the $(2n_p, n_d)$ -plane. Here, predominantly Cu holes disappear while the (large) O hole content changes only slightly. It is not apparent from the phase diagram why most parent materials can only be doped with one type of carrier. As a function of doping, T_c increases with a slope that depends on the position on the parent line, as well, and hole doping seems to be more effective in raising T_c . Another important observation concerns optimal doping, that is, the doping level for which one finds the highest T_c for a given family. According to our analysis it is related to $x = n_d + 2n_p - 1$ and not particular values of n_d and n_p . However, we do observe a slight increase of the optimal x with increasing n_p (decreasing n_d). Note that the doping level x follows from our analysis in terms of n_d and n_p inserted into (3) and is not deduced from material chemistry. Our analysis agrees with expectations also for materials doped by interstitial O_{δ} where doping level x is often derived from the T_c dome¹⁹. Interestingly, the latter materials we find located in the same group, despite significant structural differences between Hg-, Tl- and Bi-based cuprates. Also the number of close CuO2 layers in multi-layer systems does not result in significant differences in the charge distribution.

Discussion

In Fig. 2 we plotted only T_c in the $(2n_p,n_d)$ -phase diagram, but it might be of great interest to investigate whether other cuprate properties are better presented as a function of the local charge distribution, instead of the average doping level. While further analysis is beyond the scope of our paper, we shortly discuss some other cuprate properties with regard to our phase diagram.

The Néel temperature depends on the interlayer coupling and therefore is not expected to be dominated by the charge distribution in the CuO₂ plane. For example, YBa₂Cu₃O₆ has a higher T_N than Pr₂CuO₄ and La₂CuO₄. It would be interesting, however, to find out how the exchange coupling (*J*) changes along the parent line. Recently, there have been contradicting reports regarding *J* in the cuprates^{57–59}. Mallet *et al.*⁵⁸ found no correlation between *J* and $T_{c,\max}$ in R(Ba, Sr)₂Cu₃O_y, while Wulferding *et al.*⁵⁷ claimed that *J* is correlated with $T_{c,\max}$ in (Ca_xLa_{1-x})(Ba_{1.75-x}La_{0.25+x})Cu₃O_{6+y}, which was later questioned by Tallon⁵⁹.

Structural parameters of the CuO₂ plane such as distances, buckling, or disorder appear to show no clear trend with respect to n_d and n_p . However, the apical oxygen distance from the CuO₂ plane increases as one follows the parent line beginning from low n_p , similar to the maximum possible T_c . This behaviour and the concomitant change in density of states of Cu 4*s* was noted before⁶⁰.

Pressure applied to underdoped cuprates usually increases T_c , while the structural changes to even hydrostatic pressure can be complicated⁶¹. For example, specifically strained HgBa₂CuO_{4+ δ} can have almost identical CuO₆ octahedra as La_{2-x}Sr_xCuO₄, however, the large difference in their T_c values remains⁶². This might be related to the different values of n_d and n_p for these families. As a result of recent progress in anvil cell NMR^{63,64}, it is now possible to study cuprates at high pressures also with NMR⁶⁵, and it was found that the Cu splitting increases with pressure indicating changes in the planar hole contents⁶⁶. However, single-crystal studies are necessary and ongoing efforts by our group aim at providing a quantitative measure of the local charge distribution as a function of pressure.

Another important issue concerns the heterogeneity of the cuprates. We know from NMR that the static charge and spin density can vary drastically within the CuO₂ plane, in particular between different cuprate families⁶⁷. For example, the charge density in terms of the total doping x may easily vary by $\Delta x \approx 0.05$ (refs 32,68,69). Since T_c is not in a simple relation to this static inhomogeneity, only the average n_d and n_p appear to matter. From this, one would conclude that inhomogeneity is either not important for the maximum T_c, or it is ubiquitous and dynamically averaged for NMR, depending on the chemical environment.

To conclude, NMR measures the charge distribution in the bonding orbitals in the CuO₂ plane quantitatively, and since it reproduces the Uemura plot, that is, it finds the same ordering of families with respect to their maximum T_c , we now have material chemistry parameters that are responsible for setting the highest T_c and superfluid density. These findings inspired a different perspective on the cuprate phase diagram and it is likely that the complex cuprate properties might be better understood when discussed in the context of the charge distribution in the CuO₂ plane.

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Author contributions

D.R., M.J. and S.R. contributed to data gathering, analysis and writing the manuscript. All the authors discussed the results and worked on the manuscript. C.K. helped with discussion of the μ SR technique and its relation to NMR. J.H. supervised and contributed equally to the data analysis and manuscript editing, as well as providing project leadership.

Additional information

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