Cationic substitution and role of oxygen in the *n*-type superconducting T' system $Nd_{2-y}Ce_yCuO_z$

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(Received 16 August 1989)

The effect of oxygen content and cationic substitution on the structural and superconducting properties of the T' phase were studied by investigating the $Nd_{2-y}Ce_yCuO_z$ and $R_{1.85-x}R'_xM_{0.15}CuO_z$ ($R = Nd,Eu; R' = La,Y; M = Ce,Th; z \approx 4$) series. We find that solid solutions of the T' phases can be prepared, and the solubility range of La and Y depends on the sizes of the host versus the dopant ions. The transition from tetragonal T' phase to the orthorhombic T phase is quite abrupt, without evidence for the T^* phase. In this system, as for the La-Sr-Cu-O system, we find that superconductivity seems to correlate with the in-plane Cu-Cu distance, and there is a critical distance (3.95 Å) for which T_c is maximum. The as-prepared Ce- or Th-doped materials have an oxygen content per unit formula greater than four, and when heated under nitrogen they lose oxygen in two steps at temperatures of 400 and 800°C, with the amount of oxygen lost through each step being strongly dependent on sample processing. The total amount of mobile oxygen lost decreases with increasing Ce or Th content. From thermogravimetric-analysis measurements we show the crucial importance of the reducing step at temperatures greater than 850°C for inducing superconductivity as well as the importance of the cooling rate on the superconducting properties of the T' phases. The importance of oxygen disorder to the normal-state properties is demonstrated.

INTRODUCTION

Since the first discovery of high- T_c superconductivity¹ in the $La_{2-x}Ba_xCuO_4$ system that has a distorted K_2NiF_4 structure (or the so-called T phase), superconductivity has also been found in two other K₂NiF₄related structures, T' and T^* phases (Refs. 2 and 3, respectively). The T' phase R_2 CuO₄ (R = Pr-Gd) is exhibited by most rare earths, whereas the cuprate T phase is found only when R = La. One major difference between the T and T' phases is the coordination of copper. Copper in the T phase has a sixfold coordination with four of the oxygen atoms in the CuO₂ plane, and the remaining two oxygen atoms (called apical oxygen) above and below the Cu atoms. In contrast, copper in the T'phase is in fourfold coordination (i.e., no apical oxygen). Intermediate between the T and T' phases is the T^* phase, $La(Pr)_{2-x-y}R_xSr(Ba,Ca)_yCuO_4$ (*R* denotes rare earth), in which Cu is fivefold coordination with oxygen atoms occupying both the 4d (apical oxygen) and 4e(fluorite-type oxygen) sites.⁴ Here we determine, as a function of the rare earth, the structural stability of the T' phase with respect to the T and T^* phases. We report our studies of the Laand Y-substituted $R_{1.85}R'_{x}M_{0.15}CuO_{4-y}$ (R = Nd,Eu; R'=La,Y; М =Ce,Th) systems.

The superconductors with the T^* or T structure are p type (i.e., the carriers are holes), whereas the superconductors with the T' structure are n type (i.e., the carriers are electrons). This difference is indirectly revealed from the way in which superconductivity is induced in these

phases. Superconductivity is induced in the T and T^* phases by increasing the oxidation state of Cu either by divalent dopant (e.g., Sr) and/or by means of an oxidizing atmosphere. On the other hand, to promote superconductivity in the T' phase, it is necessary not only to lower the oxidation state of copper by a tetravalent dopant but also to anneal the as-grown sample in a reducing atmosphere. It is therefore important to determine how oxygen content affects superconductivity behavior in the T' phase by studying the Nd_{2-y}Ce_yCuO_z system along with other series in which trivalent Nd ions are replaced partially by La or Y, completely by Eu, or in which the tetravalent Ce ions are replaced by Th.

Finally the T^* phase, whose unit cell can be viewed as containing half of the unit cell of the *p*-type *T* phase and half of the unit cell of the *n*-type *T'* phase, appears as an attractive system in which one could expect, depending on the nature of the chemical doping, to achieve either *p*or *n*-type superconductivity. Doping studies on the T^* phase were performed, and our results will be briefly mentioned here. Fluorine substitution or tetravalent substitution have failed to produce *n*-type superconductivity in the T^* materials.

EXPERIMENTAL

To simultaneously study the structural stability of the T' phase with respect to the T and T', as well as its T_c dependence as a function of the rare earths, we first undertook a survey of several series of samples of general formula $R_{2-x-y}R'_xCe_yCuO_z$ to determine the Ce content

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(y) at which T_c is maximum. We found that for a fixed rare-earth R/R' ratio, T_c was always maximum for Ce content close to 0.15. Thus, in the following we focused only on the series with y = 0.15.The $R_{1.85-x}R'_{x}M_{0.15}$ CuO_z solid solutions were prepared by heating the appropriate oxides in the required stoichiometry in air for 20 h at 900 °C with two grindings and making into pellets before annealing at 1100 °C for 12 h or longer (step 1). A second annealing at 900 °C under N_2 (step 2) was required in order to induce superconductivity in these samples. We found that the superconducting properties of Ce-doped samples are strongly affected by the cooling rate in both synthesis steps 1 and 2, and the annealing temperature in step 2. Unless otherwise stated, the samples were cooled in air from 1100 °C in 2 h (step 1), and in nitrogen from 900 °C in 4 h (step 2).

Fluorination was carried out on two of the reported T^* phases, and samples of La_{1.08}Dy_{0.72}Sr_{0.2}CuO_{4-x}F_x and La_{0.84}Sm_{0.96}Sr_{0.2}CuO_{4-x}F_x with (x = 0-0.35) were prepared by heating the corresponding oxides and DyF₃ and SmF₃ at 920 °C for 15 h, with one regrinding. The powders were then made into pellets and fired at 1050 °C for 18 h in air and annealed in N₂ at 890 °C for 12 h.

Oxygen stoichiometry was obtained by thermogravimetric analysis (TGA) measurements under various atmospheres (N₂,O₂ or 4% H₂ in Ar) with a heating rate of 5 °C/min. X-ray-diffraction measurements with Cu K α radiation were used to determine the range of existence of the solid solutions. The upper limit of solubility range was taken as the highest value of x (R' content) or y (Ce or Th content) for which the samples did not show a trace of the second phase. Lattice parameters for singlephase materials were refined using the Bragg peaks over the θ range, 20° < 2 θ < 80°, as described in Ref. 8. Superconductivity transition temperatures were determined by ac-susceptibility measurements (13 Hz), and the resistivity temperature dependence was measured with a standard four-probe configuration.

RESULTS

Solid solutions of $Nd_{2-x}La_xCuO_z$ can be synthesized in the range 0 < x < 0.9. With Ce doping, the range was extended to 0 < x < 1.2 in $Nd_{1.85-x}La_xCe_{0.15}CuO_4$ series [Fig. 1(a)]. Mixtures of T and T' phases were found in the range 1.2 < x < 1.7 for the $Nd_{1.85-x}La_xCe_{0.15}CuO_4$ series and in the range 0.9 < x < 1.65 for the $Nd_{2-x}La_{x}CuO_{4}$ series. Pure T phase was formed for x > 1.7 and x > 1.65 for the Nd-La-Ce-Cu-O and Nd-La-Cu-O series, respectively. The transition from tetragonal T' phase to the orthorhombic T phase with increasing La doping is guite abrupt (i.e., that the two-phase region is narrow). For both series, no peaks due to the T^* phase was detected in the powder x-ray diffraction through the two-phase region. This, in conjunction with the recent report⁴ on new T^* phases containing Ca^{2+} , Ba^{2+} , or Sr^{2+} , indicates that the stability of the T^* phase is probably more sensitive to alkaline earth than rareearth dopant. This is also evident from our failure to synthesize T' phases of $Nd_{2-x-y}La_xSr_yCuO_4$. As soon



FIG. 1. Variation of the tetragonal lattice parameters a,c and volume V as a function of the x for the (a) $Nd_{2-x-y}La_xCe_yCuO_4$ series and (b) $Eu_{2-x-y}La_xCe_yCuO_4$ series.

as Sr doping is introduced, mixtures of T' and T^* phases formed.

When Nd was replaced by the smaller Eu ion in $Eu_{1.85-x}La_xCe_{0.15}CuO_4$, the solubility range of La increased to 0 < x < 1.35 [Fig. 1(b)], and as for the Nd system already described, one finds that the Ce doping extends the solubility range of La. A small decrease in the solubility range of La (0 < x < 1.1 instead of 0 < x < 1.2)was only observed when Ce was replaced by a Th ion in the $Nd_{1.85-x}La_xTh_{0.15}CuO_4$ series [Fig. 2(a)]. These solid solutions illustrate the importance of size of the host (Nd_2CuO_4) versus the dopant ions (Ce and Th) in stabilizing the T' phase. When Nd³⁺ (1.25 Å) is partially substituted by the smaller Ce⁴⁺ (1.11 Å) dopant, more La³⁺ is substituted for Nd^{3+} in $Nd_{1.85-x}La_xCe_{0.15}CuO_4$ than in $Nd_{2-x}La_{x}CuO_{4}$. Similarly, one finds the solubility range of La in M_{2-x} La_xCe_{0.15}CuO₄ to increase when Nd is replaced by a smaller ion, Eu³⁺ (1.21 Å). In contrast, the size effect is not very pronounced when comparing $Nd_{1.85-x}La_xCe_{0.15}CuO_4$ [Fig. 1(a)] with $Nd_{1.85-x}La_xTh_{0.15}CuO_4$ series [Fig. 2(a)]. This is because Th^{4+} (1.13 Å) and Ce^{4+} (1.11 Å) have a similar ionic radius.

In the case of the Y-substituted series, one observes that the $Nd_{1.85-x}Y_xCe_{0.15}CuO_4$ series [Fig. 2(b)] have a smaller solubility than the $Nd_{1.85-x}La_xCe_{0.15}CuO_4$ series, since Y^{3+} (1.16 Å) is smaller than La^{3+} or Nd^{3+} . Comparison of the $Nd_{1.85-x}Y_xCe_{0.15}CuO_4$ with $Nd_{2-x}Y_xCuO_4$ series [Fig. 2(b)] shows that the addition



FIG. 2. Variation of the tetragonal lattice parameters a,c and volume V as a function of the x for the (a) $Nd_{2-x-y}La_xTh_yCuO_4$ series and (b) $Nd_{2-x-y}Y_xCe_yCuO_4$ series.

of Ce does not enhance the Y solubility, in contrast to the La-doped Eu and Nd series. Indeed, the solubility range is about the same for both series, and at large Y concentratio peaks due to $Y_2Cu_2O_5$ were detected by powder x-ray diffraction. Thus, large Y concentrations tend to destabilize the T' structure and form the $Y_2Cu_2O_5$ phase, confirming that the T' or T phase cannot be synthesized for rare earths with small ionic radii.⁵

In all the La substituted series, $R_{1.85-x} La_x M_{0.15} CuO_{4-v}$ (R = Nd, Eu; M = Ce, Th), the lattice parameters a, c, and the volume of the tetragonal unit cell increase with increasing La concentration. The opposite was observed for the Y-subsituted T' phases. This is because La is larger than Nd and Eu, whereas Y is smaller than Nd and Eu. A general observation is that independent of the doping element, the a and c lattice parameters either increase or decrease simultaneously. In fact the a and c parameters for all compounds with a Ce content of 0.15 are almost perfectly correlated, meaning that all compounds with a given a have the same c and hence the same volume. As an example, a = 3.95 Å, c = 12.129 Å, and V = 189.5 Å³ compared to a = 3.953 Å, c = 12.116 Å, and V = 189.36 Å³ for the $Nd_{1.55}La_{0.3}Ce_{0.15}CuO_z$ and $Eu_{0.95}La_{0.9}Ce_{0.15}CuO_z$ compounds, respectively. Finally for the as-grown materials and reduced materials, the change in the lattice parameters were very small and in the limit of our resolution.

The superconducting critical temperature were determined for all the members of the Ce- or Th-doped series by the inductive method. In Fig. 3 we plot the T_c 's



FIG. 3. T_c (onset) as determined by ac susceptibility vs the *a* lattice parameter. For the Eu series, two sets of samples were investigated. Samples represented by the triangles were prepared as described in the text. The samples represented by a star were prepared by removing nitrogen at 1000 °C, putting back oxygen at 840 °C, and removing nitrogen at 900 °C

(determined as the onset of the ac signal for quenched samples) as a function of the *a* crystallographic distance (i.e., Cu-Cu distance within the CuO_2 plane). Note a correlation between the lattice parameter a and the superconducting onset temperatures (T_c) . The optimum onset temperature $(T_c = 25 \text{ K})$ for the lanthanum Cedoped samples is found for a between 3.945 and 3.955 Å. A dramatic decrease in T_c is observed when a becomes either greater than ≈3.96 Å or less than ≈3.94 Å. Interestingly, La doping seems to raise the onset of T_c from 12.5 K in $Eu_{1.85}Ce_{0.15}CuO_4$ ($a \approx 3.947$ Å) to about 17 K $Eu_{0.9}La_{0.9}Ce_{0.15}CuO_4$ ($a \approx 3.953$ Å). In the in $Nd_{1.85-x}Y_xCe_{0.15}CuO_4$ series, superconductivity was observed only for a greater than ≈ 3.94 Å. These results show that in the T' phases there is a very critical and narrow range of a distances to obtain high T_c . In support of this correlation, it is worth mentioning that the *a* lattice parameter of the highest- T_c fluorine-doped material,⁶ $Nd_2CuO_{3,7}F_{0,3}$ falls also into the maximum of Fig. 3 with $a \cong 3.945$ Å.

The temperature dependence of resistivity for the members of each series that showed the highest T_c 's are shown in Fig. 4. The T_c values are in good agreement with those deduced from ac-susceptibility measurements. Above T_c , the resistivity is semiconductinglike, as usually found for polycrystalline ceramics of this system, and explained in terms of composition inhomogeneities.⁷ In the Nd_{2-x}La_xCe_{0.15}CuO₄ series, with increasing La content beyond x = 1, one destroys superconductivity and promotes semiconducting behavior. This result contrasts with the superconducting to metallic transition observed for the La-Sr-Cu-O system or Nd-Ce-Cu-O systems upon increasing Ce or Sr content, respectively.^{8,9}

Meissner and shielding measurements performed on



FIG. 4. Temperature-dependent resistivities of selective T' phase compounds which have shown the highest T_c by acsusceptibility measurements.

several members of the Ce-doped Eu and Nd series, by means of a SQUID magnetometer, reveal that the superconducting volume fraction does not exceed 25%. This volume fraction is independent of the cation ratio.

A depression in T_c by magnetic ions is commonly observed in a conventional superconductor. Since the rare earths are closer to the CuO₂ plane in the T' phase than in the T phase (1.81 Å as compared to 2.41 Å), one would expect a greater sensitivity of T_c to rare-earth doping in the T' phase, and thereby the T_c of Nd compound to be smaller than that of the Eu compound [in which Eu⁺³ is a Van Vleck ion (J=0)]. We note, however, the opposite; T_c is smaller for the Eu compound than for the Nd compound. Therefore, as for the La_{2-x}M_xSr_{0.2}CuO₄ (T) system with M = La to Eu,¹⁰ we conclude the magnetism is obviously not the explanation for the depression in T_c observed in the T' phases when Nd is replaced successively by Pr, Sm, and Eu.^{11,12}

Finally, we find that among the T^* phases investigated, fluorination is only possible within the La_{1.08}Dy_{0.72}Sr_{0.2}CuO_{4-x}F_x, system and single phases are obtained only for values of x ranging from 0 to 0.35. Refinement of the lattice parameters (Table I) shows a slight contraction of the *c* axis and slight expansion of the *a* axis, as observed in the Nd₂CuO_{4-x}F_x series.⁶ Independent of the cooling rate or ambient used, none of the fluorinated samples show superconductivity.

OXYGEN DOPING

In addition to the Ce content, another critical parameters which affects T_c in the T' phases $(Nd_{2-x}Ce_xCuO_z \text{ or } R_{2-x-y}R'_xCe_yCuO_z)$ is the oxygen content (z). For example, when $Nd_{1.85}Ce_{0.15}CuO_z$ is annealed in N₂ below 700 °C, no superconductivity is detected. Reannealing the sample at 820 °C, a weak and incomplete superconducting transition is observed with a T_c (onset) $\cong 16$ K. On further annealing, the sample at 900 °C, bulk superconductivity is lost on further annealing at temperatures greater than 1000 °C. Thus, we investigated the role of the nitrogen annealing treatment and more precisely how the oxygen stoichiometry affects superconductivity in these compounds by the TGA technique.

The effect of cooling rate on the oxygen content of the as-grown or reduced material was studied on the compound $Nd_{1.85}Ce_{0.15}CuO_z$. Two pellets of this compound were annealed at 1100° C for 14 h. Then one pellet (denoted Q) was quenched from 1100 °C and the other pellet (denoted S) was slowly cooled (8 h) to 300 °C. Figure 5 shows the TGA trace for the nonsuperconducting Qand S samples when annealed N_2 (a), cooled in N_2 and reannealed in (b), oxygen and finally cooled in oxygen and reannealed in (c) nitrogen. The difference in Q and S is apparent from the curves in (a). Oxygen removal occurs mainly at T = 400 °C for the quenched sample (Q) and for the slowly cooled sample (S), the maximum oxygen loss occurs for temperatures above 800° C. The total amount of oxygen loss is about 5% smaller for the Q sample than for the S sample. After the first nitrogen annealing, the Q and S samples behave identically whether reannealed under oxygen [curves in (b)] or nitrogen [curves in (c)], and the TGA traces are similar to those obtained on single crystals.⁷ The difference between Q and S, with respect to the oxygen removal during the first reducing step, suggests that the excess oxygen in the Q sample is more mobile and can escape from the sample at a lower temperature than that in the S sample. We propose that at high temperatures, because of entropic effects, one favors oxygen disorder. Thus, perhaps in the S samples these oxygens order at some temperature below 1100 °C (possibly forming a superlattice that should be observed at room temperature), so that their mobility is reduced. Evidence for such a superlattice has been detected by transmission electron microscopy on a slowly cooled Cedoped sample⁷ or F-doped sample.¹³ In contrast, in the Q samples, which are cooled too fast for this order to develop, the oxygens remain disordered and more mobile superlattice was detected [no by transverse-

TABLE I. Lattice parameters of $La_{1.08}Dy_{0.72}Sr_{0.2}CuO_{4-x}F_x$ (x =0-0.35) series.

Compounds	a (Å)	<i>c</i> (Å)	V (Å ³)	
x = 0	3.8487±0.0008	12.5310±0.0030	185.6160±0.0873	
x = 0.1	$3.8483 {\pm} 0.0010$	12.5229 ± 0.0039	185.4569±0.1141	
x = 0.2	3.8491±0.0013	$12.5230{\pm}0.0057$	185.5376±0.1511	
x = 0.35	$3.8508 {\pm} 0.0011$	12.5044 ± 0.0043	185.6222±0.1247	

electromagnetic mode on quenched samples]. After the removal of the mobile oxygens from the structure during the first reducing treatment, the samples behave identically with respect to their ability to insert or deinsert oxygen (the anomaly at 400 °C disappears in both the Q and S samples). This feature is not only typical of the Nd_{1.85}Ce_{0.15}CuO₄ sample, but it is also observed in all the other $R_{1.85-x}R'_{x}M_{0.15}$ CuO_z samples investigated. Figures 5(d) and 5(e) show the TGA data for the Eu_{1.85}Ce_{0.15}CuO_z and Eu_{0.95}La_{0.9}Ce_{0.15}CuO_z compounds, respectively. As before, after the first oxygen removal under N₂ the step present at 400 °C disappears on subsequent oxygen removal, and further oxygen loss occurs only for temperatures greater than 800 °C. The



FIG. 5. Effect of processing (cooling rate and ambient) on the oxygen content for $Nd_{1.85}Ce_{0.15}CuO_z$, (a) sample annealed under nitrogen, (b) sample annealed under oxygen after quenching, (c) sample reannealed under nitrogen after quenching. The solid curve is for a sample quenching from 1100 °C in air, and the dashed curve is the slowly cooled sample. In (d) and (e) the TGA curves for $Eu_{1.85}Ce_{0.15}CuO_z$ and $Eu_{0.95}La_{0.9}Ce_{0.15}CuO_z$, respectively, are shown. The samples were heated in N₂ (1), annealed in O₂ (2), and finally reannealed in N₂ (3).

weight gain and loss then becomes identical. We have already discussed that the drop in weight at 400 °C in Fig. 5(a) could be from disordered oxygen in interstitial sites and the drop at 800 °C from ordered oxygen in the same sites. The small dip at 400 °C in Fig. 7 may indicate that the ordering is not complete in samples cooled in air from 1100 °C.

To investigate the effect of cooling rate on T_c , the nonsuperconducting pellets Q and S were each broken into two pieces. All four pieces were annealed under nitrogen at 900 °C for 10 h and either guenched from 900 °C or slowly cooled (8 h). The resulting samples are denoted QQ, QS, SQ, and SS where QQ stands for a sample that has been quenched after both step 1 (treatments at 1100 °C) and step 2 (reducing treatment at 900 °C), whereas QS stands for the pellet that has been quenched after step 1 and slowly cooled after step 2, and vice versa for the SQ and SS samples. Figure 6 shows the resistive temperature dependence for these four samples. The samples QQ, QS, SQ, and SS superconduct with onset temperatures 23, 25.3, 23.2, and 22.5 K and zeroresistance temperatures of 16.5, 19.5, 19, and 18 K, respectively. The room-temperature resistivity values increase in going from the SS to the QQ samples (uncertainty in the relative resistivity values are less than 10%). The temperature dependence of the resistivity is semiconductinglike for the samples that were quenched in step 1 (QQ and QS), whereas it is flat or even slightly metalliclike for T > 150 K for samples that were slowly cooled in step 1 (SQ and SS). Note that the QQ and SQ samples look just like QS and SS, respectively, but shifted to higher resistivity. These similarities in temperature dependence suggest that the first step (at 1100 °C) changes



FIG. 6. Effect of cooling rate on the resistivity on Nd_{1 85}Ce_{0.15}CuO₄. The notation is Q for quench and S for slow cool. The first letter represents the cooling rate from 1100 °C and the second, the cooling rate after anneal at 900 °C. Note that the processing from high-temperature determines the shape of the resistivity temperature dependence and the processing from 900 °C the room-temperature value.

Compounds	Oxygen content (y) after step (a)	Oxygen content (y) after step (b)	Change in oxygen ΔY between step (a) and step (b)	[O] _{N2}	[O] ₀₂
Nd ₂ CuO ₂	4.1097	4.044	0.066	0.1144	0.068
$Nd_{1.95}Ce_{0.05}CuO_{z}$	4.1228	4.028	0.095	0.91	0.0364
$Nd_{1.9}Ce_{0.1}CuO_2$	4.065	4.0136	0.0514	0.1116	0.0519
$Nd_{1.85}Ce_{0.15}CuO_z$	4.032	3.991	0.041	0.0908	0.0415
$Nd_{1.8}Ce_{0.2}CuO_z$	4.031	3.977	0.054	0.0778	0.0389
$Nd_{1.55}La_{0.3}Ce_{0.15}CuO_z$	4.001	3.974	0.027	0.0595	0.0440
$Nd_{1.55}Y_{0.3}Ce_{0.15}CuO_{z}$	4.033	3.989	0.044	0.0573	0.0174
$Nd_{1.55}La_{0.3}Th_{0.15}CuO_z$	4.034	3.998	0.036	0.0615	0.0321

TABLE II. TGA results for the $Nd_{2-x}Ce_xCuO_z$ and $Nd_{1.55}R_{0.3}M_{0.15}CuO_z$ series. (R' = La,Y; M = Ce,Th). H_2 reduction data taken up to 870 °C. N₂ and O₂ data taken up to 950 °C.

the term that controls the temperature dependence of the resistivity, whereas the second step (at 900 °C), changes a constant term in the resistivity. Resistivity measurements are sensitive to disorder with resistivity values increasing with increasing structural disorder. Thus our data suggests that there are two types of disorder (disorder may be cationic or anionic including phases with different oxygen ordering) produced by the different treatment and that the degree of structural disorder is greater for the QQ than for the SS samples. This is in agreement with our model derived from the TGA results; that quenching favors structural disorder. From the TGA measurements, one would expect that superconductivity could be induced in a quenched sampled by annealing it under nitrogen at 600 °C. However, annealing a quenched sample at 600 °C for 12 h did not produce superconductivity. We find that independent of the cooling rate after the reducing treatment, superconductivity can only be induced in these materials for reducing temperatures greater than 850 °C. Similarly, we find that superconductivity can only be destroyed by reannealing them under oxygen at T > 800 °C. Thus, like the La-Sr-Cu-O system, the changes in oxygen and T_c in these T' materials are reversible.

Table II displays the total oxygen content, as deduced by TGA measurements using a reducing atmosphere (less than 5% H_2 in argon), for several as-grown and nitrogen annealed members of the $Nd_{2-y}Ce_yCuO_z$ and $Nd_{2-x}La_xM_{0.15}CuO_z$ series (M = Ce, Th). Using this technique, an accurate determination of the oxygen content requires a perfect knowledge of the reduced species. For Th, the only reduced species is ThO_2 . In contrast, for cerium oxide the the reduced species CeO_{2-x} has stoichiometries that can vary over a wide range of x (0 to 0.5). Therefore, pure CeO_2 was reduced under similar conditions as the as-grown material and the reduced species was found to be Ce_2O_3 . Thus, we were able to deduce z for the Ce-doped samples with an accuracy of about 0.5%. Independent of the Ce content, the oxygen content (z) per formula unit $Nd_{2-\nu}Ce_{\nu}CuO_{z}$ is always greater than 4 (i.e., interstitial oxygens) for the as-grown samples, but decreases with increasing Ce concentration (y). In the ideal T' structure the oxygen apical positions (i.e., oxygen above and below the Cu) are not occupied.

These results suggest a deviation from the T' structure in that the extra oxygens [determined by both TGA analysis (as earlier) or by chemical titration¹⁴] may sit on the apical 4d sites. In the following, we will consider that three types of oxygen (oxygen in the CuO₂ plane, fluorite type, and apical) exist in the $Nd_{2-\nu}Ce_{\nu}CuO_{4}$ compounds. Annealing under nitrogen at 900 °C lowers the oxygen content per formula unit, and for the compound with a Ce content of 0.15, the oxygen content becomes lower than 4 (i.e., no extra oxygen). It might be purely a coincidence, but it is for this Ce concentration that the compound becomes superconducting (we stress that caution must be exercised in interpreting these results, since our maximum accuracy in determining the oxygen content is $\pm 0.5\%$). Quite interesting also is the Δz decreases with increasing Ce concentration. In other words, this indicates that the insertion of Ce reduces the ability of these materials to reversibly change oxygen stoichiometry.

To determine how the reversibility of oxygen content varies with Ce content in the $Nd_{2-y}Ce_yCuO_z$ series, or with the R/R' ratio within the $R_{2-x-y}R'_yCe_{0.15}CuO_z$ series, we monitored the weight loss or gain of the sample when heated to 900 °C at a rate of 5 °C/min in a nitrogen ambient [step (a)], quenched in nitrogen, and reannealed in oxygen to 950 °C at a rate of 5 °C/min [step (b)], respectively. The weight loss during step (a) or gain during step (b) is summarized in Table II for several members of the $Nd_{2-y}Ce_yCuO_z$ series. Note that the weight loss is always greater than the weight gain, suggesting a slight loss of one element (not yet determined) when the material is heated under nitrogen. Also, the obtained values for the uptake of oxygen agree well with the Δz value, determined previously.

Figures 7(a) and 7(b) only show the TGA traces for the as grown $Nd_{2-y}Ce_yCuO_z$ and several $Eu_{2-x-y}La_xCe_yCuO_z$ phases, respectively, when annealed under N₂. The typical TGA curve can be divided into three regions: Region I spans from room temperature to about 760 °C, region II is from \approx 760 to \approx 950 °C, and region III is above \approx 950 °C (data not completely shown). The amplitude of weight loss in oxygen through these regions changes with both the cerium content and with the R/R' ratio: Both Figs. 7(a) and 7(b) show that the weight loss in region I decreases as Ce doping in-



FIG. 7. TGA data for several samples when heated to 850° C at a rate of 5°C/min in a N₂ ambient In (a) the TGA traces are shown for the Nd_{2-y}Ce_yCuO₄ series and in (b) for the Eu_{1.85-x}La_xCe_{0.15}CuO₂ series.

creases. In contrast, note that upon partial La substitution for Eu, there is no noticeable change in the TGA curves. The weight loss in region II is slightly dependent on the Ce concentration in the $Nd_{2-y}Ce_yCuO_4$ series, but is not dependent on the type of rare earth substituted for Nd.

The oxygen lost through regions I and II may arise from the interstitial oxygens (i.e., locally ordered or disordered apical oxygen sites). However, one should bear in mind that the T' phase contains fluorite-type oxygen sandwiched between the rare-earth layers, and that fluorite-type oxides are prone to oxygen defects.¹⁵ Thus it is likely that some of the oxygen loss comes from the fluorite-type layer in the T' phase. We cannot presently distinguish between these two possibilities, and only further neutron work may resolve this issue (apical versus fluorite-type oxygens) but we can state that it is the oxygen that is removed at 800 °C from the structure, which is crucial for superconductivity. The oxygen loss in region III for temperatures greater than 950 °C probably comes from the CuO_2 layer, since we observed that T_c is suppressed when the compounds are annealed under N_2 at T > 950 °C, and have previously reported that vacancies or disorder in this layer destroy superconductivity. Simultaneously, above 1100 °C phase decomposition takes place, as observed by high-temperature x-ray diffraction, discussed in the following. We summarize [Figs. 7(a) and 7(b)] by noting that the total amount of mobile oxygen loss Δz in the $R_{1.85-x}R'_xCe_yCuO_z$ series is decreasing with increasing the Ce content (y) independent of R, and remains constant $\Delta z \approx 0.04$ for a fixed Ce content (y = 0.15), independent of the R/R' ratio. We conclude that the role of the nitrogen treatment is twofold: (1) to remove extra oxygens (either "apical" or fluorite type) within the structure and (2) to increase the number of carriers (electrons) through the reduction of Cu^{2+} to Cu^{1+} .

Another interesting feature, common to most of the investigated samples, is the change in slope (i.e., kink) in the TGA curves occurring at T = 880 °C during the anneal under nitrogen (i.e., removal of oxygen). Such a slope change may be due to a structural phase transition or a beginning of phase decomposition. To distinguish between these two possibilities, high-temperature x-ray diffraction measurements under nitrogen ambient have been performed on as-grown material. No evidence for such a transition was observed (Fig. 8) up to temperatures of 1100 °C. However, note that the spectrum at 1100 °C exhibits an extra peak at $2\theta = 28^{\circ}$. This peak, which is still present after cooling the sample to room temperature, is suggestive of a slight amount of phase decomposition. These results clearly indicate the absence of a major structural transition occurring at this temperature, but do not rule out the possibility of an oxygen orderdisorder transition that cannot be detected by x rays.

DISCUSSION

We have shown that within the $Nd_{2-y}Ce_yCuO_z$ or $R_{2-x-y}R'_xM_yCuO_y$ ($R = Nd_xEu$; R' = La,Y; M = Ce,Th) systems there is a subtle and delicate balance between composition, structure and superconducting properties.



FIG. 8. High-temperature x-ray powder diffraction of $Nd_{1.85}Ce_{0.15}CuO_4$ under nitrogen ambient. The inset shows the variation of the lattice parameters *a* and *c* as a function of temperature.

For instance, we showed that for a material with an appropriate cation concentration $(Nd_{1.85}Ce_{0.15}CuO_z)$, the superconducting properties can be varied (sharp transition and high T_c to no T_c at all), by changing the oxygen content (thereby the Cu oxidation state) through processing conditions (ambient and cooling rate). We also showed that within these new high- T_c oxides, the T_c 's are independent of whether the rare earth is magnetic or not, which is just what has been found so far for all the high- T_c oxide systems.

In Fig. 9 we compare the variation of the a and c lattice parameters for the Sr-doped La system (La-Sr-Cu-O), the Ce-doped Nd system (Nd-Ce-Cu-O) and the La doped Nd system (Nd-La-Cu-O) system. Note that the lattice parameters a and c both increase in the La-doped Nd series. Within the same T' phase when Nd is replaced by Ce, a completely different behavior is observed; an increase in a and decrease in c with increasing Ce content. Finally, in the La-Sr-Cu-O system, like for the Ce-doped Nd system a and c change in opposite direction. Howev-



FIG. 9. The variation of the *a* and *c* lattice parameters are shown as a function of *x* for the $La_{2-x}Sr_xCuO_4$, $Nd_{2-x}Ce_xCuO_4$ and $Nd_{2-x}La_xCuO_4$ series.

er, a decreases and c increases when the trivalent La ions are replaced by divalent Sr ions. The variation of the caxis in these three systems can simply be understood based on the ionic radius of the dopant versus the substituant. In the T phase La is located at a nine-coordinated site. The ionic radii of a nine-coordinated La^{3+} and Sr^{2+} are 1.3 and 1.45 Å, respectively. Thus replacement of La by Sr is expected to increase the c and a axis. Only the increase in c is observed. In the T' phase, Nd is located at a eight-coordinated site and the ionic radii of eightcoordinated Nd³⁺ and Ce⁴⁺ are 1.25 and 1.11 Å, respectively. Thus a shortening of both a and c is expected when Nd is replaced by Ce. We find that c decreases but a increases. Finally, when Nd is replaced by La in the T'phase, one should expect an increase of the c axis and aaxis as observed. Thus based on the ionic radius, one can explain the variation of the c lattice parameter for the three systems and that of the *a* lattice parameter for the La-Nd doped system but not that of a for the Ce or Sr doped system. A key for understanding these observations is that either Ce or Sr substitution affects the electron or hole concentration, whereas La substitution does not. Thus, this charge transfer might be responsible for the apparently unexpected variation of a in the Sr-doped La- or Ce- doped Nd systems. Whangbo et al.¹⁶ have shown that the Cu—O bonds have antibonding character in the CuO₂ layer x^2-y^2 bands. The substitution of La³⁺ by Sr²⁺ corresponds to add a hole or equivalently removing an electron from the $x^2 - y^2$ bands. Thus the removal of an electron from the antibonding band orbital would shorten the in-plane Cu-O bond and first the in-plane Cu-Cu distance (i.e., the a axis) as experimentally observed. In contrast, doping with Ce corresponds to adding electrons into the antibonding band orbital and thereby lenghtening the in-plane Cu-O bond and the Cu-Cu distance as observed. Therefore, the variation of the lattice parameters is quite consistent with the fact that the Tphase is doped by holes (p type) and that the T' phase is doped by electrons (*n* type).

In the La-Sr-Cu-O system, superconductivity can be achieved either by divalent dopant or oxygen uptake and T_c is maximum when the formal valence of Cu³⁺ (i.e., holes) is close to 2.15. For the Nd-Ce-Cu-O system, Ce doping and oxygen removal (both of which donate electrons to the CuO₂ layers) are required to promote superconductivity. If there is a critical electron concentration to induce superconductivity it is puzzling why this limit cannot be achieved by further increasing Ce doping only. We have tried, but no superconductivity is observed, without nitrogen reduction, to the limit of Ce solubility $(\cong 0.2)$. This illustrates the crucial role of the annealing treatment that removes some oxygen and thereby increases the number of carriers to reach the threshold carrier concentration. TGA analysis has shown, in agreement with the work of Moran et al.,14 that independent of the Ce content the oxygen content in the as-grown material is always greater than four. We have suggested that these extra oxygens occupy the 4d positions (i.e., apical positions). Upon nitrogen reduction, the extra oxygens are partially removed, and according to our data the Ce-doped Nd compound becomes superconductor when the formal valence of Cu is 1.85 (i.e., 0.15 electrons/Cu), which occurs when the oxygen content becomes lower than four. It is interesting to recall that within the La-Sr-Cu-O system a maximum in T_c was found for a formal valence of 2.15 for Cu (i.e., 0.15 holes). The accuracy of the TGA measurements is not sufficient to speculate whether or not the presence of apical oxygens is crucial for superconductivity. Future neutron studies should clarify this point.

For La-Sr-Cu-O, the reversible removal of 0.03 oxygen atoms per formula unit occurs in one step at temperatures close to 500 °C. For the Nd system the amount of mobile oxygen is similar (0.04 per formula unit) but its removal occurs at higher temperatures (800 °C). In both systems the changes in oxygen content and T_c are reversible.

For all the copper oxide superconductors, a correlation between T_c and the in-plane Cu-O distance was proposed.¹⁷ For instance, within the La-Sr-Cu-O system there is an optimum bond length (1.888 Å) for which T_c is maximum¹⁷ and this optimum bond length changes by slightly affecting the structure (i.e., by replacing Sr by Ba or Ca). In Nd-Ce-Cu-O system we found that a similar correlation exists, and clearly shown for the Eu system that there is a critical distance of about 1.95 Å at which T_c is optimum (16 K). Away from this optimum length, T_c is decreased. This again suggests the close similarity between the *n*-type and *p*-type and superconductors. In the $Eu_{1.85-x}La_xCe_{0.15}CuO_4$ system, substitution of La^{3+} for Eu^{3+} does not change the electron concentration, nevertheless its T_c versus Cu-Cu correlation shows a maximum. The only thing that make sense is that the Cu—O bond strength changes as a function of x, as pointed out by Whangbo et al., 17 for the old copper oxide superconductors. We have shown that the a and c parameters are almost perfectly correlated so that there is a correlation between T_c and c, but this is just a byproduct of the in-plane Cu-Cu distance correlation. The replacement of Eu by larger ions such as La can, to a certain extent, be viewed as a chemical pressure effect. Thus one would expect hydrostatic pressure to lead to the same effect. In fact recent studies¹⁸ on the pressure dependence of T_c for several T' phases $M_{1.85}$ Ce_{0.15}CuO₄ have shown that a maximum in T_c is observed with increasing pressure for Eu_{1.85}Ce_{0.15}CuO₄. By increasing the pressure the Cu-O distance (i.e., half of Cu-Cu distance) is changed, and at a certain pressure its bond strength reaches the critical value, thereby leading to a maximum T_c .

The optimum bond distance for T_c in the La-Sr-Cu-O and Nd-Ce-Cu-O systems are 1.88 and 1.95 Å, respectively. Thus one might expect for compounds of the T or T' phase, with Cu-O distances close to the critical values, to exhibit superconductivity if one could find a chemical substitution which will optimize the in-plane Cu—O bond length. In this respect, the Gd₂CuO₄ system which has a Cu-O distance close to the optimal one is an interesting candidate. Experiments are presently in progress.

From this study it appears that the *n*- or *p*-type systems behave similarly from a chemical point of view. The only difference is that the nature of the carriers is different (electrons and not holes). The sign of the Hall coefficient, R_H , which so far has been found to be extremely sample sensitive, ^{7,19} with either positive or negative values, still raises some questions. Only better materials will help in solving this issue.

ACKNOWLEDGMENTS

We wish to thank B. G. Bagley, J. Barner, P. F. Miceli, R. Ramesh, J. M. Rowell, J. H. Wernick, and M.-H. Whangbo for valuable discussions.

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