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# PHYSICAL PROPERTIES OF Nd<sub>1.85</sub>Ce<sub>0.15</sub>CuO<sub>4-y</sub> AT HIGH PRESSURE AND WITH DIFFERENT OXYGEN CONTENTS

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We have measured initial susceptibility on Nd<sub>1.85</sub>Ce<sub>0.15</sub>CuO<sub>4-y</sub> with y varing from 0.02 to 0.32. T<sub>c</sub> was quite insensitive to the different oxygen contents. The pressure dependence  $dT_c/dp$  for a compound with y = 0.03 was determined to be -0.04 K/kbar.

## Introduction

Since the discovery of electron-doped superconductivity in high-T<sub>c</sub> cuprates [1], each model on high-T<sub>c</sub> superconductivity has had to suffer a checking whether it is symmetric upon doping of either holes or electrons. A single – band model can be thought of to accomplish this symmetry. The theories based on holes with spin S = 1/2 fail to explain superconductivity with spin-holes on the Cu<sup>1+</sup> ions. However, from the beginning Wachter and Degiorgi [2] and de Jongh [3] have proposed a polaronic model in which they postulated a singlet state of the Cu<sup>3+</sup> ion. The S = 0 state of Cu<sup>3+</sup> had been an issue but now from the NdCeCuO-compounds it turns out anambigiously that it is a spin-hole responsible for superconductivity.

# Experiment, Results

We have prepared the NdCeCuO compound by mixing appropriate amounts of CeO<sub>2</sub>, Nd<sub>2</sub>O<sub>3</sub> and CuO to achieve a composition of Nd<sub>1.85</sub>Ce<sub>0.15</sub>CuO<sub>4</sub>. The mixed oxides were calcined in air at 950° C for 10 hours, pressed into pellets and sintered in air at 1150° C for 12 hours. The samples were quenched in air to room temperature. The as prepared material was not superconducting. Then the samples were placed into a Perkin–Elmer thermo–analyzer and quantitative amounts of oxygen were removed by heating up to 1140° C in an atmosphere of pure argon gas for different times. When the samples had reached the desired oxygen content they were quenched with a cooling rate of 10° C/min. Beginning from an oxygen deficiency of y = 0.02 per formula unit superconductivity was observed with T<sub>c</sub> near 23 K. Up to y = 0.22 the samples were single phase with a T<sub>c</sub> not changing appreciably. Typical curves of the ac–susceptibility are shown in Fig. 1. The sample with y = 0.32 was investigated by X–ray diffraction to have two phases. Probably one of them (nonsuperconducting) is responsible for the magnetic response at 8 K. The lattice parameters of a specific superconducting compound with y = 0.04 were a = 3.947(± 0.001)Å and c = 12.08(± 0.002)Å.

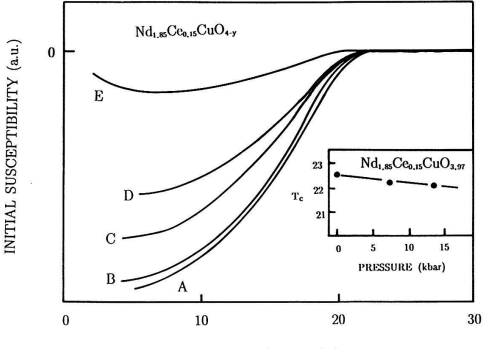
Further we have measured the pressure dependence of  $T_c$  up to 15 kbar as described in a previous paper [4]. There was only a weak pressure dependence of  $dT_c/dp = -0.04$ K/kbar (see inset of Fig.1).

# Discussion

It is amazing that the variation of the oxygen content does not influence the transition temperatur. A too large loos of oxygen (y > 0.22) results in a transformation to a red-brown phase. We believe that the outgoing oxygen comes from the (Nd,Ce)O-planes between the superconducting  $CuO_2$  sheets. As Ce can exist in two valence states the charge balance can be achieved by the transition  $Ce^{4+} \rightarrow Ce^{3+}$ . Thus, the  $CuO_2$  plane is stable and even more two-dimensional than in other superconducting cuprates.

Let us look at the pressure dependence. Assuming a band model, pressure causes a broadening of the bandwidth. However, all cuprates have more or less the same superconducting  $CuO_2$ -planes and thus a similar band structure. Therefore, upon applying pressure, they should behave in a similar way. But the pressure dependence  $dT_c/dp$  of the superconducting cuprates differs within the large range of -0.05 to 0.6 K/kbar which is not very consistent with the band-picture.

We suggest a model of exchange coupled bipolarons as described by Wachter and Degiorgi [2]. In this model the concentration of polarons (i.e. the concentration of  $Cu^{3+}$  resp.  $Cu^{1+}$ ) is crucial. At high pressure the apical oxygen ion is shifted towards the  $CuO_2$ -plane and can serve as a doping reservoir. The more apical O-ions available the more effective will be an applied pressure to adjust the optimal concentration. From this point of view it seems reasonable that the  $(La,Ba)_2CuO_4$  family with two apical O-atoms and the YBa\_2Cu\_4O\_8 and YBa\_2Cu\_{3.5}O\_7 [5] compounds with two chains resp. alternativly two and one chain as intercalation exhibit a strong pressure dependence. The YBa\_2Cu\_3O\_7 composition with only one apical oxygen is to show a lower  $dT_c/dp$  as is really the fact. For the NdCeCuO-compound with no apical oxygen above Cu(2), it is not surprising that we even have detected a negativ  $dT_c/dp$ .



**TEMPERATURE** (K)



Initial susceptibility of  $Nd_{1.85}Ce_{0.15}Cu_{04-y}$  with the different oxygen contents y : A: 0.03, B: 0.07, C: 0.02, D: 0.21, E: 0.32. The inset shows the pressure dependence of  $T_c$  of  $Nd_{1.85}Ce_{0.15}Cu_{03.97}$ .

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- [5] The pressure dependence of  $YBa_2Cu_{3.5}O_7$  was determined to be  $dT_c/dp = 0.45 \text{ K/kbar}$  (to be published elsewhere)