High- T_c superconductivity in three-fluorite-layer copper oxides. II. (Cu,Mo)Sr₂(Ce,Y)₃Cu₂O_{11+ δ}

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We have synthesized high-quality samples of the s=3 member of the (Cu,Mo)-12s2 homologous series of layered copper oxides. The (Cu,Mo)-1232 phase forms as a single phase for the cation composition (Cu_{0.75}Mo_{0.25})Sr₂(Ce_{0.67}Y_{0.33})₃Cu₂O_{11+ δ} through solid-state synthesis in air, but the samples as synthesized are not superconductive. Superconductivity is induced in the air-synthesized samples by means of high-pressure oxygenation carried out at 5 GPa and 500°C in the presence of Ag₂O₂ as an excess oxygen source. With increasing ratio of Ag₂O₂ to the (Cu,Mo)-1232 phase the *c*-axis lattice parameter gradually decreases and *T_c* increases. The highest *T_c* value of 53 K reached for samples oxygenated in the presence of 100 mol% Ag₂O₂ exceeds the values reported for Cu-based two-fluorite-layer compounds.

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I. INTRODUCTION

The known high- T_c superconductors and related layered copper-oxide phases may all be imagined as members of two structure categories: category A comprises the phases that consist of perovskite (P) and rocksalt (RS) type layers piled up with a sequence of $AO[RS]-(MO_{1\pm\delta/m})_m[P/RS]$ - $AO[RS]-CuO_2[P]-(Q-CuO_2)_{n-1}[P]$, whereas the category B phases contain additional fluorite (F) type layers inserted between two adjacent CuO₂ planes, i.e., AO[RS]- $(MO_{1\pm\delta/m})_{m}[P/RS]-AO[RS]-CuO_{2}[P]-[B-(O_{2}-B)_{s-1}][F]-$ CuO₂[P].^{1,2} From the given layer sequences the chemical formulas of the phases of the two categories, A and B, are derived at $M_m A_2 Q_{n-1} Cu_n O_{m+2+2n\pm\delta}$ and $M_m A_{2k} B_s \operatorname{Cu}_{1+k} \operatorname{O}_{m+4k+2s\pm\delta}$, respectively, or in short, $M - m^{(A)} 2(n-1)n$ and $M - m^{(A)} (2k) s(1+k)^{2.2}$. Then, a group of phases for which the $AO-(MO_{1\pm\delta/m})_m$ -AO portion is common but the number of CuO_2 planes *n* (category A), or the number of fluorite layers s (category B), varies, comprises a "homologous series".2

The s=1 phases of category B may be equally well imagined as the corresponding n=2 phases of category A. Many of these phases are well-established superconductors such as $CuBa_2YCu_2O_{7\pm\delta}$ or $Cu-1^{(Ba)}212$ (or "Y-123"). Among the (*p*-type doped³) s=2 phases of category B, $Cu(Ba, R)_2(Ce, R)_2Cu_2O_{9\pm\delta}$ (*R*=rare-earth element) or $Cu-1^{(Ba,R)}222$ was the first to show superconductivity. The highest T_c of 43 K was achieved for an R=Eusample synthesized under ambient pressure but oxygenated under elevated O₂ pressures.⁴ For these first Cu-1222 samples with Ba as the main A-site constituent, the apparent role of the trivalent R substituent was to stabilize the Cu-1222 structure. With Sr at the A site, the Cu-1222 structure is formed even without "structure-stabilizing" substituents, viz., $CuSr_2(Ce, Y)_2Cu_2O_{9+\delta}$ ⁵ though only under high O₂ pressures. Note that at ambient pressure the $(Cu, Ce)Sr_2(Ce, Y)_2Cu_2O_{9\pm\delta}$ phase rather than $CuSr_2(Ce, Y)_2Cu_2O_{9\pm\delta}$ forms in the Cu-Sr-Ce-Y-O system.⁶ Actually the $(Cu, Ce)Sr_2(Ce, Y)_2Cu_2O_{9\pm\delta}$ phase is just an example of the (Cu, M')-1^(Sr)222 phases stabilized as a result of partial substitution of the charge-reservoir Cu either by other metal atoms, i.e., M' = Ti, V, Cr, Fe, Co, Ga,Ge, Mo, W, Re, Ce, etc.,⁷⁻¹⁰ or by oxy-anions, such as BO₃³⁻, SO₄²⁻, and PO₄^{3-,11} For the Cu-12*s*2 series the third (*s*=3) member has been stabilized as well, both in multiphase samples of CuSr₂(Ce,Ho)₃Cu₂O_{11±δ} and nearly singlephase samples of (Cu,M')Sr₂(Ce,Ho)₃Cu₂O_{11±δ} (M'=Pb,Fe,Al).¹²⁻¹⁴ None of these *s*=3 samples, however, showed superconductivity.

In the present study, by taking advantage of the highpressure oxygenation (HPO) technique previously utilized in "superconductorizing" the $CoSr_2YCu_2O_{7+\delta}$ phase,¹⁵ we have been able to induce superconductivity in $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ samples of the $(Cu,Mo)-1^{(Sr)}232$ phase. Moreover, the T_c value has been successfully controlled (in terms of the amount of Ag₂O₂ used as an oxygen source in the HPO treatment) up to 53 K—just to exceed those reported for various Cu-1222-structured compounds.

II. EXPERIMENTAL

(Cu, Mo)-1^(Sr)232 А master sample of $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ was synthesized by solid-state reaction from an appropriate mixture of highpurity powders of CuO, MoO₃, SrCO₃, CeO₂, and Y₂O₃. The mixed powder was calcined at 950°C and sintered at 1020°C in air with several intermediate grindings. Portions of this as-air-synthesized (AS) sample were then highpressure oxygenated at 5 GPa and 500°C for 30 min in a cubic-anvil-type high-pressure apparatus in the presence of Ag_2O_2 as an excess-oxygen source. The amount of Ag_2O_2 mixed with the $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ powder was systematically varied from 12.5 to 100 mol% accordingly to obtain the samples of HPO-12.5, HPO-25, HPO-50, HPO-75, and HPO-100.

Powder x-ray diffraction measurements (Rigaku: RINT2550VK/U; Cu K_{α} radiation) were performed to check the phase purity and determine the lattice parameters of the

samples. The AS and HPO-100 samples were further characterized by high-resolution transmission-electron microscopy (HRTEM) and electron diffraction (ED) techniques using an ultrahigh-voltage transmission electron microscope (Hitachi: H-1500) operated at an accelerating voltage of 820 kV. For these experiments a specimen was prepared by crushing a portion of the sample into fine fragments, which were ultrasonically dispersed in CCl_4 and transferred to carbon microgrids.

The amount of "excess" or removable oxygen was estimated for the samples on the basis of thermogravimetric [(TG); MAC Science: TG/DTA 2000S] runs carried out for $a \sim 10$ mg specimen in four steps: (1) heating in N₂ at a rate of 1° C/min up to 700°C, (2) cooling down to room temperature in N₂ at a rate of 20°C/min, (3) heating in O₂ at a rate of 1° C/min to 900°C, and (4) cooling to room temperature at a rate of 1° C/min. For the AS sample, all four steps were employed, whereas the HPO samples were investigated only up to the second step.

Superconductivity and magnetic properties were measured for all the samples down to 5 K in both fieldcooled (FC) and zero-field-cooled (ZFC) modes under 10 Oe using a superconducting-quantum-interference-device (SQUID) magnetometer (Quantum Design: MPMS-XL). Resistivity measurements were carried out using a four-pointprobe apparatus (Quantum Design: PPMS).

III. RESULTS AND DISCUSSION

From the x-ray powder diffraction patterns shown in Fig. 1 for the $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ samples, both the as-air-synthesized (AS) and the high-pressure oxygenated (HPO) samples were confirmed to be of high quality; the only visible impurity diffraction peaks were due to a trace amount of nonreacted CeO₂ (for samples in which the amount of Ag₂O₂ used as an excess-oxygen source in HPO did not exceed 75 mol%). In other words, the (Cu,Mo)-1^(Sr)232 phase was found as the only Cu-containing phase [within the detection limit of x-ray diffraction (XRD)]. Naturally each HPO sample should also contain decomposition residue(s) of Ag₂O₂, i.e., Ag and/or Ag₂O. For the HPO-12.5 and HPO-25 samples, the main diffraction peak due to Ag and/or Ag₂O is clearly distinguished. With a further increase in the amount of Ag₂O₂, rather than a gradual increase in the intensity of this peak seen is that an amorphouslike background appears in the 2θ range of $30^{\circ} - 35^{\circ}$.

The diffraction peaks originating from the (Cu,Mo)- $1^{(Sr)}232$ phase were readily indexed with space group *P4/mmm*.¹³ The expected crystal structure was further confirmed for the AS and HPO-100 samples by HRTEM and ED. In Fig. 2 shown is an HRTEM image for the HPO-100 sample. The layer repetition of SrO-(Cu,Mo)O_{1+δ}SrO-CuO₂-(Ce,Y)-O₂-(Ce,Y)-O₂-(Ce,Y)-CuO₂ that is characteristic to the (Cu,Mo)- $1^{(Sr)}232$ structure is clearly revealed. No sign of extensive stacking faults is seen. From the ED patterns (Fig. 3) taken for both of the two samples, AS and HPO-100, with the incident electron beam along the [001], [010], and [1-10] directions, it is concluded that the high-pressure oxygenation does not change the basic structure of



FIG. 1. X-ray diffraction patterns for the synthesized $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ samples. The indices given are for the (Cu,Mo)-1232 structure on the basis of space group *P*4/*mmm*.

the (Cu,Mo)- $1^{(Sr)}232$ phase. Moreover, the tetragonal unit cell with the *a*:*c* lattice parameter ratio approximately at 1:4 is confirmed from the ED data.

In Fig. 4 we show the evolution of the lattice parameters a and c upon the high-pressure oxygenation as refined from the x-ray diffraction data. With an increasing amount of Ag₂O₂ used in the high-pressure oxygenation, the *a*-axis lattice parameter seems to remain quite constant, whereas the



FIG. 2. HRTEM image for the HPO-100 sample.



FIG. 3. ED patterns for the AS (a) and HPO-100 (b) samples with the incident beam along the directions, [001], [010], and [1-10] (from the top to the bottom).

c-axis lattice parameter gets systematically shortened. The shortening of the *c* parameter occurs rather rapidly up to 25 mol% of the amount of Ag_2O_2 , then slows. This suggests that upon HPO, the oxygen content of the (Cu,Mo)-1^(Sr)232 phase increases strongly only up to 25 mol%, and beyond that only a moderate change in the oxygen content is likely to occur.

To gain a better understanding of the changes in oxygen content upon the HPO, we compared the amount of depleted oxygen, $\Delta \delta$, upon an N₂ annealing up to 700°C carried out in a thermobalance under the same conditions for both the AS and the HPO samples. First shown in Fig. 5(a) are the deoxygenation (in N₂) and backoxygenation (in O₂) TG curves for the AS sample. Upon annealing in N₂ the deoxygenation occurs in three steps with onset temperatures about 200°C, 370°C, and 420°C. Such a multistep behavior suggests that accordingly the bonding strength of the oxygen



FIG. 4. Evolution of lattice parameters, a and c, upon increasing the amount of Ag₂O₂ in the high-pressure oxygenation.



FIG. 5. TG curves for (a) N_2 annealing (solid line) and subsequent O_2 annealing (broken line), of the AS sample, and (b) for the N_2 annealing of the AS (dotted line), HPO-25 (broken line), and HPO-75 (solid line) samples.

atoms being depleted slightly changes, though not yet confirmed from a detailed crystal structure analysis. The overall weight loss corresponds to ~ 0.32 oxygen atoms per $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ formula unit, i.e., $\Delta \delta_{\rm AS} = 0.32$. Then upon cycling in O₂, oxygen is incorporated into the phase again such that the original oxygen content of the AS sample is not only completely recovered but even slightly exceeded. (This is natural, since the TG annealing was done in O_2 , not in air.) In Fig. 5(b) we show two representative TG curves recorded in N₂ for the HPO samples, i.e., HPO-25 and HPO-75, together with that for the AS sample. The total oxygen loss is much larger for the HPO samples than for the AS sample, presumably due to the higher oxygen contents of these samples in comparison to that of the AS sample. However, it should be noted that for the HPO samples the total oxygen evolution consists of contributions from both the (Cu, Mo)- $1^{(Sr)}232$ phase and Ag₂O. From previous studies^{16–18} it is well known that the sharp weight and oxygen loss about 280~330°C (only faintly distinguished for HPO-25 but profoundly seen for HPO-75) is due to the reduction of Ag_2O to Ag. We therefore subtract that from the total oxygen evolution to estimate the amount of removable oxygen in the (Cu, Mo)- $1^{(Sr)}232$ phase for the HPO-25 and HPO-75 samples at $\Delta \delta_{\text{HPO-25}} = 0.84$ and $\Delta \delta_{\text{HPO-75}} = 0.91$, respectively. Now comparing the values of $\Delta \delta_{AS}$ (=0.32), $\Delta \delta_{HPO-25}$ (=0.84), and $\Delta \delta_{\text{HPO-75}}$ (=0.91) it is revealed that upon the high-pressure oxygenation the oxygen content of the (Cu, Mo)-1232 phase indeed increases considerably, i.e., by 0.84-0.32=0.52 oxygen atoms for HPO-25 and 0.91-0.32=0.59 for HPO-75. From Fig. 5(b) it may also be concluded that the onset temperatures for the oxygen-loss steps are shifted to somewhat lower temperatures for the HPO samples in comparison to those for the AS sample as an indication of weaker bonding



FIG. 6. Temperature dependence of magnetization for the AS sample (upper) and the HPO-50 sample (lower).

in the HPO samples. Moreover, it should be noted that taking an analogy to the situation seen for the related Co-1^(Sr)232 phase¹⁹ it is appropriate to assume that the oxygen-content variation in (Cu,Mo)-1^(Sr)232 occurs within the (Cu_{0.75}Mo_{0.25})O_{1+ δ} layer only.

Magnetization measurements were carried out for all the samples. The AS sample did not show any sign of superconductivity (see Fig. 6). For the HPO samples superconductivity appeared when the amount of Ag_2O_2 (used as an oxygen source) was 25 mol% or higher. The volume fraction of superconductivity (at 5 K) reaches 26% for the HPO-50 sample. The ZFC and FC curves for this sample are shown in Fig. 6. The onset of the diamagnetic signal is seen at 53 K. Figure 7 displays the resistivity (ρ) versus temperature (T) curve for the same sample that shows a clear down turn in resistivity about 50 K. The fact that the zeroresistivity is not achieved is attributed to the apparent existence of partly amorphous decomposition residue(s) of Ag_2O_2 at the grain boundaries of the superconductive (Cu, Mo)-1^(Sr)232 grains. To determine the superconductivity transition temperature properly from the ρ -T curve that is a superposition of the superconductive- and semiconductive-type behaviors, we



FIG. 7. Temperature dependence of resistivity for the HPO-50 sample.



FIG. 8. T_c values for the present (Cu,Mo)-1^(Sr)232 samples plotted against the amount of Ag₂O₂ used for the high-pressure oxygenation.

employed $d^2\rho/dT^2 \approx 0$ as a criterion for the T_c . This yielded a value of ~54 K in good agreement with the one from the magnetization measurement.

In Fig. 8 we plot the value of T_c against the amount HPO for of Ag_2O_2 used in the our $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ samples of the (Cu, Mo)-1^(Sr)232 phase. It is seen that T_c first increases strongly with an increasing amount of Ag₂O₂ and then saturates beyond \sim 50 mol%. Here one should recall that a similar saturation trend was seen for the *c*-axis lattice parameter (cf. Fig. 4). Also from the TG data we concluded that upon increasing the amount of Ag₂O₂ the amount of excess oxygen incorporated in the (Cu, Mo)-1^(Sr)232 structure increased rather fast initially but then more slowly. Thus it seems that there is a certain limit for the phase to incorporate excess oxygen, at least under the conditions presently used for the oxygen loading. Assuming that the *c*-axis lattice parameter provides us with a reasonable measure of the amount of excess oxygen in (Cu, Mo)-1^(Sr)232 and thus the overall holedoping level, we plot the value of T_c against it in Fig. 9. For comparison, also given in Fig. 9 are similar data for the first two members of the (Cu,Mo)-1^(Sr)2s2 homologous series: filled circles for our $(Cu_{0.75}Mo_{0.25})Sr_2YCu_2O_{7+\delta}$ (s=1) and $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.25}Y_{0.75})_2Cu_2O_{9+\delta}\,(s\!=\!2)$ samples additionally synthesized and high-pressure oxygenated under conditions essentially identical to those for the $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$ samples of the s=3 phase and open circles for data reported by Ono⁸ for $(Cu_{0.8}Mo_{0.2})Sr_2YCu_2O_{7+\delta} (s=1)$ and $(Cu_{0.8}Mo_{0.2})Sr_2(Ce_{0.375}Y_{0.625})_2Cu_2O_{9+\delta} (s=2)$ samples. It is seen that for all the three phases of the (Cu,Mo)-1^(Sr)2s2 series T_c increases with decreasing c-axis parameter and increasing oxygen content. None of the three phases shows overdoped-type behavior or even saturation for the T_c value with decreasing *c*-axis length and increasing oxygen content. This suggests that if the amount of oxygen could be increased further (through improvements in the HPO technique) or if the hole-doping level could be increased by some other means, e.g., control over the aliovalent Y^{III}-for-Ce^{IV} substitution level,²⁰ it might be possible to increase the T_c 's of all the three phases. Unfortunately for the presently studied $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{1-x}Y_x)_3Cu_2O_{11+\delta}$ system, our pre-



liminary efforts to tune the Y content from that of x=0.33 all ended up with multiphase samples. From the comparison of the maximum T_c values among the three members of the (Cu,Mo)-1^(Sr)2s2 homologous series (Fig. 9), we conclude that even though the values reached for the higher members are considerably lower than those obtained for the s=1 phase (88 K), the trend of decreasing T_c with increasing s, i.e., thickness of the fluorite-structured layer-block separating the two CuO₂ planes, does not continue beyond s=2, but rather the T_c value recorded here for the s=3 phase (53 K) slightly exceeds the one previously reported for the s=2 phase (50 K).⁸ We also acknowledge that the purely Mo-based $MoSr_2RCu_2O_{7+\delta}(s=1)$ and $MoSr_2(Ce, R)_2Cu_2O_{9+\delta}(s=2)$ phases were recently high-lighted due to their interesting interplays between magnetism and superconductivity.^{21,22} So far the MoSr₂(Ce, Y)₃Cu₂O_{11+ δ} (s=3) phase has not been synthesized.

Finally we emphasize that the successful induction of high- T_c superconductivity in the three-fluorite-layer copperoxide phases,²³ *M*-1232, potentially provides us with possibilities in terms of "intrinsic atomic-layer engineering".²⁴ Generally—as a consequence of its layered crystal structure—a high- T_c superconductor may be considered as a stack of intrinsic superconductor-insulator-superconductor Josephson junctions.^{25,26} Now we have superconductors for which the spacing (insulating) block between two superconductive CuO₂ planes (i.e., basal planes of the CuO₅ squarepyramid layers) has been extended from those of the single FIG. 9. T_c values for differently oxygenated samples of the three first members of the (Cu,Mo)-1^(Sr)2s2 homologous series as plotted against the *c*-axis lattice parameter: filled circles for presently synthesized samples, open circles for samples reported in Ref. 8.

R-cation layer in *M*-1212 and the "double-fluorite-layer" block in *M*-1222 to the 8-9 Å-thick "triple-fluorite-layer" block in *M*-1232.

IV. CONCLUSION

The 1232-structured phase with Cu as the main chargereservoir constituent had previously been detected in multiphase samples only. In the present work samples of the phase were obtained in essentially single-phase form for the composition, $(Cu_{0.75}Mo_{0.25})Sr_2(Ce_{0.67}Y_{0.33})_3Cu_2O_{11+\delta}$, through ambient-pressure solid-state synthesis. Moreover, we successfully induced bulk superconductivity in the (Cu,Mo)- $1^{(Sr)}232$ samples by loading the phase with ~0.5 additional oxygen atoms by means of a high-pressure-oxygenation technique. The highest T_c value of 53 K reached in this work for the (Cu,Mo)- $1^{(Sr)}232$ phase exceeded the value previously reported for the (Cu,Mo)- $1^{(Sr)}222$ phase [which has a shorter inter-CuO₂-plane distance than the (Cu,Mo)- $1^{(Sr)}232$ phase].

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- ¹T. Wada, A. Ichinose, H. Yamauchi, and S. Tanaka, J. Ceram. Soc. Jpn. **99**, 420 (1991).
- ²M. Karppinen and H. Yamauchi, Mater. Sci. Eng., R. **26**, 844 (1999).
- ³Some of the category B copper oxides show *n*-type superconductivity.
- ⁴H. Sawa, K. Obara, J. Akimitsu, Y. Matsui, and S. Horiuchi, J. Phys. Soc. Jpn. **58**, 2252 (1989).
- ⁵A. Ono, Jpn. J. Appl. Phys., Part 1 **32**, L1599 (1993).
- ⁶A. Ono and S. Horiuchi, Physica C **216**, 165 (1993).
- ⁷T. Wada, A. Nara, A. Ichinose, H. Yamauchi, and S. Tanaka,

Physica C 192, 181 (1992).

- ⁸A. Ono, Jpn. J. Appl. Phys., Part 1 **32**, 4517 (1993).
- ⁹M. Tamura, M. Sato, T. Den, and J. Akimitsu, Physica C **303**, 1 (1998).
- ¹⁰N. C. Hyatt, G. B. Peacock, I. Gameson, K. L. Moran, M. Slaski, M. O. Jones, A. J. Ellis, Y. E. Gold, R. Dupree, and P. P. Edwards, Int. J. Inorg. Mater. 1, 87 (1999).
- ¹¹P. R. Slater, C. Greaves, and M. Slaski, Physica C 235-240, 741 (1994).
- ¹²T. Wada, A. Ichinose, H. Yamauchi, and S. Tanaka, Physica C 171, 344 (1990).
- ¹³T. Wada, A. Ichinose, F. Izumi, A. Nara, H. Yamauchi, H. Asano, and S. Tanaka, Pure Appl. Chem. **179**, 455 (1991).
- ¹⁴H. W. Zandbergen, T. Wada, A. Nara, H. Yamauchi, and S. Tanaka, Physica C 183, 149 (1991).

- ¹⁵Y. Morita, H. Yamauchi, and M. Karppinen, Solid State Commun. 127, 493 (2003).
- ¹⁶T. Ito, H. Suematsu, K. Isawa, M. Karppinen, and H. Yamauchi, Physica C **308**, 9 (1998).
- ¹⁷M. Karppinen, H. Yamauchi, Y. Morita, M. Kitabatake, T. Motohashi, R. S. Liu, J. M. Lee, and J. M. Chen, J. Solid State Chem. **177**, 1037 (2004).
- ¹⁸M. Karppinen and H. Yamauchi, in *International Book Series: Studies of High Temperature Superconductors*, Vol. 37, edited by A. V. Narlikar (Nova Science Publishers, New York, 2001), pp. 109–143.
- ¹⁹Y. Morita, M. Karppinen, and H. Yamauchi (unpublished).
- ²⁰T. Watanabe, M. Karppinen, T. Motohashi, T. Nagai, Y. Matsui, and H. Yamauchi, preceding paper, Phys. Rev. B **70**, 174514 (2004).

- ²¹I. Felner and E. Galstyan, Phys. Rev. B **69**, 024512 (2004).
- ²²I. Felner and E. Galstyan, Phys. Rev. B 68, 064507 (2003).
- ²³ In two published works [Chen Xianhui, Ding Zhongfen, Qian Yitai, Chen Zuyao, and Chen Zhaojia, Phys. Rev. B **48**, 9799 (1993); Y. Akagi, H. Sasakura, S. Noguchi, T. Oka, M. Adachi, and S. Tsukui, Physica C **388-389**, 387 (2003)] possible super-conductivity in an *M*-1232 type phase has been claimed and/or mentioned. However, both the works lack the most crucial experimental evidence required to verify bulk superconductivity of the given phase.
- ²⁴S. Ikegawa, K. Nakayama, Y. Motoi, and M. Arai, Phys. Rev. B 66, 014536 (2002).
- ²⁵R. Kleiner, F. Steinmeyer, G. Kunkel, and P. Müller, Phys. Rev. Lett. 68, 2394 (1992).
- ²⁶R. Kleiner and P. Müller, Phys. Rev. B 49, 1327 (1994).