**Interfacial-Redox-Induced Tuning of Superconductivity in YBa2Cu3O7-δ**

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Many of the properties of the high- copper oxide superconductors are strongly influenced by charge doping1,2. The ability to control the doping level in these materials is therefore important not only for the development of experimental platforms that enable studies of correlated electron physics, but for multifunctional device applications as well. While traditionally3–6 the doping level is fixed during synthesis via chemical substitution or post-growth annealing, recently several techniques have demonstrated approaches which allow for on-demand control. By leveraging electrolytic double layer techniques, gating experiments7,8 on RBa2Cu3O7-δ (R=Y, Nd) thin films have achieved control over the Cu-site doping level, which determines the dominant electronic order, by introducing oxygen vacancies into the film under electric fields. In these materials, O2- ions can migrate under the influence of an externally applied electric field to eventually escape through the film surface, resulting in the formation of oxygen vacancies. To maintain charge neutrality electrons are returned to the Cu ions within the film, resulting in a reduction in Cu valence and reducing the hole concentration of the oxide. Signatures of the effects on the electronic order in these materials are profound, and point to the effectiveness of oxygen migration and vacancy formation in manipulating the properties of the cuprate superconductors. Recently, we demonstrated a different, solid-state approach to manipulate ionic distributions in oxide thin films by using a reactive Gd capping layer and thus achieve ionic control of magnetism.9–11 Leveraging the reactivity of Gd, these capping layers can extract oxygen from an adjacent oxide film, with the level of oxygen depletion controlled by the thickness of the Gd and the ion mobility in the oxide, often at room-temperature. Superconducting YBa2Cu3O7-δ (YBCO) is a prototypical example of the high- cuprates, with crystal structure and electronic ordering sensitive to oxygen stoichiometry. Combined with its high ionic conductivity,12,13 these properties make it an ideal candidate material to explore with this method.

In this study Gd capping layers of varying thicknesses are deposited on 100 nm thick YBa2Cu3O7‑δ films, dramatically altering the oxygen distribution throughout the underlying film without any annealing. As the Gd layer thickness () is increased the YBCO layer becomes progressively more oxygen deficient, demonstrating how appropriate tuning of can precisely control the remaining oxygen content of the underlayer. The extraction of oxygen from the YBCO induces a structural transition and augments the superconductivity by reducing and broadening the superconducting transition temperature . Superconductivity is completely extinguished for the sample with thickest capping layer – a corresponding change in hole doping level from optimal doping in the as-grown film to less than 0.05 holes/Cu. X-ray absorption spectra indicate a reduction of Cu valence state consistent with the removal of oxygen from the YBCO film, particularly from within the CuO chains. Since superconductivity in YBCO relies on charge transfer from the CuO chains to the CuO2 planes, disruption of the chains is likely the root cause of the disappearance of superconductivity.

Films of YBa2Cu3O7-δ (100 nm) were grown on (001) SrTiO3 (STO) substrates using pulsed laser deposition [CITATION: IEEE Trans. Appl. Supercond. 11 3209 (2001)]. The high-symmetry STO substrate (cubic, lattice parameter of 3.905 Å) is expected to grow with a degenerate epitaxy, forming 90° twinned domains [CITATION: Journal of Applied Physics 79, 1829 (1996)], with up-to 2% tensile strain exerted at the interface due to lattice mismatch [CITATION: Phys. Rev. B 39, 12355 (1989), Appl. Phys. Lett. 76, 3469 (2000)] (bulk lattice parameters of YBCO is exptected to be *a* = 3.827 Å, *b* = 3.893 Å, and *c* = 11.70 Å [CITATION Nature 328, 13 (2987)]). While the strain is known to suppress *TC*, the films are expected relax away from the substrate/film interface [CITATION: *Appl. Phys. Lett.* **72,** 2966–2968 (1998), Phys. Rev. B 57, 6173 (1998)] and show near bulk-like superconducting behavior [CITATION: Appl. Phys. Lett. 76, 3469 (2000)]. The films were then sputter coated with Gd layers of varying thickness ( 3 nm, 7 nm, 20 nm) and an Au (5 nm) protective cap, with one film kept in the as-grown state for comparison. High resolution X-ray diffraction (XRD) scans of the as-grown film, measured along the out-of-plane direction (Fig. 1) show only the family of peaks, confirming the epitaxial growth and giving a *c*-axis lattice parameter of 11.6780 ± 0.0004 Å (stat), similar to the bulk values [CITATION Nature 328, 13 (2987) and/or others]. The slightly smaller *c*-axis lattice parameter may be due to the tensile strain from the substrate [CITATION Appl. Phys. Lett. 91, 172509 (2007), or our LSCO paper, eventually]. As is increased, a monotonic shift towards lower is observed in the YBCO diffraction peaks indicating expansion in the *c*-axis. Similar lattice expansion has previously been noted in other perovskite systems14 and is a signature of oxygen depletion. In addition to the out-of-plane direction, the in-plane structure of the films was probed using reciprocal space maps (RSMs) taken near the STO substrate reflection (Fig. 2). Coordinate axes of the RSMs identify the *h* (x-axis) and *l* (y-axis) Miller indices relative to the STO substrate. When grown on cubic STO substrate, nominally orthorhombic YBCO is expected to form a twinned crystal structure, resulting in a splitting of the in-plane Bragg reflections [CITATION: Phys. Rev. B 39, 12355 (1989)]. The RSMs show a bright substrate peak near the top of each map, with a lower-intensity YBCO peak near , elongated in the in-plane direction, and consistent with the presence of two overlapping Bragg reflections from and YBCO crystal planes. Importantly, the spread of the YBCO film peak along the direction, even in the as-grown sample, indicates the films were relaxed in-plane and possess the expected orthorhombic distortion before the Gd capping layers were deposited. As is increased, the film peak broadens along the (00*l*) direction and shifts to lower , consistent with an expansion in the *c*-axis lattice parameter as seen in the symmetric scans discussed above. In addition to simple lattice expansion with increasing , a second broader feature emerges at lower angles alongside the peaks, particularly visible in the scans, associated with the optimally doped phase (OP) for nm. This secondary, oxygen-deficient (OD) phase becomes dominant for nm.

The Gd capping layer is expected to extract oxygen from the YBCO, resulting in the observed structural changes observed in XRD patterns. The extraction of oxygen was further probed by polarized neutron reflectometry (PNR), which provides a depth-resolved mapping of the nuclear scattering centers within the film. The converged depth profiles (Fig. 3a) confirm the expected 100 nm thickness of the as-grown structure, while the nuclear scattering length density (SLD, ) of the as-grown film is similar to the calculated value for YBa2Cu3O7 of . With increasing the YBCO layer increases in thickness, in agreement with the unit cell expansion observed in XRD. Commensurate with the progressive increase in thickness, the nuclear SLD decreases, particularly near the YBCO/Gd interface. The decrease of is consistent with the removal of oxygen from the YBCO. The suppression of is likely a signature of disorder near the sample surface, as has been reported15 in other YBCO thin films. With increasing the features in the SLD profiles corresponding to the protective Au layer at the top surfaces of each Gd-capped film (features a, b, c) broaden and decrease in height, indicative of interfacial roughening of the underlying layers. as the oxygen migration through the underlying films increasingly disrupts the structure. In addition to the standard nuclear profile shown in Fig. 3a, the Gd layer is explicitly identifiable by the imaginary component of its SLD (Fig. 3b), which corresponds causes neutron absorption. The imaginary component of the Gd SLD allows us to rule out the possibility of YBCO/Gd interdiffusion.

While the PNR results are consistent with the extraction of oxygen from the YBCO, fluorescence yield (FY) measurements of the X-ray absorption spectra (XA) performed at the Cu *L*2,3-edges directly confirm a change in the Cu valence resultant from the oxygen extraction. The XA result (Fig. 4) show a shift in the absorption resonance to lower energies with increasing *tGd*.. Similar spectral shifts reported in other oxygen-deficient perovskite systems16,17 have been attributed to a decrease in the average Cu valence, the result of electrons returning to the Cu ions as oxygen is leached from the film. The shoulder at eV in the as-grown YBCO spectra, a feature characteristic of lower-valence ligand states present in CuO chains8,18–20, is suppressed for greater , confirming the loss of oxygen within the chains. A second resonance associated with the Cu1+ valence state emerges at eV for 7 nm and 20 nm. This new absorption peak, appearing in conjunction with the emergent low angle feature observed in XRD, again points to the formation of a stable OD phase. The bulk sensitivity of FY measurements (70% X-ray transmission through 100 nm of YBCO at Cu-*L*2 resonance) infers that the observed oxygen depletion comes from ionic migration from deep within the film despite the interfacial origin of the leaching effect. The high ionic conductivity required for such long-range oxygen migration is in agreement with previous reports.12,13 The known sensitivity of the YBCO superconductivity to the oxygen stoichiometry suggests that this approach may therefore be used as a mechanism to design the superconducting transition.

To examine the effects of Gd deposition on the YBCO superconducting properties, magnetometry and Van der Paw resistivity measurements were performed between 5 K - 100 K. Zero-field cooled (ZFC) measurements of the magnetic moment (Fig. 5a, b) show a sharp phase transition in the as-grown YBCO film at K, typical of YBCO films grown on STO substrates21, indicating rejection of the magnetic field due to the Meissner effect, and indicating the superconducting transition. Accompanying the magnetic transition is a precipitous drop in resistivity, further confirming the superconducting transition. With increasing Gd capping layer thickness the magnetic transition shows a smooth reduction in the transition temperature , with complete suppression of the Meissner effect for nm, implying a reduction of hole doping level from optimal doping to below holes/Cu. In contrast to the magnetometry, resistivity measurements initially show only a small shift toward lower for nm. However, for nm and nm the superconducting transition is completely suppressed, with no apparent transition down to the lowest measured temperature (5 K).

The differences between the superconducting transitions observed in magnetometry and resistivity can be explained by the presence of a low-, high-resistivity OD phase stabilized alongside the OP phase YBCO film. Specifically, for the as-prepared and nm samples, a majority of the film possesses the initial structure and nominal stoichiometry of YBa2Cu3O7-δ where 0 < d< 0.5. By comparison, the nm sample shows a transition in the magnetometry, but not in the resistivity. The magnetometry signal arrises from the Meissner effect and will scale directly with the superconducting volume fraction of the film, while the superconducting transition in the resistivity necessitates a continuous superconducting pathway to exist through the sample. Therefore, the nm sample possesses some fraction which is still superconducting, as evidenced by the magnetometry, but is beyond the percolation limit for the resistive OD phase, as evidenced by the absence of a transition in the resistance. Finally, for the thickest nm sample the the absence of a transition in either the resistance or magnetic data suggests few, if any, domains remain which undergo a superconducting transition. The distinctly different superconducting properties of the OD phase may be the result of structural changes - including ordering or strain - or electron doping, both of which are consequences of the oxygen leaching; the XRD shows emergent new peaks in the nm and 7 nm samples, suggesting oxygen vacancy ordering, resulting in a structural phase transtion, may motivate the superconducting transition. Electron microscopy of the reduced film is used to confirm the local atomic ordering.

[STEM]

In summary, Gd capping layers deposited at room temperature on optimally doped YBCO thin films have been shown to remove oxygen from the underlying film via an interfacial redox reaction, with the amount of oxygen removed dependent on capping layer thickness. The superconducting transition temperature is significantly reduced, and for sufficient thickness of Gd is extinguished completely. X-ray spectroscopy measurements indicate the CuO planes become progressively more oxygen deficient with increasing Gd thickness, pointing to the reduction in hole doping level as the mechanism behind the suppression of superconductivity. Remarkably, the changes to the superconducting properties throughout the entire 100 nm thick YBCO films are induced by the migration of oxygen towards the YBCO/Gd interface, extending the viability of this approach for controlling the hole-doping level to the quasi-bulk regime.

**Supplementary Information**

**Experimental Methods**. Commercially available 100 nm thick YBCO films grown on STO substrates were purchased and subsequently sputter coated with Gd (3 nm, 7 nm, 20 nm) and a Au (5 nm) protective capping layer using Ar gas at Pa working pressure in a chamber with a base pressure of Pa. XRD characterization, including both symmetric scans and reciprocal space maps, was performed on a X-ray diffractometer equipped with parallel beam optics and Cu monochromator. Polarized neutron reflectometry was measured at the NIST Center for Neutron Research on the PBR and MAGIK beamlines. The experiment used 5 Å neutrons, and were carried out at a temperature of 6 K. Fitting of the PNR data was performed using the Refl1d software package, following a Markov-chain Monte Carlo fitting algorithm [CITATION: Curr. Opin. Colloid. Interface Sci. 17, 44–53 (2012)]. The calculated SLD was determined by the calculating the sum of the volume-scaled atomic scattering lengths. XA measurements were performed at the Advanced Light Source on beamline 4.0.2 at room temperature in a grazing incidence (30°) geometry. Both fluorescence and electron yield (EY) modes were measured, but due to the capping layers no appreciable signal was measured in the EY mode. Magnetometry measurements, FC and ZFC, were performed in a field of 1 mT. To reduce stray magnetic fields, the magnet was purged before each measurement. Resistivity was measured using a four-contact Van der Pauw geometry. The current used was at frequency of 173 Hz. The magnetic field at the sample was set to zero before the measurement by setting the temperature to the midpoint of the superconducting transition before adjusting the applied field to minimize the resistivity.

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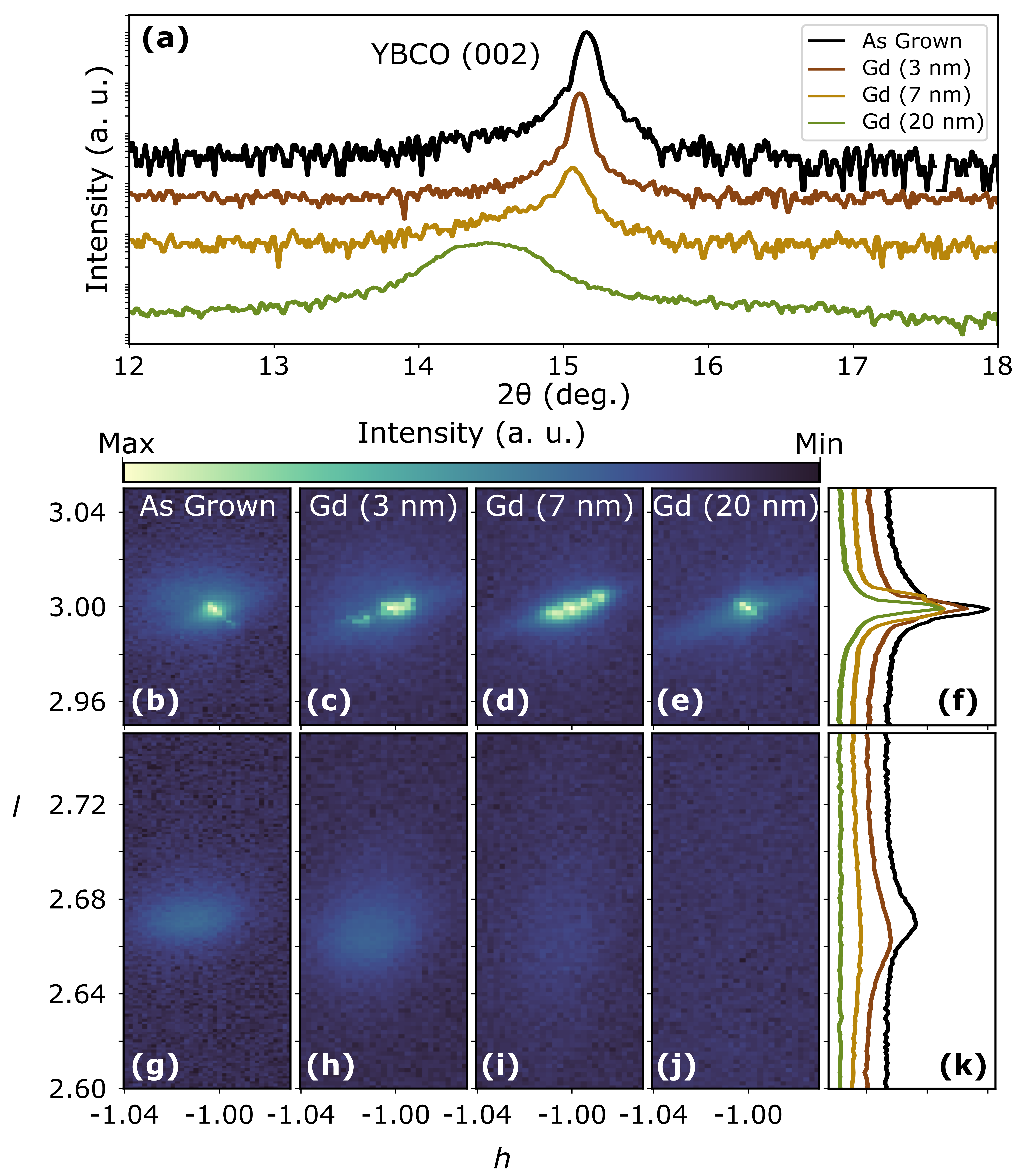
**Figures**

**Figure 1**. XRD scan (a) near the (002) YBCO peak measured with Cu radiation. Reciprocal space maps (b-e) showing the ( STO substrate peak at top, with ()/() YBCO peak at bottom (g-j), and projections (f, k) of each map along the -direction.

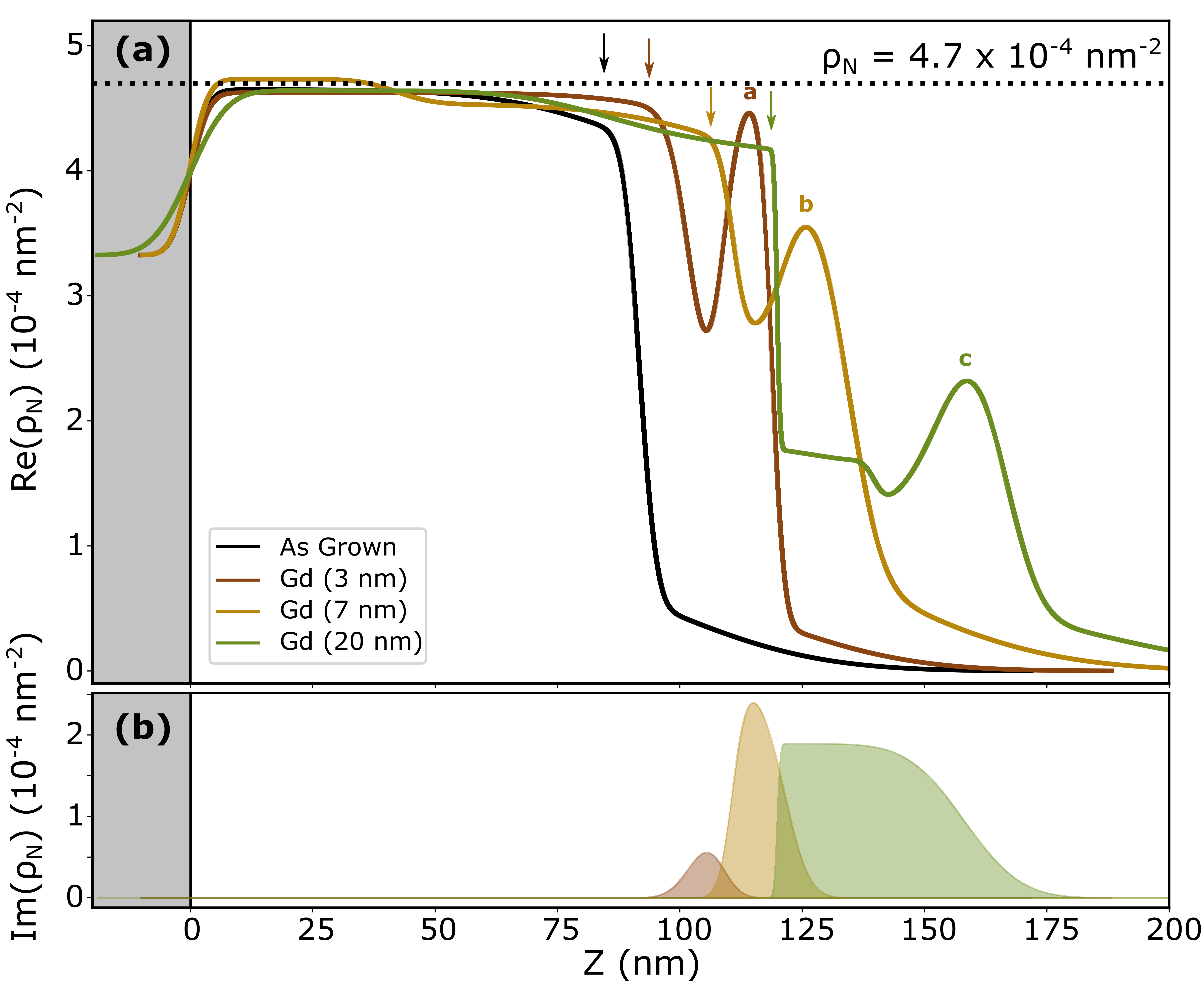
**Figure 2**. Real (a) and imaginary (b) part of the nuclear SLDs as a function of depth through the sample as measured by PNR. Shaded region () corresponds to the substrate, with () corresponding to the film. Arrows denote the top surfaces of the YBCO layers, identified by the steep drop in Re(. The peaks in Re() labeled a, b, c mark the locations of the Au capping layers.

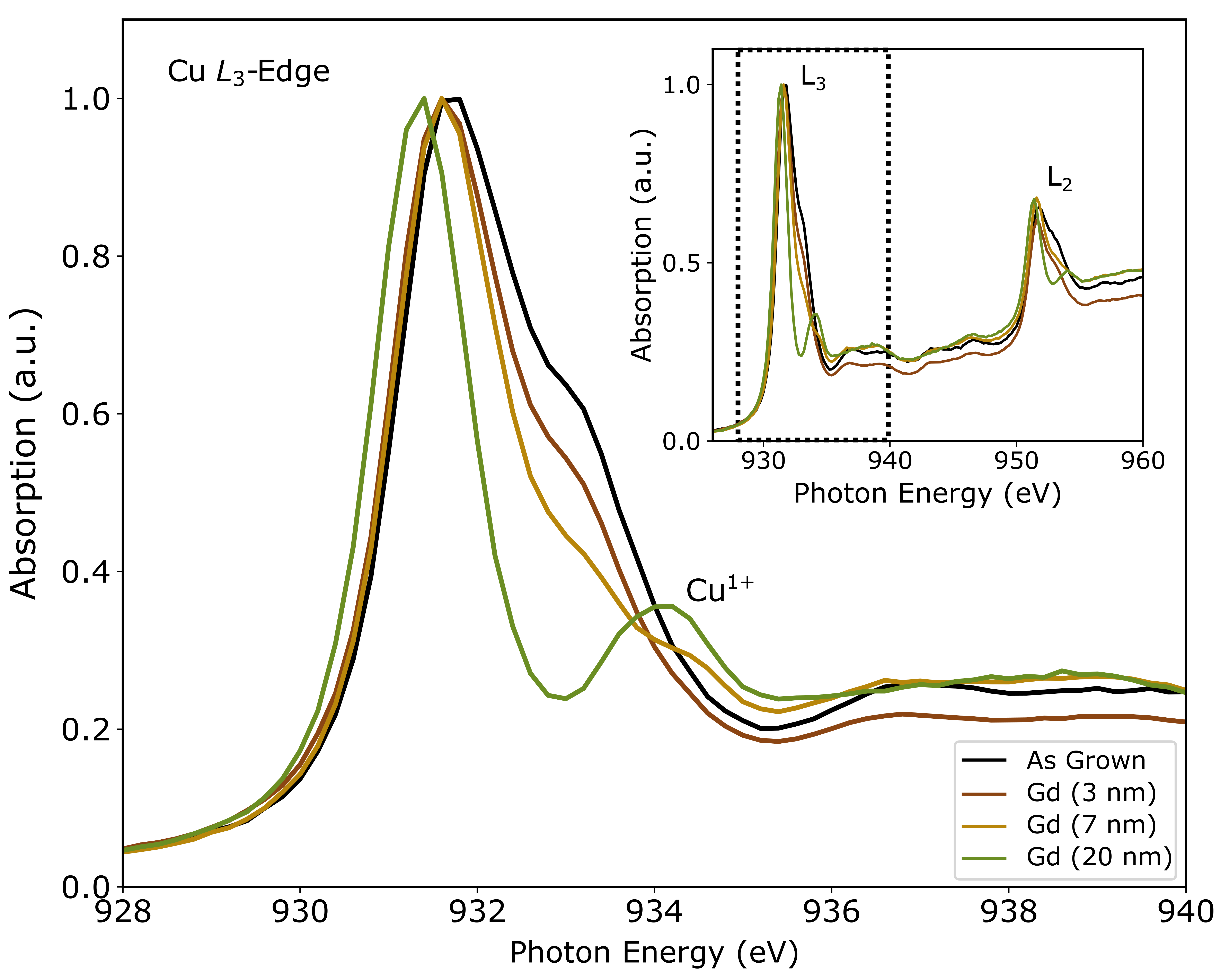
**Figure 3**. Close-up view of the normalized absorption spectra as a function of incident X-ray photon energy near the Cu L3-edge, as measured in fluorescence yield mode. The full spectral range measured including the Cu L2-edge is shown in the inset, with the shaded region corresponding to the close-up view.

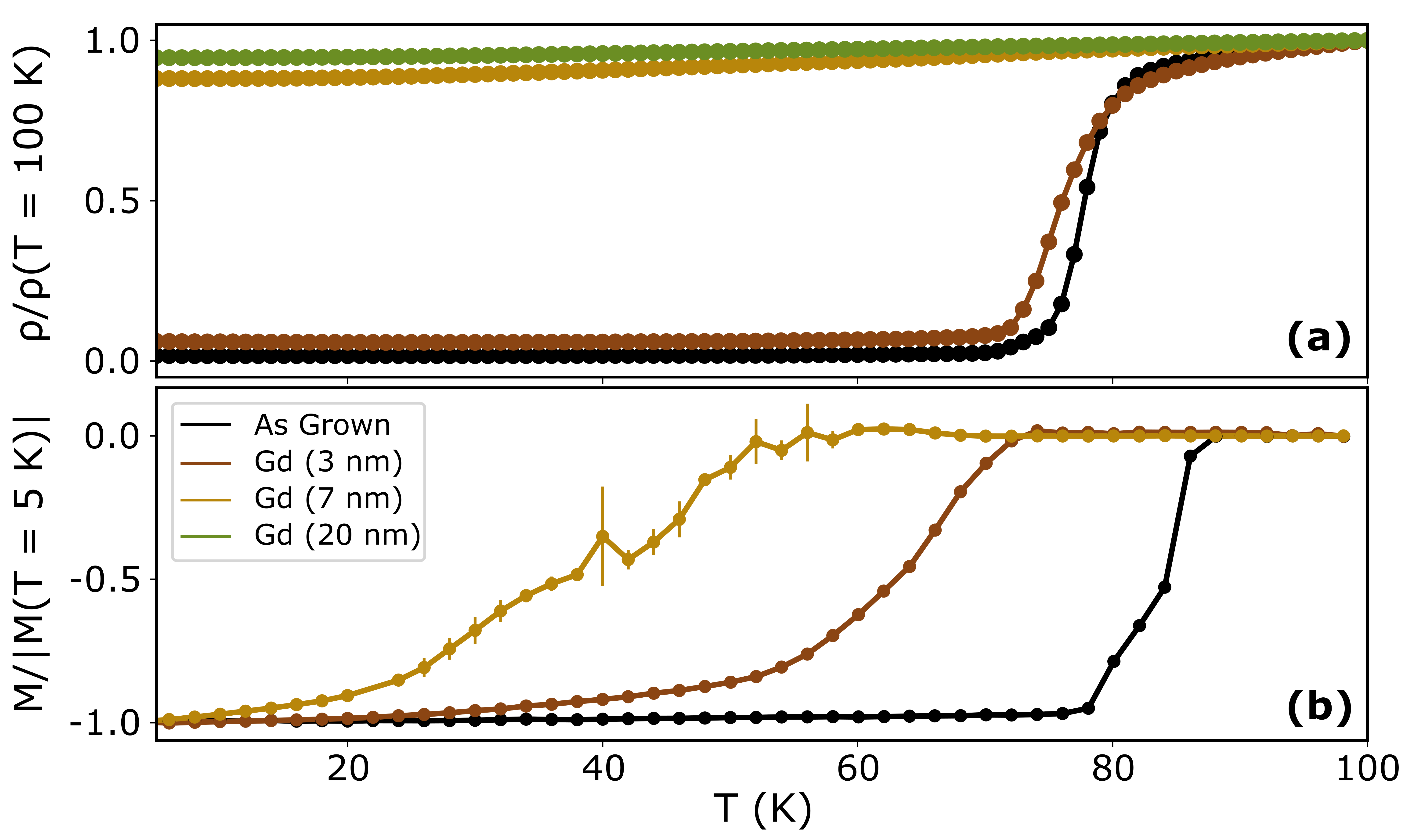
**Figure 4**. Normalized resistivity (a) and magnetization (b) as functions of temperature. The Gd (20 nm) sample has been omitted from (b), as no magnetic moment was detected at any temperature measured (to within the experimental noise).



**Figure 1**



**Figure 2Figure 4**

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**Figure 5**