Common origin of the two types of magnetic fluctuations in iron chalcogenides

Songxue Chi,1,2,* J. A. Rodriguez-Rivera,1,2 J. W. Lynn,1 Chenglin Zhang,3 D. Phelan,1 D. K. Singh,1,2 R. Paul,4 and Pengcheng Dai1,3

1NIST Center for Neutron Research, National Institute of Standards and Technology, Gaithersburg, Maryland, 20899-6102, USA
2Department of Materials Science and Engineering, University of Maryland, College Park, Maryland, 20742, USA
3Department of Physics and Astronomy, The University of Tennessee, Knoxville, Tennessee 37996-1200, USA
4Materials Measurement Laboratory, National Institute of Standards and Technology, Gaithersburg, Maryland, 20899-6102, USA
5Beijing National Laboratory for Condensed Matter Physics, Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

(Received 19 September 2011; revised manuscript received 13 November 2011; published 5 December 2011)

We use inelastic neutron scattering to study the low-energy spin excitations in moderately doped nonsuperconducting Fe1+δTe1−δSex samples. The spin excitations in this system contain components near (0.5,0,0) and (0.5,0.5,0) in the a-b plane reciprocal lattice units using tetragonal unit cell notation (a = b = 3.772 Å and c = 6.061 Å). At low energies the scattering is centered around (0.5,0,0). With increasing energy, the spectral weight of low-energy spin excitations centered around (0.5,0,0) abruptly shifts around 3 meV to the incommensurate spin excitations centered around (0.5,0.5,0). However both types of spin fluctuations exhibit the identical temperature dependence. These results indicate that the (0.5,0,0)-type spin excitations and the incommensurate excitations around the (0.5,0.5,0) position have a common origin and both must be taken into account to understand the nature of magnetism and superconducting pairing in the iron chalcogenides.

I. INTRODUCTION

The iron chalcogenides Fe1+δTe1−δSex have the α-PbO structure, which contains layers of Fe squares with the chalcogen atoms residing alternately above and below the centers of the squares, the same way in which FeAs layers are formed in the iron pnictides.1,2 The superconductivity arises when sufficient Se replaces Te in the antiferromagnetically ordered parent phase Fe1+δTe.2 Despite the similarities in crystal structure and Fermi surface topology,3,4 the parent compounds of the iron pnictides and iron chalcogenides have very different magnetic structures. The pnictides have a single-stripe in-plane collinear [C type, shown in Fig. 1(a)] antiferromagnetic (AFM) structure5,6 characterized by the wave vector (0.5,0.5,0) in the notation of the tetragonal lattice as highlighted by the shaded area in Fig. 1(a), which coincides with the Fermi surface (FS) nesting wave vector7 and the wave vectors of the neutron spin resonance in superconducting samples.8,9 The nonsuperconducting Fe1+δTe has a diagonal double-stripe bicollinear order [E-type, Fig. 1(b)] modulated along the (0.5,0,0,5) direction10,11, whose in-plane component is 45° away from the FS nesting wave vector (0.5,0.5,0) where, surprisingly enough, the spin resonance of superconducting Fe1+δTe1−δSex is found.12,13 The disparity between the static order and Fermi surface nesting as well as the large ordered moment1,14,15 in iron chalcogenides, have fueled the already heated debate about the nature of magnetism in the iron-based superconductors. The controversy over the magnetism in the iron chalcogenides has been mainly centered on whether it originates from itinerant electrons,16–18 localized moments19–21 or both.22–26

As a good probe to magnetism, the spin dynamics in the iron chalcogenides has been extensively studied with inelastic neutron scattering (INS) measurements. In undoped Fe1+δTe, the low-energy excitations start exactly at or closely to the (0.5,0,0) position depending on the amount of excess Fe.27–29

However, this dispersion around the magnetic zone center disappears at higher energies, and new rings of excitations emerge above 60 meV around integer positions such as (1,0,0). The ring centered at (1,0,0) disperses inward and eventually becomes a spot before disappearing above 275 meV.27 In the 27% Se-doped nonsuperconducting (NSC) sample,30 the magnetic response around (1,0,0) starts from the lowest measured energy and forms incommensurate magnetic (ICM) quartets instead of a ring, but with increasing energy it evolves into a ring. The dispersion is still steep and persists to energies beyond 250 meV. The normal state magnetism in superconducting FeTe1−δSex is very similar to that of the non-superconducting compounds except that the quartets never become a ring at high energies.30 The observed a-b plane spin excitations in Fe1+δTe1−δSex are summarized in Fig. 1(c) as schematics in reciprocal space.

In most Se-doped compounds the (0.5,0,0) excitations are still present at low energies. Their intensity diminishes with increasing Se-doping and becomes very weak in superconducting samples.30,34,35 It is therefore believed that the (0.5,0,0) type spin correlations have a deleterious effect on the superconducting pairing. Since chemical inhomogeneity, impurity, and phase separation exist in these materials,36–38 it is unclear whether these (0.5,0,0) spin fluctuations are intrinsic to the system or are a result of an undesirable phase or domains segregated from the primary magnetism.

To address the relationship between these two types of magnetic fluctuations, we have carried out INS measurements on moderately doped nonsuperconducting Fe1+δTe0.72Sex0.28. Our results indicate that the (0.5,0,0)-type excitations dominate the lowest-energy spectral response, but then the strength of the scattering abruptly shifts to the ICM excitations centered about (0.5,0.5,0), while the two types of spin fluctuations exhibit identical temperature dependence. Taken together, we conclude that the two types of magnetic excitations have a common origin.

DOI: 10.1103/PhysRevB.84.214407 PACS number(s): 75.47.—m, 71.70.Ch

1098-0121/2011/84(21)/214407(7) 214407-1 ©2011 American Physical Society
II. EXPERIMENT

High-quality single crystals of $\alpha$-phase Fe$_{1+y}$Te$_{1-x}$Se$_x$ were prepared with nominal composition of $x \approx 0.3$ using the flux method and co-aligned with neutrons. Neutron diffraction was carried out on a small piece of single crystal from the same batch as the 20 gram co-aligned crystals used for the INS measurements. The actual compositions of the samples were determined with prompt gamma activation analysis (PGAA) on beamline NG-7 at the NIST Center for Neutron Research (NCNR). Neutron scattering measurements were conducted on the cold neutron, multi-axis crystal spectrometer (MACS), and spin-polarized inelastic neutron spectrometer (SPINS) and the thermal triple axis spectrometer (TAS) BT-7 at NCNR. Pyrolytic graphite (PG) was used as monochromator and analyzer for all the measurements and as filter for BT-7 measurements. A BeO filter was used on SPINS with fixed final energy $E_f$ of 3.7 meV and horizontal collimations of open-80°-S-80°-open. Double-focusing monochromator, Be filter, and all 20 channels of the detection system were employed on MACS with $E_f$ fixed at 5 meV. An empty sample holder in the same sample environment was also measured and used as background subtraction for the MACS measurements. Fixed $E_f$ of 35 meV and open-25°-S-25°-120° collimations were used for INS measurements on BT-7. Open-25°-S-25°-50° with $E_f = 14.7$ meV was used for elastic scattering on BT-7. The momentum transfer $Q$ at $(q_x, q_y, q_z)$ is defined as $(H, K, L) = (q_x/a, 2\pi q_y/b, 2\pi q_z/c)$ reciprocal lattice units (r.l.u.) in the tetragonal unit cell ($P4/nmm$ space group). The lattice parameters of the tetragonal unit cell are $a = b = 3.772$ Å and $c = 6.061$ Å at $T = 1.5$ K. Elastic measurements were taken in both the $(H, K, 0)$ and $(H, 0, L)$ scattering planes, while the INS measurements were concentrated in the $(H, K, 0)$ plane. The error bars shown in the figures are statistical in nature and represent one standard deviation.

III. RESULTS

We first determine the actual stoichiometry of our Fe$_{1+y}$Te$_{1-x}$Se$_x$ crystals since these can differ significantly from the nominal compositions. For this purpose, prompt gamma activation analysis (PGAA) was carried out on a small piece of single crystal. PGAA is a nondestructive technique using neutron absorption to simultaneously determine the presence and accurate quantities of various elements in a compound. We find that the Fe:Te:Se molar ratio is 1.01:0.72:0.28. Relative expanded uncertainties for PGAA data are estimated at <5%.

The crystal was aligned in the $(H,0,L)$ scattering plane for the neutron diffraction measurements. $H$ scans were performed at various temperatures. The static magnetic order survives in this compound as short-range static order centered at the wave vector $(0.47,0,0.5)$. This is consistent with the previous reports that Se-doped bulk nonsuperconducting samples have short-range order centered at incommensurate wave vector $(0.5 - \delta, 0, 0.5)$. The $\delta$ value, 0.03 in this case, can be tuned by both the Se and Fe concentration. $H$ scans at some typical temperatures are shown in the inset of Fig. 1(d). The magnetic peaks are much broader than the instrumental resolution with Lorentzian fits giving a width of 1.6 (r.l.u.) corresponding to in-plane correlation length of 3.7(8) Å at 5 K. This is in agreement with that of similar doping FeTe$_{0.7}$Se$_{0.3}$. The integrated intensity of the $H$ scans at different temperatures is plotted in Fig. 1(d). Upon warming, the peak intensity is gradually suppressed without an abrupt transition or change of peak position. The magnetic intensities cannot be detected above 60 K. The diffuse nature of the short-range order is reflected by the concave shape of the intensity-temperature curve. Constant-$Q$ scans of inelastic neutron scattering at $(0.5,0.5,0)$ at 1.4 K and 20 K are shown in Fig. 1(e). The overall spectrum shows little temperature dependence except some enhancement at 20 K at energies below 4 meV due to thermal population. Neither a spin resonance nor the development of a spin gap at low temperature can be found, confirming the absence of bulk superconductivity.

The above characterizations place our sample in the intermediate doping part of the phase diagram where the long range AFM order is suppressed but bulk superconductivity has not emerged. In this region, there are reports about weak charge carrier localization and spin glass ordering. The magnetic and superconducting properties are sensitive not only to the Te/Se ratio, but also to the variation of Fe content. The modest amount of excess Fe in our sample ensures that no significant complications arise due to the interstitial iron.
The INS measurements focused on the $(H,K,0)$ scattering plane because of the weak $L$ dependence. For those measurements we drop the $L$ coordinate for simplicity and present the data in terms of $(H,K)$ only. In order to simultaneously monitor the two excitations we need to survey a wide region of momentum space in the low-energy transfer range. MACS is ideal for this type of measurement because of the multiple detection systems. Figures 2(a)–2(f) show some typical contour plots at 1.5 K using data folded into the first quadrant of the scattering plane. At low-energy transfers the $(0.5,0)$ type excitation is dominant with a broad peak. At the equivalent position of higher Brillouin zones, such as $(1,0.5)$ and $(1.5,0)$, peaks are weaker because of the decreased magnetic form factor. As the energy increases the $(0.5,0)$ type scattering quickly diminishes in intensity while the two ICM peaks around $(0.5,0.5)$ start to appear and $(0.5,0)$ type excitations become more easily seen in Fig. 4. The constant-$E$ cuts in the transverse $[H,1−H]$ direction through $(0.5,0.5)$ were also conducted, which yield two symmetric ICM peaks, giving the dispersion of the ICM excitation. The combinations of these two cuts as a function of energy are shown in Figs. 3(a) and 3(b). The $[H,1−H]$ cuts at energies above 7 meV are not included because the low-\(\mathbf{Q}\) space could not be reached with the chosen fixed final energy. Figure 3(a) shows that the disappearance of the broad peak at $(0.5,0)$, as energy increases, is a three-stage process instead of a continuous one. The broad quasi-elastic scattering shows a sharp drop of intensity at about 1.5 meV, followed by another abrupt drop at about 3 meV before the signal disappears above 6 meV. Figure 3(a) also shows the gradually emerging excitation at $(0.5,0.5)$ and $(0.5,0.5)$. The disappearance of the $(0.5,0)$ excitation at around 6 meV is correlated with the opening of a spin gap for the ICM peaks around $(0.5,0.5)$, as shown in Fig. 3(b). In this partial gap the ICM peaks around $(0.5,0.5)$ are suppressed in intensity, but their peak profiles remain. Remarkably, the energies where the twin ICM peaks abruptly change spectral weight coincide with those where the opposite sudden change occurs for the $(0.5,0)$ spectrum. This reciprocal interplay between the two spin excitations can be more easily seen in Fig. 4. The constant-$\mathbf{Q}$ scans at these two wave vectors show an opposite energy dependence. The
presence of the (0,5,0) fluctuations is compensated by the gap opening in the ICM fluctuations near (0,5,0). The dashed lines in Fig. 4(a) indicate the energies at which the two spectra show abrupt changes. The intensity of the constant-Q scan at (1,0,5) is also plotted on a log scale in Fig. 4(a) confirming the energy dependence of the (0,5,0) correlation.

In order to obtain the integrated intensity for the (0,5,0) spectrum, we performed cuts through (0,5,0) in the $H$ direction, instead of the $K$ direction to avoid the component of the excitation near (0,5,0,5). Because of the unreachable Q space at higher energies, only half of the thereby obtained peak is integrated and plotted against energy in Fig. 4(c). The intensity of the two ICM peaks around (0,5,0,5) is also plotted in Fig. 4(d). The constant-E scans in the transverse direction through (0,5,0,5) were also performed with the thermal triple-axis instrument BT-7. The raw data up to $E = 16$ meV and lines of fits to two Gaussians are displayed in Fig. 5(b). The ICM excitation shows very little variation in dispersion and in spectral weight between 8 and 10 meV. The sum of fitted areas of the two Gaussians, together with that of the constant-E cuts from the MACS data shown in Fig. 2, is plotted as a function of energy in Fig. 4(d). Identical scans at $E = 8$ meV using the two instruments were used to normalize the overall intensities. The spectrum of the ICM excitations about (0,5,0,5) remains constant above 7 meV. This is important because it means that the gap of the ICM excitations below 7 meV is not compensated by spectral gain at higher energies at the same wave vector as is the case in the superconducting compounds. This again indicates that spectral weight is transferred between the two wave vectors and that the (0,5,0) spectrum is at the cost of the ICM spectrum. Clearly, the spectral weight for these two types of excitations is inversely correlated, which rules out electronic phase separation or magnetic inhomogeneity as the origin of the two types of magnetic correlations.

Another conspicuous feature in Fig. 3(b) is the dispersion toward (0,5,0,5) below 3 meV before its steep outward dispersion. This hourglass-shaped dispersion has been observed in both superconducting and overdoped nonsuperconducting systems. For this underdoped sample the inward dispersion is less pronounced and is more of a bell shape. To confirm this inward dispersion and the intense quasielastic scattering below 1.5 meV, we repeated the $[H,1-H]$ scans on the cold neutron TAS instrument SPINS, which, with $E_f = 3.7$ meV and horizontally flat monochromator, offers better energy and Q resolution. Figure 5(a) shows these transverse scans through (0,5,0,5) at various fixed-energy transfers. The low background and distinct profile of the two peaks at $E = 1$ meV in Fig. 5(a), in contrast to the broad quasielastic scattering extending to $E = 1.5$ meV in Figs. 3(b) and 4(b), is the result of improved resolution. The peaks are fit with two Gaussians convoluted with the instrumental resolution. The Q positions obtained are overplotted in Fig. 3(b) with open circles. The results are consistent with the MACS data and the bell-shaped dispersion is clearly visible. The inward dispersion stops around 3 meV. Comparing our data with the studies of other compositions, it seems the increasing Se doping pushes the saddle point to higher energies. It should be also noted that in the energy range where the bell shape of the ICM excitation occurs, there is an abnormal change of the (0,5,0) spin spectrum as shown in Fig. 3(a). The change happens to both the linewidth [inset of Fig. 4(c)] and the peak intensity [Fig. 4(a)]. These anomalies are confirmed by a similar Q-E plot (not shown) for the cut through (1,0,5), which shows identical anomalies in intensity and linewidth.

We now turn to the temperature dependence of the spin excitations. The $(H, K, 0)$ planar maps have been obtained at different temperatures between 1.5 K and 308 K for four typical energy transfers: 1 meV in the quasielastic region, 4.5 meV in the spin gap, 7.0 meV at the verge of the gap, and 10 meV above the gap. Figure 6 shows the combined $(0.5, K)$ cuts at $E = 1$ meV, and the $[H, 1-H]$ cuts at $E = 1$ meV and 10 meV as a function of temperature. The integrated intensity
of these cuts at all the above-mentioned energies are plotted in Figs. 7(a) and 7(b). Both types of magnetic correlations are so robust that they maintain their well-defined features up to the highest measured temperature for all the energy transfers. On warming the (0.5,0) spectrum at $E = 1$ meV starts to gain intensity at about 60 K, where the static order disappears, reaches its maximum at about 80 K and gradually decreases at higher temperatures. The ICM spectrum around (0.5,0.5), however, is clearly gapped below the transition temperature of the static AFM order, as shown in Figs. 6(b) and 7(b). As the system is heated from the short-range static ordered phase into the paramagnetic phase, the static component also transfers to the background, resulting in the abrupt broadening of the linewidth of the (0.5,0) spectrum, as shown in Fig. 6(c). At $E = 7$ meV the intensity of the ICM spectrum is less affected by the static order below 60 K. At $E = 10$ meV [Fig. 6(d)], the ICM intensity remains unaffected by the static order and gradually increases monotonically all the way above 300 K.

The neutron scattering intensity $S_T(Q,\omega)$ at temperature $T$ is related to the imaginary part of the dynamic susceptibility $\chi(Q,\omega)$ through $S_T(Q,\omega) = |n(\omega, T) + 1|\chi''(Q,\omega)$, where $n(\omega, T)$ is the Bose factor. In order to investigate the temperature dependence of the dynamic susceptibility in the paramagnetic phase, the $S_T(\omega)$ intensities for $T > 80$ K in Figs. 7(a) and 7(b) are converted to the values of dynamic susceptibility $\chi''(Q,\omega)$ and plotted in Figs. 7(c) and 7(d). The scattering at energy transfers of 7 meV and 10 meV increases with increasing temperature as expected due to the thermal factor, but the susceptibility decreases with $T$ indicating that the intrinsic strength of the magnetic scattering decreases as the scattering evolves to higher temperature. The $T$ dependencies of the dynamic susceptibility in the paramagnetic phase ($T > 60$) were fit to the power law $\chi'' \propto C T^\beta$. The $\beta$ values obtained for wave vectors (0.5,0) and (0.5,0.5) at $E = 1$ meV, denoted by black squares in Figs. 7(a) and 7(b) (solid and empty), are $-1.28 \pm 0.02$ and $-1.26 \pm 0.03$, respectively.

Similarly the $\beta$ values at $E = 4.5$ meV (red circles) for these two vectors are $-0.68 \pm 0.01$ and $-0.65 \pm 0.01$. Note that the dynamic susceptibility of the two excitations have the identical temperature dependence. This is further evidence that both types of excitations have a common origin. At an energy of 7 meV [Fig. 7(b)] where the (0.5,0) spectrum is almost completely depleted, the $\chi''$ for (0.5,0.5) continues to follow the same relationship with $\beta = -0.66 \pm 0.01$. At an energy transfer of 10 meV, $\beta$ becomes $-0.54 \pm 0.03$.

**IV. CONCLUSION**

For the superconductors, the high-energy magnetic excitations remain unchanged when cooled from the normal state to below $T_C$. It is the low-energy part of the magnetic excitation spectrum that responds to the formation of superconducting pairs, as would be expected. These changes include the opening of a spin gap and the development of a spin resonance. In our NSC system, the gap still develops for the ICM spectrum centered around the same (0.5,0.5) position, but without the development of superconductivity or a spin resonance. Instead, we have the (0.5,0) type excitations, which apparently correspond to that part of magnetism that is needed for the spin resonance in the superconducting state, as only the (0.5,0)-type excitations are suppressed when the superconductivity and the associated spin resonance develop. Now that we know that the two types of excitations have a common origin, suppressing the (0.5,0)-type correlations through doping Se cannot be simply understood as eliminating
a coexisting phase. Rather one has to treat the two excitations as one problem when trying to reveal the driving force for magnetism and superconductivity. In that sense, the itinerant electrons alone may not be able to provide a complete answer. The interband Fermi surface nesting describes the main features of the ICM excitations such as the incommensurate excitations and the hourglass dispersion near (0.5,0). In this study, the nesting scenario for the (0.5,0) excitation has been supported by the reciprocal interplay as the Se content increases well into the superconducting region. This scenario is supported by the reciprocal interplay between the two types of excitations and the temperature dependence of the low- \( E \) dynamic susceptibility presented in this study. However, it is not possible to understand the low-energy features such as the hourglass dispersion and the abnormal change of intensity and linewidth for the (0.5,0) spectrum with just a local spin picture. Our results call for a more unified mechanism that reconciles these features and embraces the two types of excitations as having a common origin.

**ACKNOWLEDGMENTS**

The work at NIST is supported by the US Department of Commerce. SPINS and MACS utilized facilities supported in part by the NSF under Agreement No. DMR-0944772. The work at the University of Tennessee is supported by the US DOE, Basic Energy Sciences (BES), through DOE Grant No. DE-FG02-05ER46202. The work at the Institute of Physics is supported by the Chinese Academy of Science and by the Ministry of Science and Technology of China 973 program (2012CB821400).
COMMON ORIGIN OF THE TWO TYPES OF MAGNETIC . . .

27 O. J. Lipscombe, G. F. Chen, Chen Fang, T. G. Perrin, D. L. Abernathy, A. D. Christianson, Takeshi Egami, Nanlin Wang, Jiangping Hu, and Pengcheng Dai, Phys. Rev. Lett. 106, 057004 (2011).

28 Igor A. Zaliznyak, Zhijun Xu, John M. Tranquada, Genda Gu, Alexei M. Tsvelik, and Matthew B. Stone, Phys. Rev. Lett. 107, 216403 (2011).

29 C. Stock, E. E. Rodriguez, M. A. Green, P. Zavalij, and J. A. Rodriguez-Rivera, Phys. Rev. B 84, 045124 (2011).

30 M. D. Lumsden, A. D. Christianson, E. A. Goremychkin, S. E. Nagler, H. A. Mook, M. B. Stone, D. L. Abernathy, T. Guidi, G. J. MacDougall, C. de la Cruz, A. S. Sefat, M. A. McGuire, B. C. Sales, and D. Mandrus, Nature Phys. 6, 182 (2010).

31 S.-H. Lee, Guangyong Xu, W. Ku, J. S. Wen, C. C. Lee, N. Katayama, Z. J. Xu, S. Ji, Z. W. Lin, G. D. Gu, H.-B. Yang, P. D. Johnson, Z.-H. Pan, T. Valla, M. Fujita, T. J. Sato, S. Chang, K. Yamada, and J. M. Tranquada, Phys. Rev. B 81, 220502 (2010).

32 D. N. Argyriou, A. Hiess, A. Akbari, I. Eremin, M. M. Korshunov, Jin Hu, Bin Qian, Zhiqiang Mao, Yiming Qiu, Collin Broholm, and W. Bao, Phys. Rev. B 81, 220503 (2010).

33 Shiliang Li, Chenglin Zhang, Meng Wang, Hui-qian Luo, Xingye Lu, Enrico Faulhaber, Astrid Schneiderwind, Peter Link, Jiangping Hu, Tao Xiang, and Pengcheng Dai, Phys. Rev. Lett. 105, 157002 (2010).

34 Zhijun Xu, Jinsheng Wen, Guangyong Xu, Qing Jie, Zhiwei Lin, Qiang Li, W. Ratcliff, Genda Gu, and J. M. Tranquada, Phys. Rev. B 80, 104506 (2009).

35 T. J. Liu, J. Hu, B. Qian, D. Fobes, Z. Q. Mao, W. Bao, M. Reehuis, S. A. J. Kimber, K. Proke, S. N. Gvasaliya, R. Khasanov, E. Pomjakushina, and B. Roessli, J. Phys. Condens. Matter 22, 142202 (2010).

36 Hefei Hu, J. M. Zuo, J. S. Wen, Z. J. Xu, Z. W. Lin, Q. Li, Genda Gu, W. K. Park, and L. H. Greene, New J. Phys. 13, 053031 (2011).

37 D. C. Johnston, Adv. Phys. 59, 803 (2010).

38 B. Joseph, A. Ladecola, A. Puri, L. Simonelli, Y. Mizuguchi, Y. Takano, and N. L. Saini, Phys. Rev. B 82, 020502(R) (2010).

39 J. A. Rodriguez, D. M. Adler, P. C. Brand, C. Broholm, J. C. Cook, C. Brocker, R. Hammond, Z. Huang, P. Hundertmark, J. W. Lynn, N. C. Maliszewskyj, J. Moyer, J. Orndorff, D. Pierce, T. D. Pike, G. Scharfstein, S. A. Smee, and R. Vilaseca, Meas. Sci. Technol. 19, 034023 (2008).

40 J. W. Lynn, Y. Chen, S. Chang, Y. Zhao, S. Chi, W. Ratcliff, B. G. Ueland, and R. W. Erwin, J. Res. NIST (in press).

41 J. W. Lynn and Pengcheng Dai, Physica C 469, 469 (2009).

42 R. Paul, R. Lindstrom, and A. Heal, J. Radioanal. Nucl. Chem. 215, 63 (1997).

43 Jinsheng Wen, Guangyong Xu, Zhijun Xu, Zhi Wei Lin, Qiang Li, W. Ratcliff, Genda Gu, and J. M. Tranquada, Phys. Rev. B 80, 104506 (2009).

44 P. L. Paulose, C. S. Yadav, and K. M. Subhedar, Europhys. Lett. 90, 27011 (2010).

45 P. Babkevich, M. Bendele, A. T. Boothroyd, K. Conder, S. N. Gvasaliya, R. Khasanov, E. Pomjakushina, and B. Roessli, J. Phys. Condens. Matter 22, 142202 (2010).

46 Naoyuki Katayama, Sungdae Ji, Despina Louca Seunghun Lee, Masaki Fujita, Taku J. Sato, Jinsheng Wen, Zhijun Xu, Genda Gu, Guangyong Xu, Zhiwei Lin, Masanori Enoki, Sung Chang, Kazuyoshi Yamada, and John M. Tranquada, J. Phys. Soc. Jpn. 79, 113702 (2010).

47 M. Bendele, P. Babkevich, S. Katrych, S. N. Gvasaliya, E. Pomjakushina, K. Conder, B. Roessli, A. T. Boothroyd, R. Khasanov, and H. Keller, Phys. Rev. B 82, 212504 (2010).

48 E. E. Rodriguez, C. Stock, P.-Y. Hsieh, N. Butch, J. Paglione, and M. A. Green, Chemical Science 2, 1782 (2011).

49 Zhijun Xu, Jinsheng Wen, Guangyong Xu, Songxue Chi, Wei Ku, Genda Gu, and J. M. Tranquada, Phys. Rev. B 84, 134409 (2011).

50 T. M. McQueen, A. J. Williams, P. W. Stephens, J. Tao, Y. Zhu, V. Ksenofontov, F. Casper, C. Felser, and R. J. Cava, Phys. Rev. Lett. 103, 057002 (2009).

51 X. B. He, G. R. Li, J. D. Zhang, A. B. Karki, R. Jin, B. C. Sales, A. S. Sefat, M. A. McGuire, D. Mandrus, and E. W. Plummer, Phys. Rev. B 83, 220502(R) (2011).

52 K. Nakayama, T. Sato, P. Richard, T. Kawahara, Y. Sekiba, T. Qian, G. F. Chen, J. L. Luo, N. L. Wang, H. Ding, and T. Takahashi, Phys. Rev. Lett. 105, 197001 (2010).