Unusual temperature dependence of band dispersion in Ba(Fe$_{1-x}$Ru$_x$)$_2$As$_2$ and its consequences for antiferromagnetic ordering

R. S. Dhaka,$^1$ S. E. Hahn,$^1$ E. Razzoli,$^2$ Rui Jiang,$^1$ M. Shi,$^2$ B. N. Harmon,$^1$
A. Thaler,$^1$ S. L. Bud’ko,$^1$ P. C. Canfield,$^1$ and Adam Kaminski$^2$[1]

$^1$Ames Laboratory and Department of Physics and Astronomy, Iowa State University, Ames, Iowa 50011, USA
$^2$Swiss Light Source, Paul Scherrer Institut, CH-5232 Villigen PSI, Switzerland

(Dated: May 1, 2014)

We have performed detailed studies of the temperature evolution of the electronic structure in Ba(Fe$_{1-x}$Ru$_x$)$_2$As$_2$ using Angle Resolved Photoemission Spectroscopy (ARPES). Surprisingly, we find that the binding energy of both hole and electron bands changes significantly with temperature in both pure and Ru substituted samples. The hole and electron pockets are well nested at low temperature in unsubstituted (BaFe$_2$As$_2$) samples, which likely drives the spin density wave (SDW) and resulting antiferromagnetic order. Upon warming, this nesting is degraded as the hole pocket shrinks and the electron pocket expands. Our results demonstrate that the temperature dependent nesting may play an important role in driving the antiferromagnetic/paramagnetic phase transition.

PACS numbers: 74.25.Jb,74.62.Dh,74.70.-b,79.60.-i

The phase transition from antiferromagnetic to paramagnetic order is believed to play a crucial role in the emergence of high temperature superconductivity in iron arsenic high temperature superconductors$^{[1-5]}$. The associated transition temperature can be controlled by tuning parameters such as chemical substitution$^{[6-17]}$ or pressure$^{[18-20]}$. The superconducting dome exists in the phase diagram surrounding the point, where the AFM/PM phase transition temperature intersects T=0 axis and it is believed that the AF fluctuations are the glue pairing the electrons. Understanding the microscopic origin of this phase transition and the role of electronic structure is a prerequisite to validating the pairing mechanism and will likely aid in search for new superconducting materials. Our earlier studies$^{[21]}$ indicated that the AFM/PM phase transition is associated with a two- to three-dimensional change of the electronic structure. Fermi surface nesting is believed to play an important role in the emergence of the SDW and AF order, however detailed understanding is still lacking. Data from chemical substitution yields only a partially consistent picture here. It is now well established that substitutions of Fe with Co, Ru, Ni or Pd increases the carrier concentration and shift the chemical potential higher (see for example Refs.$^{[9]}$ and$^{[22]}$). This of course would worsen the FS nesting conditions, and naturally lead to weakening of the SDW. The case of isoelectronic substitution is much less clear. It is still debated whether or not substitution of As with isoelectronic P changes the carrier concentration$^{[23,24]}$, however there is sufficient evidence for changes in dispersion of individual bands, which may affect nesting in a way similar to the previous case. A curious case is substitution of Fe with isoelectronic Ru, which results in a phase diagram similar to other substitutions (albeit for much higher Ru concentrations), but it was demonstrated to have negligible effects on the band dispersion up to $40\%$.$^{[25]}$. It appears that the SDW in this case is suppressed without significant change of the nesting condition and we speculated that magnetic dilution by less magnetic Ru may play a role in such case$^{[25]}$.

In ordinary materials, in absence of phase transitions, temperature has negligible effect on the band structure, and is mostly limited to changing the occupation close to the chemical potential (Fermi function) and changes of the quasiparticle lifetime reflected in the width of the spectral function measured by ARPES.

In this Letter, we demonstrate that the band structure of one of the prototypical iron arsenic materials is changing significantly with temperature; in both pure and Ru substituted BaFe$_2$As$_2$ samples we observe significant shifts in the energy position of the bands. Upon increasing temperature the hole band moves to lower binding energy and eventually sinks below $E_f$ close to room temperature. The electron band, on the other hand, moves to lower binding energies with increasing temperature. This rather strange behavior destroys the nesting present at low temperatures upon warming the sample. Increasing temperature has therefore very similar effect to non-isoelectronic chemical substitution, where the nesting is lifted by changes of the chemical potential. Our finding explains the nearly linear decrease of the AFM transition temperature with increasing chemical substitution. It also points to a scenario where the AFM transition is driven by weakening of nesting with temperature rather than thermal fluctuations alone.

Single crystals of Ba(Fe$_{1-x}$Ru$_x$)$_2$As$_2$ were grown out of self-flux using conventional high-temperature solution growth techniques$^{[16]}$, where FeAs and RuAs were synthesized in the same manner as in Ref$^{[27]}$. The ARPES measurements were performed at 10.0.1 beamline (linear horizontal polarization, LHP) of the Advanced Light Source (ALS), Berkeley, California and at the SIS beamline (linear vertical polarization, LVP) of Swiss Light
Source (SLS), Switzerland, using a Scienta R4000 electron analyzer in ultrahigh vacuum below $4 \times 10^{-11}$ mbar and at various sample temperatures. The data were also acquired using a laboratory based system consisting of a Scienta SES2002 electron analyzer and a GammaData helium ultraviolet lamp where we use the He II line ($h\nu = 40.8$ eV) for the band dispersion maps. The samples were mounted on an Al pin using UHV compatible epoxy and in situ cleaved perpendicular to the $c$-axis, yielding single layer surfaces in the $a$-$b$ plane. The measurements carried out on several samples and temperature cycling yielded similar results for the band dispersion and Fermi surface. The energy and angular resolution was set at 15 meV and $0.3^\circ$, respectively. High symmetry points were defined the same way as in Ref. [22]. Ab initio calculations were performed with VASP [28, 29] using PAW pseudo-potentials [30, 31] and utilizing the LDA and PBE (Perdew-Burke-Ernzerhof) exchange-correlation functionals [32]. Experimental lattice parameters of $a = 3.9624$, $c = 13.0168$ and $z_{\text{As}} = 0.3545$ are used from Ref. [33].

Figs. 1(a–d) show the band dispersion maps for the parent compound BaFe$_2$As$_2$ measured at various temperatures using $35$ eV ($k_z \simeq 2\pi/c$) photon energy i.e. the upper edge of the first Brillouin zone (Z point) [34]. The second set of measurements were performed using 63 eV photon energy and the results are shown in Figs. 1(g–m). All data were divided by resolution broadened Fermi functions to reveal band behavior in the proximity to the chemical potential. At low temperature the hole band at the center of the BZ crosses the chemical potential at roughly $\pm 0.2 \pi/a$ resulting in hole like Fermi surface sheet (see also Fig. 2). With increasing temperature this band moves down, to larger binding energies and around 200 K the top of the band moves very close to the chemical potential. Corresponding momentum distribution curves (MDC’s), radii of the FS sheets and MDC widths are plotted in panels on the right. Here the two well separated MDC peaks present at low temperature move closer together and merge into flat top peak signifying significant (factor of 2) reduction of the size of the hole pocket. Those changes are accompanied by increase in the MDC width, evident from raw data as well results of fitting in panels f and m. Whereas the increase of the MDC width may be associated with paramagnetic/antiferromagnetic-orthorhombic(tetragonal/orthorhombic) transition, the
increasing temperature. Therefore nesting present at low
hole pocket decreases and electron pocket increases with
un-substituted BaFe

hole band moves to higher binding energies, while the
is fully suppressed and all data are taken in paramag-
tron band in Fig. 4(e-h); for x=0.36 the AF transition
are shown for hole band in Fig. 4(a-d) and for elec-
ning ARPES intensities measured at several temperatures
Ba(Fe

transition, we performed similar measurements using
band structure are not caused by an AFM phase
verify that the temperature induced changes of the
structure of Ru substituted samples, for x
ence of the antiferromagnetic order. Since the electronic
low temperatures will of course be affected by the pres-
ment due to thermal expansion. It is clear from these
data that size of this FS sheet decreases with increasing
temperature, which will affect nesting in a very profound
way. The above changes are strongest close to the top and
bottom edges of the BZ along the k_z direction. In Fig.
we plot the ARPES intensity along the (1,1,0) direc-
tion as a function k_z momentum. The large hole pocket
present at bottom and top edge of the zone (k_z ~ 18 \pi/c )
shrinks with temperature, whereas the dispersion close
to center of the zone remains mostly unaffected.

In the pure BaFe_2As_2 samples the band structure at
low temperatures will of course be affected by the pres-
ence of the antiferromagnetic order. Since the electronic
structure of Ru substituted samples, for x \leq 0.40 are al-
most identical [25], we can access low temperatures in the
paramagnetic state by choosing finite x values with sup-
pressed magnetic/structural phase transitions[19]. To
verify that the temperature induced changes of the
band structure are not caused by an AFM phase
transition, we performed similar measurements using
Ba(Fe_{1-x}Ru_x)_2As_2 for x=0.28 and 0.36. The result-
ing ARPES intensities measured at several temperatures
are shown for hole band in Fig. 4(a-d) and for elec-
tron band in Fig. 4(e-h); for x=0.36 the AF transition
is fully suppressed and all data are taken in paramag-
netic/tetragonal state. With increasing temperature, the
hole band moves to higher binding energies, while the
electron band moves to lower binding energies, similar to
un-substituted BaFe_2As_2. As a result, the diameter of
hole pocket decreases and electron pocket increases with
increasing temperature. Therefore nesting present at low
temperatures between hole and electron pocket weakens
significantly upon warming up. To quantify this effect,
we extracted the diameter of both pockets as a function
of temperature by fitting the MDC’s. Results are plotted
in Fig. 5. At 50K the diameters of both the hole and elec-
tron pockets are close to 0.4 \pi/a, which corresponds to
near nesting condition. Upon warming up to 300 K, the
diameter of the hole pocket decreases to ~0.32 \pi/a, while
the diameter of the electron pocket increases to 0.48 \pi/a,
which obviously destroys the nesting. We repeated the
measurements, while cycling the sample temperature to
ensure that this is an intrinsic effect and not due to aging
of the sample surface. Similar data was also obtained for
x=0.28 (not shown) demonstrating that this effect does
not change with Ru substitution. We should emphasize
that, at these Ru substitution levels, the samples do not
display AFM order at low temperatures despite presence
of apparent nesting. The suppression of AFM order in
these samples most is most likely due to magnetic dilu-
tion [25]. We note that the AFM nesting vector has a com-
ponent along the z direction. We can compare size of
both pockets at the same value of k_z because the elec-
tron pocket is quasi 2D and its diameter does not change
significantly along the z direction [21].

The large temperature induced changes of the band
dispersion are indeed very unusual and unexpected. To
the best of our knowledge, no such changes were reported
in any of the studied systems before. The most obvi-
ous explanation of this effect is subtle changes in rela-
tive atomic positions due to thermal expansion. It is

FIG. 3. (Color online) The hole Fermi surface maps of
BaFe_2As_2 (at various sample temperatures) along the k_{||}-k_z
plane measured around the center of the BZ by changing h\nu.

FIG. 4. (Color online) The band dispersion plots for
Ba(Fe_{1-x}Ru_x)_2As_2, x = 0.36 sample as a function of tem-
perature. (a-d) hole pocket at the center of the BZ.x (e-h)
electron pocket at the planar corner of the BZ. Data measured
with photon energy h\nu = 40.8 eV.
well known that the energy positions of the bands are very sensitive to the Fe-As distance. If the FeAs distance changes significantly with the temperature, this may explain our observations. We have therefore performed a band structure calculation using density functional theory (DFT) for the parent compound BaFe$_2$As$_2$ in order to understand the changes of the diameter of the pockets ($\Delta k_F$) with temperature. We considered the tetragonal structure, and thermal expansion data[35]. Room temperature parameters were used for $a$ and $c$. Next, we reduced $a$ by the amount measured in the thermal expansion data. Initial calculations using the measured atomic positions show a band with $d_{x^2-y^2}$ character shifting downward by 17 and 15 meV for LDA and PBE exchange correlation functionals. This is approximately half the observed change (35±5 meV) from the ARPES measurements. A band with $d_{x^2-y^2}$ character shifts downward by about 5 meV. At the electron pocket, bands with $d_{xz}/d_{yz}$ character shift downwards by about 5 meV.

Using room temperature parameters for $a$ and $c$, we allowed $z_{As}$ to relax such that all forces on the atoms equaled zero. We than reduced $a$ by the amount measured in the thermal expansion data and again determined $z_{As}$ associated with this value of $a$. Near the Z ($\Gamma$) point the band with $d_{x^2-y^2}$ character shifts downward by 28 and 25 meV with LDA and PBE exchange-correlation functionals, respectively. This is fairly close to the observed change from the ARPES measurements. For the electron pocket the changes are small and almost no observable shift is present in the upper band ($d_{x^2-y^2}$ character), while, the lower bands have $d_{xz}/d_{yz}$ character and shift downwards by about 6 meV. This approach points to changes of the atomic positions due to thermal expansion as a possible explanation of our results.

In conclusion, we demonstrated that the band structure of pure and Ru substituted BaFe$_2$As$_2$ changes appreciably with temperature. The nesting conditions leading to SDW and present at low temperature become weaker upon warming up. This likely contributes to destruction of AFM order at higher temperatures. In essence, increasing temperature has similar effect on the band structure and nesting as Co substitution. This explains why the AFM/PM transition in the phase diagram is almost a straight line. At higher Co concentration, the nesting and AFM order is suppressed by smaller increase of the temperature.

We would like to thank Rafael Fernandez, Andy Millis and Andrey Chubukov for very useful discussions, Sung-Kwan Mo at the ALS and staff at SLS for their excellent instrumental support. The work at the Ames Laboratory was supported by the Department of Energy, Basic Energy Sciences under Contract No. DE-AC02-07CH11358.

* kaminski@ameslab.gov

1. Y. Kamihara et al., J. Am. Chem. Soc. 130, 3296 (2008).
2. X. H. Chen et al., Nature 453, 761 (2008).
3. T. Y. Chen et al., Nature 453, 1224 (2008).
4. G. F. Chen et al., Phys. Rev. Lett. 100, 247002 (2008).
5. H. Takahashi et al., Nature 453, 376 (2008).
6. M. Rotter et al., Phys. Rev. Lett. 101, 107006 (2008).
7. A. S. Sefat et al., Phys. Rev. Lett. 101, 117004 (2008); Phys. Rev. B 78, 104505 (2008).
8. K. Sasmal et al., Phys. Rev. Lett. 101, 107007 (2008).
9. P. C. Canfield and S. L. Bud’ko, Annu. Rev. Condens. Matter Phys. 1, 27 (2010).
10. D. C. Johnston, Adv. Phys. 59, 803 (2010).
11. S. Jiang et al., J. Phys.: Condens. Matter 21, 382203
(2009); L. E. Klintberg et al., J. Phys. Soc. Jpn. 79, 123706 (2010).
[12] Yu-Zhong Zhang et al., Phys. Rev. B 80, 094530 (2009).
[13] S. Sharma et al., Phys. Rev. B 81, 174512 (2010); W. Schnelle et al., Phys. Rev. B 79, 214516 (2009).
[14] M. Tropeano et al., Phys. Rev. B 81, 184504 (2010).
[15] H. Hodovanets et al., Phys. Rev. B 83, 094508 (2011).
[16] A. Thaler et al., Phys. Rev. B 82, 014534 (2010).
[17] F. Rullier-Albenque et al., Phys. Rev. B 81, 224503 (2010).
[18] M. S. Torikachvili, S. L. Bud’ko, N. Ni, and P. C. Canfield, Phys. Rev. Lett. 101, 057006 (2008).
[19] S. A. J. Kimber et al., Nature Materials 8, 471 (2009).
[20] E. Colombier, et al., Phys. Rev. B 79, 224518 (2009).
[21] Chang Liu et al., Phys. Rev. Lett. 102, 167004 (2009).
[22] Chang Liu et al., Nature Physics 6, 419 (2010); Phys. Rev. B, 84, 020509(R) (2011).
[23] S. Thirupathaiah et al., Phys. Rev. B 84, 014531 (2011)
[24] Z. R. Ye, et al., arXiv:1105.5242v1 (2011).
[25] R. S. Dhaka et al., Phys. Rev. Lett., 107, 267002 (2011).
[26] N. Ni et al., Phys. Rev. B 78, 214515 (2008).
[27] G. Kresse and J. Hafner, Phys. Rev. B 47, 558 (1993); ibid 49, 14251 (1994).
[28] G. Kresse and J. Furthmüller, Phys. Rev. B 54, 11169 (1996); Comput. Mat. Sci. 6, 15 (1996).
[29] P. E. Blöchl, Phys. Rev. B 50, 17953 (1994).
[30] G. Kresse and D. Joubert, Phys. Rev. B 59, 1758 (1999).
[31] J. P. Perdew et al., Phys. Rev. B 23, 5048 (1981); Phys. Rev. Lett. 77, 3865 (1996); Phys. Rev. Lett. 78, 1396 (1997).
[32] M. Rotter et al., Phys. Rev. B 78, 020503 (2008).
[33] T. Kondo et al., Phys. Rev. B 81, 060507(R) (2010).
[34] S. L. Bud’ko et al., Phys. Rev. B 79, 054525 (2009).