Neutron powder diffraction study on the iron-based nitride superconductor ThFeAsN

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Neutron powder diffraction study on the iron-based nitride superconductor ThFeAsN

Huican Mao1,2, Cao Wang3, Helen E. Maynard-Casely4, Qingzhen Huang5, Zhicheng Wang6, Guanghan Cao6,7, Shiliang Li1,2,8 and Huiqian Luo1(a)

1 Beijing National Laboratory for Condensed Matter Physics, Institute of Physics, Chinese Academy of Sciences Beijing 100190, China
2 University of Chinese Academy of Sciences - Beijing 100049, China
3 Department of Physics, Shandong University of Technology - Zibo 255049, China
4 Australian Centre for Neutron Scattering, Australian Nuclear Science and Technology Organisation Lucas Heights NSW-2232, Australia
5 NIST Center for Neutron Research, National Institute of Standards and Technology Gaithersburg, MD 20899-6102, USA
6 Department of Physics and State Key Lab of Silicon Materials, Zhejiang University - Hangzhou 310027, China
7 Collaborative Innovation Centre of Advanced Microstructures - Nanjing 210093, China
8 Collaborative Innovation Center of Quantum Matter - Beijing 100190, China

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Abstract – We report neutron diffraction and transport results on the newly discovered superconducting nitride ThFeAsN with \( T_c = 30 \) K. No magnetic transition, but a weak structural distortion around 160 K, is observed by cooling from 300 K to 6 K. Analysis on the resistivity, Hall transport and crystal structure suggests that this material behaves as an electron optimally doped pnictide superconductor due to extra electrons from nitrogen deficiency or oxygen occupancy at the nitrogen site, which, together with the low arsenic height, may enhance the electron itinerancy and reduce the electron correlations, thus suppressing the static magnetic order.

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Introduction. – Unconventional superconductivity in iron pnictides or chalcogenides has been intensively investigated since the ZrCuSiAs-type crystalline \( \text{LaFeAsO}_{1-x}\text{F}_x \) (1111 family) with transition temperature \( T_c = 26 \) K was discovered in 2008 [1]. Usually, the iron-based superconductivity emerges from the proximity to a three-dimensional antiferromagnetism [2], for example, \( \text{LaFeAsO}_{1-x}\text{F}_x \) [3,4], \( \text{BaFe}_{2-x}\text{Co}_x\text{As}_2 \) (122 family) [5], \( \text{NaFe}_{1-x}\text{Co}_x\text{As} \) (111 family) [6], \( \text{FeTi}_{1-x}\text{Se}_x \) (11 family) [7,8] and \( \text{Ca}_{1-x}\text{La}_x\text{FeAs}_2 \) (112 family) [9,10], etc. In some special cases, the conductivity is very sensitive to the ion deficiency, such as \( \text{LaFeAsO}_{1-x}\text{F}_x \) [11], \( \text{Li}_{1-x}\text{FeAs} \) [12] and \( \text{K}_{0.8}\text{Fe}_{2.4}\text{Se}_2 \) [13], or the stoichiometric composition is naturally superconducting, such as \( \text{Sr}_2\text{VO}_3\text{FeAs} \) (21311 family) [14], \( \text{Ca}_{10}(\text{Fe}_3\text{Pt}_8)(\text{Fe}_2\text{As}_2)_{5} \) (10-3-8 family) [15], \( \text{KFe}_2\text{As}_2 \) [16], \( \text{RbEuFe}_4\text{As}_4 \) (1144 family) [17], \( \text{FeSe} \) [18], etc. Even so, in most of the above families, spin fluctuations persist and intimately interplay with superconductivity, while in most parent compounds, a magnetic phase transition always occurs beneath the symmetry-breaking structural transition at low temperature except for the FeSe system [19].

Specifically for the 1111 family, superconductivity can be induced among the antiferromagnetically ordered oxide (e.g., \( \text{LaFeAsO} \)) [1], fluoride (e.g., \( \text{CaFeAsF} \)) [20,21] and hydride (e.g., \( \text{LaFeAsO}_{1-x}\text{H}_x \)) [22,23] under the chemical substitution on any atomic site. Recently, the first nitride iron pnictide superconductor ThFeAsN, containing layers with nominal compositions \( \text{[Th}_2\text{N}_2 \) and \( \text{[Fe}_2\text{As}_2 \) (fig. 1(a)), has been discovered, with \( T_c = 30 \) K.

(a)E-mail: hqluo@iphy.ac.cn
for a nominally undoped compound [24]. Although the first-principle calculation of ThFeAsN indicates that the lowest-energy magnetic ground state is the stripe-type antiferromagnetic state [25,26], the normal-state resistivity shows no obvious anomaly but rather a metallic behavior. In principle, the N-N bond covalency may lower the effective nitrogen valence and lead to an internal charge transfer. Such self-doping effect may be responsible for the lowest-energy magnetic ground state is the stripe-type antiferromagnetic state [25,26], the normal-state resistivity shows no obvious anomaly but rather a metallic behavior. In principle, the N-N bond covalency may lower the effective nitrogen valence and lead to an internal charge transfer. Such self-doping effect may be responsible for the lowest-energy magnetic ground state is

\[
\begin{align*}
\chi & = 30 K \\
\lambda & = 2.41 \text{ Å} \\
\theta & = 15-136 \text{ degrees}
\end{align*}
\]

Table 1: Crystallographic data of ThFeAsN at 6K.

<table>
<thead>
<tr>
<th>Atom</th>
<th>Wyckoff</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>Uiso</th>
</tr>
</thead>
<tbody>
<tr>
<td>Th</td>
<td>2c</td>
<td>0.25</td>
<td>0.25</td>
<td>0.1380(3)</td>
<td>0.3768</td>
</tr>
<tr>
<td>Fe</td>
<td>2b</td>
<td>0.75</td>
<td>0.25</td>
<td>0.5</td>
<td>0.0934</td>
</tr>
<tr>
<td>As</td>
<td>2c</td>
<td>0.25</td>
<td>0.25</td>
<td>0.6522(5)</td>
<td>0.0884</td>
</tr>
<tr>
<td>N</td>
<td>2a</td>
<td>0.75</td>
<td>0.25</td>
<td>0</td>
<td>0.7005</td>
</tr>
</tbody>
</table>

angle 2θ range 15–136 degrees. All these diffraction patterns were refined with the Rietveld method within the program FullProf [28], and the temperature dependence of the structure parameters, such as lattice constant, full width at half-maximum (FWHM) of the (112) peak, As-Fe-As bond angles, bond length and the ionic (Th, Fe) height from the As layer, was obtained by assuming 100% occupancy of ThFeAsN. The temperature-dependent resistivity from 2 K to 300 K was measured by the standard 4-probe method, and the Hall coefficient ($R_H$) was measured by the transverse resistance under sweeping magnetic fields from −6 T to +6 T over the temperature range 50K–300 K on a Quantum Design Physical Property Measurement System (PPMS).

**Result and discussion.** The raw data of neutron diffraction patterns are presented in fig. 1. To quantitatively compare the neutron diffraction peaks, we have normalized the maximum neutron counts at 300 K to be the same as 6 K and the difference between 6 K and 40 K is obtained by direct subtraction from the raw data after normalization by monitor counts (fig. 1(c)). Most of the reflections can be indexed by a tetragonal phase in a ZrCuSiAs-type structure with the space group $P4/nmm$ ($a = b = 4.0414 \text{ Å}, c = 8.5152 \text{ Å}$), except for a very small amount of Fe₃As₁₂ and FeAs impurity phases (< 5 %), which are not detected in the X-ray diffraction experiment [24,27]. Figure 1(d) shows the Rietveld refinement of the neutron diffraction patterns at 6 K by assuming 100% occupancy of ThFeAsN. The parameters for the quality of this fitting are: profile factor $R_p = 6.47(2)$%, weighted profile factor $Rwp = 6.84(3)$%, and reduced $\chi^2$ of 2.93(1). All crystallographic parameters listed in table 1 are mostly consistent with previous X-ray diffraction results [24]. Since there is no difference between 6 K and 40 K data sets and all reflections are identified arising from the nuclear structure from 6 K up to 300 K, we thus conclude that there is no magnetic order in ThFeAsN, consistently with the Mössbauer spectroscopy results [27].

We also performed Rietveld refinements with the FullProf program considering other possibilities. The best fitting results are found in three cases: fully occupied compound (ThFeAsN), (2.7 ± 0.8)% N deficiency...
Neutron powder diffraction of ThFeAsN

Fig. 2: (Colour online) Temperature dependence of (a) lattice constant $a$, (b) lattice constant $c$, (c) FWHM of (112) Bragg peak, (d) As-Fe-As bond angles $\alpha$ and $\beta$, (e) anion (Th, Fe) height from the As layer, (f) bond length of Th-As and Fe-As in ThFeAsN.

(ThFeAsN$_{0.97}$) or $(7 \pm 2\%)$ O occupancy at N site (ThFeAsN$_{0.93}$O$_{0.07}$), among which there is no remarkable difference of the reliability of factors between ThFeAsN. For ThFeAsN$_{0.97}$, the factors are $R_p = 6.45(2)\%$, $R_{wp} = 6.82(2)\%$, and $\chi^2 = 2.92(2)$, and for ThFeAsN$_{0.93}$O$_{0.07}$ they are $R_p = 6.45(1)\%$, $R_{wp} = 6.82(2)\%$, and $\chi^2 = 2.93(1)$. Other cases such as Th or As deficiency give much worse factors, even failure of the fitting. Although the N deficiency or O occupancy as well as the exact proportions of them cannot be distinguished precisely within the data quality and instrument resolution, the N deficiency is more possibly indicated by the refinements due to a larger neutron cross-section of nitrogen.

To search for a possible structural transition in ThFeAsN, we have collected diffraction patterns over the temperature range from 300 K to 6 K. All of them can be well refined with the tetragonal phase of ThFeAsN, we have collected diffraction patterns over the temperature range from 300 K to 6 K. All of them can be well refined with the tetragonal phase of ThFeAsN, as shown in fig. 3. The resistivity data shows a metallic behavior above $T_c$ without any clear anomaly expected for a magnetic or structural transition. It can be fitted by an empirical power law $ho = \rho_0 + AT^n$ in a wide temperature range (40 K–150 K) with the exponent $n$ approximate to 1.30, suggesting possible non-Fermi-liquid behaviors governed by quantum fluctuations similar to systems around the optimal doping level [5,31–33]. Interestingly, the derivative $d\rho/dT$ (insert of fig. 3(a)) also shows a broad hump around 160 K, corresponding to the lattice distortion found in fig. 2. The Hall coefficient ($R_H$), determined by the slope of the field dependence of the Hall resistivity, is always negative and increases with temperature. By comparing with (La, Sm)FeAsO$_{1-x}$F$_x$ [34,35], the magnitude of $R_H$ in ThFeAsN suggests it is already doped by electrons, which are probably introduced by the N deficiency or O occupancy shown in the powder diffraction refinement, or the reduced valence of nitrogen as discussed before.

By summarizing the literature that has noted the arsenic height $h_{FeAs}$, As-Fe-As bond angle $\alpha$ and optimized $T_c$ for each system of iron-based superconductors, it is found that there is a close relationship between the local structure of the FeAs$_4$ tetrahedron and superconducting $T_c$, as shown by the light blue belt in fig. 4 [36–38]. Obviously, the maximum $T_c$ was achieved when the FeAs$_4$ tetrahedron is perfectly regular, with the bond angle of 109.47 degrees [39]. Interestingly, the data obtained from ThFeAsN agrees very well with other optimally doped compounds (red stars in fig. 4). Again, these results suggest ThFeAsN is nearly in an optimized superconducting state with lots of itinerant electrons and away from the “parent” compound. Another fact should be
noticed, i.e., the $h_{Fe-As}$ of ThFeAsN (1.2964 Å) is lower than those of LaFeAsO (1.3166 Å) [3,4] and SrFeAsF (1.3710 Å) [20,21]. The closer distance of Fe-As will surely favor the electron hopping, thus reducing the electron correlations and orbital order controlled by the Hund coupling $J_H$ within one atomic site [40,41]. This is a reasonable explanation for the absence of magnetic order, structural transition, and resistivity anomaly in ThFeAsN.

**Summary.** — In summary, we have carried out neutron diffraction experiments on synthesized ThFeAsN over a temperature range from 6 K to 300 K. It is seen that there is neither structural transition nor magnetic transition existing in ThFeAsN, but a structural distortion may occur around 160 K. By comparing with other iron pnictides, we conclude that ThFeAsN may be placed near the optimal doping level and have reduced electron correlations.

**REFERENCES**


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Fig. 4: (Colour online) (a) Anion height dependence of optimal $T_c$ for Fe-based superconductors. (b) Optimal $T_c$ vs. bond angle $\alpha$. The light cyan belts are from other optimally doped systems [38], and the red stars indicate our ThFeAsN compound, respectively.