# scientific reports

Check for updates

## **Study on interaction mechanism OPEN of diferent atomic ratio of neodymium, arsenic and iron**

**Juncheng Mao1,2, Run Huang1,2**\***, Chenghui Fu1,2, Xiaodong Lv3 , Lihua He1,2 & Jinzhu Zhang1,2**\*

**In this study, neodymium and arsenic were sealed into industrial pure iron cylinders at a temperature of 1223 K for 50 h. The interaction mechanism of the Nd–Fe–As system at various atomic ratios was investigated by optical microscopy, X-ray difractometry, and scanning electron microscopy. Binary compounds Fe12As5, NdAs, Fe2As, and Fe17Nd2 were the main products formed, with traces of NdFeAs compounds. In addition, at high temperatures, As content afected the difusion of Fe atoms; the difusion of Fe increased with an increase in the atomic ratio. Furthermore, the difusion ability of Nd was weaker than that of As. The major difusion mechanism of Nd was through the Fe atomic vacancy mechanism. As mainly bind to Fe to form Fe and As compounds. The formation of ternary compounds was confrmed by laboratory experiments and mismatch calculations.**

The source of residual arsenic in steel is mainly primary iron ore. Compared with iron, arsenic has a weaker oxidation potential, so it is difcult to remove arsenic from steel in the entire steelmaking process. China not only has a large amount of iron ore containing arsenic, but with the development of social economy, the scrap cycle of steel is gradually shortened. It is estimated that by 2050, the comprehensive utilization ratio of scrap steel will reach 80%<sup>1</sup>. Arsenic, as a deleterious element commonly occurring in steel, enriched at the grain boundary, which significantly deteriorate impact toughness, cold brittleness, and hot-working property of steel<sup>[2](#page-6-1)</sup>. In actual production, high-quality iron ore or molten iron can be mixed<sup>3</sup>, based on the reduction theory, steel companies add calcium to remove arsenic from molten steel<sup>[4–](#page-6-3)[8](#page-6-4)</sup>. But the former is only an emergency solution, the latter is easy to form large spherical oxides, both production processes have drawbacks. Rare earth elements (including 15 lanthanide elements, scandium, and yttrium<sup>9-[11](#page-6-6)</sup> in the third subgroup of the periodic table) are widely used for the purification, metamorphosis, and alloying of metals<sup>12,13</sup>. The unsaturated outer electronic layer of rare earth elements exhibit unique structure and strong chemical activity, which can burst out a variety of electron energy levels and exhibit high "vitality" in its external performance. Due to their active chemical properties, unique electrical and magnetic properties, rare earth elements can react with As (with a low melting point) to form compounds with high melting point, thus improving the thermoplastic and mechanical properties of steel. Rare earth elements are regarded as a treasure of new materials.

According to the iron-neodymium (Fe-Nd) binary phase diagram<sup>14</sup>, the stable compounds that can be formed between Fe and Nd include  $Fe_{17}Nd_2$  and  $Fe_2Nd$ , and according to the Fe-As binary phase diagram, the stable compounds that can be formed between Fe and As include  $Fe_2As$ ,  $Fe_3As_2$ ,  $FeAs$ , and  $FeAs_2^{15}$  $FeAs_2^{15}$  $FeAs_2^{15}$ , in a study on Nd and As compounds, the binary diagram<sup>[16](#page-6-11)</sup> shows that  $Nd<sub>3</sub>As$ , NdAs and NdAs<sub>2</sub>. Generally, the maximum solubility of As in Fe is approximately 10% at 1113  $K^{17}$  $K^{17}$  $K^{17}$ . However, the solubility decreases with a decrease in temperature and reduces to below 5% at room temperature<sup>18-20</sup>. For Re-Fe-As ternary system, the main products including REFe<sub>4</sub>As<sub>12</sub>, REFe<sub>2</sub>As<sub>2</sub> (RE=La, Nd, Sm) have been widely reported. Different atomic ratios of cerium (lanthanum), Fe, and As form the ternary compound  $RE_{12}Fe_{57.5}As_{41}$  (RE = La, Ce) and FeAs at 1173 K<sup>[21](#page-7-1),[22](#page-7-2)</sup>, whereas the Re-Fe-As ternary system forms  $La_{10}Fe_{50}As_{40}^{23}$  $La_{10}Fe_{50}As_{40}^{23}$  $La_{10}Fe_{50}As_{40}^{23}$  at 1223 K. In recent years, one of products of RE-Fe-As ternary system named  $EuFe_2As_2^{24-28}$  has attracted significant attention. Xie $^{29}$ and Fu $^{30}$  investigated the interaction of Nd–Fe–As system at high temperatures and found that the formation of the ternary compound NdFeAs depends on the formation of NdAs and FeAs<sub>2</sub>.

<sup>1</sup>School of Materials and Metallurgy, Guizhou University, Guiyang 550025, People's Republic of China. <sup>2</sup>Guizhou Province Key Laboratory of Metallurgical Engineering and Energy Process Saving, Guiyang 550025, People's Republic of China. <sup>3</sup>College of Materials Science and Engineering, Chongqing University, Shapingba District, No.174 Shazheng Street, Chongqin 400044, People's Republic of China.<sup>⊠</sup>email: rhuang@gzu.edu.cn; jzzhang@ gzu.edu.cn



<span id="page-1-0"></span>Figure 1. Schematic of the barrel-shaped cylinder and the screw plug.



<span id="page-1-1"></span>**Table 1.** Masses of Nd and As and the atomic ratio of Nd:As for sample preparation.



<span id="page-1-2"></span>**Table 2.** Experimental heating process.

Therefore, in this study, the different atomic ratio of Nd:As were sealed in a cylinder block, which was specially processed using industrial pure Fe by melting, infiltration, and diffusion. The interaction between Nd, Fe, and As at high temperature and the mechanism for the generation of ternary compounds was investigated using metallographic microscope, scanning electron microscope (SEM), and X-ray diffraction (XRD). The generation of ternary compounds was partly confrmed by calculations and laboratory experiments.

#### **Materials and methods**

Figure [1](#page-1-0) shows the barrel-shaped cylinder composed of industrial pure Fe; its principal chemical composition (mass fraction) is as follows: 0.002% C, 0.02% Mn, 0.006% P, 0.004% S, 0.005% Al, and 99.95% Fe. Before the flling operation, the oxide layer on the Nd surface was removed, Nd metal block (purity>99.9%) and As block (diameter < 1 mm, see Table [1](#page-1-1)) were flled into the industrial pure Fe cylinder block at various atomic ratios (1:1,1:2,1:3), the screw plug was welded by arc welding, and a high temperature sealant was applied to the weld to ensure it is properly sealed. Subsequently, The industrial pure Fe cylinder block was placed in a closed SRJK-2-9 tube resistance furnace and heated under high purity argon atmosphere. The experimental heating process is shown in Table [2,](#page-1-2) and it depended on the vapor pressure of As. Afer the heating process, the temperature of the furnace was reduced to room temperature (30 °C). Subsequently, argon fow into the furnace was stopped and the cylinder sample was taken out. Then, the outer side of the cylinder block was marked away from its bottom (at a distance of 16 mm); in the radial direction, it was sawed and divided into two parts, one of which was processed into metallographic samples and the other part was used for XRD analysis. The phase composition of the samples was analyzed using a PHILIPS X'-Pert PRO difractometer, and the test parameters are as follows: Copper target,  $\lambda$  = 0.154056 nm, 40 kV operating voltage, 2°/min scanning speed.

#### **Results**

**Metallographic analysis.** Figure [2](#page-2-0) shows the metallographic images of the samples at various atomic ratios under an optical microscope. At a constant heating temperature and holding time, three types of contrast (grayish, gray, and black) were observed at diferent atomic ratios. Te grayish part indicates the collective part of the cylinder, most of which was the industrial pure Fe, the gray part has a higher proportion of As, and the

2





 $(a)$ 

 $(b)$ 

<span id="page-2-0"></span>**Figure 2.** Metallographic pictures of atoms at diferent atomic ratio, (**a**) Pictures of the cylinder block's neighboring area, (**b**) Pictures around its core area.



 $\triangle$  NdAs (PDF#: 01-074-0623)  $\nabla \text{Fe}_{12}\text{As}_{5}$  (PDF#: 01-077-0104)  $\blacktriangledown$  Fe<sub>17</sub>Nd<sub>2</sub> (PDF#: 00-048-1751)  $\bullet$  Fe<sub>2</sub>As (PDF#: 00-009-0095)  $\circ$   $\alpha$ -Fe (PDF#: 03-06504150)

<span id="page-2-1"></span>

black part has a higher proportion of Nd. Figure [2](#page-2-0)a shows the neighboring area of the cylinder block, which indicates that the entire area was divided into two diferent parts: the cylinder matrix on the lef and the core component on the right. In addition, inhomogeneous granular structures were formed at all atomic ratios (the right area), and they contained the same white liner as the cylinder matrix. With an increase in the atomic ratios, the symmetry of the entire granular structure increased and then decreased. In addition, with an increase in the difusion of the grayish area, the difusion moved farther away from the edge, which consequently reduced the difusion. Particularly, with a decrease in the distance between the gray area and the edge of the boundary, the proportion of the gray area reduced. At high temperatures, the Fe atom difused into the core area of the sample, while the Nd atom difused into the matrix area, and the Fe and As atoms difused together, and consequently, Fe atoms gradually formed a circular structure.



**Figure 4.** Backscattered electron maps of the sample at various atomic ratios (**a**) 1:1, (**b**) 1:2, (**c**) 1:3.

<span id="page-3-0"></span>

<span id="page-3-1"></span>**Table 3.** EDS analysis results of diferent atomic ratio samples (at/%).

**Phase analysis.** For the phase analysis, the sample from the core area was ground in a mortar into a pow-der. Subsequently, the phase of the samples was characterized by XRD analysis, as shown in Fig. [3.](#page-2-1) The main difraction peaks observed in the XRD spectra could be attributed to the formation of fve types of compounds  $(F_{e_12}As_5, Fe_2As, NdAs, Fe_1rNd_2, and \alpha-Fe)$  in the ternary system of the high-temperature fusion samples. When the atomic ratio of Nd and As was 1:1 and 1:2, the intensity of the difraction peaks was stable. During the experiment, As sublimed at high temperatures, while the highly reactive Nd reacted with As. Consequently, the number of compounds (NdAs) increased. As the experiment progressed, a small amount of As difused into the external matrix of the cylinder block, and Fe difused to its core area and reacted with As, forming As compounds (Fe<sub>2</sub>As).

XRD spectra of Nd–Fe–As powder compounds in the ternary system were not found in relevant literature. However, the energy dispersive spectroscopy (EDS) analysis suggests the formation of the ternary compounds (NdFeAs).

SEM analysis was carried out on the reduced sample, and EDS analysis was employed to analyze the image at diferent contrasts. As shown in Fig. [4](#page-3-0), the samples are mainly distributed in four types of contrast phase (black, dark gray, light gray, and white), which are identifed by A, B, C, and D, and each contrast phase tissue was investigated using EDS analysis; the results are shown in Table [3.](#page-3-1) The microstructure of the black, dark gray, and light gray contrast had an irregular shape. In addition, the gray and light gray components were mainly composed of Nd and As, combined with the XRD spectra of Nd and As, this result indicates that the contrast phase structure was a NdAs crystal structure, whereas the black contrast phase structure was mostly saturated As α-Fe solid solution.

As discussed above, the products of Nd and As at different atomic ratios (1:1, 1:2, 1:3) contained  $Fe_{12}As_{5}$ ,  $Fe<sub>2</sub>As, NdAs, and Fe<sub>17</sub>Nd<sub>2</sub> in all the ternary systems. However, the amount of different phases changes with an$ increase in the atomic ratios, and the various contrast phase structures show a symbiotic relationship.

**Diffusion analysis of samples.** Figure [5](#page-4-0) shows the line-scanning atlas analysis of the transition areas of Nd and As at diferent atomic ratios. According to the graph, the three elements (Nd, Fe, As) were in a continuous distribution in a banded gradient. In addition, the amount of Nd in the white contrast region was signifcantly higher than that in other regions. Furthermore, the amount of Fe in the black contrast region was the highest, and As existed in all the contrast phases.

During the experiment, Fe difused into the central region of the cylinder block, whereas the difusion activation energy of As was lower than that of  $Fe^{31}$ . In addition, As diffused via the formation of Fe and As compounds, and Nd difused via its vacancy mechanism and the formation of the Fe and As compounds. Because the chemical potential of Nd increased with an increase in its content, and the solubility of As was maintained at 10% during experimental temperature, vacancies were produced during the sublimation process. The interaction between As and Nd produced NdAs compounds, and the remaining Nd and Fe formed  $Fe_{17}Nd_2$  compounds to fill the vacancies. In addition, Fe vacancies were formed by thermal vibration; consequently, these vacancies provided a condition for the diffusion of  $Nd^{32}$ . Furthermore, the grain size gradually decreased, thus increasing the grain boundary area. Simultaneously, the grain boundary vacancies increased accordingly. The combined effect of the generated vacancies, the Nd-generated solute-vacancy compounds, and the enhancement of the thermal difusion rate leads to the segregation of Nd at the grain boundary.

4



<span id="page-4-0"></span>**Figure 5.** Sweep spectrum of the edge line of the cylinder samples at diferent atomic ratio.



<span id="page-4-2"></span>**Figure 6.** Crystallographic relationships of the  $Fe<sub>12</sub>As<sub>5</sub>$  (0001) and NdAs (111) planes.

**Planar mismatch calculation and analysis.** According to the planar mismatch theory, the calculation result of the mismatch of two planes should be less than 6% to achieve good heterogeneous nucleation; heterogeneous nucleation is expected to occur if the mismatch is greater than 12%. The planar mismatch can be calculated using Eq.  $(1)$  $(1)$  [33.](#page-7-10)

<span id="page-4-1"></span>
$$
\delta_{(hkl)_n}^{(hkl)_s} = \frac{1}{3} \sum_{i=1}^3 \left[ \frac{d_{[uvw]_s}^i \cos \theta - d_{[uvw]_n}^i}{d_{[uvw]_n}^i} \right] \times 100 \tag{1}
$$

where  $\delta$  is the average of the three mismatches between the (hkl), and (hkl)<sub>n</sub> planes;  $d_{[uvw]_s}$  and  $d_{[uvw]_n}$  is the interatomic spacing along  $[uvw]_s$ , and  $[uvw]_n$  respectively; and  $\theta$  is the angle between two corresponding direction.

The atom matching diagram of the surface between the  $Fe_{12}As_{5}$  (0001) and NdAs (111) planes is illustrated in Figs. [6](#page-4-2) and [7](#page-5-0). The former is indicated by "X" and the latter is indicated by "O". The crystal parameters involved in the calculations are listed in Table  $4^{34}$  $4^{34}$  $4^{34}$ , and the specific calculation data are shown in Table [5.](#page-5-2) The calculation



<span id="page-5-0"></span>**Figure 7.** Diagram of plane and crystal direction of NdAs (111).

		<b>Lattice parameters</b> (25 °C, nm)		
Compounds	<b>Crystal</b> system	a <sub>0</sub>	b <sub>0</sub>	c <sub>0</sub>
Fe <sub>12</sub> As <sub>5</sub>	Hexagonal	0.6786		1.6301
<b>NdAs</b>	Cubic	0.5987		
Fe <sub>2</sub> As	Tetragonal	0.3632		0.5981
$Fe_{17}Nd_{2}$	Hexagonal	0.8574		1.2464
<b>NdFeAs</b>	Tetragonal	0.39655		0.8575

<span id="page-5-1"></span>**Table 4.** Crystallographic parameters of possible compounds of Nd and As in steel.



<span id="page-5-2"></span>Table 5. Calculation details of the lattice between Fe<sub>12</sub>As<sub>5</sub> and NdAs compounds.

result of the two mismatched planes was 17.86%, indicating that  $Fe_{12}As_5$  cannot be used as the effective heterogeneous nucleation core in the formation NdAs.

The calculation results of the mismatch are listed in the Table [6](#page-6-14). The mismatch between Fe<sub>2</sub>As and NdFeAs was 8.33%, indicating the high probability for Fe<sub>2</sub>As to effectively act as the heterogeneous nucleation cores for the formation of ternary NdFeAs compound.



<span id="page-6-14"></span>

### **Conclusions**

- (1) When the atomic ratios of Nd and arsenic were between 1:1 and 1:3 at a maximum temperature of 1223 K for 50 h,  $Fe<sub>12</sub>As<sub>5</sub> Fe<sub>2</sub>As, NdAs, and Fe<sub>17</sub>Nd<sub>2</sub> were formed. In addition, with an increase in the atomic ratios$ of Nd and As, the formation of the NdAs compounds decreased and the formation of Fe<sub>2</sub>As compounds increased (Fe<sub>2</sub>As was generated during the diffusion of As toward the cylinder block).
- (2) In the ternary system (Nd–Fe–As), the difusion of Fe was dependent on the amount of As. In addition, the Fe atom difused toward the core of the cylinder block, and its amount decreased with increase in the depth of its difusion. Furthermore, with an increase in the proportion of Nd and As, the difusion of As into the external matrix of the cylinder increased.
- (3) The EDS spectra suggest the formation of ternary compounds (NdFeAs), and the mismatch calculation indicated that the Fe<sub>2</sub>As can act as effective heterogeneous nucleation cores for the formation of ternary compounds (NdFeAs).

Received: 12 November 2020; Accepted: 5 February 2021 Published online: 19 February 2021

#### **References**

- <span id="page-6-0"></span>1. Pauliuk, S., Wang, T. & Müller, D. B. Moving toward the circular economy: the role of stocks in the Chinese steel cycle. *Environ. Sci. Technol.* **46**, 148–154. <https://doi.org/10.1021/es201904c>(2011).
- <span id="page-6-1"></span>2. Cheng, H. J., Wang, F. M., Li, C. R., Tan, Y. P. & Wang, J. J. Efect of arsenic on microstructure and mechanical properties of 45 steel. *Heat Treat. Met.* **35**, 33–38. <https://doi.org/10.13251/j.issn.0254-6051.2010.02.033> (2010).
- <span id="page-6-2"></span>3. Björkman, B., Samuelsson, C. Recycling of steel. In: *Handbook of Recycling: state-of-the-art fot practitioners, analysts, and scientists. Elsevier, Amsterdam* (eds Worrell, E and Reuter M. 65–83. [https://doi.org/10.1016/B978-0-12-396459-5.00006-4\(](https://doi.org/https://doi.org/10.1016/B978-0-12-396459-5.00006-4 )2014).
- <span id="page-6-3"></span>4. Nakamura, Y., Tokumitsu, N., Harashima, K. & Segawa, K. Refining of 18% Cr–8% Ni steel with Ca–CaF<sub>2</sub> solution. *Trans. ISIJ.* **16**, 623–627. <https://doi.org/10.2355/isijinternational1966.16.623>(1976).
- 5. Kitamura, K., Takenouchi, T. & Iwanami, Y. Removal of impurities from molten steel by CaC2. *Tetsu-to-Hagané.* **71**, 220–227. [https](https://doi.org/10.2355/tetsutohagane1955.71.2_220) [://doi.org/10.2355/tetsutohagane1955.71.2\\_220](https://doi.org/10.2355/tetsutohagane1955.71.2_220) (1985).
- 6. Li, W. B., Bao, Y. P., Wang, M. & Lin, L. Experimental study on arsenic removal of molten steel with diferent Ca alloy. *J. Eng. Sci.* **38**, 484–493.<https://doi.org/10.13374/j.issn2095-9389.2016.04.006> (2016).
- 7. Fu, B., Xun, Z. L., Wu, G. L., Wu, L. J., Wu, Y. J., Ke, C. Experimental study on the dearsenization of hot metal with  $CaC_2-CaF_2$ slag. *Chin. J. Process Eng.* **10**, 146–149 (2010).
- <span id="page-6-4"></span>8. Dong, Y. C., Shi, Z. P., Zhang, L. M., Peng, Y. Q. & Hong, Y. R. Study on dearsenization of molten iron. *Iron Steel.* **19**, 1–7. [https://](https://doi.org/10.13228/j.boyuan.issn0449-749x.1984.09.001) [doi.org/10.13228/j.boyuan.issn0449-749x.1984.09.001](https://doi.org/10.13228/j.boyuan.issn0449-749x.1984.09.001) (1984).
- <span id="page-6-5"></span>9. Zhang, F., Li, Y. & Li, X. Current status of rare-earth resources development and utilization at home and abroad and suggestions for rare-earth management in China. *Mod. Min.* **12**, 11–14. <https://doi.org/10.3969/j.issn.1674-6082.2018.12.003> (2007).
- 10. Dutta, T. *et al.* Global demand for rare earth resources and strategies for green mining. *Environ. Res.* **150**, 182–190. [https://doi.](https://doi.org/10.1016/j.envres.2016.05.052) [org/10.1016/j.envres.2016.05.052](https://doi.org/10.1016/j.envres.2016.05.052) (2016).
- <span id="page-6-6"></span>11. Zhang, Y., Liu, C. S., Gao, L. F., Zhang, Z. G. & Zhang, P. Marine authigenic deposits mineral-new felds for the development for the development of rare-earth resources. *Adv. Mater. Res.* **291–294**, 1748–1751. [https://doi.org/10.4028/www.scientifc.net/](https://doi.org/10.4028/www.scientific.net/AMR.291-294.1748) [AMR.291-294.1748](https://doi.org/10.4028/www.scientific.net/AMR.291-294.1748) (2011).
- <span id="page-6-7"></span>12. Li, W. C., Lin, Q., Ye, W. & Zhang, C. Y. Kinetics of rare-earth efect on the low carbon steel containing arsenic. *J. Beijing Univ. Iron Steel Technol.* **2**, 61–67. <https://doi.org/10.13374/j.issn1001-053x.1983.02.025>(1983).
- <span id="page-6-8"></span>13. Du, T. Physical-chemistry efect of rare-earth elements on metallic material. *Act Metall. Sin.* **33**, 69–77 (1997).
- <span id="page-6-9"></span>14. Zhang, F. Z. Te prospect of the application of rare-earth in iron and steel from the phase diagram of rare-earth. *Iron Steel.* **8**, 66–75 (1986).
- <span id="page-6-10"></span>15. Brandes, E. A. & Brook, G. B. *Smithells Metals Reference Book* (Butterworth-Heinemann, Oxford, 1983).
- <span id="page-6-11"></span>16. Kobzemko, G. F., Chernogorenko, V. B., L'vov, S. N., Lesnaya, M. I. & Lynchak, K. A. Equilibrium diagram and properties of alloys of the As–Nd system. *ZhurnalNeorgancicheskoiKhimii.* **20**(8), 2169–2173 (1975).
- <span id="page-6-13"></span><span id="page-6-12"></span>17. Yin, G. Te distribution of arsenic in steel. *Gangtie* **16**, 20–28.<https://doi.org/10.13228/j.boyuan.issn0449-749x.1981.02.004>(1981). 18. Duan, S., Chen, X., Yang, B., Liu, D. & Dai, Y. Ab-initio molecular dynamics simulation of thermal decomposition of arsenic-iron
- compounds in vacuum. *Chin. J. Vac. Sci. Technol.* **34**, 1128–1134. <https://doi.org/10.13922/j.cnki.cjovst.2014.10.23> (2014). 19. Hu, Q. Q., Zhou, J., Zhang, X. Y., Yan, Z. & Yang, Y. T. Comparison of As(V)removal by three diferent types of ferric iron adsor-
- bents. *Water Purif. Technol.* **33**, 71–74.<https://doi.org/10.15890/j.cnki.jsjs.2014.06.013>(2014).

7

- <span id="page-7-0"></span>20. Todorov, I. et al. CaFe<sub>4</sub>As<sub>3</sub>: a metallic iron arsenide with anisotropic magnetic and charge-transport properties. *J. Am. Chem. Soc.* **131**, 5405.<https://doi.org/10.1021/ja900534h> (2009).
- <span id="page-7-1"></span>21. Stoyko, S. S., Blanchard, P. E. R., Mar, A. Ternary rare-earth iron arsenide RE<sub>12</sub>Fe<sub>575</sub>As<sub>41</sub>(RE=La, Ce). *Inorganic Chemistry*. **49**, 2325–2333. <https://doi.org/10.1021/ic902231b>(2010).
- <span id="page-7-2"></span>22. Zhang, J. Z., Dou, S. T. Study on interaction between Cerium and Arsenic. *Advanced Materials Research.* **194–196**, 1231–1234. [https://doi.org/10.4028/www.scientifc.net/AMR.194-196.1231](https://doi.org/10.4028/www.scientific.net/AMR.194-196.1231) (2011).
- <span id="page-7-3"></span>23. Liu, X., Zhang, J. & Zhu, S. Microstructure and compound developed from La–As–Fe system at 1223 K. *Adv. Mater. Res.* **702**, 145–148. [https://doi.org/10.4028/www.scientifc.net/AMR.702.145](https://doi.org/10.4028/www.scientific.net/AMR.702.145) (2013).
- <span id="page-7-4"></span>24. Sklyarova, A. *et al.* Mössbauer study of hyperfine interactions in EuFe<sub>2</sub>(As<sub>1</sub><sub>x</sub>P<sub>x</sub>)<sub>2</sub> and BaFe<sub>2</sub>(As<sub>1<sub>→x</sub>P<sub>x</sub>)<sub>2</sub>. *J. Magn. Magn. Mater.* **378**,</sub> 327–332. <https://doi.org/10.1016/j.jmmm.2014.11.054>(2015).
- 25. Munevar, J. et al. Magnetism in superconducting EuFe<sub>2</sub>As<sub>14</sub>P<sub>0.6</sub> single crystals studied by local probes. *Solid State Commun*. **187**, 18–22.<https://doi.org/10.1016/j.ssc.2014.02.001>(2014).
- 26. Wu, T. *et al.* Magnetic phase diagram of Eu1−x LaxFe2As2 single crystals. *J. Magn. Magn. Mater.* **321**, 3870–3874. [https://doi.](https://doi.org/10.1016/j.jmmm.2009.07.043) [org/10.1016/j.jmmm.2009.07.043](https://doi.org/10.1016/j.jmmm.2009.07.043) (2009).
- 27. Terashima, T. et al. Pressure-induced antiferromagnetic bulk superconductor EuFe<sub>2</sub>As<sub>2</sub>. *Physica C* 470, S443–S444. [https://doi.](https://doi.org/10.1016/j.physc.2009.10.064) [org/10.1016/j.physc.2009.10.064](https://doi.org/10.1016/j.physc.2009.10.064) (2010).
- <span id="page-7-5"></span>28. Drief, F., Zaoui, A., Kacimi, S. & Merabet, B. Ab-initio investigation of the electronic structure in the superconducting EuFe2(As1−xPx)2. *Physica C Supercond. Appl.* **512**, 22–27. <https://doi.org/10.1016/j.physc.2015.02.045>(2015).
- <span id="page-7-6"></span>29. Xie, W. H., Huang, R., Zhang, J. Z., Li, W. & Yang, Y. Study on the interaction of rare earth element neodymium, iron and arsenic at 1173 K. *J. Funct. Mater.* **49**, 1134–1138 (2018).
- <span id="page-7-7"></span>30. Fu, C. H., Huang, R., Xie, W. H., Luo, J. X. & Zhang, J. Z. Study on high-temperature interaction mechanism of Nd–Fe–As system. *Materials.* **12**, 3060.<https://doi.org/10.3390/ma12193060>(2019).
- <span id="page-7-8"></span>31. Boi, B. I. & Lui, R. J. Difusion in iron-arsenic alloys. *J. Mater. Sci.* **11**, 887–891.<https://doi.org/10.1007/bf00542306> (1976).
- <span id="page-7-9"></span>32. Li, Y. L. *et al.* High-temperature interaction of Ce–Fe–As ternary system. *Mater. Res. Express.* [https://doi.org/10.1088/2053-1591/](https://doi.org/10.1088/2053-1591/ab1198) [ab1198](https://doi.org/10.1088/2053-1591/ab1198) (2019).
- <span id="page-7-10"></span>33. Bramftt, B. L. Te efect of carbide and nitride additions on the heterogeneous nucleation behavior of liquid iron. *Metall. Trans.* **1**, 1987–1995.<https://doi.org/10.1007/BF02642799>(1970).
- <span id="page-7-11"></span>34. Villars, P. Pauling fle. In *Inorganic Solid Phases, SpringerMaterials (Online Database)*, Springer: Heidelberg, Germany, 2012.

#### **Acknowledgements**

Tis research was funded by the Nation Natural Science Foundation of China (Grant No. 51764005 and 52064010), Key Nurturing Projects of Guizhou University (No. [2019]07).

#### **Author contributions**

R.H. conceived and designed the experiment; C.F., L.X., and L.H. contributed to retrieve documents; J.M. collected the data and made the graph; J.M. analyzed the data and wrote the paper. Tis article was checked and revised by J.Z. All authors reviewed the manuscript.

#### **Competing interests**

The authors declare no competing interests.

#### **Additional information**

**Correspondence** and requests for materials should be addressed to R.H. or J.Z.

**Reprints and permissions information** is available at [www.nature.com/reprints.](www.nature.com/reprints)

**Publisher's note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional afliations.

**Open Access** Tis article is licensed under a Creative Commons Attribution 4.0 International  $\boxed{0}$ License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit<http://creativecommons.org/licenses/by/4.0/>.

 $© The Author(s) 2021$