Solid-state synthesis, magnetic and structural properties of interfacial B2-FeRh(001) layers in Rh/Fe(001) films

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Here we first report results of the start of the solid-state reaction at the Rh/Fe(001) interface and the structural and magnetic phase transformations in 52Rh/48Fe(001), 45Rh/55Fe(001), 68Rh/32Fe(001) bilayers from room temperature to 800 °C. For all bilayers the non-magnetic nanocrystalline phase with a B2 structure (nfm-B2) is the first phase that is formed on the Rh/Fe(001) interface near 100 °C. Above 300 °C, without changing the nanocrystalline B2 structure, the phase grows into the low-magnetization modification $\alpha''$ ($M_{S} \approx 825$ emu/cm$^3$) of the ferromagnetic $\alpha'$ phase which has a reversible $\alpha' \leftrightarrow \alpha''$ transition. After annealing 52Rh/48Fe(001) bilayers above 600 °C the $\alpha'$ phase increases in grain size and either develops into $\alpha''$ with high magnetization ($M_{S} \approx 1,220$ emu/cm$^3$) or remains in the $\alpha'$ phase. In contrast to $\alpha'$, the $\alpha'' \leftrightarrow \alpha''$ transition in the $\alpha''$ films is completely suppressed. When the annealing temperature of the 45Rh/55Fe(001) samples is increased from 450 to 800 °C the low-magnetization nanocrystalline $\alpha''$ films develop into high crystalline perfection epitaxial $\alpha''(001)$ layers, which have a high magnetization of ~ 1,275 emu/cm$^3$. $\alpha''(001)$ films do not undergo a transition to an antiferromagnetic $\alpha''$ phase. In 68Rh/32Fe(001) samples above 500 °C non-magnetic epitaxial $\gamma(001)$ layers grow on the Fe(001) interface as a result of the solid-state reaction between the epitaxial $\alpha'$ (001) and polycrystalline Rh films. Our results demonstrate not only the complex nature of chemical interactions at the low-temperature synthesis of the nfm-B2 and $\alpha'$ phases in Rh/Fe(001) bilayers, but also establish their continuous link with chemical mechanisms underlying reversible $\alpha' \leftrightarrow \alpha''$ transitions.

An intriguing feature of the equilibrium diagram of the Fe-Rh system is the existence of the low-temperature transition at $T_{K}^{\alpha'' \rightarrow \alpha'} \approx 100$ °C between antiferromagnetic (AFM) $\alpha''$ and ferromagnetic (FM) $\alpha'$ phases in a narrow (0.48 < x$_{Rh}$ < 0.56) concentration interval in chemically ordered B2-FeRh alloys. This transition is accompanied by an isotropic volume expansion of ~ 1% of the B2-FeRh unit cell, which not only radically changes the magnetic properties, but also changes the entropy and resistivity. The magnetostuctural $\alpha'' \rightarrow \alpha'$ (AFM-FM) transition gives existence to the giant magnetostriction$^5$ large magnetoeresistance$^6,7$ and magnetocaloric effects$^8$ in B2-FeRh alloys. Numerous studies of bulk samples indicate that the starting temperature $T_{K}^{\alpha'' \rightarrow \alpha'}$ and characteristics of the $\alpha'' \rightarrow \alpha'$ transition may be modified by variations the magnetic field$^9,10$, microstructure$^{11}$, thermal treatment$^{12}$, energetic ion irradiation$^{13}$, stress$^{14}$ and hydrostatic pressure$^{15,17}$ and can also be tuned over a wide temperature range (100–600 K) by chemical substitution$^{9,18}$. Although bulk B2-FeRh samples exhibit a sharp transition with a small thermal hysteresis, thin films and nanoparticles often have an incomplete and broad asymmetrical hysteresis AFM-FM transition$^{1,19–34}$. The starting transition temperature $T_{K}^{\alpha'' \rightarrow \alpha'}$ and magnetic properties in B2-FeRh films depend not only on compositional variations$^{26,27,34}$, but are also highly sensitive to the type of substrate$^{18,24–26,29,33}$, the thickness of the thin film samples$^{20,24,30}$, the nature underlayers$^{33,36}$ and the capping layers$^{31}$. Significant stresses arise at the film/substrate interface$^{20,24}$, which makes it possible to control

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the magnetic and electrical transport properties of B2-FeRh films on ferroelectric substrates by varying the external electric field. Many groups using magnetron sputtering or molecular beam epitaxy achieved epitaxial B2-FeRh(001) films grown on MgO(001) crystalline substrates. In most studies high-temperature heat treatments above 600 °C were used to obtain the high crystalline quality of chemically B2-ordered phase. However, chemical interactions of Rh with Fe and solid-state synthesis of the Rh-Fe phases at the Rh/Fe interface remain completely uninvestigated.

In this work, we describe the solid-state reactions between polycrystalline Rh with epitaxial Fe(001) films in 52Rh/48Fe(001), 45Rh/55Fe(001) and 68Rh/32Fe(001) bilayers with 52Rh:48Fe, 45Rh:55Fe and 68Rh:32Fe atomic ratios, respectively. Under these conditions, the reaction products of 52Rh/48Fe(001) fall into the (0.48 < x_Rh < 0.56) concentration interval and the reaction products for 45Rh/55Fe(001) and 68Rh/32Fe(001) samples lie in Fe-rich and Rh-rich regions of the Fe–Rh system. The main purpose of this article is to show the complex and intricate nature of the B2-FeRh phases synthesis, which begins to form on the Rh/Fe interface at ~ 100 °C.

Results

Structural and magnetic phase transformations in 52Rh/48Fe(001) bilayer during annealing up 800 °C. Figure 1a shows a schematic diagram of the phase transformations consistently occurring in 52Rh/48Fe(001) bilayers on a MgO(001) substrate during annealing from room temperature to 800 °C which builds on the X-ray diffraction (XRD) analysis (Fig. 1b) and magnetic measurements (Fig. 1c,d). The transformations consist of the formation of a nonferromagnetic B2-FeRh phase (nfm-B2) thin layer at the Rh/Fe interface of the as-deposited sample, which above 300 °C turns into the αl phase. As will be shown below, the ferromagnetic αl phase may have two B2 modifications with similar or equal lattice parameters: a low-ferromagnetic αl phase having a magnetization ~ 825 emu/cm² and a fully reversible αl ↔ αh (AFM-FM) transition and a high-ferromagnetic αl phase with a magnetization ~ 1,220 emu/cm² and with a completely depressed αl ↔ αh transition. As the annealing temperature increases above 600 °C the reaction products contain a mixture of epitaxial αl(001) and αh(001) grains. Figure 1b shows the XRD profiles of the as-deposited 52Rh/48Fe(001) film and after annealing at temperatures from room temperature to 800 °C. As-deposited samples show a strong Fe(002) peak, proving the epitaxial Fe(001) film growth on the MgO(001) substrate, and wide and very low diffraction peaks (001), (002) located at ~ 30° and ~ 62°, which are a signature of significant mixing, a reaction between the Rh and Fe(001) layers and the synthesis of a very thin epitaxial layer of nanocrystalline B2-ordered FeRh phase. These (001), (002) reflections were greatly broadened, split and their intensity did not change up to 600 °C. This strongly suggests that the reaction starts with the formation of αl-textured nanograins containing B2—ordered FeRh phases with close lattice parameters and significant lattice distortions. After annealing above 600 °C the Fe(002) peak decreases significantly and practically disappears at 800 °C, which points to the complete termination of the reaction between the Fe and Rh layers. At the same time the (001), (002) peaks grow insignificantly, which is due to the annealing-induced coarsening of the B2—ordered FeRh nanograins. Using Scherrer’s formula the average grain size was estimated S = 0.90 ± 0.02 for the synthesized B2-FeRh samples at temperatures in the 600–800 °C range. Figure 1c shows the dependence of the in-plane relative magnetic anisotropy constant K4(Ta)/K4 relative magnetization as a function of the annealing temperature T_a (Fig. 1c) has an unusual shape and convincingly shows complex and intricate scenarios of various phase formation and their mutual transformations in the Rh/Fe(001) bilayer during annealing to 800 °C. Within experimental error the relative anisotropy constant K4(Ta)/K4 and the relative magnetization M_s(Ta)/M_s decrease identically as the annealing temperature increases up 300 °C. The synchronous decrease of the K4(Ta)/K4 and M_s(Ta)/M_s values clearly proves the onset of the slow mixing of the Rh and Fe layers and the synthesis of the nonferromagnetic B2-FeRh(001) phase (nfm-B2) on the Rh/Fe(001) interface, whose volume increases with an increase in annealing temperature to 800 °C. This means that the first magnetic anisotropy constant K4 of the B2—ordered FeRh phase is zero or less than the measurement error (0.2 × 10⁴ erg/cm³). A value of K4 ~ 0 is consistent with the FeRh single films exhibiting an in-plane easy axis of magnetization with no measurable magnetocrystalline anisotropy. Therefore, the K4(Ta)/K4 dependence can be used to find the thickness, magnetization and magnetic moment of the reacted Fe(001) layer after annealing at T_a (Supplementary Note 1). The dependence of the relative magnetization M_s/M_s as a function of annealing temperature T_a (Fig. 1c) has an unusual shape and convincingly shows complex and intricate scenarios of various phase formation and their mutual transformations in the Rh/Fe(001) bilayer during annealing to 800 °C. Within experimental error the relative anisotropy constant K4(Ta)/K4 and the relative magnetization M_s(Ta)/M_s decrease identically as the annealing temperature increases up 300 °C. The synchronous decrease of the K4(Ta)/K4 and M_s(Ta)/M_s values clearly proves the onset of the slow mixing of the Rh and Fe layers and the synthesis of the nonferromagnetic B2-FeRh(001) phase (nfm-B2) on the Rh/Fe(001) interface, whose volume increases with an increase in annealing temperature to 300 °C. Above 300 °C the magnetization increases to M_s ~ 1, which indicates a phase transformation of the nfm-B2 to the ferromagnetic αl phase. From Fig. 1c follows that the trilayer system Rh/αl/Fe(001) has the magnetization of the initial sample M_s ~ 825 emu/cm² (M_s/M_s ~ 1) in the temperature range (350–500 °C), which means that only the Fe atoms contribute to the saturation magnetization of αl-FeRh. Surprisingly, after annealing at 550 °C, the saturation magnetization decreases to 660 emu/cm² (~ 20%), but the reaction product layer participates completely in the reversible αl ↔ αh transition (Fig. 1d). At annealing temperatures above 500 °C, the films begin to partially peel off from the substrate due to the strong stresses arising during the synthesis of the epitaxial αl-FeRh(001) film at the αl ↔ αh(Fe(001)/Fe(001)) interface. Therefore, a possible explanation is the deformation of some αl—grains with a loss of ferromagnetic order. However, the question about the exact origin of the decrease of M_s/M_s at 550 °C remains open. All of the 52Rh/48Fe(001) samples had the same temperature dependencies of the magnetization M_s(T_a) up to 600 °C. However, after annealing above 600 °C the magnetizations of M_s sharply increased and
assumed values that centered either around $M_{s1} \approx 1,220 \text{ emu/cm}^3$ ($M_{s1}/M_0^0 \approx 1.47$, B-samples) or $M_{s2} \approx 940 \text{ emu/cm}^3$ ($M_{s2}/M_0^0 \approx 1.15$, A-samples) (Fig. 1c).

Temperature dependence of the reversible AFM-FM transition in the 52Rh/48Fe bilayer. In experiments, all 52Rh/48Fe(001) samples, after annealing each in a temperature range of 100–800 °C, were subjected to an investigation of the $\alpha' \leftrightarrow \alpha''$ transition, consisting of measurements of the magnetic moment $m(T_a)$ of the synthesized layer B2-FeRh in the Rh/B2-FeRh/Fe (001) trilayer by the torque method at room temperature, after annealing at temperature $T_a$ and after cooling in liquid nitrogen $m(N)$ for which $\eta = 0$ is absence of transition and $\eta = 1$ is perfect transition (see “Methods”). Figure 1d shows the temperature dependence of the degree $\eta(T_a)$ of the $\alpha' \leftrightarrow \alpha''$ transition which has values (200–300 °C) $\eta = 0$, (400–600 °C) $\eta > 0.95$ and (700–800 °C) $\eta = 0$ for the (B) samples and $\eta = 0.95–0.8$ for the (A) samples in the specified temperature ranges. In the temperature interval 200–300 °C, the equality $\eta = 0$ is determined by the formation of a non-magnetic nfm-B2 phase. Although the $\alpha'_1$ (001) layer is located between the Rh and Fe (001) layers and grows epitaxial on the surface of Fe (001) in the (400–600 °C) interval it has a reversible $\alpha'_1 \leftrightarrow \alpha''$ transition with a low residual magnetization (see Supple-
In the (650–800 °C) interval the reaction ends and only the (A) samples reveal a reversible $\alpha_1' \leftrightarrow \alpha_2''$ transition, while this transition does not occur in the (B) samples. We hypothesize that the $\alpha_1'$ phase has two B2-ordered polymorphic modifications having similar lattice parameters, but different magnetic properties. In the first modification the $\alpha_1'$ phase has a low magnetization about 825 emu/cm$^3$, which is defined only by the Fe atoms with the Rh atoms not contributing to the magnetization, and the $\alpha_1'$ undergoes a complete reversible $\alpha_1' \leftrightarrow \alpha_2''$ transition (Supplementary Fig. 3a). In contrast, in the second $\alpha_1''$ modification, which is primarily in the (B) samples, the Fe atoms supposedly polarize the Rh and the Rh atoms make the contribution to the saturation magnetization 1,220 emu/cm$^3$ and in the $\alpha_1''$ phase the $\alpha_1'' \leftrightarrow \alpha_2''$ transition is completely suppressed (Supplementary Fig. 3b). Under such conditions, with an increase in temperature above 650–800 °C, some low-magnetic $\alpha_1'$ nanograins turn into high-magnetic $\alpha_1'$ and therefore, the samples have a magnetization between 825 and 1,220 emu/cm$^3$ and exhibit an imperfect reversible transition with a residual magnetization. To summarize, the chemical reaction between Fe and Rh arises on the Fe/Rh interface at a low annealing temperature (~ 100 °C) and with an increase in annealing temperature the phase sequence 52Rh/48Fe $\rightarrow$ (300 °C) nfm-B2 $\rightarrow$ (300 °C) $\alpha_1' \rightarrow$ (600 °C) $\alpha_1'$ or $\alpha_2''$ is formed as shown in Fig. 1a.

The phase and magnetic evolution in the 45Rh/55Fe(001) bilayer during annealing up 800 °C. The schematic diagram in Fig. 2a is based on the results presented in Fig. 2b–d and shows the same
nonferromagnetic nfm-B2 phase which was formed in the 45Rh/55Fe and 52Rh/48Fe bilayers during annealing from room temperature to 300 °C. Above 300 °C the nfm-B2 phase consistently turns into ferromagnetic low-magnetization α′, which above 450 °C grows into the high-magnetization α′ phase. As shown in Fig. 2b after annealing above 400 °C the (001) and (002) peaks start to grow and become very strong above 500 °C, which indicates the formation of high structural quality epitaxial α′(001) layers on the MgO(001) surface. The K(T)/K0 and M4/M40 dependencies (where for 45Rh/55Fe(001) samples K(T)/K0 = 2.5 × 105 erg cm3 and M40 = 875 emu/cm3, see “Methods”) presented in Fig. 2c show that the nanocrystalline phase α′(001) with low magnetization (M4/M40 ~ 1.0) grows into the high structural quality epitaxial phase α′(001) with high magnetization (M4/M40 ~ 1.47) above 450 °C. The high quality of the chemical ordering of the α′(001) films after annealing at 500 °C and 800 °C supports the order parameter S = 0.96 ± 0.02, which is more than the S = 0.90 ± 0.02 for α′ in 52Rh/48Fe films. This result is unexpected, since the α′ exists in a narrow composition range of nearly equiatomic concentration and must have a more complete B2 order than the Fe-rich αh phase. The α′(001) and α′(001) films have the same orientation relationship with the substrate MgO(001) (Supplementary Fig. 1). Above 500 °C the Fe phase has completely reacted with the Rh as evidenced by the K(T)/K0 dependence (Fig. 2c). As can be seen from Fig. 2d the degree of the AFM-FM transition η ~ 0.9, which means the α′ phase exists in a narrow temperature range (300–450 °C) and the AFM-FM transition has a relatively low magnetization (Supplementary Fig. 4). Figure 2d shows the absence of the AFM-FM transition (η = 0), which indicates the formation of a α′ phase from the α′ phase after annealing above 450 °C (see also Supplementary Fig. 3). From these facts it transpires that the Fe-rich α′ phase formed from the equiatomic α′ compound above 450 °C by the solid-state reaction α′ + Fe → (450 °C) α′. Finally, we proved that even a slight Fe doping of α′ causes a chemical reaction between Fe and α′ and the start of the synthesis of the α′ phase. This suggests that the high magnetization α′ phase occurring in the 52Rh/48Fe(001) bilayer (B samples) also has more Fe content than α′ and explains the compositional heterogeneity arising as a result of the phase evolution. Figure 3a shows a schematic of the phase sequence formation in the nanocrystalline nonferromagnetic nfm-B2 and the low-magnetization α′ phases during annealing up to 500 °C in the 68Rh/32Fe(001) bilayer system (Fig. 3b, see Supplementary Fig. 6). Annealing above 500 °C the epitaxial γ(001) layer grows on the α′(001) surface originating from the solid-state reaction Rh + α′ → (500 °C) γ. As shown in Fig. 3b the reaction is fully completed after 800 °C and the final reaction product is the γ(001) layer which has a cube-on-cube orientation relationship γ(001)[110][MgO(001)][100] with the MgO(001) substrate. (Supplementary Fig. 7). The relative magnetization M4(T)/M40 and the K(T)/K0 dependence presented in Fig. 3c confirm the sequential formation of nonferromagnetic nfm-B2 at about 100 °C, ferromagnetic α′ above 300 °C and the paramagnetic γ phases above 500 °C. As follows from Fig. 3d, the ferromagnetic α′ phase exists and undergoes a reversible AFM-FM transition (η ~ 0.9) in the 300–500 °C temperature range (Supplementary Fig. 6). Finally, we revealed the phase formation sequence 68Rh/32Fe → (~ 100 °C) nfm-B2 → (~ 100 °C) α′ → (450 °C) α′ is induced by the solid-state reaction method and the final reaction product is a highly B2-ordered phase α′, which has a high magnetization of 1,270 emu/cm3 and in which the reversible α′↔α transition is completely suppressed.

Phase formation sequence, magnetic and structural development in the 68Rh/32Fe(001) bilayer during annealing up 800 °C. Figure 3a shows a schematic of the phase sequence formation in the 68Rh/32Fe(001) bilayer based on data presented in Fig. 3b–d. Figure 3a shows the same phase sequence including the sequential formation of the nanocrystalline nonferromagnetic nfm-B2 and the low-magnetization α′ phases during annealing up to 500 °C in the 68Rh/32Fe, 52Rh/48Fe (Fig. 1a) and 55Rh/45Fe (Fig. 2a) bilayers. After annealing above 500 °C the epitaxial paramagnetic γ(001) phase starts to grow at the Rh/α′(001) interface (Fig. 3a). The XRD patterns show that after annealing at 400 °C the Fe layer forms an epitaxial α′(001) layer in the Rh/α′(001) bilayer system (Fig. 3b, see Supplementary Fig. 6). Annealing above 500 °C the epitaxial γ(001) layer starts to grow on the α′(001) surface originating from the solid-state reaction Rh + α′ → (500 °C) γ. As shown in Fig. 3b the reaction is fully completed after 800 °C and the final reaction product is the γ(001) layer which has a cube-on-cube orientation relationship γ(001)[110][MgO(001)][100] with the MgO(001) substrate. (Supplementary Fig. 7). The relative magnetization M4(T)/M40 and the K(T)/K0 dependence presented in Fig. 3c confirm the sequential formation of nonferromagnetic nfm-B2 at about 100 °C, ferromagnetic α′ above 300 °C and the paramagnetic γ phases above 500 °C. As follows from Fig. 3d, the ferromagnetic α′ phase exists and undergoes a reversible AFM-FM transition (η ~ 0.9) in the 300–500 °C temperature range (Supplementary Fig. 6). Finally, we reveal the phase formation sequence 68Rh/32Fe → (~ 100 °C) nfm-B2 → (~ 100 °C) α′ → (~ 300 °C) γ during thermal annealing up to 800 °C and the final reaction product contains only an epitaxial γ(001) layer on the MgO(001) surface.

Kinetic growth of the nanocrystalline nfm-B2 layer during thermal aging at 110 °C. To gain a better understanding of the origin and formation of the very thin nanocrystalline nfm-B2 layer at the Rh/Fe(001) interface at the temperature of the α′ ↔ α transition, the synthesis kinetics of the nfm-B2 layer during isothermal aging at 110 °C was investigated. As stated above, deposition of Rh on Fe(001) at room temperature at any ratio of thicknesses between Rh and Fe leads to the formation of a thin interfacial nanocrystalline nfm-B2 layer. However, slight heating to 100 °C during sputtering cannot be excluded, therefore, aging of as-deposited samples was investigated at 110 °C. XRD data from the as-deposited Rh/Fe(001) bilayer and after aging up to 360 h shows a small broadened (001) superlattice and the fundamental (002) peaks of the nfm-B2 phase which did not change from aging time to 360 h (Fig. 4a). This means that long-time aging does not lead to the nfm-B2 grains coarsening and clearly demonstrates the inherent nanocrystalline nature of nfm-B2, which is also inherited by the α′ phase. Figure 4b shows the TEM image and EDX line scan results of the Rh/Fe(001) bilayer aged at 110 °C for 360 h. The elemental profile obtained by the EDX line scan across the Rh/nfm-B2/Fe(001) trilayer shows an interfacial nfm-B2 thin film beside the initial Rh and Fe layers. The TEM image displays a compositional contrast that is proportional to the difference in the average atomic number of the respective areas. As a consequence, the reaction product nfm-B2 can be easily distinguished in the interfacial region from the initial Rh and Fe(001) areas. This allows one to estimate the thickness of the interfacial nfm-B2 layer α′ (001) ~ 15 nm (Fig. 4b). Figure 4c shows the thickness d(nfm-B2) of the Fe(001) layer, which entered into a reaction with Rh at 110 °C, as a function of aging time. There is a large experimental error in the thickness d(nfm-B2) measurements and therefore the dependence d(nfm-B2)(t) is impossible to describe by an unambiguous kinetic equation, which suggests a possible growth mechanism. As shown in Fig. 4c the thickness d(nfm-B2) after aging for 360 h at 110 °C is ~ 8 nm, which corresponds to the thickness of α′(001) ~ 17 nm of the nfm-B2 layer (Supplementary Fig. 8a). This value agrees well with the estimated d(nfm-B2) ~ 15 nm obtained from TEM-EDX analysis. These findings are in good agreement with...
a previous report\textsuperscript{42} which showed and discussed the tendency towards intermixing and interfacial Rh–Fe alloy formation at the Rh/Fe interface at room temperature.

**Phase transition from nfm-B2 to $\alpha'_l$ at 300 °C.** For a further understanding of the reactivity of Fe and Rh during the transition from nfm-B2 to $\alpha'_l$ around 300 °C, we investigated the evolution of the temperature dependencies of the magnetization and the thickness of the interfacial nfm-B2 layer in the 52Rh/B2-FeRh/48Fe(001) trilayer after annealing at 280 °C, 300 °C and 350 °C. Figure 5a shows the M–T curves for the 52Rh/nfm-B2/48Fe(001) trilayer after annealing at 280 °C which do not contain changes in either the forward or the reverse direction. This clearly demonstrates that the B2-FeRh interlayer remains a non-magnetic nfm-B2 layer after annealing in the 110–280 °C temperature interval. From the cross-sectional TEM image and the EDS line-scan (Fig. 5a), it can be seen that the $d_{\text{nfm-B2}}$ interlayer in the 48Rh/nfm-B2/52Fe(001) trilayer increases to a thickness around $d_{\text{nfm-B2}} \approx 30$ nm after annealing at 280 °C. (Supplementary Fig. 8b). As shown in Fig. 5a,b slight increase in the annealing temperature to 300 °C leads to the appearance a road FM-AFM transition with a temperature hysteresis of magnetization about 80 K and a small increase in the $d_{\text{B2-FeRh}}$ thickness of the B2-FeRh interlayer to ~ 40 nm (Supplementary Fig. 8c). The change in magnetization at the AFM-FM transition gives a rough estimate of the thickness $d(\alpha'_l) \sim 20$ nm of the $\alpha'_l$ layer in the B2-FeRh interlayer. This clearly proves that after annealing at 300 °C the B2-FeRh interlayer contains a mixture of 50%
Figure 4. Direct evidence of the low-temperature synthesis of the thin interfacial nanocrystalline nfm-B2 layer in 52Rh/48Fe bilayers. (a) XRD pattern of the 52Rh/48Fe bilayer after aging at 110 °C between 60 and 360 h. (b) TEM image of the cross-section and the EDS element mapping of the Fe and Rh in the 52Rh/48Fe bilayer after aging at 110 °C for 360 h. (c) The dependence of the reacted layer thickness $h_{Fe}$ during aging up to 360 h at 110 °C.

nfm-B2 and 50% α$^{'}$ phases (Supplementary Note 2). Figure 5c shows that the thickness $d_{B2-FeRh}$ grows quickly to 135 nm as the annealing temperature increases to 350 °C (Supplementary Fig. 8d). This value is in good agreement with the estimate obtained from the $K_\chi(T)/K_\chi^0$ dependence (Fig. 1c, see Supplementary Note 3). EDS anal-
ysis gives the average value of the atomic concentration ratio of Fe and Rh at about 1.0, which confirms that the reaction product layer is the B2-FeRh phase. Pores are visible on the Fe/nfm-B2 interface (Fig. 5c, Supplementary Fig. 8d), which are well known as Kirkendall voids. The explanation of the Kirkendall effect is based on the difference in rates of the reacting atoms and it proves that Fe is the dominant diffusing species during B2-FeRh formation. Figure 5c shows the M–T curve having a road FM-AFM transition with large thermal hysteresis width about 100 K, which often occurs in thin film system. This may be due to inhomogeneities of composition, stresses and nanocrystalline growth as a result of non-equilibrium synthesis of the B2-FeRh phases. Thus, this is proof that the reactivity of Fe and Rh is very low at 110 °C and begins to increase greatly during the transition nfm-B2 → αlʹ above 300 °C. This causes strong temperature dependencies up to 500 °C of the relative magnetic anisotropy constant $K_4(T)/K_4^0$ (Figs. 1, 2, 3) and electrical resistance (Supplementary Fig. 9).

Discussion
Numerous experiments have proved that as temperature rises, only one phase appears on the interface of bilayer film systems, which is called the first phase. The initiation temperatures $T_{in}$ of the first phase for most bilayers lie below 400 °C. However, many thin-film reactions are initiated near room temperature and even occur at cryogenic temperatures. At such low temperatures diffusion is extremely small and cannot provide atomic transfer in a solid state. This suggests an alternative view, in which not diffusion, but chemical interactions play a crucial role in the initiation and kinetics of solid state interfacial reactions. Under the influence of chemical interactions that arise above the initiation temperature $T_{in}$, the chemical bonds break in the reactants, and the reacting atoms migrate into the reaction zone to synthesize new compounds. Previously, it was shown that the initiation temperatures $T_{in}$ are close to or coincide with the solid-state transformation temperatures $T_k$ of several reagent-based binary systems, such as order–disorder transitions, the superionic transition, the spinodal decomposition, martensitic transformations and others. This suggests that the same chemical interactions underlie and control both the solid-state thin-film reactions and the corresponding solid-state transformations. The equality $T_{in} = T_k$ indicates that low-temperature solid-state thin-film reactions in A/B bilayers occur only in A–B binary systems, which have corresponding low-temperature solid-state transformations. Therefore, the study of reactions in A/B bilayers with different layer ratios is a study of the low-temperature part of the A–B phase diagram.

It is well established that ordered B2 alloys, such as NiTi, AuCd, NiAl have reversible low-temperature martensitic transformations, in which the high-temperature austenite B2-phase develops into a low-temperature martensitic phase through a complex process of the formation of intermediate phases. We have shown earlier the initiation temperatures $T_{in}(Ti/Ni) < 150 °C$, $T_{in}(Ni/Al) ~ 180 °C$, $T_{in}(Cd/Au) = 67 °C$ in Ti/Ni, Ni/Al, Cd/Au bilayers, respectively. These temperatures are close to or coincide with the reverse martensitic transformation starting temperatures $A_s(Ti-NiTi) ~ 100 °C$, $A_s(B2-NiAl) ~ 180 °C$, $A_s(B2-CdAu) = 67 °C$. The reversible α’ ↔ α'' transition in B2-FeRh possesses all the characteristics of a martensitic transformation (α’-austenite, α''-martensite), because it can pass at high speeds, has isotropic volume changes at the transition, can be

Figure 5. The nfm-B2 to αlʹ phase transformation. TEM images of the cross-section, the EDS element mapping and the temperature dependent magnetization curves of the 52Rh/48Fe bilayer after aging (a) at 280 °C, (b) 300 °C, (c) 350 °C. The appearance of a loop in the M–T curve in (b) indicates the start of the ferromagnetic αlʹ phase formation at 300 °C.
induced by the application of stress and magnetic field, has martensitic instabilities and the α’ and α” lattices have a cube-on-cube orientation relationship. According to the phase diagram, the transition α’ ↔ α” has a minimum temperature 100 °C among other structural transformations in the Fe-Rh system. From the above, we have concluded that the initiation temperature of the reaction  in the Rh/Fe bilayer coincides with the martensitic-like transition temperature 100 °C in B2-FeRh. The coincidence of the starting temperature of the reaction between Fe and Rh and the temperature of the magnetostuctural AFM-FM transition suggests common chemical mechanisms behind both phenomena, but that connection remains to be confirmed by additional experimentation.

As mentioned above, the formation of compounds at the Rh/Fe interface starts in the temperature range 100–300 °C from the synthesis of the non-ferromagnetic nfm-B2 phase. Since this phase has only (001) and (002) B2-FeRh reflections, this means that nfm-B2 is either a martensitic-like antiferromagnetic α’-phase or a non-ferromagnetic martensitic variant which is stabilized by strains resulting from the non-equilibrium synthesis of the nfm-B2 phase. B2-FeRh, similar to other B2 phases of NiTi, AuCd, NiAl alloys, experiences premartensitic instabilities with the subsequent formation of the different structural variants of martensite. Our hypothesis is that the amorphous phase appears above 100 °C at the Rh/Fe interface, which is then transformed into a nanocrystalline state containing B2 nanograins of martensite variants with lattice parameters close to α” martensite. This is consistent with the possible existence of various structural phases of FeRh, predicted by ab initio calculations and found in the experimental study. Competition between the α’ and α” phases during the partial crystallization suppresses grain growth and stabilizes the nano-grained B2 structures in an amorphous matrix. The formation of an amorphous phase is a quite common phenomenon in the initial stage of solid-state reactions in bilayers and multilayers, although the nature of this phenomenon is still a subject of dispute. It is interesting to note the general features of the initial stage of the synthesis of B2 phases in Ti/Ni and Rh/Fe thin films. In Ti/Ni multilayers, the amorphous phase starts near the martensitic transformation temperature (450 °C) which turns into B2-NiTi at annealing temperatures above 350 °C. Similar to Ti/Ni, the reaction in the Rh/Fe(001) bilayer starts at ~100 °C with the formation of the interfacial amorphous phase, which partially crystallizes. From this point the as-deposited films consist of nanocrystalline B2 grains dispersed in the amorphous matrix. This strongly suggests that both amorphous phases are amorphous martensite, which may be a universal phenomenon of the solid-state synthesis of martensitic phases. This scenario is different from the synthesis of B2-NiAl and B2-AuCd, which begin to form in Al/Ni and Cd/Au bilayers at martensitic transformation temperatures without the formation of an intermediate amorphous martensite.

Analysis of the general thermodynamic characteristics and features of B2-FeRh and B2-NiTi suggests the possibility of a fabrication of the B2-FeRh compound by self-propagating synthesis (Supplementary Note 3). Our approach assumes ~100 °C is the starting temperature of the formation of the B2 phase, which is associated with the reversible AFM-FM transition, and the synthesis temperatures of the α” phase coincide with the phase transition temperatures in the Fe-rich and Rh-rich regions of the Fe–Rh system, respectively. Such an approach is justified by us for the well-studied Fe–Ni system and made it possible to predict phase transformations in other binary metallic systems. Therefore, further study of solid-state reactions in Rh/Fe films, depending on the composition, will make it possible to specify the low-temperature part of the Fe–Rh phase diagram, which still remains unknown.

In conclusion, we have uncovered that regardless of the Rh and Fe thicknesses the thin nanocrystalline B2-FeRh layer starts to form at ~100 °C and grows up to 300 °C on the Rh/Fe interface. Above 300 °C the amorphous phase is converted into a low-ferromagnetic B2 α’ phase with a magnetization ~825 emu/cm3 without changing the nanocrystalline structure. Above 500 °C the α” phase reacts with Fe and formed B2 α” phase with a magnetization ~1,270 emu/cm3 in the 45Rh/55Fe(001) samples and the α” phase reacted with Ph and formed the non-ferromagnetic γ phase in the 68Rh/32Fe(001)samples. Magnetic analysis has revealed that only the α” undergoes the complete reversible α’ ↔ α” transition and there is no transition in the α” samples. Thus, our work not only provides an idea of how phase sequences start and develop depending on the composition of the Fe/Rh bilayers, but also suggests the interesting possibility that a similar chemical mechanism may be at play behind the low-temperature reaction of Fe and Rh and the AFM-FM transition in B2-FeRh.

Methods
52Rh/48Fe(001), 45Rh/55Fe(001) and 68Rh/32Fe(001) bilayers preparation and characterization. At first the epitaxial Fe(001)/MgO(001) films were grown on single-crystal MgO(001) substrates by a thermal evaporation method in a vacuum chamber at a pressure of 10–6 mbar. To obtain high-quality Fe(001) films, the substrates were previously outgassed at 300 °C for 1 h and the Fe layers were deposited at 250 °C. Epitaxial Fe(001) films had the orientation ratio Fe(001)[100]∥[MgO(001)[110] with the MgO(001) substrate and the magnetocrystalline anisotropy constant K1, which coincided with the value of bulk iron K1 = 4.9 × 105 erg/cm3. The constant K1 in the (001) plane was determined by a torque magnetometer in a magnetic field H = 12 kOe. The torque curve L(φ) in the (001) plane was determined according to the equation

\[ 2L(\phi) = K_V \sin 4\phi + 2K_1 \sin(2\phi + \gamma) \]

In addition to the dominant 4ϕ-term, there is a minor 2ϕ-term, which is due to the surface roughness of the MgO(001) substrate and a slight misorientation of the Fe(001) grains. In the equation, K1 is the uniaxial anisotropy constant, V is the volume of the film, φ is the angle between the easy axis of fourfold anisotropy and the magnetization M, γ is the angle between the easy axis of fourfold anisotropy and the axis of the uniaxial anisotropy. The K_V value was calculated from the torque curve L(φ) at the maximum of the 4ϕ-term: 2L_{max} = K_V. High quality epitaxial Fe(001)/MgO(001) films also can be obtained by various other methods as reported in the literature.

The starting Rh(001) bilayers were obtained by the evaporation of the Rh layers on Fe(001)/MgO(001) samples using dc sputtering in a magnetron sputtering system. The base pressure of the chamber was less...
than $1 \times 10^{-6} \text{ mbar}$, and a working pressure of $\sim 1 \text{ mTorr}$ Ar was used during sputtering. To prevent a reaction between Rh and Fe, the Rh layer was deposited at room temperature. Under such deposition conditions the polycrystalline Rh layer was formed on the Fe(001) surface. Three different types of samples were prepared for the experiments, namely Rh/Fe (001) bilayers with Rh-rich, approximately 1Fe:1Rh and Fe-rich atomic ratios, each with a total thickness about 300 nm. The elemental chemical composition determined by energy dispersion X-ray (EDX) analysis showed sample compositions of Rh$_{68}$Fe$_{32}$, Rh$_{52}$Fe$_{48}$ and Rh$_{45}$Fe$_{55}$ respectively. The saturation magnetization $M_s^0$ and the magnetic fourfold anisotropy constants $K_4^0$ were determined for the total volume of the 52Rh/48Fe, 45Rh/55Fe and 68Rh/32Fe bilayers, which turned out to be $M_s^0 = 825 \text{ emu/cm}^3$, $K_4^0 = 2.2 \times 10^5 \text{ erg/cm}^3$ for the 52Rh/48Fe bilayers, $M_s^0 = 875 \text{ emu/cm}^3$, $K_4^0 = 2.4 \times 10^5 \text{ erg/cm}^3$ for the 45Rh/55Fe bilayers and $M_s^0 = 450 \text{ emu/cm}^3$, $K_4^0 = 1.25 \times 10^5 \text{ erg/cm}^3$ for the 68Rh/32Fe bilayers.

The reversible AFM-FM phase transitions were checked using a superconducting quantum interference device (SQUID) magnetometer. Magnetic fields of $H = 1.0 \text{ kOe}$ were applied along the in-plane [100] MgO direction, which coincides with the easy axis of the Rh/Fe (001) bilayers, at all measurements in the 77–400 K temperature interval. The saturation magnetization $M_s$ and the coercivity $H_c$ were measured with a vibration magnetometer in magnetic fields up to 22 kOe. All saturation magnetization measurements were monitored using the torque method.

The formed phases were identified with a DRON-04-07 diffractometer (CuKa radiation). The epitaxial relationships between MgO(001) and the B2, $\gamma$ layers that formed in the reaction products were X-ray studied with a PANalytical X′Pert PRO diffractometer with a PIXcel detector. CuKa radiation monochromatized by a secondary graphite monochromator was used in the instrument. The order parameter for the synthesized samples at temperatures in the 600–800 $^\circ\text{C}$ range was estimated using the equation $S = (I_{001}/I_{002})^{1/3}/1.07$, where $I_{001}$ and $I_{002}$ are experimental integrated intensities of superstructural (001) and fundamental (002) reflections.

The cross-sectional samples for TEM studies were prepared with a focused ion beam (single-beam FIB, Hitachi FB2100) at 40 kV. In order to protect the surface of interest from milling by the Ga + ion beam during sample preparation, a Ge layer was deposited onto the Fe-Rh film before cross-sectional sample preparation by FIB. TEM studies were carried out using a Hitachi HT7700 TEM (acceleration voltage 100 kV, W source) equipped with a STEM system and a Bruker Nano XFlash 6T/60 energy dispersive X-ray (EDX) spectrometer. The imaging and EDX spectroscopy line scans and mapping were carried out in STEM mode with an electron probe of diameter $\sim 30 \text{ nm}$.

**Characterization of solid-state phase transformations.** The starting 52Rh/48Fe, 45Rh/55Fe and 68Rh/32Fe bilayers were annealed at temperatures ranging from 50 to 800 $^\circ\text{C}$ in increments of 50 $^\circ\text{C}$. The samples were held at each temperature at a pressure of $10^{-6} \text{ Torr}$ for 1 h. To characterize the phase transformations the crystal structure, the magnetic moments $m(T)$ (where $m(300 \text{ K}) = M_s^0$), the magnetic fourfold anisotropy constants $K_4(T)$ and the degree of the $\alpha^\prime \rightarrow \alpha^\prime$ transition $\eta(T)$ were determined for all bilayers after annealing at each temperature $T_a$. In order to fully characterize the phase transformations, cross-sectional SEM images and an elemental analysis of the phases of several samples using EDX were carried out.

**Measurement of the $\alpha^\prime \rightarrow \alpha^\prime$ transition degree.** In experiments, the degree $\eta(T)$ of the FM-AFM phase transition was determined for all 52Rh/48Fe, 45Rh/55Fe and 68Rh/32Fe samples after annealing at each temperature in the range of 100–800 $^\circ\text{C}$. The magnetic moments $m(T)$ were measured by the torque method, after annealing at temperature $T_a$ and cooling in liquid nitrogen $m(T_0)$, to find the value of the degree $\eta(T)$. The magnetic moment $m(T)_{\text{FeRh}}$ of the synthesized $\alpha^\prime$—FeRh layer in the Rh/$\alpha^\prime$—FeRh/Fe(001) trilayer after annealing at temperature $T_a$ is equal to the difference $m(T)_{\text{FeRh}} = m(T)_{\text{FeRh}} - K_4(T)_{\text{FeRh}}m(T)/K_4$. The magnetic moments of the $m(T)$ film and the $K_4(T)m^2/K_4$ untreated layer of the Fe(001) layer, where $K_4(T)m^2/K_4$ is the magnetic moments of the untreated Fe(001) layer. After placing the sample in liquid nitrogen, only the ferromagnetic $\alpha^\prime$—FeRh phase in the Rh/$\alpha^\prime$—FeRh/Fe(001) trilayer is subjected to the transition into the antiferromagnetic $\alpha$” phase and the magnetic moment $m(T)$ is reduced to $m(T)$. The quantity $\eta = 1 - m(T)/m(T_0)$ is a quantitative characteristic of the degree of the $\alpha^\prime \rightarrow \alpha^\prime$ transition, where $\eta = 0$ and $\eta = 1$ mean the absence of and the complete FM$\rightarrow$AFM transition (residual magnetization is zero), respectively.

Received: 17 July 2019; Accepted: 10 June 2020
Published online: 02 July 2020

**References**

1. Lewis, L. H., Marrows, C. H. & Langridge, S. Coupled magnetic, structural, and electronic phase transitions in FeRh. *J. Phys. D Appl. Phys.* 49, 323002 (2016).
2. Annaorazov, M. P. et al. Anomalously high entropy change in FeRh alloy. *J. Appl. Phys.* 79, 1689–1695 (1996).
3. Cooke, D. W. et al. Thermodynamic measurements of Fe–Rh alloys. *Phys. Rev. Lett.* 109, 255901 (2012).
4. Mankovsky, S. et al. Temperature-dependent transport properties of FeRh. *Phys. Rev. B* 95, 155319 (2017).
5. Ibarra, M. R. & Algarabel, P. A. Giant volume magnetostriction in the FeRh alloy. *J. Phys. D Appl. Phys.* 49, 295001 (2016).
6. Suzuki, L., Naito, T., Roh, M., Sato, T. & Taniyama, T. Clear correspondence between magnetoresistance and magnetization of epitaxially grown ordered FeRh thin films. *J. Appl. Phys.* 109, 07C717 (2011).
7. Chen, X. Z. et al. Tunneling anisotropic magnetoresistance driven by magnetic phase transition. *Nat. Commun.* 8, 449 (2017).
8. Nikitin, S. A. et al. The magnetocaloric effect in Fe49Rh51 compound. *Phys. Lett. A* 114, 363–366 (1985).
9. Kouvel, J. S. Unusual nature of the abrupt magnetic transition in FeRh and its pseudobinary variants. *J. Appl. Phys.* 37, 1257–1258 (1966).
10. Onodera, R. et al. Observation of a metamagnetic transition of FeRh alloy under high magnetic fields and high temperatures. *J. Phys. Inst. Met. Mater.* 80, 186–191 (2016).
11. Chirkova, A. et al. The effect of the microstructure on the antiferromagnetic to ferromagnetic transition in FeRh alloys. Acta Mater. 131, 31–38 (2017).
12. Takahashi, M. & Oshima, R. Annealing effect on phase transition of equiatomic FeRh alloy. Mater. Trans. JIM 36, 735–742 (1995).
13. Hashimoto, A. et al. Effect of high temperature annealing on non-equilibrium phase transitions induced by energetic ion irradiation in FeRh and Ni,V intermetallic compounds. Jpn. J. Appl. Phys. 53, 05FC08 (2014).
14. Takahashi, M. & Oshima, R. Stress induced phase transition of iron–rhodium alloys. J. Phys. IV 05, C8-491–C8-496 (1995).
15. Zakharov, A. I., Kadomtseva, A. M., Levitin, R. Z. & Ponyatovskii, E. G. Magnetic and magnetoelastic properties of a magnetic iron–rhodium alloy. Sov. Phys. JETP 19, 1348–1353 (1964).
16. Heeger, A. J. Pressure dependence of the FeRh first-order phase transition. J. Appl. Phys. 41, 4751–4752 (1970).
17. Vinokurova, L. I., Vlasov, A. V. & Pardavi-Horváth, M. Pressure effects on magnetic phase transitions in FeRh and FeRh alloys. Phys. Status Solidi B 78, 353–357 (1976).
18. Barua, R., Jiménez-Villacorta, F. & Lewis, L. H. Predicting magnetostructural trends in FeRh-based ternary systems. Appl. Phys. Lett. 103, 102407 (2013).
19. Xie, Y. et al. Effect of epitaxial strain and lattice mismatch on magnetic and transport behaviors in metamagnetic FeRh thin films. AIP Adv. 7, 056314 (2017).
20. Barton, C. W. et al. Substrate induced strain field in FeRh epilayers grown on single crystal MgO (001) substrates. Sci. Rep. 7, 44397 (2017).
21. Chen, I., Gao, Y., Wu, L., Ma, J. & Nan, C.-W. A magnetic glass state over the first-order ferromagnetic-to-antiferromagnetic transition in FeRh film. Mater. Res. Lett. 5, 329–334 (2017).
22. Suzuki, I., Hamasaki, Y., Itoh, M. & Taniyama, T. Controllable exchange bias in Fe/magnetic FeRh bilayers. Appl. Phys. Lett. 105, 172401 (2014).
23. Yap, Q. J. et al. Phase ordering and its effect on magnetic and structural properties of FeRh ultrathin films. J. Appl. Phys. 116, 043902 (2014).
24. Ceballos, A. et al. Effect of strain and thickness on the transition temperature of epitaxial FeRh thin films. Appl. Phys. Lett. 111, 172401 (2017).
25. Bennett, S. P. et al. Giant controllable magnetization changes induced by structural phase transitions in a metamagnetic artificial multiferroic. Sci. Rep. 6, 22708 (2016).
26. Han, G. C. et al. Suppression of low-temperature ferromagnetic phase in ultrathin FeRh films. J. Appl. Phys. 113, 123909 (2013).
27. Jiang, M. et al. Influence of film composition on the transition temperature of FeRh films. J. Cryst. Growth 438, 19–24 (2016).
28. Gatel, C. et al. Inhomogeneous spatial distribution of the magnetic transition in an iron–rhodium thin film. Nat. Commun. 8, 15703 (2017).
29. Ne, J. et al. Magnetic properties of ordered polycrystalline FeRh thin films. RSC Adv. 7, 44097–44103 (2017).
30. Suzuki, I., Koike, T., Itoh, M., Taniyama, T. & Sato, T. Stability of ferromagnetic state of epitaxially grown ordered FeRh thin films. J. Appl. Phys. 105, 07E501 (2009).
31. Loving, M. G. et al. Strain-tuning of the magnetocolor transition temperature in model FeRh films. J. Phys. D Appl. Phys. 51, 024003 (2018).
32. Yamada, S. et al. Low-temperature B2 ordering and magnetic properties of Fe_{50−x}Rh_{x} films on bcc alloys. Phys. Rev. B 92, 094416 (2015).
33. Kande, D., Pisara, S., Weller, D., Laughlin, D. E. & Zhu, J.-G. Enhanced B2 ordering of FeRh thin films using B2 NiAl underlayers. IEEE Trans. Magn. 47, 3296–3299 (2011).
34. Cher, K. M., Zhou, T. J. & Chen, J. S. Compositional effects on the structure and phase transition of epitaxial FeRh thin films. IEEE Trans. Magn. 47, 4033–4036 (2011).
35. Yamada, S. et al. Exchange coupling in metallic multilayers with a top FeRh layer. AIP Adv. 6, 056115 (2016).
36. Witte, R. et al. Epitaxial strain-engineered self-assembly of magnetic nanostructures in FeRh thin films. J. Phys. D Appl. Phys. 50, 025007 (2017).
37. Lee, Y. et al. Large resistivity modulation in mixed-phase metallic systems. Nat. Commun. 6, 5959 (2015).
38. Chen, J. et al. Strain modulated ferromagnetic to antiferromagnetic transition in FeRh/BaTiO_{3}(001) heterostructures. J. Appl. Phys. 121, 194103 (2017).
39. Fina, I. et al. Electric-field-adjustable time-dependent magnetoelectric response in martensitic FeRh alloy. ACS Appl. Mater. Interfaces 9, 15577 (2017).
40. Cherifi, R. O. et al. Electric-field control of magnetic order above room temperature. Nat. Mater. 13, 345–351 (2014).
41. Thiele, J.-U., Maat, S. & Fullerton, E. E. FeRh/FePt exchange spring films for thermally assisted magnetic recording media. Appl. Phys. Lett. 82, 2859–2861 (2003).
42. Busch, M., Gruyter, M. & Winter, H. Spin polarization, structure and chemical composition of the Rh/Fe(100) interface. Surf. Sci. 205(582), 31–41 (2005).
43. Poate, J. M. et al. (eds) Thin Films-Interdiffusion and Reaction (Willey-Interscience, New York, 1978).
44. Colgan, J. E. G. A review of thin-film aluminate formation. Mater. Sci. Rep. 5, 1–44 (1990).
45. Pretorius, K., Theron, C. C., Vantomme, A. & Mayer, J. W. Compound phase formation in thin film structures. Crit. Rev. Solid State Mater. Sci. 24, 1–62 (1999).
46. Myagkov, V. G. et al. Solid state synthesis of Mn_{5}Ge_{3} in Ge/Ag/Mn trilayers: Structural and magnetic studies. J. Solid State Chem. 246, 378–387 (2017).
47. Myagkov, V. et al. Long-range chemical interactions in solid-state reactions: Effect of an inert Ag interlayer on the formation of Fe_{50−x}Rh_{x} in epitaxial Pd_{0}/Ag(001)/Fe(001) and Fe(001)/Ag(001)/Pd(001) trilayers. Philos. Mag. 94, 2595–2622 (2014).
48. Myagkov, V. G., Zhigalov, V. C., Bykova, L. E. & Bondarenko, G. N. Long-range chemical interaction in solid-state synthesis: Chemical interaction between Ni and Fe in epitaxial Ni(001)/Ag(001)/Fe(001) trilayers. Int. J. STD 18, 117–124 (2009).
49. Myagkov, V. G., Bykova, L. E., Li, L. A., Turpanov, I. A. & Bondarenko, G. N. Solid-phase reactions, self-propagating high-temperature synthesis, and martensitic transformations in thin films. Dokl. Phys. 47, 95–98 (2002).
50. Myagkov, V. G., Bykova, L. E., Zharkov, S. M. & Bondarenko, G. N. Formation of NiAl shape memory alloy thin films by solid-state reaction. Solid State Phenom. 138, 377–384 (2008).
51. Myagkov, V. G., Bykova, L. E. & Bondarenko, G. N. Solid-state synthesis and martensitic transformations in thin films. Dokl. Phys. 48, 30–33 (2003).
52. Radu, I. et al. Laser-induced generation and quenching of magnetization on FeRh studied with time-resolved X-ray magnetic circular dichroism. Phys. Rev. B 81, 104415 (2010).
53. Unal, A. A. et al. Laser-driven formation of transient local ferromagnetism in FeRh thin films. Ultramicroscopy 183, 104–108 (2017).
54. Zakharov, A. I., Kadomtseva, A. M., Levitin, R. Z. & Ponyatovskii, E. G. Magnetic and magnetoelastic properties of a magnetic iron–rhodium alloy. Sov. Phys. JETP 19, 1348–1353 (1964).
55. Chen, L. J. Solid state amorphization in metal/Si systems. Mater. Sci. Eng. A Rep. 29, 115–152 (2000).
58. Zhu, Y., Liao, G., Shi, T., Tang, Z. & Li, M. Highly choreographed atomic motion and mechanism of interface amorphization. *Acta Mater.* **125**, 69–80 (2017).
59. Bhatt, P., Chaudhari, S. M. & Fahlman, M. Influence of Ti layer thickness on solid state amorphization and magnetic properties of annealed Ti/Ni multilayer. *J. Phys. Condens. Matter* **19**, 376210 (2007).
60. Cho, H., Kim, H. Y. & Miyazaki, S. Allopying process of sputter-deposited Ti/Ni multilayer thin films. *Mater. Sci. Eng. A* **438–440**, 699–702 (2006).
61. Zhang, L. et al. Amorphous martensite in ß-Ti alloys. *Nat. Commun.* **9**, 506. https://doi.org/10.1038/s41467-018-02961-2 (2018).
62. Myagkov, V. G., Zhigalov, V. C., Bykova, L. E. & Bondarenko, G. N. Solid-state synthesis and phase transformations in Ni/Fe films: Structural and magnetic studies. *J. Magn. Magn. Mater.* **305**, 534–545 (2006).
63. Ohnuma, I., Gendo, T., Kainuma, R., Inden, G. & Ishida, K. Phase equilibria and thermodynamic evaluation approximating short-range ordering energy in the Fe–Rh binary system. *ISIJ Int.* **49**, 1212–1219 (2009).
64. Chikazumi, S. Epitaxial growth and magnetic properties of single-crystal films of iron, nickel, and permalloy. *J. Appl. Phys.* **32**, S81–S82 (1962).
65. de Vries, M. A. et al. Hall-effect characterization of the metamagnetic transition in FeRh. *New J. Phys.* **15**, 013008 (2013).

**Acknowledgements**
The authors thank L.A. Solovyov for their help on the orientation relationships measurements and I.V. Nemtsev for assisting with the measurements of the samples’ composition. This work was supported by RFBR together with the Government of the Krasnoyarsk Territory, the Krasnoyarsk Regional Fund of Science (project № 19-43-240003). The work is partially based upon the experiments performed on Krasnoyarsk Regional Center of Research Equipment of Federal Research Center «Krasnoyarsk Science Center SB RAS».

**Author contributions**
A.A.I. and V.S.Z. prepared the samples and characterized them. V.S.Z. carried out annealing experiments. D.A.V. characterized the samples by SQUID magnetometry. M.N.V. performed TEM experiments. L.E.B. and A.A.M. planed and performed resistance measurements. G.N.B. carried out the X-ray diffraction experiments. V.G. initiated the study and wrote the manuscript with inputs from L.E.B. and A.A.M. All authors contributed to the manuscript and the interpretation of the date.

**Competing interests**
The authors declare no competing interests.

**Additional information**
*Supplementary information* is available for this paper at https://doi.org/10.1038/s41598-020-67837-2.

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