Realization of magnetostructural coupling by modifying structural transitions in MnNiSi-CoNiGe system with a wide Curie-temperature window

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The magnetostructural coupling between structural and magnetic transitions leads to magneto-multifunctionalities of phase-transition alloys. Due to the increasing demands of multifunctional applications, to search for the new materials with tunable magnetostructural transformations in a large operating temperature range is important. In this work, we demonstrate that by chemically alloying MnNiSi with CoNiGe, the structural transformation temperature of MnNiSi (1200 K) is remarkably decreased by almost 1000 K. A tunable magnetostructural transformation between the paramagnetic hexagonal and ferromagnetic orthorhombic phase over a wide temperature window from 425 to 125 K is realized in (MnNiSi)_{1-x}(CoNiGe)_{x} system. The magnetic-field-induced magnetostructural transformation is accompanied by the high-performance magnetocaloric effect, proving that MnNiSi-CoNiGe system is a promising candidate for magnetic cooling refrigerant.

Magnetostructural transformation (MST), a coupling between the structural and magnetic transition, attracts considerable attention due to various associated interesting magnetoresponsive effects, such as magnetic shape memory effect, magnetic field-induced strain, magnetic anisotropy, and magnetoelastic effect. These effects show potential applications in sensors, magneto-mechanical devices, energy-harvesting devices, magnetic cooling refrigeration, and so on. In order to realize MST in a phase-transition material, a large magnetization difference (ΔM) between the two structural phases is essential and is always pursued to increase the magnetic field-driving capacity. If MST in a given system is tuned to occur between a paramagnetic (PM)/antiferromagnetic (AFM) state and a ferromagnetic (FM) state, rather than between two FM states, a large ΔM will be gained. As an example, MST is widely observed in Heusler-type magnetic shape memory alloys, which show ferromagnetic martensitic transitions.

Based on this viewpoint, another type of magnetic shape memory material, the hexagonal MnM′X compound (M′ = Co or Ni, X = Si or Ge) is designed to obtain MST. In stoichiometric MnM′X compounds, the structural transformation takes place from a martensitic-like hexagonal Ni2In to anorthorhombic TiNiSi structure during the cooling process. The main challenge in MnM′X compounds is that the structural transformation temperature (Tt) is usually much higher than the magnetic-ordering temperatures of both hexagonal and orthorhombic phases, and the transformation occurs between two PM states, resulting in a low ΔM and low magnetic field-driving capacity.

It is known that, in MnM′X compounds, the stability of Ni2In and TiNiSi structures is highly dependent on the covalent bonds between M′ and X atoms, and that between the neighbouring Mn and Mn atoms. The stoichiometric tuning, foreign-atoms-substituting, and applied stress may alter the strength of covalent bonds, thus influence Tt. For example, by introducing interstitial B atoms or substituting Cu for Mn

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atoms, $T_t$ of MnCoGe can be reduced from 420 K to below Curie temperature of the orthorhomic phase ($T_C$) and MST from PM hexagonal to FM orthorhombic phase is realized during cooling$^{22,31}$. In the case of MnNiGe system, the stoichiometric tuning can reduce $T_t$ from 470 K to below than 200 K and obtain MST from FM hexagonal to AFM orthorhombic phase with decreasing temperature$^{20}$. Moreover, it is reported that if $T_t$ lies in the Curie-temperature window (CTW), which is the range between Curie temperatures of the hexagonal and orthorhombic phases$^{16,33,34}$, the structural transition will couple with magnetic state changes, bringing a large $\Delta M$. The CTW is expected to be broad enough, so that MST and the coupled magnetoresponsive effects can be freely tailored in a large temperature range. However, in MnM’Ge-based compounds, the relatively low magnetic ordering temperatures restrict the further enlargement of CTW.

Aimed at the potential applications with magnetoresponsive effects, a good candidate of MnM’X compound with MST should have both a large $\Delta M$ and a wide CTW. As a member of MnM’X family, MnNiSi has a high $T_C$ of 622 K$^{35}$, indicating a potential large CTW. However, $T_t$ in stoichiometric MnNiSi alloy is as high as 1200 K$^{24}$, which is almost two times higher than those in MnCoGe and MnNiGe$^{24–27}$. Compared to the previously mentioned alloy systems, to effectively tune down $T_t$ to below $T_C$ is a big obstacle in MnNiSi system before the realization of MST$^{36,37}$. Recently, the isostructural alloying opens up a new feasibility to realize the magnetostuctural coupling in MnNiSi-based compounds$^{16,33,34,38–41}$. By applying this method, the wide CTW can be further obtained. In this work, by Co and Ge co-substitution, we perform the isostructural alloying of MnNiSi and CoNiGe. In this (MnNiSi)$_{1-x}$(CoNiGe)$_x$ system, $T_t$ is successfully tuned down to below than $T_C$, and the tunable MST between PM hexagonal and FM orthorhombic phase can be obtained in a large CTW by altering the CoNiGe-content. Due to the large $\Delta M$ between two phases, the observed MST can be induced by the external magnetic field at room temperature (RT). The effect is accompanied by a large magnetocaloric effect, indicating the potential applications in magnetic cooling refrigerator.

**Results**

**Structural characterization.** Figure 1(a) shows X-ray diffraction (XRD) patterns of (MnNiSi)$_{1-x}$(CoNiGe)$_x$ measured at RT. Inset: $a_{ortho}/b_{ortho}$ and $c_{hex}/a_{hex}$ ratios. (b) Temperature-dependent XRD of (MnNiSi)$_{0.66}$(CoNiGe)$_{0.34}$ from 353 to 493 K. Inset: cell volumes from 353 to 493 K for (MnNiSi)$_{0.66}$(CoNiGe)$_{0.34}$.

![Figure 1](https://www.nature.com/scientificreports/)
transits to Ni$_2$In structure with increasing temperature. According to XRD analysis, the temperature-dependent unit-cell volume is calculated, shown in the inset of Fig. 1(b). A large decrease of 2.5% in unit-cell volume is observed during the structural transformation on heating, indicating a remarkable atomic displacement during the structural reconstruction.

MST and magnetic phase diagram. In order to confirm the CoNiGe-content dependent $T_t$, DSC measurements of (MnNiSi)$_{1-x}$(CoNiGe)$_x$ ($x = 0.32, 0.33, 0.34, 0.35, 0.36, 0.37, 0.38, 0.39, 0.40, 0.41$ and $0.42$) were carried out upon cooling and heating with a ramp rate of 10 K/min, which are shown in Fig. 2. The observed large endothermic/exothermic peaks during heating/cooling cycles are associated with the latent heat of the first-order structural transitions between TiNiSi and Ni$_2$In structures. The thermal hysteresis between heating and cooling cycles signifies the first-order nature of structural transformation. For the sample with $x = 0.34$, the endothermic peak appears at 410 K, which agrees well with the temperature-dependent XRD analysis. It is also found that the endothermic and exothermic peaks shift towards lower temperatures with the increase of CoNiGe-content. This phenomenon verifies that the introduction of CoNiGe can reduce $T_t$ of MnNiSi from 1200 K to be below RT.

Since the magnetic properties of MnM′X alloys are sensitive to the Mn-Mn distances$^{44}$, the large distortion of unit-cell during the structure transformation may bring about considerable changes of magnetic states. The temperature dependences of magnetization ($M$-$T$) for (MnNiSi)$_{1-x}$(CoNiGe)$_x$ ($x = 0.36, 0.38$ and $0.41$) measured during heating and cooling with a ramp rate of 2 K/min at a magnetic field of 0.1 T are shown in Fig. 3(a) (some other representative M-$T$ curves are shown in the supporting information). A sharp magnetic transition from the high-temperature PM to low-temperature FM state is observed during cooling. The transition shifts to the lower temperatures with the increase of CoNiGe-content. The obvious thermal hysteresis, reflecting the irreversibility between cooling and heating cycles, suggests the first-order nature of the transition. $T_t$, here defined as the temperature where |$dM/dT$| is the maximum, agrees well with the DSC measurement. These results prove that the studied samples experience MST between FM orthorhombic and PM hexagonal phase as the temperature changes. While it is worth noting that Mn atoms carry the majority of magnetic moments in MnNiSi alloys$^{35}$. To investigate the saturated magnetization, $M_s$, of MnNiSi-CoNiGe system, M-$B$ curves of some samples are measured at 4.2 K in Fig. 3(b). As shown in the inset, $M_s$ decreases with increasing CoNiGe-content, and it is lower.
than that of stoichiometric MnNiSi (2.62 μB/f.u.)\(^35\). It indicates that the substitution of large-moment Mn atoms by small-moment Co atoms modifies the exchange interactions between Mn-Mn atoms\(^37\).

According to DSC and magnetic measurements, the structural and magnetic phase diagram of \((\text{MnNiSi})_{1-x}(\text{CoNiGe})_x\) system is obtained, as shown in Fig. 4. The sample with \(x = 0.32\) transits from PM hexagonal to PM orthorhombic phase at 450 K, then to FM orthorhombic phase at 425 K during cooling. Upon the further increase of CoNiGe-content to 0.43, \(T_t\) continuously decreases to 125 K. When \(x\) is higher than 0.43, MST disappears and a weak magnetic spin-glass-like state is found, similar as \(\text{Mn}_{1-x}\text{Fe}_x\text{NiGe}\)\(^16\). Thus, a CTW ranged from 425 to 125 K is established in \((\text{MnNiSi})_{1-x}(\text{CoNiGe})_x\) system, where alloys undergo a tunable MST coupled with a magnetic transition from PM to FM state.

Magnetic-field-inducing MST and magnetocaloric effect. In the case of Ni-Mn based ferromagnetic shape memory alloys, due to the large \(\Delta M\) between FM austenite and weak magnetic martensite, the austenite is energetically more favorable in the magnetic field, giving rise to the decrease of \(T_t\) and the magnetic-field-inducing MST from the martensite to austenite\(^1\). Similarly, in \((\text{MnNiSi})_{1-x}(\text{CoNiGe})_x\) system with a first-order PM-FM transition, the magnetic field will stabilize FM TiNiSi structure and lead to a MST from the hexagonal to orthorhombic phase. For the sample with \(x = 0.38\), \(T_t\) increases 4.8 K under a magnetic field of 5 T, indicating that MST can be induced by the magnetic field (Fig. 5(a)). Due to the discontinuities of spin and lattice, MST is associated with a large magnetic entropy change (\(\Delta S\)). It is known that the maximum value of \(\Delta S\) can be estimated from \(M-T\) curves measured at different constant fields (here, \(B = 0.1\) and 5 T, respectively) using the Clausius-Clapeyron equation:

\[
\frac{\Delta S}{\Delta M} = -\frac{\partial B}{\partial T} 
\]

For \((\text{MnNiSi})_{0.62}(\text{CoNiGe})_{0.38}\), the calculated maximum value of \(\Delta S\) is \(-41.8\) J/(kg·K), where \(\Delta M = 41\) A·m\(^2\)/kg, \(\Delta B = 4.9\) T and \(\Delta T = 4.8\) K (Fig. 5(a)). To further confirm the magnetocaloric effect during the transition, \(\Delta S\) in the heating process is also calculated from the isothermal magnetization curves (\(M-B\) curves in Fig. 5(b)) using the Maxwell relation. As shown in Fig. 5(c), the maximum value of \(\Delta S\) is \(-40.3\) J/(kg·K) at 295 K, which agrees with the value obtained by Clausius-Clapeyron equation. In \((\text{MnNiSi})_{1-x}(\text{CoNiGe})_x\) system, the large CTW offers the possibility to obtain large \(\Delta S\) values in the temperature range of nearly 300 K. As an example, the substitution levels of \(x = 0.40\) and 0.39 give rise to \(\Delta S\) of \(-31.7\) and \(-30.5\) J/(kg·K) for \(\Delta B = 5\) T at 245 and 270 K, respectively (Fig. 5(c)). The observed maximum \(\Delta S\) is larger than some promising magnetocaloric systems, such as some \(\text{MnM'}\text{Ge}\)-based systems\(^16,30,33\), Ni-Mn based alloys\(^7,8\) and rare earth-transition metal intermetallic compounds\(^45\).

The effective refrigeration capacity (\(R_{\text{Ceff}}\), which is calculated by subtracting the average hysteresis loss (HL) from the refrigeration capacity (RC) value, is commonly adopted to evaluate magnetocaloric effect\(^46\). For sample with \(x = 0.38\), RC value is 170.1 J/kg around room temperature, numerically calculated by integrating the area under \(\Delta S-T\) curves, using the temperatures at half-maximum of the peak as the integration limits. And HL, calculated from the area surrounded by hysteresis loops (\(M-B\) curves), is negligible, as shown in the inset of Fig. 5(c). Therefore, \(R_{\text{Ceff}}\) is about 169.8 J/kg for this sample under the field change of 0–5 T. These indicate the potential applications for the RT magnetic cooling refrigerator. Besides, as mentioned above, the MnNiSi-CoNiGe system undergoes large changes of lattice parameters and unit volume during the transition (Fig. 1(a)), which can be induced by the applied magnetic field. This may be utilized for the strain-based application\(^15\).

Discussion

The observed MST between FM orthorhombic and PM hexagonal phase is achieved by adjusting \(T_t\) into the CTW. However, as mentioned above, \(T_t\) of MnNiSi is as high as 1200 K and cannot be efficiently decreased by the conventional methods\(^36,37\). Here, the question is why Co and Ge co-substitution can sharply reduce \(T_t\) to below \(T_C\), leading to the observed MST. In \(\text{MnM'}\text{X}\) alloys, covalent bonds form between \(\text{M'}\) and \(\text{X}\) atoms and between the neighbouring Mn and Mn atoms both in TiNiSi and Ni\(_2\)In structures\(^36,28,29\). These covalent bonds act...
as skeletons, stabilizing the crystalline structure\textsuperscript{16,29,47}. The structure transformation can be understood as a competition between the strengths of covalent bonds in TiNiSi and Ni\textsubscript{2}In structures\textsuperscript{16,34,47}. Thus, altering the strength of covalent bonds can influence $T_t$. For the system with $M' = \text{Ni}$, $T_t$ of MnNiGe lies around 470 K\textsuperscript{24,25}, which is 730 K lower than that of MnNiSi\textsuperscript{24}. This suggests that altering the main-group elements can influence the strength of covalent bond, leading to the change of $T_t$. Similar phenomenon is found in Al substituted MnNiGe alloy\textsuperscript{48}. Except for altering elements which form $M'$-X covalent bond, the decrease of $T_t$ is also observed when Mn is replaced by other 3$d$-atoms, such as Mn\textsubscript{1−x}Fe\textsubscript{x}CoGe alloy\textsuperscript{49}. Based on the site-preference rule\textsuperscript{50}, Fe atoms occupy Mn sites and induce lattice change in $c$ axis. This change will influence the separation of Mn atoms, leading to the enhancement of the strength of Mn-Mn covalent bonds, which is helpful to stabilizing the Ni\textsubscript{2}In structure. For MnNiSi, $T_t$ cannot be efficiently reduced by the single-element substitution\textsuperscript{36,37}. However, when CoNiGe is introduced, Co atoms occupy 3$d$-atom Mn sites, Ge atoms occupy main-group Si sites, and $T_t$ decreases by almost 1000 K with increasing CoNiGe-content.

From the view of applications, a large CTW enables tunable magnetoresponsive effects in a wide temperature range. In this (MnNiSi)$_{1−x}$(CoNiGe)$_x$ system, the CTW is as large as 300 K, which is comparable to the previously

Figure 5. (a) M-T curves of (MnNiSi)$_{0.62}$(CoNiGe)$_{0.38}$ for applied fields $B = 0.1$ T and 5 T. (b) Isothermal M-B curves measured around $T_t$ for the sample with $x = 0.38$ (c) Temperature dependence of $\Delta S$ in the field changes of 0–2 T and 0–5 T for the samples $x = 0.38$, 0.39 and 0.40. Inset: magnetic hysteresis loss of sample with $x = 0.38$.
reported systems, such as MnCoGe-based system\textsuperscript{21}, Mn\textsubscript{1−x}Co\textsubscript{x}NiGe\textsubscript{33}, MnNiGe:Fe system\textsuperscript{16}, (Mn, Fe)Ni(Fe, Si)\textsuperscript{34} and Gd-Si-Ge alloys\textsuperscript{51}. It is known that the realization of FM-FM-type MST should meet the condition that $T_t$ can be tuned into the temperature window between the magnetic-ordering temperatures of orthorhombic and hexagonal phases. In (MnNiSi)\textsubscript{1−x}(CoNiGe)\textsubscript{x} system, $T_t$ of orthorhombic phase is around 425 K (shown in Fig. S1). However, with the increase of CoNiGe-content, a spin-glass-like state appears and the magnetic order-disorder transition of hexagonal phase is still not observed (Fig. S1). This phenomenon indicates that the magnetic ordering temperature of hexagonal phase is lower than 125 K. Therefore, a large CTW between 425 and 125 K is established in (MnNiSi)\textsubscript{1−x}(CoNiGe)\textsubscript{x} system.

**Conclusions**

To summarize, we have successfully realized a PM-FM magnetostructural coupling in (MnNiSi)\textsubscript{1−x}(CoNiGe)\textsubscript{x} system. By introducing CoNiGe which possesses a stable Ni$_3$In structure, $T_t$ decreases sharply, resulting in the first-order MST between FM orthorhombic and PM hexagonal phase. This is due to the enhancement of covalent bonds which modifies the relative stability of two structures. Besides, a large CTW of 300 K is established, which will benefit the multifunctional applications of the materials over a wide temperature range. Due to the large $\Delta M$ during the transition, MST can not only be induced by temperature, but also by magnetic field. Large, tunable magnetocaloric effect generated from MST is obtained. The magnetic entropy change of (MnNiSi)\textsubscript{0.62}(CoNiGe)\textsubscript{0.38} reaches $-40.3\text{J/(kg·K)}$ under the field change of 0–5 T around RT, suggesting the potential applications in magnetic cooling refrigerator.

**Methods**

The samples with nominal compositions of (MnNiSi)\textsubscript{1−x}(CoNiGe)\textsubscript{x} ($x = 0.32, 0.33, 0.34, 0.35, 0.36, 0.37, 0.38, 0.39, 0.40, 0.41, 0.42, 0.43$ and $0.44$) were prepared by arc-melting the appropriate amounts of raw materials in a water-cooled copper crucible under a high purity argon atmosphere for three times. As-cast ingots were annealed in vacuumed quartz tubes at 1073 K for four days before quenched into the cold water.

The crystal structures of the specimens were identified by X-ray diffraction (XRD, Bruker, D8 Advance) analysis with Cu-K$_\alpha$ radiation. The structural transitions were investigated by differential scanning calorimetry (DSC, Mettler Toledo, DSC 3).

Magnetic measurements were performed with vibrating sample magnetometer (VSM, LakeShore, 7407) and Physical Property Measurement System (PPMS, Quantum Design, Dynacool). Isothermal magnetic entropy change ($\Delta S$) was calculated from the isothermal magnetization curves using Maxwell equation (2):

$$\Delta S(T, H) = S(T, H) - S(T, 0) = \int_0^H \frac{\partial M}{\partial T} \, dH$$

(2)

To avoid the irreversibility in the magnetic-field-induced first-order MST, isothermal magnetization curves were measured using a so-called loop process\textsuperscript{36}: in the heating process, the isothermal magnetization were measured with a temperature interval of 2 K in the magnetic field variation from 0 to 5 T. For each M-B curve, the samples were initially cooled down to complete orthorhombic region (for our samples, it's around 150 K) at 5 K/min. Then the samples were heated slowly to the measuring temperature at 3 K/min. To guarantee the temperature stability of measurement, a waiting time of 300 s was held at the initial and targeted temperatures. Besides, for the samples with $x = 0.38, 0.39$ and 0.40, the highest loop temperatures were 301, 288 and 261 K, respectively.

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Acknowledgements
This work is sponsored by National Natural Science Foundation of China (Grant No: 51271093, 51571121), Fundamental Research Funds for the Central Universities (Grant No: 30920140111010), Jiangsu Natural Science Foundation for Distinguished Young Scholars (Grant No: BK20140035), Cooperative Innovation Fund of Jiangsu Province, and China Postdoctoral Science Foundation (Grant No: 2015M580495). This work is also supported by the Priority Academic Program Development of Jiangsu Higher Education Institutions.

Author Contributions
J.L. and F.X. proposed the idea, J.L. prepared the samples, G.P. carried out the magnetic measurement, I.A.S. and N.U.H. performed the DSC measurement, J.L. and Y.Y.G. analyzed the data, Y.Y.G. and G.Z.X. provided fruitful discussions.

Additional Information
Supplementary information accompanying this paper at http://www.nature.com/srep
Competing financial interests: The authors declare no competing financial interests.
How to cite this article: Liu, J. et al. Realization of magnetostructural coupling by modifying structural transitions in MnNiSi-CoNiGe system with a wide Curie-temperature window. Sci. Rep. 6, 23386; doi: 10.1038/srep23386 (2016).

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