Tuning the Magnetocaloric Properties of the La(Fe,Si)$_{13}$ Compounds by Chemical Substitution and Light Element Insertion

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Abstract: LaFe$_{13-x}$Si$_x$ compounds exhibit a giant magnetocaloric effect and are considered as a good magnetocaloric working substance for an environmentally friendly cooling technique. Nevertheless as the Curie temperature $T_C$ is around 200 K, it is necessary to tune $T_C$ near room temperature for magnetic refrigeration. In this work we present a review of the various methods of synthesis and shaping of the LaFe$_{13-x}$Si$_x$ type compounds as well as the influence of chemical substitution, light element insertion or combination of both on $T_C$, magnetic entropy and adiabatic temperature variation ($\Delta S_M$ and $\Delta T_{ad}$), and stability upon cycling. The advantages and drawbacks of each method of preparation and type of element substitution/insertion are discussed. The implementation of these NaZn$_{13}$ type materials in active magnetic refrigerator is presented and their performances are compared to that of Gd in prototypes.

Keywords: intermetallics; NaZn$_{13}$ compounds; magnetocaloric effect; magnetic refrigeration

1. Introduction

Classical refrigeration technologies are using refrigerants that deplete the ozone layer and contribute to global warming, and will be therefore forbidden by different climate protocols. Alternative refrigerants present also various drawbacks (flammable, toxic). Therefore, it is worth developing new refrigeration technologies without environmental problems [1,2], such as magnetic refrigeration that is based on the magnetocaloric effect. The development of efficient magnetocaloric materials has become challenging since the discovery of a giant magnetocaloric effect (MCE) near room temperature (RT) in Gd(Ge,Si)$_5$, MnAs$_{1-x}$Sb$_x$, and FeRh based alloys [3–11].

Since that time, intensive studies have yielded the discovery of several families of materials, among which the La(Fe,Si)$_{13}$ system, which displays a giant magnetocaloric effect, and is considered as a good magnetocaloric material for magnetic refrigeration. This system is particularly interesting, because it has relatively cheap, abundant, and non-toxic constituents. However several questions should be solved to use this material for magnetic refrigeration: (i) the synthesis and shaping of the material should be fast, not too expensive, and appropriate for an industrial scaling, (ii) the chemical composition should be adjusted to reach room temperature, as La(Fe,Si)$_{13}$ Curie temperature is close to 200 K, while keeping a giant MCE effect, and (iii) the compounds should remain stable upon a large number of thermomagnetic cycles. In this paper, we first propose a review of the main works that have been undertaken to solve these different challenges. The different methods of synthesis and shaping will be presented and then compared. The influence of both chemical substitution of La, Fe, and Si by other elements as well as the insertion of light elements (H, C) will be presented to show the advantages and drawbacks of each solutions.

We have developed a synthesis method combining high energy ball milling with a subsequent short annealing treatment in order to obtain a single-phase compound. Besides, a reactive Spark Plasma Sintering (SPS) method has been used in order to produce and
sinter a powdered sample in one single step. The carbon atoms were inserted by solid/solid reaction, while the hydrogenation of the parent intermetallic La(Fe,Si,Co)_{13} alloys was carried out using a Sievert method. A study of our own experimental results will then be presented, which combines Fe for Co substitution with the hydrogen insertion in order to give an example of a strategy developed to obtain performant magnetocaloric materials. The insertion of light elements (H, C) or/and cobalt for iron substitution are used in order to improve the Curie temperature near room temperature for magnetic refrigeration applications. The tuning of the Curie temperature is explained by a combination of the structural and electronic properties. The advantages and disadvantages of each type of element insertion are discussed and the misunderstanding on interstitial site is clarified.

2. Sample Synthesis

Condition of Synthesis of La(Fe_{1−x}Si_{x})_{13} Type Alloys

Contrary to LaCo_{13}, the binary LaFe_{13} compound does not form, but the 1:13 phase can be stabilized by a partial Si for Fe substitution. The study of the ternary La-Fe-Si ternary phase diagram at 1173, 1373, and 1573 K [12,13] showed that the LaFe_{13−x}Si_{x} compounds only exist in a very narrow range of composition (with La/(Fe+Si) =1/14) and crystallize either in the cubic NaZn_{13} type structure (phase τ_1) or the tetragonal Ce_{2}Ni_{17}Si_{9}-type structure (phase τ_2). At 1200 K, τ_1 is single phase for 0.08 ≤ x ≤ 0.21 and τ_2 for 0.25 ≤ x ≤ 0.40. Both of the phases coexist between 0.21 and 0.25. The Si concentration at the boundaries varies with the equilibrium temperature. The NaZn_{13} phase forms via a peritectic reaction at T = 1665 K:

\[ \text{LFeSi} + \text{La} + \text{LaFe}_2\text{Si}_2 \rightarrow \tau_1 \quad \text{or} \quad \alpha-\text{Fe} + \text{La} + \text{LaFe}_2\text{Si}_2 \rightarrow \tau_1 \]

The as-cast alloys are constituted by a mixture of α-Fe(Si) and LaFeSi (formed from the liquid L_{La} phase upon cooling) and an appropriate thermal treatment is necessary for obtaining the NaZn_{13} τ_1-phase via a diffusion process [14–16]. The single phase NaZn_{13} samples can be synthesized by arc or induction melting with a further annealing treatment at temperatures between 1273 and 1423 K and during several days or weeks [11,12]. It is necessary to take the risk of La oxidation into account and an excess of La is generally added in order to avoid the formation of secondary phases [17]. The chemical purity of each element, their preparation into a glove box, and the quality of the atmosphere upon melting and annealing (secondary vacuum or Argon pressure) are crucial to obtain single phase compounds. The annealing time can be shortened by increasing the annealing temperature. In ref [18] the NaZn_{13} was obtained for an arc-melted sample annealed 1 h at 1573 K, but some α-Fe was still present. However, in order to prepare a large quantity of sample for magnetic refrigeration application, the too long duration or too high temperature of the annealing treatment remains an economical drawback. Therefore, to reduce the annealing time several alternative synthesis methods have been developed. They are mainly based to a reduction of the grain size and an intimate contact between the various phases obtained as cast to shorten the diffusion path. Among these methods, one can find melt-spinning [19–23], melt extraction [24], drop-tube solidification [25], and solid state sintering [26]. Gas atomization, followed by an annealing treatment of 1 hour at 1323 K, has been developed by Erasteel Company, in order to produce batches of 500 kg of alloys for industrial applications [27]. Another industrial alternative was developed by the Vacuumschmelze Company [28,29], based on powder metallurgy using reactive sintering.

Our group has developed a high energy ball milling method starting with a mixture of LaSi alloys, α−Fe and Si. A ball milling of only 1 hour, followed by an annealing treatment of 30 min at 1423 K, were sufficient for obtaining 95% of the NaZn_{13} phase with very small quantities of α−(Fe,Si), La_{2}O_{3} and La(OH)_{3} [30]. A comparison of the results that were obtained by these different techniques can be found in [31].

Furthermore, it is also necessary to shape the magnetocaloric material in appropriate form as a function of the design of the magnetic refrigeration device. Several methods have been investigated, like extrusion after mixing with an epoxy polymer [27], thermal
decomposition and recombination (TDR) process [29], and selective laser melting [32]. Erasteel has been chosen to mix the powder that was obtained from gas atomization with epoxy, whereas Vacuumschmelze has developed the TDR process in order to shape their materials in specific form. They prepare an alloy containing a mixture of α−Fe and LaFeSi, which can be easily cut without breaking and then process to the annealing treatment in order to obtain the NaZn_{13} phase.

Our group also succeeded in obtaining a NaZn_{13} sample with a density that is very close to that of bulk sample by combining high energy ball milling with reactive spark plasma sintering (SPS). It was possible to directly obtain the NaZn_{13} phase by sintering the non-annealed BM sample at 1273 K with a heating rate of 100 K/min [31]. The best results were obtained under Ar atmosphere with a final step of 15 min at 1273 K [33].

3. Properties of the La(Fe_{1−x}Si_{x})_{13} Compounds

Table 1 reports the crystallographic data of both τ_{1} (NaZn_{13}) and τ_{2} (Ce_{2}Ni_{17}Si_{9}) phases. The crystal structure of the NaZn_{13} phase is shown in Figure 1. The evolution of the cell parameters versus the Si content have been detailed in [34]. The cubic cell parameter slightly decreases versus Si content (a ≈ 11.46 Å). The tetragonal distortion corresponds to an expansion along the c axis and an intermediate tetragonal τ’_{2} phase is found as the distortion progressively increases. The cell volume decrease becomes steeper upon the tetragonal distortion for a Si concentration larger than 20 at. %.

Table 1. Crystallographic data of the La(Fe,Si)_{13} phases [34].

| Phase         | Pearson | Space Group | Atm. | Wyck. | Sym. | Position |
|---------------|---------|-------------|------|-------|------|----------|
| NaZn_{13}     | cF112   | Fm3c        | La   | 8a    | 432  | 1/4, 1/4, 1/4 |
|               |         |             | Fe   | 6b    | m3   | 0, 0, 0   |
|               |         |             | Fe, Si | 96i | m..  | 0.01806, 0.1192 |

| Ce_{2}Ni_{17}Si_{9} | tI56 | 44/mcm     | La   | 4a    | 422  | 0, 0, 1/4 |
|                     |      |            | Fe1  | 4d    | m..m | 0, 1/2, 0 |
|                     |      |            | Fe2  | 16k   | m..  | 0.2024, 0.0691, 0 |
|                     |      |            | Fe3  | 16l   | ..m  | 0.1294, 0.6294, 0.1832 |
|                     |      |            | Si   | 16l*  | ..m  | 0.33, 0.83, 0.118 |

Figure 1. The crystal structure of the cubic NaZn_{13} phase. The La atom occupy the 8a sites (Na), whereas the Fe atoms are located on the 8b and 96i sites (Zn). In the right figure: one Fe (8b) atom is surrounded by 10 Fe (96i) and 8 La (8a). The Si atoms are substituted in the 96i position.

All of the La(Fe_{1−x}Si_{x})_{13} compounds are ferromagnetic. The Curie temperature \( T_C \) monotonously increases from 200 to 260 K for \( 0.1 < x < 0.21 \), and then decreases abruptly reaching 60 K for \( x = 0.34 \) [34]. The spontaneous magnetization decreases linearly versus Si content with a discontinuity at the \( \tau'_{2} - \tau_{2} \) transition. The evolution of the magnetic entropy variation \( \Delta S_M \) at the transition of the \( \tau_{1} \) phase was studied in [30]. It showed that the \( \Delta S_M \) peak was larger and higher for \( x = 0.108 (\Delta S_M^{\text{max}} = 32.6 \text{ J/kgK}) \) and
broadens progressively as the Si content increases. This indicates a diminution of the first order character of the transition. As the magnetic properties of the $\tau_2$ phase are not suitable for magnetocaloric application and they will not be discussed in the following.

Thermal expansion measurements for the $\tau_1$ phase showed that the cell volume decreases sharply at $T_C$, thus confirming the first order character of the ferromagnetic-paramagnetic transition [35]. Above $T_C$, an itinerant electron metamagnetic (IEM) behavior is observed. The linear decrease of $T_C$ under applied pressure confirms its strong dependence versus cell volume variation, characteristic of the magnetovolume behavior of IEM compounds.

The NaZn$_{13}$ compounds show hysteresis that yields energy loss upon magnetic cycles due to the first order character of the metamagnetic FM-PM transition. Therefore, several studies have been undertaken in order to reduce the hysteresis [36–39]. It was found that the hysteresis increases with the field sweep rates, but it can be reduced by playing on the sample shape to lower the demagnetization coefficient. Indeed, the hysteresis is smaller for thin or porous samples and fragments when compared to bulk samples. The hysteresis can be also significantly reduced when the applied field is parallel to the smaller dimension of the sample. The presence of local chemical inhomogeneity’s also play a role in the local magnetic properties [40].

Therefore, La(Fe$_{1-x}$Si$_x$)$_{13}$ compounds with the NaZn$_{13}$ structure are very interesting for their giant magnetocaloric effect and other advantages (moderate cost, environmental friendly), but their Curie temperature remains too low for room temperature magnetic refrigeration. Therefore, many works have been undertaken to reach an operating room temperature range while keeping a giant MCE: the substitution of La or Fe, insertion of light elements (H, C), or combination of both. The main and most interesting results will be described in the following part of this paper.

4. Effect of Iron Substitution in La(FeSiM)$_{13}$ Compounds

The Curie temperature of La(Fe$_{13}$Si)$_{13}$ alloys was found to be much lower than the LaCo$_{13}$ alloy (1318 K) [41]. The main reason is that Fe-Fe interactions are much weaker than the Co-Co interactions in NaZn$_{13}$-based systems. This fact can explain the increase of $T_C$ in the Co-substituted La-Fe-Si alloys, which was attributed to the strong Co-Fe and Co-Co exchange interactions.

Hu et al. [42], Liu et al. [21], Katter et al. [28], Bjork et al. [43], and Hansen et al. [44] have substituted cobalt for iron atoms, while preserving the first-order magnetic phase transition from paramagnetic to ferromagnetic state at $T_C$ in order to increase the Curie temperature of LaFe$_{13-2x}$Si$_x$. The obtained compound, LaFe$_{11.2}$Co$_{0.7}$Si$_{1.1}$ presents a very large magnetic entropy change $\Delta S_M = 20.3$ J/kgK under a magnetic field change $\Delta H = 5$ T at $T_C = 274$ K. The main reason of this large $\Delta S_M$ is the huge negative lattice expansion at $T_C$. In addition, the LaFe$_{11.2}$Co$_{0.7}$Si$_{1.1}$ compound exhibits a small temperature hysteresis.

At the Curie temperature, the unit cell parameter decreases sharply as the structure stay cubic with $Fm\bar{3}c$ space group. The unit cell parameter in the paramagnetic state was smaller than in the ferromagnetic one, with a change of 0.43% for this magnetic state transition.

For LaFe$_{11.2}$Si$_{1.8}$ compound Hu et al. [45] substituted Fe by Co and studied La(Fe$_{1-x}$Co$_{x}$)$_{11.2}$Si$_{1.8}$ ($x = 0 – 0.08$) compounds, that they prepared by arc melting and subsequent annealing at 1323 K. They found that the Curie temperature derived from the ZFC-FC magnetization increases from 222 K for $x = 0$ to 307 K for $x = 0.08$, while the magnetic entropy change decreases from 13 J/kg K for $x = 0$ to 8 J/kg K for $x = 0.08$ under magnetic field variation of 0 – 5 T. The later value of $\Delta S_M$ is just smaller than Gd at the same magnetic field changes 0 – 5 T. Another interesting result observed is the disappearance of the field induced itinerant electron metamagnetic transition with cobalt content; indeed, Hu et al. [45] underlined that the substitution of iron by cobalt induced a disappearance of the asymmetrical broadening of $\Delta S_M$ peak.

Figure 2 highlights the fact that Si content has much less influence on $T_C$ when considering a Co substituted compound. It shows that the increase of $T_C$ is linear with the substitution rate and it reaches about 310 K for $x = 1$. The increase of $T_C$ is due to
the addition of new 3d electrons by the cobalt, which modifies the density of states of the material.

![Figure 2](image_url)

Figure 2. Curie temperature evolution of compounds for which Fe has been partially substituted by Co (a) [46], (b) [47], (c) [48], (d) [49], (e) [18], (f) [50], (g) [21], and (h) [51].

Figure 3 shows a passage from first-order toward second-order magnetic phase transition visible through the quick decrease of $\Delta S_{\text{max}}$ with $T_C$. An increase in $T_C$ of 100 K due to the Co substitution results in a decrease of $\Delta S_{\text{max}}^\text{M}$ of 13 J/kg·K. The decrease of $\Delta S_{\text{max}}^\text{M}$ is much less than in the case of the silicon content change. Therefore, it is preferable to use Co substitution to increase $T_C$. The decrease of $\Delta S_{\text{max}}$ due to a second-order transition is compensated by the widening of $\Delta S_M$. In this case, the Relative Cooling Power (RCP) appears to be unchanged. On the other hand, a more spread out transition in working temperature can be efficient for applications.

![Figure 3](image_url)

Figure 3. Relative magnetic entropy evolution as a function of $T_C$ for materials in which Fe iron has been substituted by Co, compared to the compound without Co. (a) [46], (b) [47], (c) [48], (d) [49], (e) [18], and (f) [50].

Paul-Boncour et al. [52] studied the influence of cobalt substitution for LaFe$_{11.5-x}$Co$_x$Si$_{1.5}$ compounds ($x \leq 2.2$), prepared by ball milling and one of either short annealing treatment or reactive Spark Plasma Sintering (SPS). Their magnetic properties have been investigated while using magnetic measurements, differential scanning calorimetry, and Mössbauer spectroscopy. Figure 4a shows the cell volume at 293 K plotted versus Co content. The annealed and sintered powder display close cell volumes for a given x Co content. We observe a critical range with a cell volume jump, which can be related to the change of
magnetic order, depending on whether samples are above or below $T_C$, in the ferromagnetic state as the cell volume decreases versus Co content.

The Curie temperature was estimated either from the maxima of $dM/dT$ or $dV/dT$ derivative curves or by the maxima of the DSC peaks. Figure 4b shows the good agreement obtained between these different methods. $T_C$ increases linearly versus $x_{\text{Co}}$ content according to the following relation: $T_C = 226 + 97 \times x_{\text{Co}}$.

$\Delta S_{\text{M}}^{\text{max}}$ curves versus temperature for LaFe$_{11.5-x}$Co$_x$Si$_{1.5}$ compounds with $\mu_0\Delta H$ of 1 T are presented in Figure 5. $T_C$ shift to larger temperature and a decrease of $\Delta S_{\text{M}}^{\text{max}}$ are clearly observed as the Co content increases.

The $57$Fe Mössbauer spectra that were collected at room temperature, for $x = 0.8, 1, 1.2$, and 2.2, have been refined with several ferromagnetic sextets for both inequivalent $8b$ and $96i$ iron sites. The obtained weighted average hyperfine field $<B_{\text{HF}}>$, together with $T_C$, versus $x$ Co content, are shown in Figure 6. As the Co content becomes larger, an increase of the mean Fe local moment is observed. It is worth noticing that the Co substitution increases both $T_C$ and the iron local moment.

Gebara et al. [53] have studied LaFe$_{11.8-x}$Co$_x$Si$_{1.2}$ alloys ($x = 0.52, 0.66, 0.8, 0.94, 1.08$) composites that were prepared by high energy ball milling showing the evolution of lattice constant, the Curie temperature, magnetic entropy change, and the evolution of the Fe Mössbauer spectra. Surprisingly, the magnetic entropy changes are not very sensitive to the Co content in their composites.

The substitution of Fe by the $Z$ elements ($Z = V, Cr, Ni, Cu$) in the La(Fe$_{1-x}Z_x$)$_{11.4}$Si$_{1.6}$ system can lead to a strong influence on the electron concentration on the magnetic and magnetocaloric properties [54]. This is due to the properties of itinerant metamagnets that are very sensitive to the $3d$ electron density at the Fermi level. For this motivation, Pathak et al. [54] have studied the possibility of substituting Fe by other $3d$ metals (V, Cr, Ni, Cu).
Figure 6. Evolution of the Curie temperature $T_C$ and the weighted average hyperfine field $<B_{HF}>$ versus $x$ Co content [52].

For $Z = V$ and at low $Z$ substitution rates, in $\text{La(Fe}_{1-x}Z_x)_{11.4}\text{Si}_{1.6}$, a loss of the magnetocaloric effect and an increase in $T_C$ are observed; $T_C = 211$ K for $x = 0$ increases up to 223 K for $x = 0.01$, and $\Delta S_m^{\text{max}}$ decreases from 15 to 8 J/kg.K. On the contrary, a substitution of Fe by Mn in $\text{La(Fe}_{1-x}\text{Mn}_x)_{11.7}\text{Si}_{1.3}$ [55] shows a decrease in $T_C$, as well as a decrease of $\Delta S_m^{\text{max}}$.

5. Effect of La Substitution in $\text{La}_{1-z}R_z(\text{Fe}_{0.88}\text{Si}_{0.12})_{13}$

The substitution of La by another rare earth atom influences the structural and magnetic properties of the material. Anh et al. [56] performed a systematic study on the effect of Nd substitution on magnetocaloric effect in $\text{La}_{1-z}\text{Nd}_z\text{Fe}_{11.4}\text{Si}_{1.6}$ ($z = 0 - 0.4$). They found a decrease of the unit cell parameter and $\Delta S_m^{\text{max}}$ with an increasing Nd content.

Fujita et al. [57,58] and Fujieda et al. [59,60] studied the effect of partial substitution of La in $\text{La(Fe}_{0.88}\text{Si}_{0.12})_{13}$ compound. Figure 7 (Left) shows that the unit cell parameter and $T_C$ linearly decrease with Ce/Pr content; these results were mainly attributed to the magnetovolume effect [59,60].

In addition, $\Delta S_m^{\text{max}}$ increases with the substitution rate (Figure 8). We can see that the effect is more contrasted with Ce atoms, which can be tetravalent. In an alloy, its average valence is therefore greater than three and its average ionic radius is smaller than for other rare earths.
Figure 8. $\Delta S_{\text{M}}^{\text{max}}$ vs. R content $z$, for $\text{La}_{1-z}\text{R}z\text{Fe}_{11.5}\text{Si}_{1.5}$ (R = Nd, Pr) [61,62].

The effect of substituting La by another rare earth atom on the unit cell parameter, $T_C$, and MCE was also studied by Shen et al. [61,62] for (R = Ce, Pr, Nd) and by P. Gebara and J. Kovac [63,64] for (R = Dy, Ho).

The maximum value of $\Delta S_{\text{M}}^{\text{max}}$ was found to increase from 23.7 J/(kg K) for $z = 0$ to 32.0 J/(kg K) for $z = 0.3$ in $\text{La}_{1-z}\text{Nd}z\text{Fe}_{11.5}\text{Si}_{1.5}$, for a field change $\mu_0\Delta H$ of 0-5 T, but this comes with a change of the hysteresis loss from 21.2 J/kg to 77.5 J/kg. Moreover, the relative cooling power (RCP) is enhanced by the partial substitution of Nd [61,62]. On the other hand, the substitution of La by Ho or Dy causes a decrease of magnetic entropy change [63,64].

6. Effect of Light Element Insertion in La(FeSi)$_{13}X$

6.1. Hydrogen Insertion

Fujieda et al. [65] and Fujita et al. [35] have shown that the insertion of hydrogen atoms (H) into $\text{La(Fe}_{0.88}\text{Si}_{0.12})_{13}$ significantly increases the Curie temperature, which increases linearly with the hydrogen content while maintaining the first order character of the transition. They have controlled the hydrogen concentration by changing both the annealing temperature and the hydrogen gas pressure. Fujieda et al. [65] adjusted $T_C$ to around room temperature (278 K) by hydrogen absorption. Fujita et al. [35] showed that, due to the increase of $T_0$, temperature at which the IEM transition disappears, this phenomenon appears above the Curie temperature, Figure 9 (Right). The disappearance of the IEM transition reflects the suppression of the renormalization effect due to the cell volume expansion of the $\text{La(Fe}_{0.88}\text{Si}_{0.12})_{13}$H$_y$ compounds through magnetovolume effects.

Figure 9. (Left) Unit cell parameter $a$ vs. hydrogen content [66,67]. (Right) Magnetic phase diagram of $\text{LaFe}_{11.44}\text{Si}_{1.5}\text{H}_y$. The IEM transition occurs between $T_0$ (the temperature at which the IEM transition disappears) and $T_C$ [35].
Phejar et al. [68] found that the unit cell parameter $a$ increases linearly with hydrogen content, independent of the composition of the parent compound; this result is in agreement with previous results [66,67] (Figure 9-Left). In order to follow the evolution of the crystallographic and magnetic parameters as a function of the temperature, they have performed NPD experiments between 1.5 and 300 K. In order to have a smaller incoherent background and, therefore, a better signal/noise ratio, the hydrogen is replaced by deuterium (D). The La(Fe$_{0.88}$Si$_{0.12}$)$_{13}$D$_{0.7}$ compound exhibited a magnetostrictive effect around $T_C$, resulting in a contraction of the cell parameter at the transition, which is the result of the IEM transition, as shown in Figure 10 (left). The cell parameter of the $\alpha$-(Fe,Si) inclusions increases, as expected, from a thermal expansion behavior without anomaly at $T_C$. A different behavior has been found by Gebara et al. [69], where an anomalous behavior of $\alpha$-Fe impurity cell parameter was observed at $T_C$ in La(Fe$_{13}$Si)$_{13}$ compounds with La substituted by Ce, Pr, Ho, or Fe by Mn. This means that micro-strains are negligible in the deuteride. Figure 10 (right) shows the magnetic moments of Fe$_{96i}$ and Fe$_{8b}$ atoms in the LaFe$_{11.5}$Si$_{1.5}$D$_{0.7}$ compound.

Neutron powder diffraction (NDP) experiments have been performed to solve the structure and localize the interstitial site in order to remove ambiguity concerning the interstitial occupied by hydrogen (deuterium) or carbon (24$d$ or 48$f$ site) [68]. Paul-Boncour et al. [68,70] have refined the NDP patterns while using the Rietveld method and have found that the interstitial D(H) was located on the 48$f$ site (Figure 11); this result is in perfect agreement with previous work [67].

Mandal et al. [71] showed that they have succeeded in tuning the Curie temperature from 199 to 346 K, with the hydrogen content without changing significantly $\Delta S^\text{max}_M$ value. Figure 12 (Left) shows the evolution of the Curie temperature for materials in which hydrogen has been inserted as a function of the insertion rate $y_H$ taken from different works [35,68,71,72]. Figure 12 (Right) shows the evolution of $\Delta S^\text{max}_M$ versus the insertion rate $y_H$ for LaFe$_{13-x}$Si$_x$H$_y$ [35,68,71]. According to this figure, it can be noticed that, for the Si content 1.3 $\leq x \leq$ 1.6, we obtain a first-order magnetic transition, which results in a larger $\Delta S^\text{max}_M$ than for the compositions $x \geq$ 1.8, which represent a second-order magnetic transition. Moreover, for $x = 1.4$ and 1.6, $\Delta S^\text{max}_M$ is almost constant for the different hydrogen insertion rates $y_H$. 

![Figure 10. (Left)](image_url1)

![Figure 10. (Right)](image_url2)
6.2. Carbon Insertion

Another possibility for increasing unit cell parameters and, thus, the Curie temperature of the material by magneto-volumic effect is the insertion of carbon atoms into the interstitial sites of the structure. This carbon insertion induces a decrease in $\Delta S_M$. The insertion of carbon in the material changes the electronic state and, thus, the magnetic properties of the material. The decrease of $\Delta S_M$ as a function of $T_C$ is of the same order as that obtained by the substitution of iron by cobalt: a decrease in entropy of 19 J/kg.K for an increase in $T_C$ of 100 K. Thus, it seems that, in this case, the transition also goes from first order to second order.

Li et al. [73] studied the insertion of carbon influences the phase formation, $T_C$, and magnetic entropy change of LaFe$_{11.7}$Si$_{1.3}$. The unit cell parameter $a$ increases with C content, $T_C$ increases from 194 to 225 K, and $\Delta S_M$ was equal to 27.5 J/kg K$^{-1}$.Phejar et al. [68] studied the effect of the carbonation of LaFe$_{13-x}$Si$_x$ compounds. LaFe$_{11.5}$Si$_{1.5}$C$_{0.7}$ compound displays a magnetostrictive effect around $T_C$ and a classical thermal expansion for the $\alpha$-(Fe,Si) impurities as for the deuteride, as shown in

Figure 11. The unit cell of LaFe$_{11.5}$Si$_{1.5}$D$_{0.7}$ projected along the $(a, c)$ plane.

Figure 12. (Left) Evolution of the Curie temperature for materials in which hydrogen has been inserted vs. the insertion rate $y_H$ (a) [72], (b) [35], (c) [71], (d) [68], and (e) [70]. (Right) $\Delta S_M^{\text{max}}$ vs. the insertion rate $y_H$ for LaFe$_{13-x}$Si$_x$H$_y$ (a) [35], (b) [71], and (c) [68].
Figure 13 (left). Figure 13 (right) shows the evolution of the magnetic moments versus the temperature. The magnetic moment decrease is not as abrupt when compared to the hydrides, which is due to a weakening of the first order transition.

Figure 13. Unit cell parameter $a$ vs. temperature for LaFe$_{11.5}$Si$_{1.5}$C$_{0.7}$ (left). Magnetic moments of Fe atoms vs. temperature, in the 96$i$ and 8$b$ crystallographic sites (right). The results are obtained from neutron diffraction [68].

7. Combination of Substitution and Light Element Insertion

Hydrogen insertion in La(Fe$_{1-x}$Si$_x$)$_{13}$ compounds allow for maintaining a large magnetic entropy variation with $T_C$ at room temperature (RT). However, several studies have shown that, for H composition lower than 1.5 H/f.u. and with $T_C$ close to room temperature, there is a splitting into two hydride phases with low and high H concentrations after few hours or weeks at RT [55,74,75]. The sample is no more homogeneous and the magnetic entropy variation is split in two peaks. In order to avoid this segregation, it is safer to prepare saturated hydrides, but their Curie temperature is around 330–340 K. This temperature can be reduced to RT by partial Mn for Fe substitution for a fully hydrogenated hydride. In this case, the hydride remains stable without desorption or segregation versus time or upon magnetic cycles, despite $T_C$ being near RT. A systematic study on the influence of Mn for Fe substitution [76] has revealed that it is possible to vary $T_C$ between 260 and 350 K ($0.1 \leq x_{Mn} \leq 0.4$ Mn/f.u.; Si concentrations: 1.2 and 1.4 Si/f.u. and $1.77 \leq y_H \leq 1.97$ H/f.u.). The drawback is a significant reduction of the $\Delta S_M$ and $\Delta T_{ad}$ values as the Mn content increases significantly [77], but this Mn substitution remains interesting to vary $T_C$ in a narrow temperature range, as, in this case, the reduction of $\Delta S_M$ is limited [78].

Another alternative is to partially substitute La by Ce [79], as, beside the reduction of $T_C$, there is an increase of the MCE effect versus the rate of Ce for La substitution. However, it was found that, in this case, the hydrogenation significantly reduces the MCE effect as compared to that of the parent alloys, which is less beneficial than expected. The influence of Ce substitution for carbides was also investigated, but, despite the larger magneto-volume effect of C versus H insertion, the reduction of $\Delta S_M$ was more pronounced [50]. Hai et al. [80,81] studied the influence of both C and H insertion on Ce substituted alloys on the hydrogenation kinetic by in-situ neutron diffraction. The preferential insertion of C in 24 $d$ sites introduces a cell distortion and modifies some specific Fe-Fe distances. The presence of C is found to influence the H insertion and to slow down the hydrogenation kinetic. The presence of Ce substituted to La also modifies the kinetic of reaction due to its smaller atomic radius.

Further studies have been undertaken to combine both Ce for La and Mn for Fe substitution with hydrogen insertion [27,82,83]. An adjustment of the rate of both Ce and Mn in La$_{0.9}$Ce$_{0.1}$Fe$_{11.7-x}$Mn$_x$Si$_{1.3}$ hydrides allows for maintaining a significant MCE effect near RT. The influence of the bonding with epoxy [27,82] was found to improve the
mechanical properties and slightly reduce the MCE effect. High hydrogen pressure sintering on La$_{0.9}$Ce$_{0.1}$Fe$_{11.7-x}$Mn$_x$Si$_{1.3}$ hydrides [83] showed a beneficial effect for a pressure of 40 MPa. However, an applied pressure of 50 MPa induced the precipitation of $\alpha$-Fe reducing the MCE effect.

Beside these works, the influence of Co insertion on the magnetocaloric properties of hydrides has been investigated [70]. The aim was not to maintain $T_C$ near RT, but rather to increase it for heat pump applications above room temperature. $T_C$ is found to increase linearly versus Co content and H concentration for LaFe$_{10.5-x}$Co$_x$Si$_{1.5}$H$_y$ compounds (Figure 14 (Left)), but the $dT_C/dy$ slope changes versus Co concentration. $T_C$ can be expressed as a function of $x$ and $y$ while taking a negative mixing term into account:

$$T_C(x,y) = 210 + 100 \cdot x + 83 \cdot y - 18 \cdot x \cdot y$$

For $x = 1$ and $y = 1.5$, $T_C = 400$ K, whereas it would have been 450 K without the negative exchange term.

The Co substitution induces a reduction of the magnetic entropy variation, whereas it remain almost constant before and after hydrogenation, as shown for $x_{Co} = 0.8$ and $y = 1.4$ (Figure 14 (Right)).

8. Implementation in Active Magnetic Regenerators

The optimization of the chemical composition by the substitution or/and light element insertion in La(Fe,Si)$_{13}$ compounds has been reviewed in previous part. But magnetic refrigeration requires also the use of an active magnetic regenerator (AMR) to increase the temperature span ($\Delta T_{\text{span}}$) by generating a temperature gradient between the hot and cold sources [84]. Many prototypes have been tested while using Gd or its alloys as reference refrigerants in AMR [85]. Although Gd offer many advantages (stability, easy to machine, large $\Delta T_{\text{ad}}$, the absence of thermal hysteresis, large thermal conductivity), its elevated cost and limited global reserve constitutes a serious drawback for a broad public application. Therefore, it is necessary to test and optimize other promising magnetocaloric material, such as La(Fe,Co,Si)$_{13}$ in AMR. The performances of different regenerators using Gd and La(Fe,Co,Si)$_{13}$ material have been compared and discussed in [86–89]. In all of these prototypes, the AMR were fixed and the magnetic field mobile (rotation of permanent magnets arranged in Halbach cylinder or reciprocating movement). One of the best geometries to favor heat exchange is thin parallel plates (0.25–1 mm) that are separated by a small gap [87,90]. Compared to Gd, which presents broad $\Delta S_M$ and $\Delta T_{\text{ad}}$ peaks, La(Fe,Co,Si)$_{13}$ compounds display higher but narrower peaks due to their first order
transition. In addition, due to its smaller specific heat, Gd has a two time larger $\Delta T_{ad}$ than La(Fe,Co,Si)$_{13}$ for a field variation around 1 T. To expand the $\Delta T_{\text{span}}$, the use of a composite or several plates of La(Fe,Co,Si)$_{13}$ compounds with different $T_C$, obtained by varying the Co content, have been therefore tested. Thin plates prepared by TDR method and purchased by Vacuumshmelze GmbH and Co were used in several prototypes [86–88]. Other tests were performed with an epoxy-bonded plates [89]. Better performances were obtained while using regenerators with at least four different compositions that were arranged to optimize the thermal gradient, as compared to plates with only one or two different compositions.

Balli et al. [86] built a prototype with two AMRs that are alternatively magnetized and demagnetized in order to reduce the magnetic forces acting on the magnetic refrigerant. The applied magnetic field is parallel to the plates to limit the demagnetization field which is corrected. Working with two AMR, one with Gd and the other with a La(Fe,Co,Si)$_{13}$ composite, they obtained a comparable maximum $\Delta T_{\text{span}}$ of 14 and 16 K for LaFeSi-based and Gd AMR respectively. They have tested the corrosion behavior of La-Fe-Co-Si compounds in different solutions of water with an additional corrosion inhibitor and a silicon oil. The best resistance to corrosion was obtained with 3% of noxal anti-oxidant. Because water has a larger specific heat ($C_p = 4.2$ J/g K) than Si oil ($C_p = 1.6$ J/g K), it is more favorable as fluid heat transfer. The study of the corrosion of Gd and La(Fe,Co,Si)$_{13}$ in various heat conducting fluids is detailed in Forchelet et al. [91].

Legait et al. [88] compared the performance of Gd, a manganite oxide (Pr$_{0.62}$Sr$_{0.35}$MnO$_3$) and La(Fe,Co)$_{13-x}$Si$_x$ compounds under a wide range of fluidic and magnetic operating conditions. Their devices do not use a thermal exchanger, therefore it only provides non-load temperature spans. They have followed the influence of non-dimensional numbers (utilization ratio $U$ and volume ratio $V^*$) and physical properties, such as thermal conductivity. They observed that each regenerator is most efficient over a specific domain of utilization. A material with low thermal conductivity (manganese) is more efficient at low frequency, whereas the materials with large thermal conductivity are better at a high frequency.

Tusek et al. [87] have tested their AMR with a device that is equipped with a reciprocating magnet and a heat exchanger to investigate the cooling load under different $\Delta T_{\text{span}}$. They have observed that the maximum $\Delta T_{\text{span}}$ varies versus the utilization factor $U$ and the operating frequency $P$. It is larger for LaFeSi-based AMR when compared to Gd at low $U$ values (moderate $\Delta T_{\text{span}}$), whereas that of Gd increases for larger $U$ values. The maximum $\Delta T_{\text{span}}$ also depends on the hot side temperature and it is maximum at 35°C for Gd, 27°C for a two-layered, and 43°C for a seven-layered La(Fe,Co,Si)$_{13}$ AMR.

AMR was also prepared with epoxy-bonded plates that were constituted of a composite of La(Fe,Co)$_{13-x}$Si$_x$ powder in a polymer matrix pressed into thin plates [89]. This AMR yields smaller $\Delta T_{\text{span}}$ than with sintered plates, but it has very good mechanical properties, as it remains stable without significant changes over 90,000 cycles. Because the AMR was constituted with two-layered plates (two different $T_C$), further improvements can be expected by preparing a AMR with four-layered plates. Beside these experimental results, numerical investigations have been performed through a two-dimensional (2D) model of an AMR refrigerator for various magnetocaloric materials [92,93]. The best performance in terms of temperature spans and coefficient of performance (COP) were obtained for Gd, Gd$_2$Si$_2$Ge$_2$, La(Fe,Mn,Si)$_{13}$H$_y$, and La(Fe,Co,Si)$_{13}$ compounds as compared to MnFe(P,As) and manganite oxide.

The thickness of the plates as well as the spacing between the plates, the $T_C$ distribution, the intensity of the magnetic field, and the operating frequencies are among the important parameters for optimizing in order to obtain the best cooling performances. Therefore, the properties of AMR with La(Fe,Co,Si)$_{13}$ material should be optimized to reach as good AMR performances as Gd. These materials remain very promising for magnetic refrigeration near room temperature application due to their many advantages in term of cost and global reserve.
9. Conclusions

In this review, we have presented the main works dealing with the magnetocaloric effect of LaFe\(_{13-x}\)Si\(_x\). The different methods of synthesis and shaping were presented and compared. The influence of both chemical substitution of La, Fe, and Si by other elements, as well as the insertion of light elements (H, C), were presented to show the advantages and disadvantages of each solutions. Fe for Co substitution or H insertion are considered to be the most efficient way to use these materials as refrigerant near room temperature. A combination of metal substitution and light element insertion is another alternative to adjust their working conditions. La(Fe,Si)\(_{13}\) type compounds have been implemented in active magnetic regenerators and their performance in magnetic refrigerator prototype when compared with that of Gd.

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**Abbreviations**

| Abbreviation | Full Form |
|--------------|-----------|
| AMR          | Active Magnetic Regenerator |
| COP          | Coefficient of Performance |
| DSC          | Differential Scanning Calorimetry |
| FM           | Ferromagnetic |
| IEM          | Itinerant Electron Metamagnetic |
| MCE          | Magnetocaloric effect |
| NPD          | Neutron Powder Diffraction |
| PM           | Paramagnetic |
| RT           | Room Temperature |
| SPS          | Spark Plasma Sintering |
| \(T_C\)      | Curie Temperature |
| TDR          | Thermal Decomposition and Recombination |
| RCP          | Relative Cooling Power |
| ZFC-FC       | Zero Field Cooled - Field Cooled |

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