Full length article

Design and synthesis of a magnesium alloy for room temperature hydrogen storage

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A B S T R A C T

Safe and high-density storage of hydrogen is a key issue for development of hydrogen as a clean energy carrier. Hydride-forming materials are potential candidates for safe and high-density storage of hydrogen and among them, Mg-based alloys are the most investigated ones due to their high gravimetric capacity. The key issue for practical application of Mg-based alloys is that they do not desorb hydrogen without heating due to the strong hydrogen binding energy. In this study, we employed first-principles calculations to design a Mg-based alloy with a low hydrogen binding energy and room temperature hydrogen storage properties. The designated material, highly-homogenous Mg₄NiPd with a BCC-based CsCl-type structure, was successfully synthesized by severe plastic deformation via the high-pressure torsion (HPT) method. The alloy exhibited reversible hydrogenation and dehydrogenation at room temperature with high phase stability. This discovery introduces a rational approach to design and synthesize new alloys for hydrogen storage using the concept of binding energy engineering.

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1. Introduction

Hydrogen is currently considered to be an important future energy carrier [1]. Despite significant interest in hydrogen as a clean fuel with zero CO₂ emissions, there are still considerable issues that need to be addressed for the future application of hydrogen as a fuel. Among these issues, the design of new technology to store hydrogen safely, with high volumetric and gravimetric density and without using a heating system is a major challenging task [1]. Since hydrogen is the lightest known element, its storage in the form of gas under ambient pressure is impractical and requires a very large volume per kilogram. Hydrogen storage in the form of compressed gas, usually at several hundred bars, is being commercialized for automobiles, but compressed hydrogen gas requires special and expensive tanks to ensure hydrogen is stored safely [1]. Hydrogen storage in the form of liquid is being used in aerospace industry, but large energy levels are required for liquefaction and the hydrogen loss through evaporation is another issue [1].

One promising technique for the high-density and safe storage of hydrogen is the use of different hydrides such as Mg-based, Ti-based, La-based or complex hydrides [2,3]. Mg-based hydrides, which are known as the first introduced hydrogen storage candidate materials [4], are the most investigated metal hydrides due to their large volumetric and gravimetric storage capacity [2]. Despite high-density storage capacity of Mg-based hydrides, their main drawback is still their high thermodynamic stability due to the

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strong metal-hydrogen binding energy and thus their high dehy-
rogenation temperature (maximum ~630 K for pure MgH2 and
minimum ~460 K for Mg2NiH4, while the desirable tempera-
tures are below 373 K and ideally room temperatures) [2]. Despite
this apparent thermodynamic barrier to reduce the dehydrogenation
temperature of Mg-based materials to room temperature [5], it was
shown that the Mg-H binding can be broken at room temperature in
a few compounds based on rare-earth elements and/or Ca with the
Mg addition [6–9]. Refs. [6–9] suggest that room temperature
hydrogen storage is still achievable in Mg-based alloys, provided
that the alloys with low hydrogen binding energy can be designed
and synthesized successfully.

The main strategy currently used to reduce the dehydrogenation
temperature of hydrogen storage materials is to mix elements with
two different hydrogen affinities, specifically: an element that
readily reacts with hydrogen, such as Mg, Ti or La (component A)
with negative binding energies, and an element that has no af-

finity to react with hydrogen, such as Ni or Fe (component B) with pos-
itive binding energies [1]. This strategy has been successfully
employed to fabricate binary Ti-based (AB-type TiFe) [10] and La-
based (AB5-type LaNi5) compounds [11] that can reversibly store
hydrogen at room temperature. In the case of the Mg-based binary
systems, however, they can only function above ~460 K, with A2B-
type Mg2Ni exhibiting the lowest temperature [4,5].

In a previous study, we calculated the hydrogen binding energy
in different binary Mg-based systems from first-principles and
compared the results with experimentally measured dehydroge-
nation temperatures [12]. We found that intermetallic materials,
such as Mg2Ni and Mg2Co with dehydrogenation temperatures
lower than pure Mg, reduce the hydrogen binding energy to less
negative values. However, some other intermetallics, such as
Mg2Pd, Mg2Sn, Mg2Si, Mg2Ge, Mg2In and MgAg have positive
hydrogen binding energies and thus cannot absorb hydrogen. The
target hydrogen binding energy for room-temperature hydrogen
storage is a negative value close to zero, but none of the current Mg-
based binary systems meet this target.

As briefly summarized in Fig. 1(a), to engineer new three-
component (ternary) Mg-based compounds with a low hydrogen
binding energy close to zero, our strategy is mixing two different
types of binary compounds at the atomic scale: (A') a compound
with a negative hydrogen binding energy (such as Mg2Ni) and (B') a
compound with a positive hydrogen binding energy (such as
Mg2Pd). Although many different composition can be fabricated by
mixing the compounds with negative and positive hydrogen
binding energies, we selected Mg2NiPd (Mg2Ni + Mg2Pd) among
many other possible choices because of well-know catalytic effect
of Ni and Pd elements on hydrogen storage in Mg-based alloys [2],
which minimizes the effect of kinetic issues on the experimental
results.

Although the above-mentioned strategy seems feasible from a
theoretical perspective, there are some technical limitations to
mixing these compounds homogeneously and achieve a uniform
atomic-scale elemental distribution (i.e. a homogenous single
phase) because of the thermodynamic immiscibility of Mg in many
different systems [13]. For example, if Mg2NiPd is fabricated by
slow solidification, three phases (Mg2Ni and Mg2Pd) with
heterogeneous elemental distributions are formed, as reported in
the ternary phase diagram of Mg-Ni-Pd [14,15]. One effective so-
lution for atomic-scale mixing of Mg with other elements at the
atomic scale is the application of severe plastic deformation (SPD)
using high-pressure torsion (HPT) (see the following references for
more information on the principles of SPD [16,17] and the details
and background of HPT [18,19]). As a brief description, in the HPT
method as schematically shown in Fig. 1(b), a disc sample is
compressed between two anvils under high pressure and concurrently
subjected to severe strain by rotating one of the anvils with respect to
the other anvil for N turns. Earlier studies showed that the HPT
method is quite effective for synthesizing new Mg-based com-
 pounds [20,21], even in immiscible systems. HPT has also been used
to activate hydrogen storage materials [22,23] and to enhance the
hydrogenation kinetics in pure Mg [24,25], Mg-based composites
[26,27], and Mg alloys [28,29].

The goal of this study is to propose a strategy of designing and
synthesizing new compounds with promising functionality, i.e. a
new Mg-based alloy that can reversibly store hydrogen at room
temperature. Here, we report the successful design and fabrication
of a Mg2NiPd alloy with the BCC-based CsCl-type structure, which
is thermally stable up to 440 K and reversibly store 0.7 wt% of
hydrogen at room temperature. Despite low hydrogen storage ca-
cacity of the alloy, this study opens up a rational approach to design
and synthesize new materials beyond the scope of known equi-
librium phase diagrams for promising functions such as room
temperature hydrogen storage.

2. Materials and methods

2.1. Materials processing

Small pieces of high-purity (>99%) Mg, Ni and Pd with a nominal
composition of Mg2NiPd were melted under an argon atmosphere
and cast into a metallic mold to fabricate a rod with a 12-mm
diameter and a 16-mm length. The as-cast material was subjected
to SPD using the HPT process [16–19] to achieve a single phase with
an atomic-scale elemental distribution. To do this, the rod was first
sliced into discs 0.8 mm thick and 10 mm diameter, with each disc
being further processed by HPT under a pressure of P = 6 GPa at
room temperature for either N = 20, 100, 500 or 1500 turns with a
rotation speed of *ω = 1 rpm.

2.2. Materials characterization

Following the HPT processing, the discs were polished by me-
canical grinding and buffing to mirror-like surfaces, and the
 elemental distribution was examined at the mid-radius of the discs
by a scanning electron microscope (SEM) equipped with an energy
dispersive X-ray spectroscopy (EDS) using an accelerating voltage
of 15 kV.

The elemental mixing and the formation of new phases were
studied using X-ray diffraction (XRD) analysis with Cu Kα radiation
produced using a current of 40 mA and an accelerating voltage
of 40 kV. The XRD patterns were examined in detail using the Rietveld
method in the FullProf Suite software [30,31].

To examine the final microstructure of the alloys after 1500 HPT
turns, a thin foil was prepared 4 mm away from the disc center
using a focused ion beam (FIB) system. It should be noted that after
such large numbers of HPT turns, microstructure and properties do
not vary radially in the samples [32–34]. The foil was examined with
transmission electron microscopy (TEM) in bright-field (BF),
dark-field (DF), selected-area electron diffraction (SAED) and high-
resolution modes using an accelerating voltage of 200 or 300 kV.

The spatial distribution of the three elements was then char-
acterized after 1500 HPT turns at the atomic scale using atom probe
tomography (APT). Samples were prepared using a FIB system using
the lift-out method at a distance of 4 mm from the disc center. APT
analysis was carried out using a LEAP 4000HR instrument at a
temperature of 40 K with electric pulses with a 20% pulse fraction,
and a 200 kHz pulse repetition rate. Three dimensional re-
constructions were generated using the Integrated Visualization &
Analysis Software (IVAS) and other data processing was performed
with the GPM 3D-soft (Groupe de Physique des Matériaux 3D-soft).
2.3. Hydrogen storage

The hydrogen storage performance of the samples was examined in a Sieverts-type gas absorption apparatus. One disc sample processed using HPT for 1500 turns was broken to pieces with dimensions smaller than 0.1 mm under an argon atmosphere and subjected to evacuation at room temperature for 2 h to remove possible moistures. The pressure-composition isotherm (PCI) measurements were performed at 305 K up to a hydrogen pressure of 10 MPa for several hydrogenation/dehydrogenation cycles. After the PCI measurements were taken, each sample was examined by XRD analysis.

Finally, hydrogenation behavior of the HPT-synthesized phase and its thermal stability were examined by in situ XRD analysis using synchrotron radiation with a wavelength of 0.07149 nm either under an argon atmosphere during heating, or at room temperature under a hydrogen atmosphere at a pressure of 1.1 MPa (the maximum achievable hydrogen pressure in the facility) at SNBL, ERSF, Grenoble, France.

2.4. Calculation methods

The crystal structure of the Mg$_4$NiPd and its hydrogen binding energy were examined theoretically from first-principles. Since a BCC-based crystal structure was experimentally detected in different Mg-based alloys such as Mg-Ni \[35\], Mg$_{50}$Ni$_{45}$Pd$_5$ \[36\], Mg-Ni-V \[20\], Mg-Pd-V \[20\], Mg-Ti \[37\], Mg-Zr \[21\] and Mg-V \[20\] after SPD processing by ball milling \[35,36\] or HPT \[20,21,37\], the initial structure was modeled by a 54-atom supercell constructed from $3 \times 3 \times 3$ BCC unit cells. A random configuration of the constituent elements was simulated using a special quasi-random structure (SQS) \[38-40\], in which the center sites were occupied by Mg atoms and the corner sites were randomly occupied by either Mg, Ni or Pd (CsCl-type based on our preliminary experimental data). The SQS was obtained by application of the CLUPAN code \[41,42\]. The calculations were conducted with the plane-wave basis projector augmented wave (PAW) method \[43\] in the framework of density functional theory (DFT) within the generalized gradient approximation (GGA) of the Perdew-Burke-Ernzerhof (PBE) form \[44\], as implemented in the VASP (Vienna Ab initio Simulation Package) code \[45-47\]. A plane-wave energy cutoff of 350 eV was used. The Brillouin zone was sampled with a $\Gamma$-centered $15 \times 15 \times 15$ k-point mesh per BCC unit cell using the Methfessel-Paxton scheme \[48\] with a smearing width of 0.4 eV. The crystal structure was optimized until the energy convergences and the residual forces decreased to less than $10^{-8}$ eV and $1 \times 10^{-3}$ eV/Å, respectively. The optimized crystal structure was visualized three dimensionally by employing the VESTA code \[49\].

![Fig. 1](image-url). (a) Schematic illustration of the hydrogen binding energy effects on the hydrogen storage behavior of materials in the Mg-Ni-Pd system; and (b) schematic illustration of high-pressure torsion (HPT) method. Data for the hydrogen binding energy and dehydrogenation temperatures ($T_{Deh}$) in (a) were taken from Emami et al. \[12\].
Phonon calculations were performed using the PHONOPY code [50] in order to examine the dynamic stability of the optimized crystal structure. Force constants were obtained for the 54-atom supercell model using an atomic displacement of 1 p.m. Phonon frequencies at reciprocal-space points were calculated using Γ-centered $64 \times 64 \times 64$ meshes, and the phonon density of states was calculated using the tetrahedron method [51,52].

Hydrogen binding energies were calculated for the three primary types of octahedral sites, namely, $4\text{Mg-2Ni}$, $4\text{Mg-1Ni-1Pd}$ and $4\text{Mg-2Pd}$, since these could be found the most frequently in the supercell model. Hydrogen atoms were located in the selected sites and the structures were optimized in terms of their lattice shape and atomic positions. The theoretical hydrogen binding energies were calculated using the following equation, as described previously [12]:

$$
\Delta E = E(\text{Mg}_4\text{NiPdH}) - \left[ E(\text{Mg}_4\text{NiPd}) + \frac{1}{2} E(\text{H}_2) \right]
$$

In this equation, $\Delta E$ is the hydrogen binding energy and $E$ denotes the internal energies at 0 K.

3. Results and discussion

3.1. Experimental evidence for atomic-scale mixing and new phase formation

In order to manipulate the hydrogen binding energy, it is essential to generate an atomic-scale uniform distribution of elements and form a single phase. To do this, the as-cast material was processed by HPT for 20, 100, 500 and 1500 turns and the distribution of elements was examined by SEM-EDS, STEM and APT. Moreover, the formation of new single phase and its structure were examined by XRD and TEM.

To assess the distribution of the elements in the alloy, we analyzed the mid-radius of the discs by SEM-EDS. Fig. 2(a) shows the SEM-EDS analysis of samples. The as-cast sample ($N = 0$) contained three intermetallic phases with the compositions $\text{Mg}_8\text{Ni}_3\text{Pd}$, $\text{MgNi}_2$ and $\text{Mg}_5\text{Pd}_2$, as measured by EDS analysis quantitatively. The formation of these three intermetallics is not consistent with the reported ternary phase diagrams of Mg-Ni-Pd, in which the $\text{Mg}_4\text{NiPd}$ composition should contain Mg, $\text{Mg}_2\text{Ni}$ and $\text{Mg}_6\text{Pd}$ [14,15]. The reason for this contradiction is that the ingot was casted in a metallic mold which resulted in rapid cooling rate and formation of three metastable phases. The three intermetallic phases are well distinguished after the HPT processing for 20 and 100 turns, but their sizes become considerably smaller, as reported for many other HPT-processed multi-phase systems [20,32]. The three phases can barely be distinguished after 500, and after 1500 turns, the elemental distribution becomes homogeneous within the detection limits of SEM-EDS.

Since the SEM-EDS method with low detection limit can not show the atomic-scale homogeneity, the distribution of the three elements at the atomic scale was examined by APT analyses. The APT mappings in Fig. 2(b) show that the distribution of the three elements at the atomic level occurred after 1500 turns. The uniform atomic-scale distribution of the three elements is more apparent in the line profiles of Fig. 2(c) which show that the compositional fluctuations are ±7 at.% for Mg and ±4 at.% for Ni and Pd.
Importantly, two other volumes analyzed by APT exhibited similar features with reasonable atomic-scale elemental homogeneity (see the supporting video), which is essential to tune the atomic sites for a low hydrogen binding energy [12]. Supplementary video related to this article can be found at https://doi.org/10.1016/j.actamat.2018.02.033.

To assess the formation of a new single phase, we analyzed the materials by XRD analyses. The XRD profiles in Fig. 3(a) shows that the peaks for the three initial intermetallics (Mg₈Ni₃Pd, MgNi₂ and Mg₅Pd₂) are clearly visible for the ingot (N = 0), which is in good agreement with the SEM-EDS analysis shown in Fig. 2(a). After 20 and 100 turns, the intermetallic peaks for the three species are still visible, but with considerable peak-broadening. Such peak broadenings confirm that large number of lattice defects, including dislocations [33] and grain boundaries [34], is formed after the HPT processing. It was suggested previously that the formation of these lattice defects is one main mechanism for the atomic-scale elemental distribution and formation of new phases after SPD processing [53–55]. The peak intensity for the three intermetallics greatly decreases after 500 turns, and a new single phase with a BCC-based partly ordered CsCl-type structure and a lattice parameter of a = 0.319 nm forms after 1500 turns, as shown in Fig. 3(b). The formation of this BCC-based phase after 1500 turns, which is likely due to the effect of lattice defects on the dynamic stability of phases [53–55], is in good agreement with earlier reports on the formation of BCC-based phases in SPD-processed Mg-based alloys [20,21,35–37], and is in line with the purpose of this study to synthesize a single-phase material with high atomic-scale elemental homogeneity.

To examine the thermal stability of the BCC-based phase, in situ XRD analyses were conducted using synchrotron radiation at different temperatures. The heating rate was 12 K/min and each pattern was taken in 20 s. As shown in Fig. 3(c), the BCC-based structure is incredibly stable up to 440 K suggesting the capability of the Mg₄NiPd phase for wide applications including hydrogen storage.

To examine the nanostructural features of the BCC-based phase, TEM observations were conducted. The BCC-based phase has a nanocrystalline morphology with an average grain size of 10 ± 4 nm, as shown in the TEM bright-field (a) and dark-field (b) images in Fig. 4. The perfect ring pattern of the SAED analysis with several well-defined rings, taken from a region with a diameter of 300 nm, also confirms the presence of nano-sized polycrystals within the selected area. Examination of the TEM lattice images and the corresponding fast Fourier transform (FFT) diffractograms confirm that the BCC-based phase has a distorted structure and the atomic planes are not in an ideal geometrical condition (Fig. 4(c)), but some less-distorted structures can be partly detected (Fig. 4(d)).

### 3.2. Low hydrogen binding energy and room-temperature hydrogen storage

To examine the formation and stability of the BCC-based structure, the first-principles calculations were conducted. Fig. 5(a) shows the initial structure modeled by SQS and Fig. 5(b) shows the optimized structure. The examination of optimized lattice structure in Fig. 5(b) confirms that the BCC-based CsCl-type phase with a distorted lattice can remain stable provided that the three elements are well-mixed at the atomic scale over the two crystallographic BCC sites. The lattice parameter can be estimated from the optimized volume per atom as a = 0.325 nm (Fig. 5(b)). The ~2% overestimation from the experimental value of a = 0.319 nm is within reasonable accuracy of the GGA-PBE exchange-correlation function capabilities [56], and which suggests good correlation between the first-principles calculations and the experimental data. The plot of phonon density of states versus frequency, as shown in Fig. 5(c), shows that the BCC-based structure does not exhibit imaginary phonon frequencies, which implies that this phase is dynamically stable [50].
To examine the thermodynamics of the BCC-based structure for hydrogen storage, the hydrogen binding energies were calculated in three types of the most abundant octahedral sites, namely, 4Mg-2Ni, 4Mg-1Ni-1Pd, and 4Mg-2Pd. Fig. 5(d) summarizes the theoretical hydrogen binding energies for the three main octahedral interstitial sites in the dynamically stable BCC-based structure. For the 4Mg-2Ni sites, the hydrogen binding energy is negative, and thus, hydrogen may not be desorbed from these sites. For the 4Mg-2Pd sites, the hydrogen binding energy is positive and therefore hydrogen cannot occupy these sites. However, for the 4Mg-1Ni-1Pd sites, with a composition that matches the Mg₄NiPd alloy, the hydrogen binding energy is close to the ideal binding energy for room-temperature hydrogen storage material (see Fig. 1). These calculations indicate that hydrogen can be absorbed and desorbed in certain octahedral sites in Mg₄NiPd at room temperature.

Fig. 6(a) shows the PCI results at room temperature (305 K) for the sample processed by HPT for 500 turns, after one and five hydrogenation/dehydrogenation cycles. The sample absorbed 0.6 wt% hydrogen with 0.5 wt% reversibility in the first cycle. However, the total absorption increases to 0.86 wt% (equivalent to Mg₄NiPdH₁₂.₂₅) after the fifth cycle, with an improved reversibility of 0.7 wt%. The current results on room temperature hydrogen storage introduce a practical approach to designing and synthesizing new hydrogen storage materials that can reversibly store hydrogen at room temperature.

To examine the stability after cyclic hydrogenation and dehydrogenation, XRD analysis was performed after five PCI cycles. Fig. 6(b) shows the XRD profiles for the sample after the PCI measurements. It is apparent that the BCC-based phase remains stable after at least five hydrogenation/dehydrogenation cycles. There is no indication that trapped hydrogen was present. This confirms that the hydrogen is desorbed at room temperature, in good agreement with the PCI results shown in Fig. 6(a).

A question naturally arises now: where are the hydrogen atoms or what hydride was formed from the Mg₄NiPd alloy? In the context of this study, it was hard to identify the place of hydrogen or the structure of the hydride by ex situ analysis because of the room-temperature hydrogen storage reversibility and disappearance of the hydride phase during the ex situ analysis. To confirm that the hydrogen atoms were absorbed in the lattice, in situ XRD analysis under a hydrogen atmosphere at a pressure of 1.1 MPa (this pressure is not high enough for full hydrogenation, but it corresponds to the maximum experimentally achievable hydrogen pressure in our facility) was conducted. The results indicated a
volume expansion of 1.03% after hydrogenation (see XRD peak shifts in Fig. 6(c)). This suggests that the hydrogen atoms started occupying the octahedral sites with low hydrogen binding energy at a pressure of 1.1 MPa, which naturally results in volume expansion.

4. Conclusions

In summary, we report the design and fabrication of the Mg₄NiPd alloy, having capacity to store hydrogen at room temperature due to its low hydrogen binding energy. The Mg₄NiPd alloy with a BCC-based CsCl-type structure and a low hydrogen binding energy (a negative energy close to zero) was designed by employing first-principles calculations and synthesized by SPD using the HPT process with 1500 turns. This metastable alloy is highly homogeneous and is remarkably stable up to 440 K. It reversibly absorbs and desorbs 0.7 wt% hydrogen at room temperature (305 K). To our knowledge, this is the first successful demonstration of tuning the dehydrogenation temperature of Mg-based alloys to room temperature by applying binding energy engineering principles. More importantly, it is worth emphasizing

![Fig. 5](image_url)

**Fig. 5.** The BCC-based structure is dynamically stable and shows low hydrogen binding energy at 4Mg-Ni-Pd octahedral sites. (a) Initial lattice structure, (b) optimized lattice structure, (c) phonon density of states curve and (d) calculated hydrogen binding energy at three main types of octahedral interstitial sites obtained from first-principles calculations for the Mg₄NiPd with a BCC-based structure. Imaginary phonon frequencies are shown as negative values below the dotted line in (c); the hydrogen binding energies at the three different locations were averaged for each site type.

![Fig. 6](image_url)

**Fig. 6.** The BCC-based Mg₄NiPd phase shows room-temperature reversible hydrogen storage and remains stable after five hydrogenation/dehydrogenation cycles. (a) Hydrogen PCI results at 305 K under hydrogen pressures up to 10 MPa, (b) XRD profile after PCI measurements obtained using Cu Kα radiation, and (c) XRD profile obtained using *in situ* synchrotron radiation under a hydrogen atmosphere at a pressure of 1.1 MPa for the Mg₄NiPd processed by HPT with 1500 turns. The vertical lines and arrows in (c) show the peak shifts due to hydrogenation.
that this strategy can be universally applied to design new materials with superior functional properties beyond the scope of the equilibrium phase diagrams.

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