Divacancies and the hydrogenation of Mg-Ti films with short range chemical order

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We obtained evidence for the partial chemical segregation of as-deposited and hydrogenated Mg0.80Ti0.20 films (0 ≤ y ≤ 0.30) into nanoscale Ti and Mg domains using positron Doppler-broadening. We exclusively monitor the hydrogenation of Mg domains, owing to the large difference in positron affinity for Mg and Ti. The electron momentum distribution broadens significantly upon transformation to the MgH2 phase over the whole compositional range. This reveals the similarity of the metal-insulator transition for rutile and fluorite MgH2. Positron lifetime studies show the presence of divacancies in the as-deposited and hydrogenated Mg-Ti metal films. In conjunction with the relatively large local lattice relaxations we deduce to be present in fluorite MgH2, these may be responsible for the fast hydrogen sorption kinetics in this MgH2 phase. © 2010 American Institute of Physics. [doi:10.1063/1.3368698]

Mg-Ti alloys are promising materials for application as hydrogen storage media, metal hydride rechargeable batteries, hydrogen sensors, and smart solar collectors.1,2 Hydrogenation of Mg-Ti films with Ti-concentrations larger than ~15% (Mg0.85Ti0.15) leads to the formation of a fluorite Mg1−yTiyHx phase with substantially faster hydrogenation kinetics than the common rutile MgH2 phase. At the same time, a high hydrogen storage capacity of up to 6.5 wt % capacity is reached for Mg0.80Ti0.20.1

The Mg-Ti-H films have an intriguing microstructure. While Ti and Mg are immiscible metals on a macroscopic scale, codeposition of Mg and Ti by magnetron sputtering leads to Mg-Ti films with a coherent structure, as indicated by x-ray and electron diffraction studies,3–5 and a very high degree of intermixing of the Mg and Ti. Extended x-ray absorption fine structure (EXAFS) studies, in contrast, indicated that atomically mixed alloys are not formed. Instead, a partial chemical segregation into Mg and Ti domains on a length scale of the order of less than 10 nm was suggested.3 The stabilization of the cubic fluorite phase is thought to occur via elastic coupling of MgH2 to the nanoscale TiH2 domains. This coupling remains effective during the full (de-)hydrogenation cycle because of the similarity of the unit cell volumes of cubic TiH2 and hexagonal closed packed Mg in the intermediate state.3 A direct observation of these phase segregated domains, however, is currently lacking.

Furthermore, the fast hydrogen kinetics of these types of films—especially in the cubic hydride phase formed at Ti concentrations larger than 15%—is not well understood. The metal hydride lattice contracts upon increased incorporation of Ti, leading to reduced space for hydrogen mobility.3,4 It was suggested that the unoccupied octahedral positions in the fluorite MgH2 structure could aid the hydrogen mobility.5

Ab initio studies indicate that vacancies form a key factor in the hydrogen diffusion of both the rutile and fluorite MgH2 phases.7–9 Further insight into the occurrence of vacancies and their possible role in the hydrogenation kinetics in Mg-Ti-H films is thus strongly warranted.

In recent studies on Mg, Mg2Ni, and Mg/Si films, positron annihilation emerged as a sensitive tool to monitor the hydrogenation and phase transitions in Mg-based metal films in a depth-resolved manner.10 Here, we provide direct evidence for chemical segregation in Mg-Ti films using the positron Doppler broadening depth profiling method. Furthermore, our positron annihilation lifetime spectroscopy (PALS) study shows that divacancies are present in the metal sublattice. Such vacancies may aid to the fast hydrogen sorption kinetics associated with the cubic fluorite MgH2 phase.

Four Mg0.95Ti0.05, four Mg0.80Ti0.20, and two Ti films were deposited in a UHV system by rf magnetron cosputtering of Mg and Ti targets in argon atmosphere, on suprasil glass substrates. The films were covered with a 1–5 nm Pd capping layer to prevent oxidation and enhance hydrogen dissociation. Two samples of each composition were hydrogenated using a pressure of 90–100 mbar at room temperature. In addition, a Pd-capped Mg-layer with a thickness of ~1.3 μm was deposited on a glass substrate by dc plasma sputtering. The positron Doppler broadening of annihilation radiation (511 keV) was measured using positrons with a kinetic energy in the range of 0–25 keV. Momentum windows |p| < 3.0 × 10−3 mec and 8.2 × 10−3 mec < |p| < 23.4 × 10−3 mec for S and W, respectively, were used11 (see inset of Fig. 1). The S parameter is a measure of annihilation with valence electrons, providing sensitivity to the electronic structure and the presence of vacancies, while W is a measure for positron annihilation with semicore electrons, providing chemical sensitivity to the positron trapping site.11 PALS studies were performed using the pulsed low energy positron beam system (PLEPS).12 Positron lifetime spectra were collected at selected depths using positron implantation.
energies in the range of 0.5–18 keV, and analyzed with POSWIN (Ref. 13) using a three lifetime component analysis. Reference spectra were collected on a p-SiC reference sample to extract the instrumental time resolution function.

The results of the Doppler depth-profiling studies are summarized in the S-W diagram of Fig. 1. The depth-profiles were analyzed using the VEPFIT program,14 the fitted curves are characterized by straight lines in the S-W diagram for as-deposited and for hydrogenated Mg70Ti30, reflecting the single layer behavior of the metal (metal hydride) layer.10 The S-W cluster point characteristic for the metal Mg70Ti30 phase shows a clear and large shift to the S-W point for the fluorite Mg90Ti10H2 phase upon hydrogenation of the layer. Our previous studies10,15 demonstrated that such a large reduction in S-parameter is a direct result of the metal-insulator transition occurring during the transition from the Mg to the MgH2 phase. The transition from the free-electron-like Mg metal to the insulator metal hydride phase leads to a substantial broadening of the electron momentum distribution.15

A remarkable feature of the S-W diagram is that the cluster points for Mg70Ti30 and Mg90Ti10 lie very close to those of well-annealed (defect-free) Mg and of sputter-deposited Mg layers.16 Despite the high Ti fractions of up to 30%, the cluster points remain surprisingly far from the S-W point of well-annealed Ti. This is in a sharp contrast to Mg-based alloys such as Mg2Pd and Mg2Si, which show large shifts toward S-W points intermediate of those of Mg and Pd or Si, respectively.10 Evidently, in our case, positrons do not see an Mg-Ti alloy with an atomically mixed composition but rather detect the presence of pure Mg, as revealed from the position of the S-W cluster points. This selective detection requires the presence of chemically segregated Mg areas, into which the positron will trap during its diffusional motion which follows on the implantation and subsequent thermalization process.16 The trapping is driven by the large difference in positron affinity, by 2.1 eV,17 for Mg metal relative to Ti (Fig. 2). The Ti domains embedded in a Mg environment will act as positron potential barriers and, consequently, once a positron is injected in a Mg domain, it will not be able to re-enter neighboring Ti domains during its diffusion motion. The positron Doppler broadening studies thus provide direct evidence for the presence of Mg domains that are chemically segregated from Ti domains. The absence of a clear signature of Ti in the positron measurements further indicates that the coherent embedding of Ti-domains in Mg leads to interfaces with hardly any vacancy-related defects. Such interface defects would act as positron trapping sites, which are easily detected by the positron method16,18 via annihilation with 3d-electrons of the surrounding Ti atoms since this would lead to a distinctly higher value of the W parameter than observed for Mg (Fig. 1).

The selective probing by positrons offers the interesting possibility to monitor the metal-insulator transformation of the Mg areas in these types of Mg-Ti films directly. Upon hydrogenation, the positrons remain confined to MgH2 areas, as demonstrated by the S-W points of the Mg70Ti30, Mg50Ti50 films and of MgH2 (Ref. 10) films (open squares), Mg50Ti10H2 and Mg90Ti10H2 films (open circles), and TiH2 (open squares) films.

![FIG. 1. (Color online) S-W diagram for as-deposited and hydrogenated Mg70Ti30 films, with the cluster points surrounded by circles; the running parameter is the positron implantation energy E. The solid lines are extracted from the combined VEPFIT analysis of the S and W depth profiles. For comparison purposes, the S-W cluster points are shown for (1) defect-free bulk Mg (Ref. 10) and Ti (filled squares), and (2) Mg (Ref. 10) and MgH2 (Ref. 10) films (filled triangles), Mg90Ti10 and Mg50Ti10H2 films (open squares), and TiH2 (open squares) films.](Image)

![FIG. 2. (Color online) Schematic representation of the nanoscale chemically segregated Mg-Ti domain structure. The schematic drawing for the positron potential energy E+ shows that a Ti domain embedded in Mg acts as a positronic potential barrier with height ΔE+ = 2.1 eV (Ref. 17).](Image)

![FIG. 3. (Color online) Doppler S-parameter for the Mg70Ti30 layers in the as-deposited (open circles) and hydrogenated (filled circles) states as a function of Ti-composition.](Image)
ergies for which positrons annihilate solely within the Mg-Ti(H) layer, namely, revealed a nearly single exponential decay with a high intensity in the range 95%–100% for the second lifetime component, associated with vacancies, for each of these films. This shows that the defect concentrations are at least of the order of \(10^{-4}\) (Ref. 20) but higher values are quite likely. Figure 4 presents the extracted positron lifetimes for the Mg layer, and for the Mg-Ti layers in the as-deposited and hydrogenated states. The measured positron lifetime of 312 ± 4 ps for the Mg film is close to the value of 316 ps for the divacancy in Mg obtained in recent \textit{ab initio} calculations.\(^{21}\) The comparison with experimental and theoretical positron lifetimes for the monovacancy\(^{22,23}\) and divacancy\(^{21}\) in Mg strongly indicates that trapping occurs in divacancies. Clearly, the detected positron lifetime in the Mg\(_{1-x}\)Ti\(_x\) metal phase scales proportional to the volume per metal atom\(^{3,4}\) (Fig. 4), as expected for the divacancy.\(^{24}\) The Mg-Ti lattice contracts proportionally to the Ti-fraction\(^{3,4}\) which reduces the size of the divacancy. This leads to a larger electron density present at the positron trapping site with correspondingly higher positron annihilation rates. In contrast, Fig. 4 shows that the positron lifetime for fluorite phase Mg\(_{70}\)Ti\(_{30}\)H\(_x\) is nearly equal to the positron lifetime for rutile phase Mg\(_{90}\)Ti\(_{10}\)H\(_x\), despite the significantly smaller volume per formula unit, by 16\%, for the fluorite phase.\(^{3,4}\) This indicates that the divacancy in the fluorite MgH\(_2\) structure (inset of Fig. 4) occupies a relatively large open space leading to a low electron density. This requires local lattice relaxations of nearby hydrogen and Mg atoms, which can be facilitated by displacements to neighboring empty octahedral sites.\(^{6}\) The combined effect of divacancies present and relatively large local lattice relaxations may effectively lower the migration barriers for hydrogen diffusion in the fluorite MgH\(_2\) phase, leading to the fast hydrogen transport. Indeed, recent \textit{ab initio} modeling studies\(^{7-9}\) point to the relevance of vacancy-mediated hydrogen diffusion in MgH\(_2\) and related metal hydrides, since this provides a major contribution to the hydrogen mobility.

In conclusion, evidence for chemical segregation in Mg-Ti films into Mg and Ti domains was obtained using positron annihilation methods. Selective detection of the Mg areas in these films provide a local view on the metal-to-insulator transition upon hydrogenation to fluorite phase MgH\(_2\). Experimental evidence is provided for the presence of divacancies in the metal sublattice of both the rutile and fluorite MgH\(_2\) phases. The lifetimes of these vacancies are comparable, though their size must differ considerably. Hence, we deduce the existence of local lattice relaxations in the fluorite phase, which may play a crucial role in the enhanced hydrogen transport in fluorite MgH\(_2\).

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\(^{21}\) J. M. Campillo Robles, private communication (February 18, 2010).

