In situ X-ray Diffraction Investigation of Hydrogen Storage Alloys During Charge and Discharge

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Abstract. Metal hydride electrodes have shown to be useful both in secondary batteries, e.g. the Ni/metal hydride system, and more recently in fuel cells, especially for direct borohydride fuel cells. This work provides in situ X-ray diffraction (XRD) results aiming at characterizing the phase transitions of the LaNi4.75Sn0.25 alloy, which is usually used in Ni/metal hydride batteries. Results have shown that the method of charging leads to different phase predomination and different site occupancies, but both methods lead to almost the same discharge capacity. In the electrolysis process, γ-β phase transitions are predominant during the charge/discharge, while in the chemical charging by exposure to a borohydride solution the α-γ phase transition is more important, indicating occurrence of smaller localized stress in the alloy, with benefit on the material lifetime. Essentially the same phenomena were observed for the alloy with Pt, either with respect to the alloy structure and the electrochemical characteristics.

Introduction

Metal hydride electrodes have shown to be useful both in secondary batteries, e.g. the Ni/metal hydride system, and more recently in fuel cells, especially for direct borohydride fuel cells.[1,2] The charge and discharge processes of the metal hydride alloy are important in both of these systems. In the battery application, the extent of charging determines the discharge capacity, and the rate of diffusion of the hydride species within the structure of the alloy contributes to the limiting rate of charge or discharge. For fuel cell applications, the formation of the hydride phase may affect both the catalytic activity and the structural stability of the electrode.[3,4]

The most common Ni/metal hydride battery anode uses an AB5-type material, which is an intermetallic alloy where the A element is usually La or Mm (misch metal, which is a natural mixture of metals comprised of Ce, La, Nd and Pr). The B element may be any or a combination of transition metals. The structure of this type of alloy generally follows that of LaNi5, which crystallizes in the CaCu5-type structure (space group P6/mmm; 1 La in 1a; 0, 0, 0; 2 Ni in 2c: 1/3, 2/3, 0; 3 Ni2 in 3g; 1/2, 0, 1/2). Replacement of La and Ni by different A and B elements can dramatically improve the cycle life of the alloys and reduce problems of hysteresis, peak broadening and intrinsic degradation.[5] These problems appear, because the hydriding and dehydriding processes lead to an increase or decrease of the lattice constant of the crystal structure, respectively. Such changes may cause particle collapse leading to the formation of segregated lanthanum phases, which diminishes the electrical conductivity and increases the surface area leading to increased alloy corrosion.[5] Partial substitutions of the A and B components can also lead to improvement of the maximum discharge capacity of the alloys. For example, the substitution of molybdenum to form La0.55Ce0.45Ni5.15Co0.85–xMn0.35Al0.65Mo0.5 (x = 0.00, 0.15, 0.20, 0.25) was investigated and it was observed that the composition where x = 0.25 presented the highest discharge capacity and faster electrochemical kinetics compared to the other compositions.[7] Recently, catalytic electrodes consisting of MmNi5.25Co0.25Mn0.35Al0.65 (AB5-type alloy) and multi-walled carbon nanotubes (MWNTs) were investigated for NaBH4 electrooxidation by cyclic voltammetry. It is found that the electrode performance is significantly affected by the content of MWNTs and the optimized content of MWNTs is found to be 2 wt.%. This was attributed to MWNTs, which acts as a hydrogen adsorbent to diminish hydrogen release.[8] In situ X-ray powder diffraction (XRD), such as used in the investigation reported herein, enables the structure of the metal hydride alloy to be monitored during the processes of hydrogen absorption and desorption. Previously reported in situ XRD studies of the mechanisms of phase/structural transformations during hydride formation, decomposition, and the kinetics of the absorption/desorption processes in “fast” intermetallic hydrides such as LaNi5–Hx have shown the contributions of three hydride phases and their interconversion (α→γ→β). In situ XRD results obtained during the hydriding–dehydriding process of a LaNi4.75Sn0.25 alloy electrode showed that a lattice strain occurs in the hydride phase during the hydriding process.[11] A study of the activation of LaNi5 and LaNi4.75Al0.25 found changes in the crystallite size and evidence of lattice strain during this process.[12] For LaNi5Co it was observed that the presence of Co in the composition of the alloy generates less internal stress and leads to a higher discharge capacity compared to LaNi5.[5] Several metal alloy hydrides have been studied in our group as electrocatalysts for the oxidation of borohydride ions in alkaline media. The electrochemical data suggest that alloy hydriding occurs simultaneously with the borohydride oxidation. We
showed that LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ and LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ – Pt were able to charge when exposed to the borohydride solution and that the hydrolysis process of borohydride ions on LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ was not significant. The study reported in this work provides in situ XRD results aiming at characterizing the phase transitions of the alloys when the electrodes are exposed to borohydride solution. Results obtained for the formation of the hydride phase via electrolysis are compared to those obtained for the spontaneous formation of the hydride upon contact with the borohydride solution.

**Results and Discussion**

Fig. 1 shows the electrode potential vs. discharge capacity curves obtained during the electrochemical discharging process ($i_{\text{discharge}} = 1.0$ mA), for the electrodes charged either electrochemically applying a cathodic current (- 1.0 mA) or chemically by exposure to NaBH$_4$ $10^2$ mol L$^{-1}$ for two hours in the in situ XRD flow cell. During both of these procedures, XRD measurements were carried out every 20 seconds, with the aim of mapping the structural changes occurring during the processes of charge and discharge either by water electrolysis and by exposure to borohydride ions.

![Figure 1](image1.png)

**Figure 1.** Electrode potential as a function of discharge capacity for LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ with and without Pt when charged by the electrolysis and by exposure to NaBH$_4$ $10^2$ mol L$^{-1}$ in KOH 1.0 mol L$^{-1}$.

In agreement with our previously published works, the results of charge capacity for LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ with and without Pt show that the alloys present almost the same discharge capacity when charged by both electrolysis and exposure to borohydride solution for 2 hours. For the alloy without Pt it is seen that initial potential for both methods is similar, whilst for the material with Pt charged by exposure to borohydride, the initial potential of the discharge process following chemical charging is lower (more negative) than that obtained following the electrochemical charging. This result is explained by the fact that hydrogen oxidation and the direct oxidation of borohydride ions occurred in parallel for the Pt modified electrode.

Fig. 2 shows the in situ XRD results obtained during the electrochemical process of charge and discharge of the LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ alloy, with and without Pt, in the range 19.5° ≤ 2θ ≤ 24°. Noise in the data is attributed to the formation of bubbles during the charging process. Shifts in the background (removed) were also observed and attributed to the flowing electrolyte. Nevertheless, the quality of the XRD data obtained is excellent. Peaks associated with three different phases during the alloy hydriding and dehydriding for both samples are clearly observed in the data shown in Fig. 2. For LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ (Fig. 2A) the first phase, α (the largest diffraction peak at 2θ = 22.0°), refers to the original crystal structure of the alloy, containing only a small amount of hydride. The second phase, γ, (peak at 2θ = 21.5°) represents formation of a new crystal structure due to the insertion of hydride, thus causing an expansion in the volume of unit cell. Finally, phase β (peaks at 2θ = 20.0° and 20.5°) represents the crystal structure of either the fully hydrided alloy or with at least two-thirds of the total possible hydride content. Previous results have shown that the direct transition from phases α to β represents an expansion along both the c and a axes, corresponding to an increase of 25% in the crystal volume.

![Figure 2](image2.png)

**Figure 2.** In situ XRD results for the LaNi$_{4.7}$Sn$_{0.2}$Cu$_{0.1}$ alloy with and without Pt, during the electrochemical process of charge and discharge.
while the transition from α to γ followed by the transition from γ to β result in a smaller volume expansion (22.8%) and is thus accompanied by less localized stress. This last phenomenon is expected to occur for the alloys containing other transition metals partially replacing Ni on the B element of AB5 metal hydride alloys, as is the case of Sn.

The results obtained for the Pt modified alloy are shown in Fig. 2B. The peaks attributed to the α, β, and γ phases of the alloy are largely unchanged upon addition of Pt. There is, however, an additional peak characteristic of the Pt fcc lattice at 2θ = 22.6°. This latter peak could only be observed due to the high intensity x-ray source used in these XRD experiments and was not observed in our previous studies.\[^{13,14}\]

Fig. 3 shows plots of the intensity of the more prominent peaks for each phase as a function of the charging/discharging time for the LaNi4.7Sn0.2Cu0.1 and LaNi4.7Sn0.2Cu0.1 - Pt samples. In both cases the peaks for the α and γ phases decrease slightly during the charging and then increase during discharge. An opposite trend is observed for the peak attributed to the β phase. For the alloy without Pt, in particular, it is observed that the α and γ phases coexist in the beginning of the charge, with the γ phase most likely being present at the surface of the metal hydride particles, while the α phase is in the core. The opposite may occur at the end of discharge, explaining why the maximum peak intensity of the γ-phase is lower at the beginning of the alloy charging than at the end of discharge, when a compression of the crystal structure occurs due to the hydride oxidation.\[^{10}\]

It should be noted that the γ-β phase transformation involves the transition between a phase with low concentration of hydride species to another one initially containing low hydride content, but retaining the structure up to the insertion of about two-third of the full possible M-H content.\[^{5}\] A continuous lattice expansion and contraction occurs during this γ-β phase transition,\[^{5}\] as mentioned above.

Figs. 4 and 5 show the corresponding in situ XRD results for the LaNi4.7Sn0.2Cu0.1 alloy with and without Pt, obtained during chemical charging, followed by the electrochemical discharge. Some differences with respect to the XRD patterns obtained during the electrochemical charging are apparent. For LaNi4.7Sn0.2Cu0.1 it is observed (Fig. 4A) that the α-γ phases are apparent, but there is no significant appearance of the β-phase, as also shown in Fig. 5. This suggests lower insertion of hydrogen into the lattice structure in this later case, compared to those of electrochemical charging. This is consistent with the results presented in Fig. 1, which indicates that the total charge accumulated during chemical charging is ca. 80% of the total charge accumulated from electrochemical charging. Regarding the lines for the α-γ phases in Fig. 5, it should be noted that the trends are similar as those for the electrochemically

![Graph 1](image1)

Figure 3: Peak intensity for LaNi4.7Sn0.2Cu0.1 and LaNi4.7Sn0.2Cu0.1 – Pt during the charge and discharge process by electrolysis.

![Graph 2](image2)

Figure 4: In situ XRD results for the alloy LaNi4.7Sn0.2Cu0.1 with and without Pt deposited during the charge by exposure of BH₄⁻ 10⁻² M for 2 h followed by the electrochemical discharge.
charged/discharged alloys (Fig. 3). So, these results are consistent with the fact that, as mentioned before, the γ phase is present at the surface of the metal hydride particles, while the α phase is in the core. The results for the alloy with Pt show that, also in this case the charging and discharging only involves the α-γ phase transition. As mentioned before, the β-phase appears for levels of alloy hydriding higher than ca. 67%, meaning that the appearance of this phase would only occur by the end of the alloy chemical charging and the beginning of the discharging, but this is not confirmed. Explanation for this contradiction may be given in terms of the presence of residual borohydride covering the metal hydride particle surface or trapped into the carbon network support following the chemical charging, which may be able to replenish some of the oxidized hydride in the course of the discharging process. As mentioned above for the Pt-containing alloy, the lower potentials observed in the beginning of the discharge step provide evidence of the presence of the borohydride ions in contact with the Pt particles, because of the lower reversible potential of the borohydride oxidation compared to that of the metal hydride. In summary, the appearance of the γ-phase during the borohydride charging process confirms hydride insertion into the alloy. However, there is no formation of the β-phase and this indicate occurrence of smaller localized stress in the alloy, compared to the electrochemical charging, with benefit on the alloy lifetime, as evidenced by the results in Fig. S2 (support information). So, results show that the system essentially remains with the α and γ-phases during the whole experiments, confirming that the charging process by exposure to borohydride is in fact smaller than the electrochemical step; but this is not surprising because of the low BH$_4$ concentration used in the chemical charging process.

Figure 5: Peak intensity for LaNi$_4$Sn$_2$Cu$_0.1$ during the charge by exposure to borohydride $10^2$ mol L$^{-1}$ and discharged by electrolysis.

Conclusions

The in situ XRD results have shown that the method of charging of the alloy, electrochemical or chemical, leads to different phase predominance and different site occupancies, but both methods lead to almost the same discharge capacity. In the electrolysis process, the γ-β phase transition is predominant while in the chemical charging by the exposure to borohydride solution the α-γ phase transition is more important, indicating occurrence of smaller localized stress in the alloy, with benefit on the material lifetime. The results have also shown that the alloys with and without Pt lead essentially to the same phenomena, either with respect to the alloy structure and the electrochemical characteristics.

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In situ X-ray diffraction XRD was used to investigate the phase transitions of the metal hydride alloys when the electrodes were charged/discharged. Results have shown that the method of charging, electrochemically or chemically, leads to different phase predominance and different site occupancies, but both methods lead to almost the same discharge capacity.