



Chemical surface modification for the improvement of the hydrogenation kinetics and poisoning resistance of TiFe

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ABSTRACT

Hydrogen storage performance of the TiFe-based materials suffers from difficulties with hydrogenation and sensitivity towards impurities in hydrogen gas reducing hydrogen uptake rates and decreasing the cycle stability. In present work the surface modification of the sintered $Ti_{1.1}Fe_{0.9}O_x$ and arc-melted TiFe was performed using autocatalytic deposition of the Pd-based catalytic layers in order to achieve improvement of the H storage characteristics. Pd deposition proved to be efficient in significant facilitation of the hydrogenation ability of the materials at moderate H_2 pressures and room temperature, even after their long exposure to air. Activation performance of the sintered $Ti_{1.1}Fe_{0.9}O_x$ is superior than that for the arc-melted TiFe. This effect was associated with the presence of the oxygen-containing suboxide $Ti_4Fe_2O_{1-x}$ in the sintered sample acting as a hydrogen transfer catalyst. γ -Aminopropyltriethoxysilane pre-functionalization and subsequent Pd–P autocatalytic deposition onto the sintered $Ti_{1.1}Fe_{0.9}O_x$ intermetallic substrate resulted in a better hydrogenation kinetics compared to the samples prepared by the conventional Pd deposition.

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1. Introduction

Recently experienced difficulties in achieving uninterrupted supply of the Rare Earth metals and alloys, put a demand to develop solid hydrogen storage materials alternative to the commonly used AB_5 type alloys. Such alternatives first of all are within the alloys of Ti, Zr and Mg.

TiFe intermetallic compound is a well-studied hydrogen storage material known for more than 30 years [1]. Its reasonable storage capacity (up to 1.9 wt.% H), low pyrophoricity, abundance and low cost [2] cause interest to its applications. Improvements of the hydrogen storage performance of TiFe, including simplification of its activation and modification of the thermodynamic behaviours, can be achieved by a partial substitution of Fe with other metals, first of all, by Mn and V [2–5]. Different techniques used in the preparation of TiFe and its derivatives include smelting of the constituent elements [2,6], mechanical alloying of (Ti + Fe) or $(TiH_2 + Fe)$, followed if necessary by annealing [7–9] or sintering of (Ti + Fe) in hydrogen atmosphere [10–12]. Despite the listed methods proved to be easily upscalable, however, special precautions must be taken during the synthesis because of the ability of the TiFe-based materials to easily pick-up oxygen, and the possibility of the formation of $TiFe_2$ which does not absorb hydrogen

under acceptable pressure–temperature conditions. To avoid these complications, Ti is taken in 10–30% excess as compared to the stoichiometric Ti:Fe ratio, and/or deoxidisers like rare-earth metals are introduced. These modifications improve hydrogen storage performance, in particular two hydrogen absorption–desorption plateaux for $Ti_{1+x}Fe$ are merged thus providing high hydrogen storage capacities at reasonable hydrogen pressures [1,2,6], as well as having a benefit of an easier activation of the $Ti_{1+x}Fe/RE$ alloys [13,14]. Alternatively, the activation can be facilitated by the introduction of oxygen, to yield $Ti_{1+x}FeO_y$ ($x = 0.22–0.24$, $y = 0.007–0.04$) [6].

The main drawback of TiFe and its derivatives is in their poor activation performances and low poisoning tolerance resulting in the suppression of hydrogen absorption even by trace amounts of added to hydrogen active gases like oxygen and water vapour [15,16]. Attempts to address these problems included ball milling of the parent alloy together with additives of H_2 dissociation catalysts, Pd [17] and Ni [18]. Studies of the thin films TiFe + Pd showed that Pd coating (20 nm) of TiFe (100–200 nm thick films deposited on silicon) promotes the hydrogenation of TiFe; the rate of hydrogen uptake can be further increased by annealing of the Pd-capped TiFe in air [19]. Most probably, the reason for that is in the formation of oxygen-containing phases ($Ti_4Fe_2O_{1-x}$) which were shown to easily absorb hydrogen at mild conditions [20–22]. The formation of such phases was also observed in the course of arc-melting of titanium, iron and iron oxide [6], or sintering of Ti and Fe powders in non-purified hydrogen (99.8% $H_2 + 0.2\% O_2$), to yield TiFe as a major product [10].

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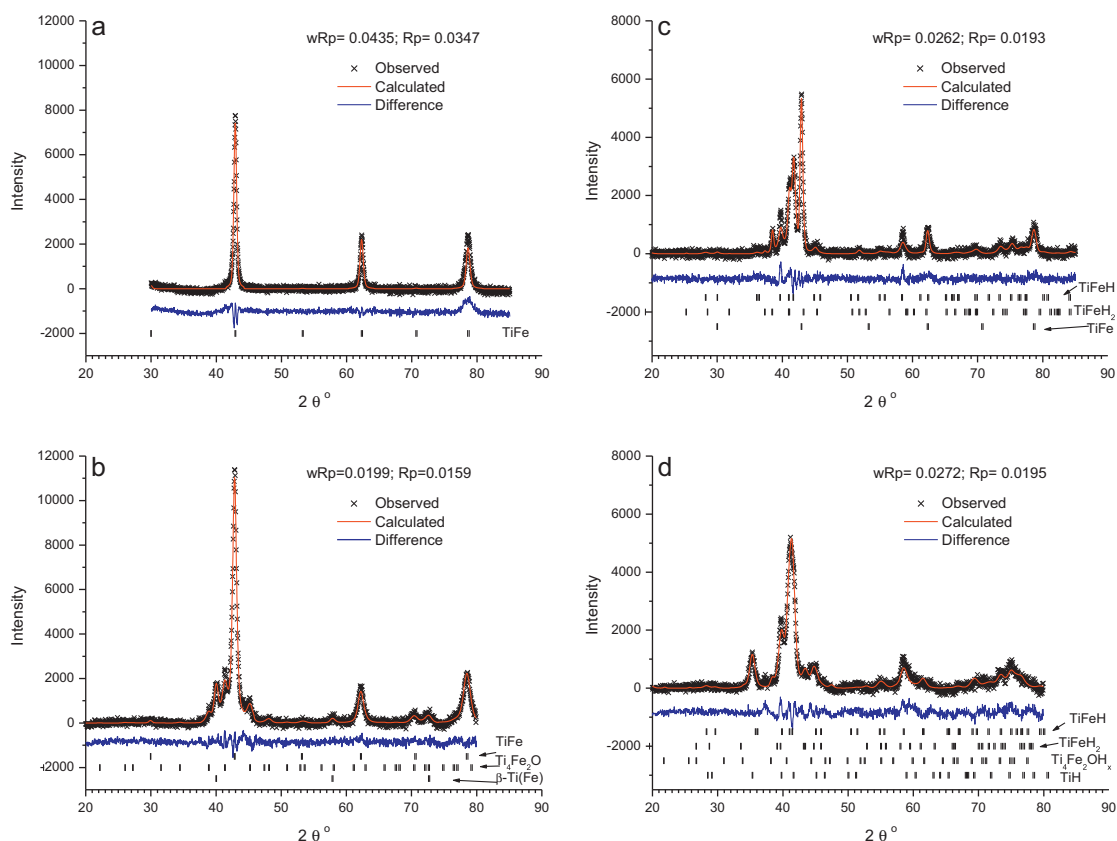


Fig. 1. XRD pattern of TiFe substrates: a – arc-melted TiFe; b – sintered $Ti_{1.1}Fe_{0.9}O_x$; c – arc-melted TiFe after hydrogenation; d – sintered $Ti_{1.1}Fe_{0.9}O_x$ after hydrogenation.

The improvement of activation and H sorption properties in the modified Ti-based intermetallics, which are very sensitive towards surface poisoning, will be more pronounced than in the case of easier activated AB_5 -type materials [23]. Thus, TiFe-based substrates are very relevant in studying the effects of their surface modification. Taking into account the mentioned above sensitivity of the activation performances of TiFe to the preparation routes, it would be also important to reveal the effects of the surface modification of the TiFe substrates prepared by different methods.

This work presents experimental results on morphological and hydrogen charging characteristics of the arc-melted and sintered oxygen-containing TiFe alloy, both were surface-modified through the autocatalytic deposition of Pd(P) [24–26]. The primary objective was in the development of the Rare Earth-free hydrogen storage materials with improved activation performance and increased poisoning tolerance.

2. Experimental

TiFe intermetallic was prepared by arc-melting (Institute for Energy Technology) of the high-purity metals (>99.99%) taken in equiatomic amounts (total mass ~5.0 g) on a water-cooled copper crucible in protective argon atmosphere. Subsequently, the prepared metal ingots were pulverised by ball-milling in argon for 10 min.

Sintered $Ti_{1.1}Fe_{0.9}O_x$ was prepared by mixing Ti and Fe powders (Alfa Aesar, purity 99.99%, particle size 44 μm) in 1.1: 0.9 atomic ratio, through grinding in a mortar, and placing the mixture (~10 g) into a quartz or tungsten sample holders positioned in a quartz tube inside a tubular furnace (South African Institute for Advanced Materials Chemistry). The tube was flushed at room temperature with Ar (Afrox Ltd., 99.999%, O_2 and H_2O impurities below 10 ppm) for 1 h to displace air. This was followed by heating to 1000 °C, maintaining the high temperature for 1.0 h, and cooling down to room temperature under Ar flow.

The excess of Ti (~22% as to the stoichiometric amount in the sintered material) was chosen to (i) avoid a formation of inactive in hydrogenation $TiFe_2$ and (ii) increase the Ti/Fe ratio to allow formation of $Ti_4Fe_2O_x$. From experiments (see next

section, Fig. 1, Table 1) we have found that chosen content of Ti and sintering conditions yielded a mixture of two hydride-forming constituents, TiFe and $Ti_4Fe_2O_{1-x}$ formed in ratio of approximately 3:1.

Surface modification of the TiFe-based substrate was achieved by autocatalytic deposition of palladium in hypophosphite-based baths. The materials were initially sensitized-activated in a $SnCl_2$ -PdCl₂ colloidal solution (30 min; 300 rpm; 20 °C) prepared by dissolving 0.3 g PdCl₂ in 2.15 g HCl and mixing with 25 g $SnCl_2 \cdot 2H_2O$ dissolved in 14.45 g HCl and 8.75 g KCl dissolved in 80 mL HCl. The activated powders were then accelerated in a 10% (w/v) solution of disodium ethylene-diamine-tetraacetic acid (Na_2EDTA) for 10 min. The materials were further plated (pH ~9.0; 50 °C; 30 min; 300 rpm) in an ammoniacal bath solution, which was constituted as follows: 1.0 g L⁻¹ PdCl₂; 27 g L⁻¹ NH_4Cl ; 160 mL L⁻¹ NH_4OH ; 4.0 mL L⁻¹ HCl; 10 g L⁻¹ NaH_2PO_2 . A customized autocatalytic deposition technique allowing to use dilute (< 1.0 g/L) solutions of Pd salts with 100% utilization of Pd [25] was also utilized in the surface modification of the sintered $Ti_{1.1}Fe_{0.9}O_x$ based on the detection of oxygen. The technique involved surface pre-functionalization of the sintered substrate using 1.0 vol.% solutions of γ -aminopropyltriethoxysilane (γ -APTES) to facilitate the surface immobilization of Pd nuclei in the sensitization-activation process, resulting in increased densities of deposited Pd(P) particles and surface Pd loading on the intermetallic compared to that prepared without the pre-functionalization step [26].

The samples were characterised by XRD (Bruker AXS D8 Advance, Cu-K α , $\lambda_1 = 1.5406 \text{ \AA}$, $\lambda_2 = 1.5444 \text{ \AA}$, $\lambda_2/\lambda_1 = 0.5$, $2\theta = 20\text{--}85^\circ$), SEM (Zeiss Ultra 55, Norwegian University of Science and Technology, 4.0 kV, secondary electrons), and volumetric measurements of hydrogen absorption kinetics ($T = 20^\circ C$, $P_0 \sim 30$ bar H_2 , for 24 h). Before the measurements of hydrogen absorption, the system was evacuated to not less than 1.0×10^{-3} mbar at room temperature/ $T = 400^\circ C$, for 1 h.

All materials were allowed exposure to air (no less than two weeks) prior to the experimental studies of their hydrogen absorption performances.

3. Results and discussions

Summary of the obtained data on the phase-structural composition of the samples is presented in Table 1 and is given together with relevant reference information on the constituent phases [20,22,27–31]. The XRD pattern of the initial alloys and their corre-

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