Synthetic nanocomposite MgH$_2$/5 wt. % TiMn$_2$ powders for solid-hydrogen storage tank integrated with PEM fuel cell

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Storing hydrogen gas into cylinders under high pressure of 350 bar is not safe and still needs many intensive studies dedicated for tank’s manufacturing. Liquid hydrogen faces also severe practical difficulties due to its very low density, leading to larger fuel tanks three times larger than traditional gasoline tank. Moreover, converting hydrogen gas into liquid phase is not an economic process since it consumes high energy needed to cool down the gas temperature to $-252.8$ °C. One practical solution is storing hydrogen gas in metal lattice such as Mg powder and its nanocomposites in the form of MgH$_2$. There are two major issues should be solved first. One related to MgH$_2$ in which its inherent poor hydrogenation/dehydrogenation kinetics and high thermal stability must be improved. Secondly, related to providing a safe tank. Here we have succeeded to prepare a new binary system of MgH$_2$/5 wt. % TiMn$_2$ nanocomposite powder that show excellent hydrogenation/dehydrogenation behavior at relatively low temperature ($250$ °C) with long cycle-life-time (1400 h). Moreover, a simple hydrogen storage tank filled with our synthetic nanocomposite powders was designed and tested in electrical charging a battery of a cell phone device at $180$ °C through a commercial fuel cell.

Hydrogen is an energy carrier, which holds tremendous promise as a new clean energy option$^{1,2}$. It is a convenient, safe, versatile fuel source that can be easily converted to a desired form of energy without releasing harmful emissions$^{3,4}$. A key advantage of hydrogen is that when burned, carbon dioxide (CO$_2$) is not produced. Hydrogen storage is one of the most crucial difficulties restricting utilization of hydrogen energy for real applications. However, storage hydrogen gas into gas cylinders under high pressure reached to 350 bar is a well-known technology, using such pressurized hydrogen gas tanks as a source of fuel in vehicles is not safe at present and still needs many intensive studies dedicated for improving the structural and mechanical properties of the materials used in tank’s manufacturing. Extreme conditions for on-road vehicle service should be defined, to demonstrate performance of storage systems demonstrated both under the stresses of normal vehicle operation and under externally imposed stresses$^5$. Likewise pressurized hydrogen gas, liquid nitrogen possesses many difficulties related to its very low density. Accordingly, the size of liquid hydrogen requires larger tanks reaches to about three times larger than the traditional gasoline tank$^6$. Practically, converting hydrogen gas into liquid hydrogen is not an economic process since it consumes large amount of energy required to cool down the gas temperature to $-252.8$ °C. For instance, liquefying 1 kg of hydrogen gas in medium-size plant requires 10 to 13 kWh of electrical energy$^7$. Moreover, liquid nitrogen is not safe since it has a high flammability range. Boil-off losses associated with the storage, transportation and handling of liquid nitrogen can consume up to 40% of its available combustion energy$^8$.

**Solid Hydrogen**

Apart from gaseous and liquidus phases of hydrogen, solid hydrogen has been considered as the most reliable and safe practical solution for providing clean energy required for different applications, using proper fuel cells such as proton-exchange fuel cells membrane (PEM)$^9$. Hydrogen can be simply stored in nanocrystalline metal...
powders such as Mg and Mg-based nanocomposite powders in the form of MgH₂. The choice of Mg is attributed to its high hydrogen capacity (7.60 wt. %, 0.11 kg H₂ L⁻¹), natural abundance, cheap price, operational cost effectiveness, and light weight. Accordingly, MgH₂ has become a potential candidate for fuel cell applications used in light-duty vehicles and mobile application²⁵.

However, nanotechnology has had an obvious impact on producing industrial scale of uniform nanocrystalline MgH₂ powders, using a room-temperature reactive ball milling technique (RBM)²⁶. The nanophase of such metal hydride system still shows serious drawbacks that should be solved first before nominating the system for real applications. Firstly, MgH₂ has a high thermal stability making the hydrogen releasing at moderate temperatures (below 300 °C) very difficult²⁷,²⁸,²⁹. Secondly, MgH₂ exhibits very slow kinetics of hydrogenation/dehydrogenation at temperatures around 350 °C³⁰,³¹. Innumerable efforts have been targeted to improve the kinetics behavior of MgH₂ by catalyzing the metal hydride powders with wide spectrum of mono, binary, and multicalytic systems. One of the earliest work proposed for improve MgH₂ powders was achieved by Prof. R. Schulz and his team work in 1999³². In their work, MgH₂ powders were catalyzed by ball milling with one of 3-d transition metal powders of Ti, V, Mn, Fe and Ni. Based on their results, Ti and V showed better catalytic effect for hydrogen absorption and desorption when compared with Ni. Furthermore, Hanada et al.³³ reported very interesting results on catalyzing of MgH₂ powders by small amount (1 mol. %) of Fe, Co, Ni and Cu nanoparticles. The as-mechanically doped MgH₂/Ni powders obtained after a very short milling time (2 h) showed excellent hydrogenation/dehydrogenation kinetics properties and enjoyed high storage capacity (~6.5 wt. %)³⁴. Since then, different schools have reported attractive results upon using pure elemental powders such as Al, Ti, Fe, Ni, Cu and Nb,³⁵ intermetallic compounds³⁶,³⁷, metastable big-cube Zr₂Ni³⁸, and metal/metal oxide binary nanocomposite³⁹,⁴⁰ for improving the kinetics of hydrogen absorption/desorption of MgH₂. More recently, an interesting study was reported by Ouyang et al.³⁵ when they successfully prepared Mg₂Ni-Ni solid solution with an Mg₂Ni-type structure. They pointed out that the introduction of In-semimetal into Mg₂Ni not only significantly improved the dehydrogenation kinetics but also greatly lowered the thermodynamic stability. This was implied by lowering the activation energy of dehydrogenation and enthalpy change to very low values of 28.9 kJ/mol and 38.4 kJ/mol H₂, respectively³⁵.

Besides metal, semimetal, and metallic metastable phases, hard metal oxide of Nb₂O₅⁴¹ and refractory material powders and, such as SiC⁴², and TiC⁴³ find a space of applications as excellent kinetics modifier used successfully for improving the hydrogen absorption/desorption behaviors of MgH₂ system. More recently, we demonstrated the first report of employing a metallic glassy Zr₇₀Ni₂₀Pd₁₀ powders for enhancing the hydrogenation/dehydrogenation properties of MgH₂ powders⁴. In general, it is agreed that mechanically-induced doping of MgH₂ with the abrasive powders of hard phases such as carbides, oxides, intermetallic and metallic glassy alloys materials lead to fast grain refining of the MgH₂ upon releasing the crystalline stored energy, leading to refine the MgH₂ grains along their grain boundaries where superfine grains are formed. Such desirable fine grains with their short-distance grain boundaries always facilitate short diffusion path⁴⁵, leading to fast diffusion of the hydrogen atoms⁴⁶–⁴⁸. Recently, Crivello et al.⁴⁹ have introduced a useful review article discussing the several ways used for improving MgH₂-based materials.

Recently, Zhu et al. investigated a powerful technique called plasma-milling (P-milling)⁵⁰ when they successfully introduced dielectric barrier discharge plasma (DBDPT) during vibratory milling of Mg(In)-MgF₂ composite powders.⁵¹ The combination between the plasma process, and the impact and shear forces generated by the milling balls led to enhance the milling process for refining and/or alloying the milled powders.⁵²,⁵³

Apart from the milling process employed for preparing MgH₂ nanocrystalline and MgH₂-based nanocomposite powders, direct current magnetron sputtering technique was successfully employed in 2004 by Ouyang et al.⁵⁴ for preparing multi-layer hydrogen storage thin films with Mg and MnMn₁₋ₓ(CoAlMn)ₓ. They reported that this technique can led to the formation of high quality nano-scale multi-layer composite with well-controlled film thickness and well-bonded interface between the multilayers⁵⁵. One advantage of this promising technique can be realized from its ability on synthesizing ultra-thin films (4 nm) with clean interfaces.

**Solid Hydrogen Tanks**

Merits of using solid hydrogen tanks are obvious for those who live in isolated communities having lack in connection to the public power grid. However, renewable energy sources that generate electricity directly through for example photovoltaics (PVs) and windmills are environmentally friendly and can provide cost-effective solutions, their energy availability varies drastically from time to day per one day. Accordingly, energy storage is necessary to meet the electricity demand with the required reliability⁵⁶. Thus, the combination of stored solid hydrogen generated from excess electricity and a fuel cell is a promising solution⁵⁷.

In general solid hydrogen storage tanks differ in their design and materials used when compared with those used to store pressurized hydrogen gas and liquid hydrogen. This because diffusion of hydrogen atoms into metal lattice (e.g. Mg) does not require the application of very high pressure since the gas-solid exothermic reaction between the two phases takes place simultaneously at relatively low pressure (below 15 bar/300 °C). Moreover, the need of using expensive cryotanks with controlled pressure (similar to those used for liquid hydrogen) is not required. Accordingly, the basic process for design, materials selections, and manufacturing of vessels or tanks contain metal hydride, as a source of hydrogen becomes inexpensive in materials science point of view. In fact, the success of using solid hydrogen tank for providing hydrogen to operate an electrical system with constant rate of hydrogen flow through a suitable fuel cell can be firstly attributed to the hydrogenation/dehydrogenation kinetics behavior and cyclability of the metal hydride materials stored in the tank.

Most the authors published fine articles related to the Mg-based hydrogen storage materials, focused mainly on materials preparations, characterization and investigation the hydrogenation/dehydrogenation behaviors of the metal hydride systems. However, there are some studies reported interesting and promising results related to designing and utilizing of solid hydrogen tank. Of these, Zu et al.⁵⁸ proposed an advanced design that can be used...
for real manufacturing of hydrogen storage tank. However, their design may improve the geometric flexibility and structural performance of composite toroidal hydrogen storage tanks; they have not examined the system with any types of metal hydride to ensure the validity of their tank model for hydrogenation/dehydrogenation processes. A numerical work proposed by Gkanas et al. showed a three-dimensional computational model regarding coupled heat and mass transfer during both the hydrogenation and dehydrogenation process. In this study, two different types of hydrides, LaNi5 and an AB2-type (Ti-Zr-Mn intermetallic system) were selected as possible candidate materials. Their results showed the possibility of hydrogen gas uptake and release with moderate kinetics at 20 °C and 15 bar of H2. A very interesting and promising model was recently introduced by Gattia et al. They used compacted powders prepared by ball milling MgH2 with Nb2O5 with mixed with 5 wt.% ENG. The pellets were coated by different type of metal coating, using sputtering and thermal evaporation techniques. The material were tested by inserting the pellets into a cylindrical tank. Unfortunately, the kinetics of the system did not show attractive characteristics since the temperature required for cyclic procedure taken place at 310 °C. On the other hand, significant number of patents related to designing hydrogen storage tanks are available. One of the earliest invention related to this regard dates back to 1969, when Lyon et al. proposed a process and an apparatus used to utilize MgH2 as a source of hydrogen for useful fuel cell applications. In their prototype, Mg reacted with hydrogen gas to form MgH2 and then decomposed at high temperature (277°C to 649°C) under pressure ranging from 1 bar to 207 bar. Applications of such high temperatures and pressure are consider as drawback of their system. In 2014, Ornath introduced an interesting idea for manufacturing a hydrogen storage tank that can lead to lower the temperature of hydrogen uptake and release. Unluckily his patent did not contain any experimental results to prove his idea. Details and more information on the employing different types of metal hydride materials on solid-state hydrogen storage are recently reviewed and published by Rusman and Dahari et al.41.

Apart from the interesting and useful solid hydrogen storage models invented by many authors, our present has been addressed to satisfy two objectives; the first is focused on synthesizing and characterizing a new nanocomposite MgH2/5 wt. % TiMn2 system with advanced kinetics behaviors and cyclability, where the second objective is focused on utilizing the as-synthesized nanocomposite powders in a self-manufactured hydrogen storage tank interfaced to 40 W/4.5 A proton exchange membrane (PEM) fuel cell. This system feeds the PEM-fuel cell with the released hydrogen gas form the tank. The converted electrical energy was utilized to charge the battery of a cell phone device through 5 V voltage regulator. As far as the authors know, this is the first time to examine the validity of a new Mg-based nanocomposite in real applications.

Results and Discussion
Structure. The general and local structure beyond the nano level for the nanocomposite MgH2/5 wt. % TiMn2 powders were investigated by means of X-ray diffraction (XRD) and field emission-high resolution transmission electron microscope (FE-HRTEM) techniques, respectively. The XRD pattern of elemental Mg powders (precursor) obtained after 200 h of RBM under 50 bar of a hydrogen gas atmosphere is shown in Fig. 1(a). The powders

Figure 1. Structural characteristics of nanocrystalline MgH2 and nanocomposite MgH2/5 wt. % TiMn2 powders milled for different RBM time. (a) XRD patterns of hcp-Mg powder obtained after ball milling under 50 bar of H2 gas atmosphere for 200 h. XRD patterns of as-synthesized MgH2 powder doped with 5 wt.% TiMn2 powder and then ball milled for (b) 3 h, (c) 12.5 h, (d) 37.5 h, and (e) 50 h. The BFI image of the end-product (50 h) is shown in (f), where the HRTEM image for zone I indexed in (f) is presented in (g). HRTEM image of zone II indexed in (f) is presented in (h). The NBDPs for zones III and IV shown in (h) are elucidated in (i,j), respectively.
consisted of a fine nanocrystalline mixture of β-MgH₂ (PDF file# 00-012-0697) and γ-MgH₂ (PDF file# 00-035-1184) phases, implied by the broad Bragg peaks shown in Fig. 1(a). Low intensity Bragg peak related to fcc-MgO (PDF file# 01-079-9866) was detected at a scattering angle of about 42° due to the oxidation of the sample during XRD sample preparation outside of the glove box.

Based on the purpose of the present study, the end-product (200 h) of as-synthesized MgH₂ powders was manually doped with 5 wt. % TiMn₂ powders inside the glove box and then charged into the milling vial together with the milling media. The vial was then pressurized with 50 bar of hydrogen gas and mounted onto a high-energy ball mill where the RBM process takes place for different milling time. After 3 h of milling, the powders revealed Bragg peaks related to MgH₂ phase, where the Bragg peaks corresponding to TiMn₂ powders were hardly seen, as shown in Fig. 1(b). This may be attributed to the effect of cold welding generated by the milling process on TiMn₂ powders, leading to sticking the powders onto the milling media (vial’s internal wall and balls). Further ball milling time led to fragmentation of TiMn₂ agglomerate powders to fall into the vial’s milling zone and then milled together with MgH₂ powders. After 12.5 h, the milled powders consisted of ultrafine particles of MgH₂ coexisted with TiMn₂ (PDF file# 01-071-9712) powders, as implied by the broad Bragg peaks related to both phases (Fig. 1(c)). This broadening manifested in the Bragg peaks raised from both refinement of the MgH₂ and TiMn₂ crystallites and accumulated macrostrain during the RBM process. It should be notified that neither reacted powders related to the formation MgTiHₓ and TiH₂ phases nor elemental phases corresponding to metallic Mg and Ti could be detected even after longer RBM time, ranging between 37.5 h (Fig. 1(d)) and 50 h (Fig. 1(e)). Moreover, the Bragg-peaks of metallic hcp-TiMn₂ maintained their peak positions after 50 h of RBM, implying the absence of solubility into the MgH₂ lattice, as elucidated in Fig. 1(e).

The bright field image (BFI) of nanocomposite MgH₂/5 wt.% TiMn₂ powders obtained after 50 h of RBM time is displayed in Fig. 1(f). The nanocomposite powders revealed Moiré-like fringes with nanocrystalline-structure morphology (5 nm to 17 nm). Moreover, the lattice fringes of MgH₂ (β and γ phases) and hcp-TiMn₂ were regularly separated with an interplanar spacing (d) matching well with the reported PDF files cited in Fig. 1(a). Based on careful analysis performed of at least 30 examined zones for three individual samples, we could not detect the existence of any other phase(s). This implies the formation of binary nanocomposite MgH₂/5 wt.% TiMn₂ powders. A different examined zone located at the top edge of the particle, zone (II) was analyzed by HRTEM (Fig. 1(h)). Two selected zones (III, and IV) in Fig. 1(h) were selected to get their corresponding nano beam diffraction patterns (NBDPs). The grains located in each zone contained ultrafine nano-grains ranging in size between 5 to 8 nm, as displayed in Fig. 1(h). It can be notified that the grains did not reveal specific orientation (Fig. 1(h)) and the development of dislocation tangles into sub-boundaries can be seen clearly in zones III and IV (Fig. 1(h)). The dislocations presented inside the individual grains suggesting the nanostructure development by severe plastic deformation created by the ball milling media.

The NBDPs taken from the zone III and IV elucidated in Fig. 1(h) are shown in Fig. 1(i) and (j), respectively. Both electron diffraction revealed Debye continuous rings corresponding to β- and γ-MgH₂ phases, plus hcp-TiMn₂. The present of these diffractions indicates the random orientation of the grains in the nanocomposite powders (Fig. 1(i)). In addition, the circular-like NBDPs of MgH₂ (Fig. 1(j)) were overlapped with numerous closely spaced spots, each from diffraction of a single crystallite for TiMn₂.

Morphology and elemental analysis. In order to understand the distribution effect of TiMn₂ powders into MgH₂ matrix on the kinetics behaviors and thermal stability of the hydride phase, careful energy-dispersive X-ray spectroscopy (EDS) elemental mapping experiments were conducted for the powders obtained after the early (1–3 h), intermediate (25 h) and final (50 h) stages of RBM processing time. The composite powders obtained after 1 h of RBM revealed a fine MgH₂ aggregates (Fig. 2(a) and (b)) doped with thick layer particles of TiMn₂ (Fig. 2(c) and (d)). After 3 h of RBM (Fig. 2(e)), TiMn₂ particles (Fig. 2(g) and (h)) were deformed into flake-like morphology and heterogeneously distributed into the MgH₂ matrix (Fig. 2(f)). The EDS analysis of the powders obtained after the early stage of milling indicated that the concentration of the kinetic-modifier phase (TiMn₂) was widely varied from particle to particle and even within the particle itself. The local chemical analysis of TiMn₂ measured with an average area of 0.3 μm was in the range between 0.7 and 64 wt. %.

Increasing the RBM time (25 h) led the large TiMn₂ flaky particles to be disintegrated into smaller particles, ranging in size between 10 to 150 nm in diameter, as shown in Fig. 2(i–l). However, these metallic particles are heterogeneously embedded into the MgH₂ matrix (Fig. 2(j)). The TiMn₂ concentration in the host MgH₂ matrix was significantly improved, as indicated by the near concentration values varied from 3.6 to 7.8 wt. %. Toward the end of the RBM time, the TiMn₂ particles (dark lenses presented in Fig. 2(m)) revealed significant changes in their shapes and possessed spherical-like morphology with fair distributions (Fig. 2(o), and (p)) into MgH₂ matrix (Fig. 2(n)). This morphological improvement of TiMn₂ particles was followed by a dramatic deduction in their sizes laid within a narrow size distribution range between 5 nm to 27 nm (Fig. 2(m,o,p)). Accordingly, and based on EDS local analysis of different locations in several particles, the concentration of TiMn₂ in the MgH₂ matrix was outstandingly improved to be in the range between 4.8 to 5.3 wt. %.

Thermal stability. Differential scanning calorimetry (DSC) performed at a constant heating rate of 20°C/min under helium gas flow of 100 ml/min was employed to investigate the effect of RBM time and TiMn₂ additive on the decomposition temperature (dehydrogenation temperature at normal pressure) of as-synthesized MgH₂ powders. The DSC trace of as-synthesized MgH₂ powders obtained after 200 h of RBM revealed a broad endothermic event laid at a peak temperature of 728 K, as shown in Fig. 3(a). This endothermic peak notably shifted to the lower temperature side (697 K) upon milling for short time (3 h), as displayed in Fig. 3(b). The peak
decomposition temperature for those samples obtained after 6 h (Fig. 3(c)) and 12.5 h (Fig. 3(d)) were 671 K and 628, respectively. The tendency of MgH₂ powders to be decomposed at lower temperature (581 K) with increasing the RBM time was continuously occurred during the intermediate stage of RBM (25 h), as displayed in Fig. 3(e). During the final stage of RBM (37.5 h) the decomposition temperature was retreated to 546 K, as elucidated in Fig. 3(f). This value did not remarkably changed (541 K) for the nanocomposite powders obtained after 50 h of RBM time (Fig. 3(g).

**Apparent activation energy of dehydrogenation.** In order to realize the effect of doping MgH₂ with 5 wt. % TiMn₂ powders on the apparent activation energy (Eₐ) of the metal hydride phase, individual DSC experiments were conducted with different heating rates (5, 10, 20, 30 and 40 °C/min). In this study the effect of RBM on Eₐ were investigated for all samples obtained after different milling stages. Figure 4(a) to (c) displayed selected DSC curves conducted at different heating rated (k) for 3 samples obtained after selected RBM times (3 h, 37.5 h, and 50 h). All the scans revealed single endothermic events related to the decomposition of MgH₂ into metallic Mg and hydrogen gas, as confirmed by XRD technique. While the peak height increased proportionally with the increasing the k from 5 °C/min to 40 °C/min, the peak temperatures (Tₑ) were significantly shifted to the higher temperature side, as presented in Fig. 4(a–c)). The Eₐ of dehydrogenation related to each sample was calculated according to the Arrhenius equation:

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E_a = -RT \ln(k/k_0)
\]

where k is a temperature-dependent reaction rate constant, R is the gas constant, and T is the absolute temperature. The Eₐ values were determined by measuring the Tₑ corresponded to the different k and then plotting ln(k) versus 1/Tₑ. The Eₐ values were then obtained from the slope of line (−E/R, where R is the gas constant).
Based on these measurements, the nanocomposite powders obtained after 3 h of RBM time showed a high $E_a$ value (138.03 kJ/mol), as shown in Fig. 4(d). This indicates a high thermal stability of the powders against decomposition. In contrast, $E_a$ of nanocomposite MgH$_2$/5 wt.% TiMn$_2$ powders obtained after 37.5 h (Fig. 4(e)) of RBM showed a lower value (117.76 kJ/mol), indicating a significant destabilization of the MgH$_2$ upon high-energy ball milling with TiMn$_2$ phase. The $E_a$ value did not show notable improvement (116.99 kJ/mol) upon increasing the RBM time to 50 h, as elucidated in Fig. 4(f). The apparent $E_a$ of our system is closed to the reported one for MgH$_2$/5 wt.% Zr$_{70}$Ni$_{30}$Pd$_{10}$ powders (92 kJ/mol)\(^4\). However, our system showed lower $E_a$ value when compared with pure MgH$_2$\(^4\) (164 kJ/mol)\(^4\), Mg$_{63}$In$_{38}$Al$_{15}$Ti$_3$ (125.2 kJ/mol)\(^4\), and Mg$_{71}$Ba$_2$ (173.92 kJ/mol)\(^7\), systems, it is well above than those corresponding values reported for MgH$_2$ powders coated by Ti-based thin films (30.8 kJ/mol)\(^4\), CeH$_{2.73}$-MgH$_2$-Ni nanocomposite (63 kJ/mol)\(^8\), Mg- TiVMn (85.2 kJ/mol)\(^9\), Mg- Li$_2$TiO$_3$ (84 kJ/mol)\(^9\), and MgH$_2$-Ta$_2$O$_5$ (74kJ/mol)\(^5\) systems.

**Figure 3.** Effect of RBM time on the thermal stability of MgH$_2$/5 wt.% TiMn$_2$ powders obtained after selected RBM time. DSC curves of MgH$_2$ powder obtained after 200 h of RBM time and then mechanically doped with 5 wt.% TiMn$_2$ powder for (a) 0 h, (b) 3 h, (c) 6 h, (d) 12.5 h, (e) 25 h, (f) 37.5 h, and (g) 50 h.

During the intermediate stage of RBM (25 h) the MgH$_2$ grain size was reduced to 38 nm (Fig. 5(b)), where $T_p$ and $E_a$ reached to lower values of 582 K (Fig. 5(a)) and 124 kJ/mol (Fig. 5(b)), respectively. Toward the end of the RBM processing time (37.5 h–50 h) the MgH$_2$ grain continued their tendency to be reduced in sizes (14 nm–7 nm) according to the double successive effects of balls- and TiMn$_2$-micro milling media. Further destabilization of MgH$_2$ taken place during this final stage of RBM, referred by the reduction occurred in $T_p$ values (546 K–540 K), as shown in Fig. 5(a). Most importantly and as a result of MgH$_2$ grain refinement, $E_a$ tended to drop into a lower value (116–117 kJ/mol), as elucidated in Fig. 5(b).
Hydrogenation/dehydrogenation kinetics behavior. The improvement of kinetics related to hydrogen absorption and desorption processes attained upon mechanical doping MgH₂ with 5 wt. % TiMn₂ powders were monitored after different RBM time, using Sievert's method

Kinetics of hydrogenation. The hydrogenation kinetics behavior investigated at 250 °C under 10 bar H₂ gas pressure for nanocomposite MgH₂/5 wt.% TiMn₂ powders obtained after selected RBM time are shown in Fig. 6(a). The shaded zone indexed in Fig. 6(a) is elucidated in Fig. 6(b) with a different time scale. After 25 h of RBM, the sample absorbed 3.5 and 4 wt. % H₂ after 1.25 and 4 min, respectively as shown in Fig. 6(b). As previously discussed in Fig. 5(b), increasing the RBM time resulting a significant decreasing in the MgH₂ grain size, allowing an easier hydrogenation process. Accordingly, the sample obtained after 37.5 h of RBM time showed a significant development on absorbing 5.1 wt. % H₂ after only 1.25 min (Fig. 6(b)). Marginal improvement (5.4 wt. % H₂) was notified with increasing the absorption time to 3 min (Fig. 6(a)). Further increasing of the absorption time (~7.75 min) did not cause any improvement on hydrogen uptake and the sample saturated at 5.4 wt. % H₂, as elucidated in Fig. 6(a). However, longer RBM time (50 h) did not show a major beneficial effect on the hydrogen uptake (5.4 wt. % H₂) and hydrogenation kinetics, the sample possessed a notable ability to absorb higher H₂ value with shorter time (~4.7 wt. %/0.6 min), as displayed in Fig. 6(b). This value is higher than the one shown by

**Figure 4.** Dependence of activation energy on the RBM time for nanocomposite MgH₂/5 wt.% TiMn₂ powders. DSC curves of the nanocomposite powders obtained after RBM time of (a) 3 h, (b) 37.5, and (c) 50 h. The apparent activation energy (Eₐ) was calculated from the slope of line shown in the Arrhenius plots, displayed in (d), (e,f) for the samples milled for 3h, 37.5 h, and 50 h, respectively.
the sample obtained after 37.5 h of RBM (4.4 wt. % H₂) at the same absorption time. It should be mentioned that Mg/5 wt.% Ta₂O₅ binary system can uptake about 6 wt.% H₂ within 60 min at 100 °C⁵². Doping MgH₂ with 5 wt.% Li₂TiO₃ led to enhance the hydrogenation kinetics conducted at 300 °C to absorb 5.5 wt. % H₂ within 5 min⁵¹. For Mg doped with 30 wt.% TiMn₁.₅, 2 min was required to perform complete hydrogenation (~4.4 wt. % H₂) at 300°C⁵⁴. MgH₂-VTiCr and MgH₂-TiMn₂ systems possessed nearly complete absorption of ~5.5 and 4.5 wt. % H₂, respectively within 15 min at 300°C⁵². It has been reported that doping MgH₂ with 5 wt.% TiMn₂ led to conduct hydrogenation of ~4.8 wt. % H₂ within 250 min at 100 °C⁵⁵. Doping MgH₂ with 20 wt % Ti₀.₄Cr₀.₁₅Mn₀.₁₅V₀.₃ led to absorb 5.7 wt.% H₂ in 100 min at low temperature (100 °C)⁵⁶.

The XRD pattern of 50 h sample taken after hydrogenation process at 250 °C is shown in Fig. 6(c). The sample revealed a domain structure of γ-MgH₂ coexisted with small molecular fractions of γ-MgH₂. Moreover, the XRD analysis did not confirm the formation of any intermediate or reacted phases (MgTi, TiH₂, MgTiH) where the TiMn₂ additive powder maintained its original hcp-structure (PDF file #01-071-9712), as displayed in Fig. 6(c). This implies that TiMn₂ additive can be considered as heterogeneous catalysts, in which it enhanced the hydrogenation/dehydrogenation kinetics of MgH₂ without entering into a chemical reaction with MgH₂ matrix.

Kinetics of dehydrogenation. In order to get the complete picture of TiMn₂ effect on enhancing the behavior of MgH₂ powders, the kinetic of hydrogen releasing for the samples obtained after 25 h, 37.5 h, and 50 h of RBM were carefully investigated at 250°C at 200 mbar H₂ gas pressure. Figure 6(d) and (e) present the effect of RBM time on the desorption kinetics of nanocomposite MgH₂/5 wt. % TiMn₂ powders. After very short desorption time (1 min) the 3 selected samples showed poor dehydrogenation kinetics, indicated by the low values of H₂ released (~0.4 to 0.8 wt. %), as displayed in Fig. 6(e). Significant improving on the hydrogenation kinetics was attained for 50 h sample that desorbed ~1.9 wt. % H₂ within 2 min (Fig. 6(e)). In contrast, a moderate improvement could be notified for the sample obtained after 37.5 h and 25 h that succeed within 2 min to discharge ~1.4 and ~1.2 wt. % H₂, as presented in Fig. 6(e). Increasing the absorption time led to enhance the ability of the 3 samples for desorb more H₂, exemplified by the H₂ wt. % absorbed at 3 min for 25 h (~1.8), 37.5 h (~2.2), and 50 h (~3), as shown in Fig. 6(d). After 7 min, the 50 h sample attained a saturated value of hydrogen released (~5.4 wt. %), where 25 h and 37.5 h samples released only ~4.5 and 4.9 wt. % H₂, respectively (Fig. 6(d)). Where
there was no remarkable change in the desorbed hydrogen could be notified for 50 h sample upon increasing the desorption time between 7 to 10 min, the 25 h and 37.5 h samples tended to getting changes in the $H_2$ desorbed values that reached to $-5.1$ and $-5.4$ wt.\% after 10 min of the desorption time (Fig. 6(d)). Previously published MgH$_2$/5 wt.\% TiMn$_2$ system possessed fast dehydrogenation ability at 300 °C to discharge 4.5 wt.\% $H_2$ within only 2 min. Different research group reported that MgH$_2$/5 wt.\% TiMn$_2$ system can desorb its storage capacity of 4.8 wt.\% $H_2$ in 6 min at 270 °C. It was demonstrated that MgH$_2$/30 wt.\% TiMn$_{1.5}$ system can achieve complete absorption/desorption processes in 20 min at 250–300 °C with rather poor hydrogen storage capacity (4.3 wt. \% $H_2$). MgH$_2$/Ti$_{0.4}$Cr$_{0.15}$Mn$_{0.15}$V$_{0.3}$ composite system was able to discharge its storage capacity (5.7 wt. \% $H_2$) within 30 min at 100 °C.

The crystal structure related to 50 h sample obtained after the dehydrogenation test was examined by XRD. The sample revealed sharp Bragg peaks corresponding to hcp-Mg (PDF file #00-004-0770) coexisted with fine hcp-TiMn$_2$ (PDF file #01-071-9712) particles having broad Bragg peaks patterns, as shown in Fig. 6(f). Neither reacted nor intermediate phases could be detected, implying the absent of any undesirable reactions during the dehydrogenation process.

Cyclability of hydrogen absorption/desorption. Measuring the cycle-life-time, which reflects the capability and performance of synthesized nanocomposite MgH$_2$/5 wt.\% TiMn$_2$ powders for achieving continuous hydrogenation/dehydrogenation cycles was investigated. For comparison, the cycle-life-time test of pure MgH$_2$ powders obtained after 200 h of RBM time (before doping with TiMn$_2$ powders) was conducted and the results are displayed in Fig. 7(a). To ensure the cyclic continuity and to improve the of hydrogenation/dehydrogenation kinetics of pure MgH$_2$, the test was achieved at high temperature (300 °C). The pressures used for hydrogenation/dehydrogenation processes were 8/0.2 bar, respectively.

However, the as-prepared nanocrystalline MgH$_2$ powders possessed excellent hydrogen storage capacity of about 6.9 wt. \% (Fig. 7(a)), the powders failed to maintain their capacity. This is indicated by a monotonic degradation on the storage capacity upon increasing the cycle-life-time, reaching a lower value (6.3 wt. \% $H_2$) after only 75 h, as shown in Figs 7(a) and 8(a). When the powders were subjected to longer cycle-life-time in the range between 75 – 275 h (Figs 7(a) and 8(a)), the hydrogen storage capacity was severely degraded to reach a lower value of 5.4 wt. \% $H_2$ after 200 h (Fig. 8(a)). Toward the end of the test (275 h), the MgH$_2$ powders were disable to store more than 5.24 wt. \% $H_2$, as elucidated in Fig. 7(a). The tendency of nanocomposite MgH$_2$/5 wt.\% TiMn$_2$ powders to manifest hydrogen storage degradation was almost absent even after 1400 h of continuous hydrogen uptake/discharge, as presented in Figs 7(a) and 8(b).

In order to realize the reasons responsible for such severe degradation, the morphology of the material powders obtained after 275 h of cycle-life-time (Mg powders) were examined by FE-SEM technique. Noticeably, the metallic powders after this test tended to agglomerate and composite larger particles with average size of 13μm in diameter, as presented in Fig. 8(c). This agglomeration behavior was a result of the continuous applications of rather high hydrogen pressure on the powders at a high temperature (300 °C). Moreover, the topology of the...
The powder was solid, where the pores facilitating the hydrogen diffusion into/out the powder were absent (Fig. 8(c)). The absence of favorable pores and cavities degrades the performance of MgH₂ powders to maintain their hydrogen storage capacity. Unluckily, and according to the application of high temperature and pleasure the metallic Mg grains revealed severe grain growth with an average diameter of 80 nm, as shown in the STEM-dark field image (DFI) in Fig. 8(e).

The beneficial effect of doping nanocrystalline MgH₂ with 5 wt. % TiMn₂ powders on the hydrogenation/dehydrogenation cyclability conducted at 250 °C under hydrogenation/dehydrogenation pressure of 8/0.2 bar is clearly realized in Fig. 7(b). The nanocomposite powders possessed very high performance for achieving 1400 cycle-life-time (~1000 continuous cycles) without severe degradation, as displayed in Fig. 7(b). During the first 400 h, the sample maintained its original hydrogen storage capacity (5.4 wt. %), that was slightly decreased to about 5 wt. % H₂ after 900 h, as shown in Fig. 7(b). Further unseasoned decreasing can be seen after 1200 h when the hydrogen storage capacity dropped a little to the level of 4.95 wt. % H₂, as presented in Fig. 7(b). The last 200 h of the cycle-life-time test is individually displayed in Fig. 8(b). During this last stage of the test, the nanocomposite powders maintained their hydrogen storage capacity at almost a constant value of 4.95 wt. %, as elucidated in Fig. 8(b). Moreover, the kinetics of hydrogenation/dehydrogenation processes remaining constant with no obvious failure or decay.

Since milling MgH₂ with TiMn₂ powders did not lead to the formation of any reacted intermediate phase (Fig. 6(c) and (f)) responsible for enhancing the kinetics and performance of charging/discharging hydrogen cyclability, the justification behind such obvious improvements can be related to a morphological reason. Figure 8(d) presents the FE-SEM of nanocomposite MgH₂/5 wt.% TiMn₂ powders after completion of 1000 hydrogenation/dehydrogenation cycles. The aggregated powders consists of ultrafine MgH₂/TiMn₂ particles,
ranging in sizes between 0.22 μm to less than 0.5 μm in diameter, as shown in Fig. 8(d). In addition, TiMn₂ fine particles (~5 nm in diameter) maintained their tendency for adhering onto the MgH₂ surface of powders, even after 1400 h of cycle-life-time, as shown in the STEM-DFI presented in Fig. 8(f). Comparing the STEM-DFI presented in Fig. 8(d) with that one for nanocomposite powders before conducting the cycle life-time test (Fig. 2(m–p)) led us to confirm the absence of any undesired grain growth for both MgH₂ grain matrix and TiMn₂ particles. The limited grain growth seen in MgH₂ grains is probably attributed to the distribution of TiMn₂ particles (grain-growth inhibitors) into the MgH₂ matrix. It was pointed out by Yao et al.57 that hydrogen diffusion is much faster when Mg grains were in the nanoscale level.

Based on these results, it can be concluded that one advantage of using hard intermetallic compounds such as TiMn₂, refractory metal carbides and amorphous alloys for improving the kinetics behavior and cycle-life-time of MgH₂ powders is related to the tendency of these hard particles to surround MgH₂ grains leading to block their attitude of growing at high temperature and pressure. Another advantage of TiMn₂ intermetallic compound additives with their nano-dimensional particle size is the improvement of the cyclability performance for MgH₂ system. This present system can be considered as one of the most reliable performance system among the most well known system, exemplified by MgH₂-5 wt. % of Ti₅₋₆Cr₄₋₅Mnₖ₋₇V₀₃ (73 cycles, 290 °C)⁵⁶, TiMn₂ (100 cycles, 300 °C)⁵⁵, VTiCr (100 cycles, 300 °C)⁵⁵, FeTi (500 cycles, 200 °C)⁵⁸, metallic glassy of Zr₇₀Ni₂₀Pd₁₀ (100 cycles, 200 °C), 10 wt. % of big-cube Zr₂Ni (2546 cycles, 250 °C)²¹, ZrNi₅ (600 cycles, 275 °C)⁴¹, 5Ni/5Nb₂O₅ (180 cycles, 250 °C)⁵⁹, and 5TiC/5Fe-12Cr (530 cycles, 275 °C)²⁵.

Integrated hydrogen storage system for fuel cell Applications. Our present work have two objectives; the first is focused on synthesizing and characterizing a nanocomposite MgH₂/5 wt. % TiMn₂ powders. The FE-SEM micrographs of the powders obtained after completion the cyclic hydrogenation tests displayed in (a,b) are shown in (c,d), respectively. STEM-DFIs of the powders obtained after completion 275 h cycle-life-time (MgH₂ powders) and 1400 h (MgH₂/5 wt.% TiMn₂ powders) are shown in (e,f), respectively.

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Figure 8. Influence of cycle-life-time and TiMn₂ additives on the morphological characteristics of the powders. The cycle-life-time of the last 200 h for (a) nanocrystalline MgH₂ powders, and (b) nanocomposite MgH₂/5 wt.% TiMn₂ powders. The FE-SEM micrographs of the powders obtained after completion the cyclic hydrogenation tests displayed in (a,b) are shown in (c,d), respectively. STEM-DFIs of the powders obtained after completion 275 h cycle-life-time (MgH₂ powders) and 1400 h (MgH₂/5 wt.% TiMn₂ powders) are shown in (e,f), respectively.
The tank was then mounted on a temperature controlled hotplate (Fig. 9(c)). The hydrogen storage tank was connected to a 40 W/4.5 A proton exchange membrane (PEM) fuel cell (Fig. 9(a)) through a pipeline allowing the released hydrogen gas from the tank to be passed to the PEM-fuel cell, as shown in Fig. 9(a). The converted electrical energy required to charge the battery of a cell phone device through a 5 V voltage regulator (Fig. 9(a)). The PEM-fuel cell system is controlled and operated with a software where the data output corresponding to hydrogen flow rate, voltage and current were obtained and stored.

Before starting the fuel-cell testing, the powders were subjected to pressure-composition-temperature (PCT) analysis to ensure the possibility of dehydrogenation process at a low temperature (180 °C) and to investigate the pressure required to achieve the decomposition process. MgH₂/5 wt.% TiMn₂ powders obtained after 50 h of milling. In the PCT experiment, the sample was firstly activated at high temperature (350 °C) that corresponding to about 35 bar of H₂ for 12 h. The powders were then charged with hydrogen at 180 °C, where the corresponding pressure reached to 0.4 bar, as shown in Fig. 10(a). Under this low temperature (180 °C) the fully charged powders tended to discharge their hydrogen storage capacity (5.43 wt.%) at 0.3 bar, as displayed in Fig. 10a. The PCT experiments were repeated 5 times under the same conditions to ensure the reproducibility of the results. It is worth to be mentioned that the presence of minimal pressure-hysteresis gap between the pressure of absorption and desorption (~0.1 bar) indicates the powders' ability to achieve long term of cycle-life-time without failure, as was previously presented (Fig. 7b). Moreover, and in part to ensure our results, new set of kinetic measurements were conducted at 180 °C for both hydrogenation and dehydrogenation reactions taking place under 8–10 bar/0.2 bar (Fig. 10(b)). The possibility of achieving both reactions at 180 °C was approved, however, both reactions showed rather slow kinetics when compared with the results obtained at 250 °C (Fig. 6(a) and (d)).

Prior to fuel cell experiments processing start, the nanocomposite MgH₂/5 wt.% TiMn₂ powders were activated at 350 °C by charging/discharging hydrogen gas atmosphere at 30 bar/1 bar, respectively for 10 continuous cycles to obtain MgO-surface free nanocomposite powders. A continuous hydrogen gas flow pressurized at 10 bar was introduced at 180 °C to the Ti-tank containing the powders. The Ti-tank was then kept under this temperature (180 °C), where the readings shown by the hydrogen pressure gauge connected to the tank (Fig. 10(c)) was monitored and recorded every 1 min. These readings were used to construct a relationship between the time required for the powders placed to be charged with hydrogen, as shown by the open red symbols presented in Fig. 10(c). Completion of the absorption process was realized when the hydrogen was completely absorbed by the powders and the tank's pressure dropped to the atmospheric level (Fig. 10(c)). The hydrogen gas cylinder was
then removed and disconnected from the tank, where the tank was kept under this 180 °C for almost 40 min. In order to construct a relation between the time required to release hydrogen gas stored in the powder in Ti tank, the readings of the pressure gauge was recorded every 1 min. The monotonical increase of hydrogen pressure inside the closed Ti-tank implies a gradual hydrogen release from the powders, as elucidated by the closed blue symbols shown in Fig. 10(c). The time required to release hydrogen from MgH₂ powders at 180 °C was about 25 min, as shown in Fig. 10(c). At this temperature, we started to open the valve of the tank shown connected to the PEM-fuel cell system (Fig. 9(a)) with nearly constant hydrogen flow of 175 ml/min (Fig. 10(d)).

Figure 10(d) displays the hydrogen gas flow transported into the PEM-fuel cell. The hydrogen released from the nanocomposite powders was used to feed the PEM-fuel cell through a pipeline, as shown in Fig. 9(a). During the battery charging time (28 min) the hydrogen gas released from the hydrogen storage tank at 180 °C passed delivered into the PEM-fuel cell at constant rate of 175 ml/min, as shown in Fig. 10(d). Providing the fuel cell with such a constant rate of hydrogen gas flow leads to generate constant values of voltage (7 V) and current (1 A), as displayed in Fig. 10(e) and (f), respectively.

In summary, we have synthesized nanocomposite MgH₂/5 wt.% TiMn₂ system with advanced storage capacity (~5.3 wt.% H₂), hydrogenation/dehydrogenation kinetics and long hydrogen charging/discharging cyclability shown at rather low temperature (250 °C). The new nanocomposite powders were charged into a self-made Ti-tank and employed as a source of hydrogen required to charge a battery of a cell-phone device through a commercial proton exchange membrane (PEM) fuel cell.

**Methods**

**Preparation of MgH₂ powders.** Elemental Mg metal powders (~80 μm, 99.8% provided by Alfa Aesar - USA), and hydrogen gas (99.999%) were used as starting materials. An amount of 5 g Mg was balanced inside a He gas atmosphere (99.99%) - glove box (UNILAB Pro Glove Box Workstation, mBRAUN, Germany) and sealed together with fifty FeCr balls into a hardened steel vial (150 ml in volume), using a gas-temperature-monitoring system (GST; supplied by evico magnetic, Germany). The ball-to-powder weight ratio was maintained at 40:1. The vial was then evacuated to the level of 10⁻³ bar before introducing H₂ gas to fill the vial with a pressure of 50 bar. The reactive ball milling (RBM) process was carried out at room temperature, using a high-energy ball mill (Planetary Ball Mill PM 400 provided by RETSCH, Germany). The RBM process was interrupted after selected milling time (3, 6, 12.5, 25, 50, 100, and 200 h) where the vial was opened inside the glove box to take a small amount (~300 mg) of the milled powders for different analysis. Then, the RBM process was resumed, using the same operational conditions shown above.
The as-synthesized MgH₂ powders were then mixed in the glove with the 5 wt. % of TiMn₂ shots, using an agate mortar and pestle. Five gram of the mixed powders for each composite system were charged together with fifty Cr-steel balls into the hardened steel vial and sealed under He gas atmosphere. The vial was then filled with 50 bar of hydrogen gas atmosphere and mounted on the high-energy ball mill. The milling process was interrupted after selected time (3, 6, 12.5, 25, 37.5 and 50 h) and the powders obtained after an individual milling time were completely discharged into 8 Pyrex vials for different analysis. Then, new MgH₂/5 wt% TiMn₂ mixed powders were charged again and ball milled under the same milling conditions. The contamination contents of Fe and Cr of the powders obtained after 50 h of ball milling were 1.19 and 0.32 wt. %, respectively.

**Powder characterizations.** XRD AND HRTEM. The crystal structure of all samples was investigated by XRD with CuKα radiation, using 9kW Intelligent X-ray diffraction system, provided by Smartlab-Rigaku, Japan. The local structure of the synthesized material powders was studied by 200 kV-field emission high resolution transmission electron microscopy/scanning transmission electron microscopy (HRTEM/STEM, supplied by JEOL-2100F, Japan), which is equipped with Energy-dispersive X-ray spectroscopy (EDS) supplied by Oxford Instruments, UK. In addition to the elemental analysis achieved by EDS approach, we employed ICP technique to get the elemental analysis by a chemical analytical approach.

Thermal stability. Shimadzu Thermal Analysis System/TA-60WS-Japan, using differential scanning calorimeter (DSC) was employed to investigate the decomposition temperatures of MgH₂ powders with a heating rate of 20°C/min. The activation energy for of the powders obtained after different RBM time were investigated, using Arrhenius approach with different heating rates (5, 10, 20, 30, 40°C/min).

The hydrogenation/dehydrogenation behaviors. The hydrogen absorption/desorption kinetics were investigated via Sievert's method⁶⁹–⁷², using PCTPro-2000, provided by Setaram Instrumentation, France, under hydrogen gas pressure in the range between 200 mbar (for dehydrogenation) to 10 bar (for hydrogenation). The samples were examined at different temperatures of 50, 100, 250, and 275°C. In the PCT measurements, the dosed pressure in absorption/desorption was gradually increased/decreased by 1000 mbar until equilibrium pressure reached to 13000 and 50 mbar, respectively. The PCT absorption/desorption kinetics were fitted in real-time by the HTPSwin software, to determine the sufficient equilibration time (the next point would start when the uptake had relaxed just 99% to asymptote). A minimum time of 30 minutes per equilibrium point and a maximum timeout of 300 minutes were set for each kinetic step in both the absorption and desorption isotherms.

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Author Contributions
M. Sherif El-Eskandarany generated the idea, designed the experiments, carried out the TEM and XRD analysis, and contributed on SEM and DSC analysis. He designed the hydrogen storage tank and tackled the experiments related to charge the battery of a cell-phone device together with F. Aldakheel and A. Alkandary. He wrote the manuscript after discussions the results with E. Shaban, F. Aldakheel, A. Alkandary, M. Behbehani, and M. Al-Saidi. E. Shaban shared M. Sherif El-Eskandarany the task related to sample preparations, tackled the kinetics and cycle-life-time measurements and shared M. Sherif El-Eskandarany XRD experiments and analysis. F. Aldakheel shared with A. Alkandary a part of sample preparations, using reactive ball milling. They shared also in the structural, morphological analysis, as well as kinetics measurements. M. Behbehani tackled the thermal analysis task together with M. Al-Saidi, using DSC technique. All experiments and analysis were done at the Nanotechnology Laboratory, Energy and Building Research Center, Kuwait Institute for Scientific Research.

Additional Information
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