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# Modelling a kinetic deviation of the magnesium hydrogenation reaction at conditions close to equilibrium

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## ARTICLE INFO

### Article history:

Received 29 December 2018

Received in revised form

2 March 2019

Accepted 3 April 2019

Available online 3 May 2019

### Keywords:

Magnesium hydrogenation

Thermochemical energy storage

Shrinking-core model

Knudsen diffusion

PCT equation

Langmuir

## ABSTRACT

A model has been derived for the magnesium hydrogenation reaction at conditions close to equilibrium. The reaction mechanism involves an adsorption element, where the model is an extension of the Langmuir adsorption model. The concept of site availability ( $\sigma_s$ ) is introduced, whereby it has the capability to reduce the reaction rate. To improve representation of  $\sigma_s$ , an adaptable semi-empirical equation has been developed. Supplement to the surface reaction, a rate equation has been derived considering resistance effects. It was found that close to equilibrium, surface resistance dominated the reaction.

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## Introduction

Research efforts to store surplus energy for concentrated solar power (CSP) plants have looked beyond latent heat technologies, to a potentially cheaper and more effective technology, thermochemical energy storage (TCES). [1,2], A potential TCES system is based on utilising the enthalpy of reaction of a reversible metal hydride reaction, where metal hydrides are selected based on several governing factors, such as operating temperature, enthalpy of reaction, plateau pressure and cost. Many hydrides have been identified for a TCES system, in part due to their versatility, for example

combining magnesium with a transition metal such as nickel or iron enables the tuning of the operating temperatures at a reasonable cost [3]. In addition, adding a destabilisation agent, such as In or Ge, enables thermodynamic modification for further tuning within a TCES system if needed [4,5]. However, this paper focuses on magnesium hydride, due to the simple case of a binary system and the suitability for a thermal oil circuit [6]. A key challenge with the design of this system are the reaction kinetics, where sufficient understanding is needed to facilitate an efficient energy store [2]. As such, this paper focuses on the kinetics, specifically the hydrogenation reaction.

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<https://doi.org/10.1016/j.ijhydene.2019.04.036>

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Analysis of the literature reveals the intricate nature of the Mg hydrogenation reaction, whereby the reaction is sensitive to temperature, particle size, gas pressure, irregularities in particle structure, concentration, and diffusion effects [7–9]. A popular choice are the nucleation-growth-impingement models known as the Johnson–Mehl–Avrami–Kolmogorov (JMAK) [10]. This model is based on the concept of an extended volume. However, there are drawbacks with this concept, as in practice the reactive surface area would decrease as more reactant depletes.

Rate equations developed for metal hydrogenation that include temperature dependency are commonly governed by an Arrhenius term, and this is widely accepted, however the pressure dependency is still being debated. These rate equations can be written in many forms, such as dimensionless [11], mass concentration [12] or mole concentration [13], of which they commonly resemble a first order equation based on the original work of Mayer and Groll [14]. Chaise et al. also attempted to fit their data using an Arrhenius term and other reaction mechanisms [15].

A pressure term typically contains the gas pressure  $P$  and equilibrium pressure  $P_{eq}$ . These range from a straight pressure

### Site availability model (SAM)

The SAM contains elements of the Langmuir's adsorption model (LAM) [19], and a shrinking core approach [20], whereby the overall reaction progresses through steps, shown below. Fig. 1 illustrates the mechanism.

- |   |  |
|---|--|
| 1 Transport of $H_2$ to Mg surface                                |  |
| a. Initial permeation through the porous bed (external diffusion) |  |
| b. Diffusion through product layer (internal diffusion)           |  |
| 2 Dissociation of $H_2$ at Mg surface:                            | $H_2 \rightarrow 2H$                   |
| 3 Reaction via adsorption and depletion of core:                  | $2H + 2S \xrightarrow{k_2} 2H \cdot S$ |

The simple case whereby step 3 is assumed the rate-determining step is shown in Adsorption reaction. Inclusion of Step 1 is discussed in section resistances. The SAM is compared with LAM below:

Langmuir (Forward rate)	Site Availability Model (SAM)
The rate of attachment to an active site (S) is proportional to rate of collisions that H make with active sites	Same
An active site is an unoccupied site.	An active site is a site that is unoccupied and available.
The collision rate to an active site is proportional to the <b>pressure over the surface (<math>P_H</math>)</b> .	The collision rate to an active site is proportional to the <b>site availability (<math>\sigma_s</math>)</b> .
As H only adsorbs on active sites, it is proportional to concentration of active/vacant sites. ( $C_v$ )	Same
$r_{H \cdot S} = k''P_H C_v$	$r_{H \cdot S} = k''\sigma_s C_v$

driving force (conventional Langmuir) [16,17], or via a pressure ratio. It has been observed that modelling  $P_{eq}$  near to the equilibrium isotherm yields a closer result to experimental data [13]. To model the equilibrium pressure, approaches have varied from entirely empirical equations, such as a partition of two separate exponential functions [13], to semi-empirical functions which model the overall shape with mathematical functions and use the Van't Hoff equation to determine the temperature dependence [12].

Switching attention to the overall format of the pressure ratio, Mayer et al. modelled a  $LaNi_{4.7}Al_{0.3}$  bed using the term  $\ln(P/P_{eq})$  [14]. Herbrigg et al. modelled a hydride-graphite composite bed using  $(P - P_{eq})/P_{eq}$  [13]. This term originated from Ron (1999), who experimented with different pressure terms [18]. A pressing question is how the pressure dependency is interpreted, whether that be by conventional Langmuir assumptions or perhaps via a new mechanistic concept, or, not at all.

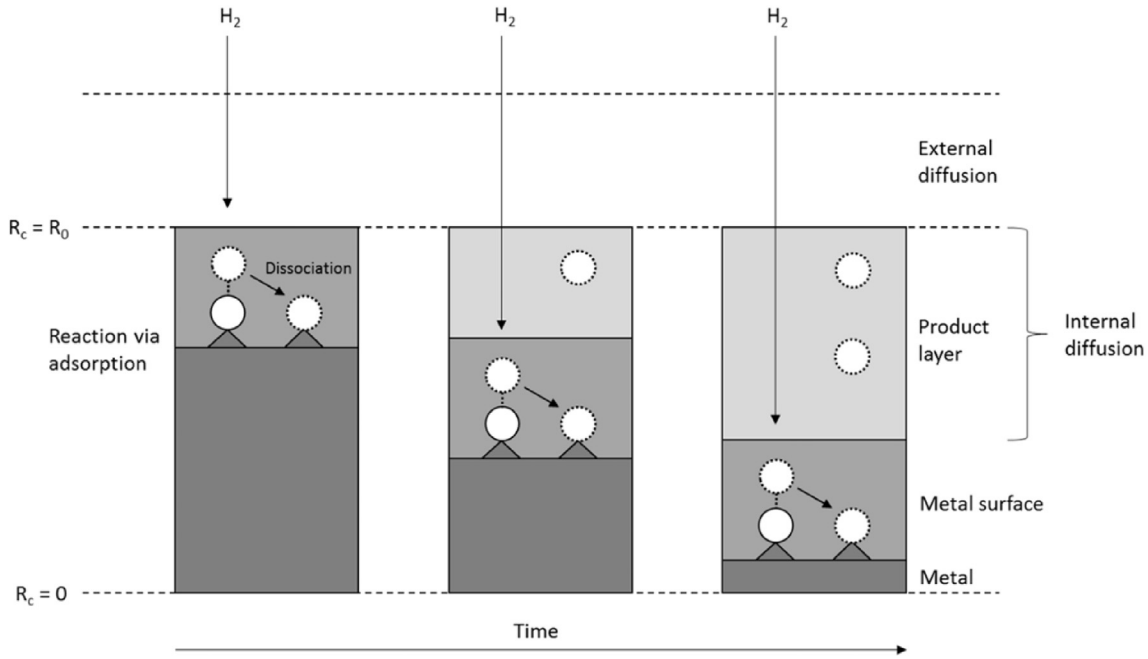
As such, this paper presents the findings of an alternative derived model that encompasses an expansion of Langmuir, temperature & pressure dependency, and diffusion & surface effects. In addition, this model is compared to experimental data.

### Site de-activation

For  $H_2$  sorption to a metal to occur, the gas pressure ( $P$ ) must be greater than the equilibrium pressure ( $P_{eq}$ ). As the exothermic reaction invokes a rise in temperature, which causes the equilibrium pressure  $P_{eq}$  to rise and approach  $P$ ; the thermodynamic driving force is reduced and so must the reaction rate. In effect,  $P_{eq}$  can be considered the thermodynamic boundary.

Expanding on site theory, if the reaction rate must fall, the rate of a successful collision to an active site must decrease. Thus, it is assumed that the rise in temperature and subsequently the reduction in pressure driving force, must temporarily de-activate a site, even if that site is unoccupied/unavailable, to reduce the amount of successful collisions, and therefore reduce the reaction rate.

Within SAM, this theory is represented by the ratio of available sites to unavailable sites. As temperature is linked to the equilibrium pressure, then across each time step,  $P_{eq}$  indicates the unavailable sites. At the next time step, the change in  $(P - P_{eq})$  will be an indication of sites that have become inactive across that time step. Therefore, the ratio that sites



**Fig. 1 – Proposed general mechanism of hydrogenation kinetics for Magnesium. This is a 2D cutout section of a sphere.  $R_0$  is the radius of the sphere,  $R_c$  is the radius of shrinking metal reactant.**

that are available to ones that are unavailable can be estimated by the site availability,  $\sigma_s$ .

$$\sigma_s \approx \frac{S_{av}}{S_{Unav}} \approx \frac{P - P_{eq}}{P_{eq}} \quad (1)$$

#### Isotherm representation

A semi-empirical isotherm equation has been developed to represent metal hydrogenation systems. The equilibrium pressure is calculated by Eq. (2).

$$P_{eq} = P_N \left\{ 1 - \frac{1}{y_1} \ln \left( \frac{1 - \theta}{\theta} \right) \right\} \quad (2)$$

$$P_N = \exp \left( \frac{\Delta H}{RT} - \frac{\Delta S}{R} \right) \quad (3)$$

$P_N$  is the normalisation pressure. For a typical PCT, this point would be where  $\theta = 0.5$ , often this position is the plateau pressure. Note, Eq. (2) does not work at  $\theta = 0$  or at  $\theta = 1$ . The slope parameter  $y_1$  can be constant or a variable, such as  $f(\theta)$ , where increasing  $y_1$  increases the slope. Eq. (2) can be modified for a system that exhibits two separate growth phases:

$$P_{eq} = P_N \left\{ 1 - \left( \frac{1}{y_1} \ln \left( \frac{1 - \theta}{y_2 \theta} \right) \right)^{\frac{1}{2}} \right\} \quad (4)$$

where ' $y_2$ ' is a constant that can shift the location of the normalisation point.

#### Adsorption reaction

If step 3 is considered the rate-determining step, the rate of forming  $H \cdot S$  is,

$$r_{H \cdot S} = k'' \sigma_s C_v \quad (5)$$

Units:  $k'' = m \ s^{-1}$  &  $r'' = mol \ m^{-2} \ s^{-1}$ . Assuming a site balance where the total sites is the sum of vacant and filled sites ( $C_T = C_v + C_{H \cdot S}$ ), and incorporating the normalised hydride fraction  $\theta = C_{H \cdot S} / C_T$ , gives a first order rate equation,

$$r_{H \cdot S} = k'' \sigma_s C_T (1 - \theta) \quad (6)$$

This is an ideal scenario, where the sample can reach full capacity. If  $C_T$  is,

$$C_T = C_{ref} x_m \quad (7)$$

where  $C_{ref}$  is the reference concentration (taken as 7.66 wt%) and  $x_m$  is the maximum capacity fraction at operating conditions. Including the hydride fraction (a helpful performance measurement),  $x = \theta x_m$  the rate equation can be modified,

$$r_{H \cdot S} = k'' \sigma_s C_{ref} (x_m - x) \quad (8)$$

#### Resistances

The model now includes surface and diffusion resistance. The SAM neglects the bulk movement of hydrogen through the bed; assumes a constant bed porosity ( $\epsilon_b$ ) and the expansion of the metal hydride forms surface cracks, grain boundaries and defects in the structure allowing the permeation of molecular hydrogen direct to the inner metal surface. This is represented by Knudsen diffusion [21]. If the reaction of  $H + S \rightarrow H \cdot S$  is considered, where the rate equation is based on gaseous atomic hydrogen which takes part in the reaction, the surface resistance balance for a spherical pellet is,

$$-\frac{dn_H}{dt} = k'' C_{Hc} \sigma_s \cdot 4\pi R_c^2 \quad (9)$$

Where  $C_{Hc}$  is the concentration of H at the core interface. Converted into effective volume terms and rearranging:

$$C_{Hc} = -\frac{1}{4\pi R_c^2} \frac{RSA}{k_e \sigma_s} \frac{dn_H}{dt} \quad (10)$$

where  $k_e = k^* RSA$  and RSA is the reactive surface area. If RSA is based on a sphere,

$$RSA = \frac{3R_c^2}{R_0^3} \quad (11)$$

The radius of the core ( $R_c$ ) is related to the constant particle size ( $R_0$ ) by:

$$R_c = R_0 \left( \frac{C_H}{C_T} \right)^{\frac{1}{3}} \quad (12)$$

For the diffusion resistance balance:

$$-\frac{dn_{H_2}}{dt} = 4\pi R^2 J_{H_2} = 4\pi R^2 D_e \frac{dC_{H_2}}{dR} \quad (13)$$

Applying  $\left( dC_{H_2} = \frac{1}{2} dC_H \right)$ , the pseudo steady-state assumption and integrating [20].

$$-\frac{dn_H}{dt} \frac{RSA}{4\pi} \left[ \frac{1}{R_0} - \frac{1}{R_c} \right] = D_e RSA (C_{Hc} - C_H) \quad (14)$$

Combining the resistances, (Eqs. 10 & 14) gives,

$$-r_H = -\frac{dn_H}{dt} \frac{RSA}{4\pi R_0^2} = \frac{C_H RSA}{\frac{R_0^2 RSA}{R_c^2 k_e \sigma_s} + \frac{R_0}{D_e} \left( \frac{R_0 - R_c}{R_c} \right)} \quad (15)$$

The surface resistance is represented by  $R_0^2 RSA / R_c^2 k_e \sigma_s$  and the diffusion resistance is  $R_0 / D_e R_c (R_0 - R_c)$ . Eq. (8) is based on the concentration of atomic hydrogen in the solid state, whereas Eq. (15) is based on the concentration of atomic hydrogen in the gaseous phase, which takes part in the reaction. E.g. if the initial gas pressure was 20 bar and the final pressure was 18 bar, Eq. (15) is based on the moles corresponding to 2 bar. Therefore, a mole balance can be used where if the hydrogen is not bound, it must be in the gas phase. This enables calculation of the reacted fraction.

## Experiment

Experiments were performed using a Sievert's apparatus, where high purity hydrogen (99.9999%) was used. The sample was atomised Mg powder (26  $\mu m$ ) by SFM - FluorsidGroup Company. The bulk density was 1000 kg/m<sup>3</sup>, thus giving an assumed porosity of  $(1 - 1000/1740) \approx 0.4$ . Kinetic runs were at several constant initial temperatures and when the initial overpressure  $([P - P_{Pl}]/P_{Pl}) = 1$  ( $P_{Pl}$  is the plateau pressure). We assume that when  $([P - P_{Pl}]/P_{Pl}) = 0$ , it is at equilibrium, when the overpressure is  $\leq 1$ , it is close to equilibrium, and when  $> 1$ , we are far from equilibrium for the temperature range 330–360 °C.

The equivalent volume method was used [22], with a manifold volume of 62.9 cm<sup>3</sup> and an effective sample cell volume of 11–13 cm<sup>3</sup> depending on the temperature. The sample temperature was measured with a thermocouple in the sample.

Each sample was activated at 360 °C and 40 bara over 4 cycles. These cycles were typically 2–3 days each. For the first sample, the capacity successfully recovered by the fourth cycle and underwent experiments in triplicates at 330 °C and 360 °C, where the total number of cycles was kept low at

approximately 10 cycles, in order to minimise the possibility of sintering. To further minimise sintering, both samples were always kept in the hydrogenated state overnight, and once fully dehydrogenated (all dehydrogenation was at 360 °C and took approximately 1 h), hydrogenated again within an hour. The second sample was activated under identical conditions as the first sample, whereby capacity was recovered by the fourth cycle. Experiments of sample two at 345 °C were also performed in triplicates.

## FEM model

### Geometry

The sample sat at the base of the sample holder hole where a thermocouple was placed 15 mm from the edge of the sample. It was observed to remain relatively steady at the set point. Thus, a constant temperature boundary condition was assumed. Fig. 2 illustrates this.

### Sample dimensions

$L_2$  was assumed to be  $L_2 = 0.2L_1$ . The sample mass was 0.2 g, and assuming the magnesium's maximum capacity,  $wt_m = 0.0766$  the reference mass of hydrogen at reaction completion ( $m_{ref}$ ) is given by

$$m_{ref} = \frac{wt_m m_{Mg}}{1 - wt_m} \quad (16)$$

To account for the changing density, an assumed average porous matrix density ( $\rho_p$ ) was used. A simple average density of Mg and MgH<sub>2</sub> gave a  $\rho_p = 1590$  kg/m<sup>3</sup>. This created a pseudo value for  $L_1$  and  $L_2$ , equating to an effective sample domain volume ( $V_{de}$ ).  $V_{de}$  is calculated by considering the average maximum uptake of hydrogen fraction (constant) at operating conditions  $x_m^{av}$  (i.e. the experimental end capacity).

$$V_{de} = \frac{m_{Mg} + m_{ref} x_m^{av}}{\rho_p (1 - \epsilon_b)} \quad (17)$$

### Balance equations

To model the experimental data, the finite element method was used and achieved by using COMSOL Multiphysics 5.3a.

### Mole balance

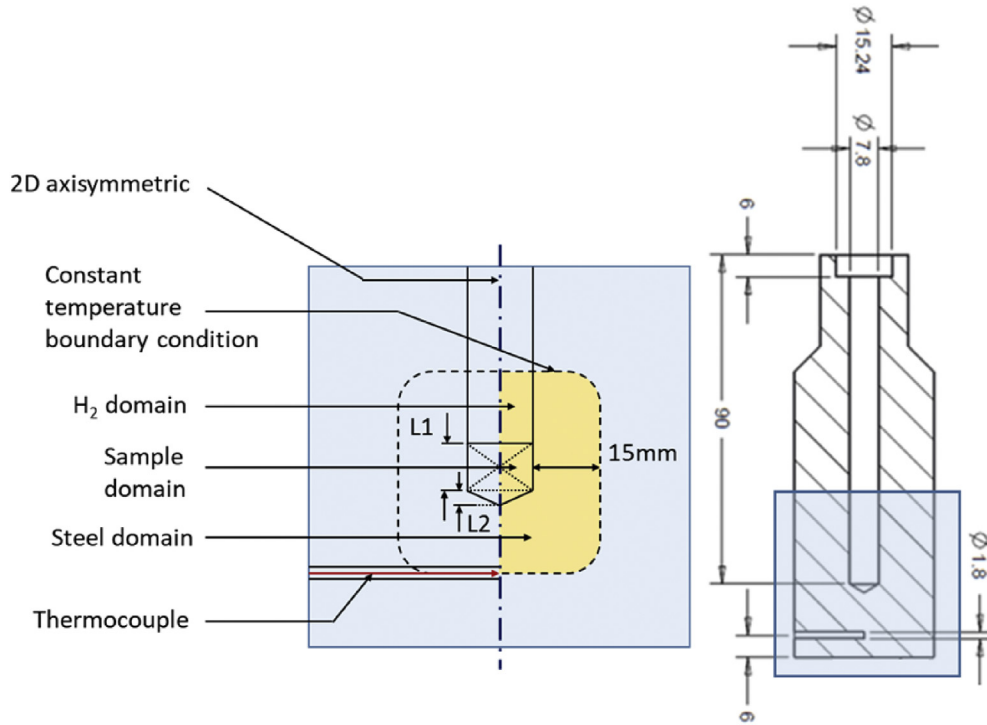
External diffusion effects are neglected, and the balance is based on the gas phase H concentration taking part in the reaction considering the bed void volume.

$$-\frac{\partial C_H}{\partial t} = r_H \quad (18)$$

The Arrhenius equation calculated the effective specific rate constant  $k_e$ , with activation energy  $E_e$  and frequency factor  $A_e$ .

$$k_e = A_e \exp(-E_e/RT) \quad (19)$$

As the maximum capacity ( $x_m$ ) varies with temperature, an empirical relation was used, with  $a_1$  and  $a_2$  being constants. The equation for  $x_m$  changes for each operating condition.



**Fig. 2 – (left) 2D axisymmetric geometry (yellow zone) (right) Schematic of sample holder, (all dimensions in mm and material of construction 316ss).**

$$x_m = a_1 T + a_2 \quad (20)$$

#### Energy balance

Conductive heat transfer was expected to dominate, thus advective heat transfer was neglected and there was thermal equilibrium between the porous matrix and hydrogen [12].

$$(\rho c_p)_e \frac{\partial T}{\partial t} - \nabla(\lambda_e \nabla \cdot T) = -r_H \Delta H \quad (21)$$

With  $\Delta H$  being the enthalpy of reaction. Considering a porous matrix and gas domain, the volumetric heat capacity  $(\rho c_p)_e$  and thermal conductivity  $\lambda_e$  were expressed in effective terms.

$$(\rho c_p)_e = (1 - \varepsilon_b) \rho_p c_{p_p} + \varepsilon_b \rho_{H_2} c_{p_{H_2}} \quad (22)$$

$$\lambda_e = (1 - \varepsilon_b) \lambda_p + \varepsilon_b \lambda_{H_2} \quad (23)$$

where  $c_{p_{H_2}}$  and  $\lambda_{H_2}$  were governed by an empirical equation which was a  $f(T, P)$ , using data from NIST database at the system pressure ranges [23].

$$\lambda_{H_2}(T[K], P[\text{bar}]) = 4.805E^{-4}T + 3.983E^{-2} + 4.344E^{-5}P \quad (24)$$

Uncertainty =  $\pm 2.698 \times 10^{-5}$  @ 1 bar,  $\pm 8.347 \times 10^{-5}$  @ 50 bar

$$c_{p_{H_2}}(T[K], P[\text{bar}]) = 1.599E^{-6}T^2 - 1.429E^{-3}T + 14.83 + 5.231E^{-4}P \quad (25)$$

Uncertainty =  $\pm 5.231 \times 10^{-4}$  @ 1 bar,  $\pm 2.926 \times 10^{-3}$  @ 50 bar.

The specific heat capacity ( $c_{p_p}$ ) and porous matrix thermal conductivity ( $\lambda_p$ ) were assumed to be constant.  $\lambda_p$  was selected

to reach the desired value of  $\lambda_e$ . At 300 K and 1 bar, the measured  $\lambda_e$  of  $MgH_2$  under air and argon was  $0.116$  and  $0.088 \text{ W m}^{-1} \text{ K}^{-1}$  respectively [24]. Together with the gas thermal conductivities, linear interpolation at the same conditions (but in  $H_2$ ) resulted in an  $\sim \lambda_e = 0.7 \text{ W m}^{-1} \text{ K}^{-1}$ . Even though  $\lambda_{H_2}$  is higher at operating conditions, this conservative estimate of  $\lambda_e$  was used.

## Results & discussion

### Isotherm modelling

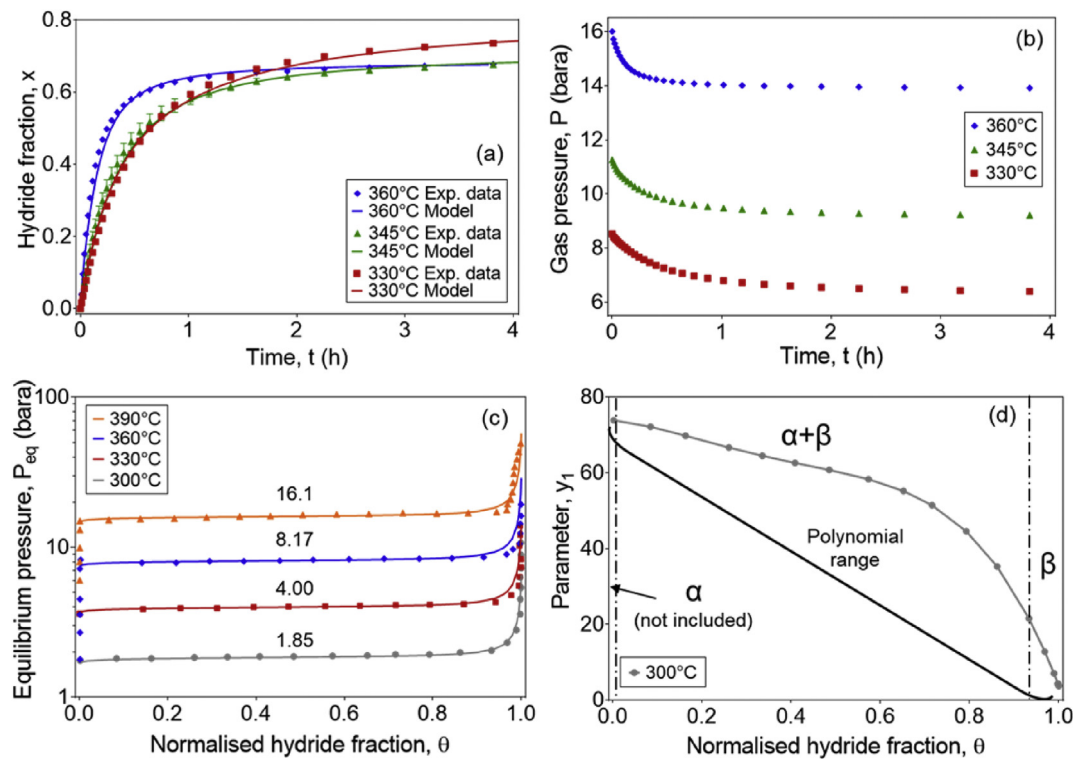
The calculated polynomial for  $y_1$  from experimental magnesium sorption PCTs between 300 and 390 °C [25] and used within COMSOL simulations is shown in Eq. (26). The paper calculated a  $\Delta H = -76.07 \text{ kJ (mol } H_2)^{-1} \pm 1.21$  and a  $\Delta S = -137.89 \text{ JK}^{-1} (\text{mol } H_2)^{-1} \pm 1.97$ . The values of  $\Delta H$  &  $\Delta S$  calculated by these authors are comparable to Refs. [26,27]. This equation sufficiently represents the plateau and  $\beta$  phase but does not include the  $\alpha$  phase. This is because the total  $\alpha$ -phase reaction time was less than 10 s, and thus could be omitted. The PCT is shown in Fig. 3, where  $y_1$  was regressed using data at 300 °C and assumed not to be a function of temperature.

$$y_1 = -245.09\theta^4 + 310.95\theta^3 - 123.74\theta^2 - 12.268\theta + 73.943 \quad (26)$$

### Kinetics

Using the equations outlined in previous sections, enabled the calculation of the hydride fraction as the reaction progressed, shown in Fig. 3. The calculated effective activation energy





**Fig. 3 – (a) Model results versus experimental uptake & initial pressure data at 1 overpressure. Averaged results of triplicates. (b) Experimental gas pressure data. (c) Equilibrium pressure fits of magnesium PCT's [25], (d) Regressed parameter  $y_1$  @ 300 °C.**

( $E_e = 172 \text{ kJ mol}^{-1}$ ,  $A = 4.5 \times 10^{11} \text{ s}^{-1}$ ) was within the error of one standard deviation to the values determined by DSC dehydrogenation under argon via Kissinger plots of the same magnesium batch ( $169 \pm 9 \text{ kJ (mol H}_2\text{)}^{-1}$  &  $A = 4.5 \times 10^{11} \text{ s}^{-1}$  (lower limit  $-1.8 \times 10^{11} \text{ s}^{-1}$  and upper limit  $+6.8 \times 10^{11}$ ) [28]. To calculate  $E_e$ , we started with the initial estimate of the experimental data. Then  $A_e$  was kept constant with  $E_e$  as a regression parameter within the COMSOL simulation. The reaction at 360 °C has progressed further compared to 330 °C and 345 °C (of which no reaction reached completion) and results indicate that lower temperatures result in higher capacities at conditions close to equilibrium. Altering the y-intercept of Eq. (20) is required with a change in initial temperature.

The gas pressure experimental data was directly used within the rate equations and shown in Fig. 3 for reproducible purposes. However, for a typical Sievert's apparatus kinetic experiment, if one plots  $P$  versus  $\theta$ , it is a straight line. Therefore, it is possible to calculate the gas pressure alongside  $\theta$  as the reaction progresses with a suitable expression.

#### Diffusion & surface resistance

Eq. (15) enables the resistances to be plotted independently. In the first case, the diffusion resistance is assumed to have a product layer porosity of 0.3 (expansion by 30%) with an assumed pore radius of 1 nm. In case two, the diffusion resistance is maximised by minimising the porosity and pore radius. Here, it is assumed a practical minimum of porosity = 0.1 and the pore radius = 0.5 nm ( $\approx 2 \times$  the diameter

of 1 hydrogen molecule) to give a maximum Knudsen diffusion of  $10^{-8} \text{ m}^2/\text{s}$ . From Fig. 4, it is apparent that the diffusion resistance does not exceed  $8000 \text{ s m}^{-1}$  for when the effective diffusion coefficient is maximised.

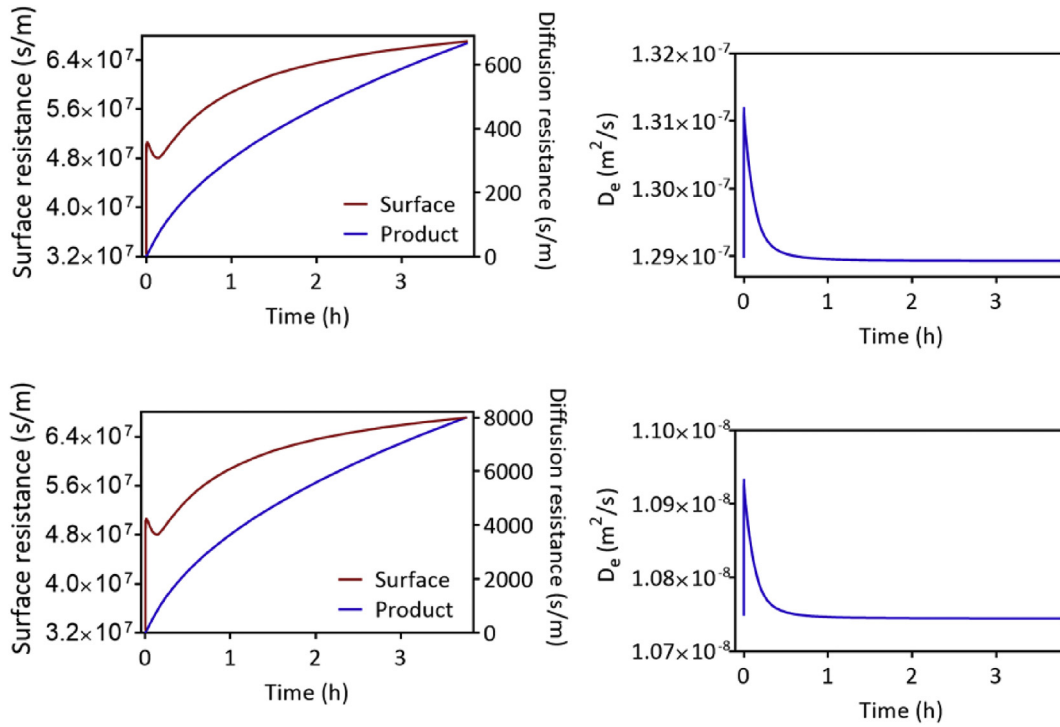
In comparison, the surface reaction resistance is in the order of  $10^7$ . In effect, a small diffusion resistance implies that hydrogen diffusing through the hydride has little effect on limiting the kinetics, if based on Knudsen diffusion. However, one cannot eliminate the possibility that diffusion resistance becomes influential at far from equilibrium operating conditions, where the additional overpressure could change the process in how the hydride forms, minimising cracks and forming an encasing shell, i.e. no longer entirely Knudsen diffusion.

#### Surface resistance

Consequently, if it is assumed that the diffusion resistance is negligible, Eq. (15) simplifies to Eq. (27):

$$-r_H = k_e \sigma_s C_H \left( \frac{R_c}{R_0} \right)^2 \quad (27)$$

Analysing Eq. (27) shows that the effective rate constant ( $k_e$ ) dominates in comparison to  $(R_c/R_0)^2$  and  $\sigma_s$ . As  $k_e = k' \text{RSA}$  where  $\text{RSA} = 3R_c^2/R_0^3$ , an increase in RSA would increase  $k_e$  and thus raise the reaction rate, which is achieved by a reduction in particle size ( $R_0$ ). This mathematically represents that reducing the particle size, is an effective method of increasing the reaction rate.



**Fig. 4 – The averaged volume resistances and effective molecular diffusion coefficient. Initial conditions at 360 °C & 16 bar ( $\sigma \approx 1$ ) over the duration of 4 h (Top)  $\varepsilon = 0.3$ , Pore radius = 1 nm. (Bottom)  $\varepsilon = 0.1$ , Pore radius = 0.3 nm. The temporary improvement of the surface resistance is due to site re-activation.**

If Eq. (27) is further simplified by substituting (12), this gives the result in terms of concentration only ((28)), and if expressed in terms of the normalised hydride fraction gives (29). This result indicates that if there is a sufficient reduction in the RSA, feasibly a high wt% material, where the rate determining step is a reaction occurring at a surface of a sphere at conditions close to equilibrium, the reaction order is 5/3. If in contrast, the material is low wt%, where there is not a significant reduction in  $R_c$ , thus  $R_c/R_0 \approx 1$ ; Eq. (27) simplifies to a first order rate Eq. (6) – similar to kinetics currently used for low wt% materials based on the work of Mayer and Groll [14] and popularised by Jemni & Nasrallah [12]. In effect, a low wt% material potentially contains a low total amount of available sites for hydrogen relative to the total surface, so the metal reactant surface does not shrink. In contrast, a high wt% material contains a high total amount of available sites, resulting in a shrinking metal reactant and therefore, the analysis indicates inclusion of the reacting surface is important when deriving a suitable rate equation for a high wt% material.

$$-r_H = k_{eH} \sigma_s C_H \left( \frac{C_H}{C_T} \right)^{\frac{5}{3}} \quad (28)$$

$$-r_H = k_{eH} \sigma_s C_T (1 - \theta)^{\frac{5}{3}} \quad (29)$$

In addition, if  $\sigma_s$  is assumed a constant, Eq. (28) is suitable for integration:

$$\left( \frac{3}{2} \right) \left( \left( \frac{C_T}{C_H} \right)^{\frac{2}{3}} - 1 \right) = k_{eH} \sigma_s t \quad (30)$$

However, there are issues with the integral method. Firstly, reaction conditions should strictly be isothermal, to reliably assign a single 'k' value at a given temperature. This can be problematic with metal hydride samples, due to low effective thermal conductivities. Further, if the inclusion of a pressure term is a valid assumption, then the site availability is not constant due to the variation in  $P$  and  $P_{eq}$ . Thus, the accuracy of the calculated effective activation energy is mainly dependent on the estimated value of  $\sigma_s$ . Even if partial differentiation was applied to Eq. (28), regarding  $\sigma_s$ , an estimation over time would still be required. In effect, the analysis indicates the integral method is not recommended to completely model  $H_2$  hydrogenation of metals. Comparably, non-linear regression involves solving the reaction rate differential equation. If coupled to an energy balance, this would better represent the key changing parameters within the system, significantly  $\sigma_s$ ; thus, mitigating the issues of the integral method.

#### Site availability

Analysis of the site availability results in several key observations. If the equilibrium pressure cannot be greater than the gas pressure, then site de-activation would limit the initial temperature spike. This indicates that with a system of exclusively hydrogen and metal, thermal

runaway cannot occur. As the rise in temperature increases site de-activation, an improvement in heat transfer would improve site-reactivation and thus increase the reaction rate. Further, as temperature varies with space and time, the wall region would experience relatively no site de-activation, whereas in the reactor centre, the opposite would occur. This important observation implies that regions experiencing a higher rate of heat transfer will transition to slower phases sooner, such as the transition from  $\alpha+\beta$  to  $\beta$  phase. Thus, modelling localised phase transitions sufficiently is required.

However, once the reaction reaches isothermal conditions (latter reaction stage), site de-activation is no longer influential. It is proposed that the site availability is still necessary, but  $\sigma_s$  is now representing phase behaviour effects. As the metal reacts, an increase in H-H interactions occur. If these interactions cause sites to be blocked, rendering these sites unavailable – then the reaction rate reduces & capacity drops. Therefore, the site availability could be a generalised measure of the phase behaviour (and temporary site de-activation) hence; to give a good approximation of the site availability throughout the reaction, it is speculated that an accurate representation of the PCT or “phase transition curve”, which is related to H-H interactions, at operating conditions is required.

We note that although  $\sigma_s$  can recognise the change in phase behaviour across each time step, it can only describe the reaction endpoint (full capacity) based on the equation used to model the equilibrium pressure. Within the equations in this paper, the capacity is only determined through a simple empirical relation (Eq. (20)) and thus this is a pitfall. Given that Eq. (20) estimates the maximum capacity when the reaction has completed, this can be influenced by the presence of a surface oxide. Although most of the magnesium activated, there is a possibility that a certain fraction did not, blocking potential sites and reducing capacity. However, we are confident that the oxygen content does not significantly influence the reaction rate as the empirical relation (4.5) mainly governs the reaction endpoint, and has little effect on the kinetic curve trajectory, which is primarily governed by the rate-determining step.

## Conclusion

Non-linear regression of a derived rate law considering both diffusion and surface effects resulted in a similar effective activation energy determined through DSC desorption experiments under Argon. At conditions close to equilibrium, the analysis suggests that the reaction is surface resistance dominating with a reaction order of 5/3. There is confirmation within the rate equation that the reactive surface area increases the reaction rate. In addition, it is known that increasing the heat transfer rate improves the kinetics, where this term is explained with the concept of site de-activation. When the reaction enters isothermal conditions, it is proposed that the site availability represents the potential influence of solid phase behaviour, which can affect the reaction rate and helps to explain the reason for variation in capacity.

## Acknowledgements

This work was supported by the Engineering and Physical Sciences Research Council (EPSRC), United Kingdom [grant number EP/L015749/1].

## Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.ijhydene.2019.04.036>.

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