Recommended Best Practices for the Characterization of Storage Properties of Hydrogen Storage Materials

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Summary of Document Structure

This Recommended Practices for the Characterization of Hydrogen Storage Materials document provides an introduction to and overview of the recommended best practices in making measurements of the hydrogen storage properties of materials. Due to the breadth of the subjects covered, material will be presented in its most concise and accessible form. The authors will use examples from literature to add clarity to key topics and to provide the reader with avenues for further detailed inquiry into a specific subject. The Recommended Practices document is divided into an introductory section and will be followed by four or more chapters that cover the topic areas of: concentration and capacity, kinetics, thermodynamics and cycle life measurements of hydrogen storage materials as well as other important hydrogen storage materials properties.

The introductory section provides the reader with the background information needed for making hydrogen storage measurements. The introduction covers the following main topics: 1) evaluating the experimental setup and procedures with respect to the ultimate purpose for making a set of measurements, 2) the relationships between direct and indirect (measureable vs. derived) quantities (which we will refer to as variables) used to describe hydrogen storage materials properties, and 3) an overview of common methods used to measure these hydrogen storage properties.

The introduction will be followed by separate chapters covering the recommended practices associated with the measurement of concentration and capacity, kinetics, thermodynamics and cycle-life properties among others. These chapters will each include: a review of theory, an evaluation of the purpose of measurements for the selection of appropriate methods, and detailed experimental considerations that are of particular importance to each property being measured.
Preface

1 Mission
The ultimate goal of the United States Department of Energy’s Hydrogen Storage Program is the development of hydrogen storage materials that meet or exceed the DOE’s targets for onboard hydrogen storage in a hydrogen-powered vehicle. The recent rapid expansion of research efforts in this field has brought the talents of a wide range of researchers to bear in solving the grand challenge of hydrogen storage. There is a need for common metrics and recommended practices for measuring the practical hydrogen storage properties of new materials that are being developed within the US DOE Hydrogen Storage Program as well as at an international level. A clear and comprehensive resource that will provide guidelines to common metrics and recommended practices in the measurements of hydrogen storage properties is critical to the success of the Hydrogen Storage Program.

1.1 Objective
The objective of this Recommended Practices for the Characterization of Hydrogen Storage Materials Project is to create a reference guide of common methodologies and protocols for measuring critical performance properties of advanced hydrogen storage materials. This document is designed to serve as a resource to the hydrogen storage materials development community to aid in clearly communicating the relevant performance properties of new materials as they are discovered and tested.

1.2 Benefit to the DOE
The benefit of this series of Recommended Practices reference guides to the DOE is to facilitate the transfer of knowledge and experience in making these critical measurements from some of the world’s experts in this field to the entire DOE and world-wide hydrogen storage community. The short-term benefit will be to provide a clear knowledge base in the form of a published resource to aid those just entering this rapidly expanding field. The long-term benefit will be to create strong ties between government, university, and small and large business entities that will lead to more open communications and ideally the establishment of uniform measurement practices and reporting.

2 Definitions
A recommended reference for detailed materials, chemistry and measurement definitions can be found in publications by IUPAC the International Union of Pure and Applied Chemistry.¹
2.1 Mechanisms
For all sorption mechanisms, the suffix -ate (sorbate) refers to the substance taken up while -ent (sorbent) refers to the sorbing substance. The suffix -tive (sorptive) refers to the sorbate substance in its natural state, i.e. the bulk gas state. The coupled terms are generally considered counterparts.\(^2\)

2.2 Sorption and Desorption
Sorption is the non-specific term that describes both adsorption and absorption. It is most often used when it is difficult or impossible to differentiate between adsorption and absorption or when both processes occur simultaneously. Desorption is the inverse process of sorption. It is important to note that the sorption and desorption of the sorbate by the sorbent is a dynamic process.

2.3 Adsorption and Absorption
Adsorption is the enrichment or depletion of one or more components in an interfacial layer. Absorption occurs when the adsorbates are incorporated into the phase of the adsorbent. In absorption, the structure and/or the chemical nature of the absorbate and absorbent may be modified. It is important to keep in mind that adsorption and absorption refer to the location of the sorbed species with respect to the sorbent, i.e. surface or bulk respectively.

2.4 Chemisorption and Physisorption
Chemisorption, or chemical sorption, includes adsorption and absorption and occurs when the forces involved are valence forces of the same kind as those operating in the formation of chemical compounds. Physisorption, or physical sorption, is restricted to adsorption and occurs when the forces involved are weak intermolecular forces of the same kind as those responsible for the imperfection of real gases and the condensation of vapors. Weak intermolecular forces are also known as van der Waals forces. Physisorption does not involve a significant change in the electronic orbital patterns of the species involved; however no absolutely sharp distinction can be made between chemisorption and physisorption. It is useful to make a distinction between molecular physisorption, in which the H-H bond in the gas phase is preserved in the sorbed state, and chemical sorption, in which the H-H bond is broken during the sorption process. For more details on features useful in distinguishing between chemisorption and physisorption, the reader is encouraged to review Everett.
2.5 Spillover

Spillover involves the dissociation of hydrogen on a metal and subsequent transference, or spillover, to the metal's support via a chemical bridge. Spillover is highly dependent on the metal dissociator and the bridges linking the metal and the support. Typically graphitic or nano-structured carbon is used as support because of its high specific surface area.³

![Figure 1. Diagram of spillover material structure.](image)

3 Materials

Materials for hydrogen storage can be divided into two families: traditional bulk or chemisorbed materials⁵, which include hydriding alloys, complexes, ammonia-based compounds, and hydrocarbons and surface or physisorbed nano-structured materials⁶ such as carbon fullerenes, nano-tubes and highly porous media like metal-organic frameworks and aerogel. The family tree in Figure 2 illustrates the relationships between different types of hydriding alloys and complexes.

![Figure 2. Family tree of hydriding alloys and complexes.](image)
3.1 Metal Alloy - Hydrides

The host hydrogen storage materials may consist of a solid solution alloy. These are characterized by varying composition and disordered substitution of one element for another on crystal lattice sites; they are formed by dissolving one or more minor elements into a primary element. Solid solutions based on Pd, Ti, Zr, Nb and V form hydrogen storage materials with some attractive features such as moderate reversible storage capacities near ambient conditions but cost and the heat released on hydrogen absorption deter the use of these materials for transportation applications. For example, (V_{0.9}Ti_{0.1})_{0.95}Fe_{0.05} has a reversible storage capacity of \( \Delta wt.\% \) = 1.8 at \( T = 36^\circ C \) and atmospheric pressure but costs but are relatively expensive for hydrogen storage. Vanadium-based solid solutions (V-SS) are compared to other metal hydrides in Table 1.

3.2 Metal Hydride Intermetallic Compounds

Intermetallic compounds (ICs) are characterized by homogenous composition and crystal structure. ICs for hydrogen storage combine a strong hydriding element A with a weak hydriding element B to create a compound with the desired intermediate thermodynamic affinities for hydrogen. Hydrogen absorbs atomically into the host metal lattice as a solid solution at low concentrations and via hydride-forming metal/hydrogen bonds at higher concentrations. The host intermetallic compound elements A and B are typically present in an integer or near-integer stoichiometric relationship. Element type A and/or B may be an ordered or disordered mixture of several metal elements. The total compositional variation has a strong impact on hydriding properties. This ability to form \( A_xB_y \) ICs containing up to 10 or more elements has been extensively studied and exploited in the commercialization of metal hydrides for hydrogen storage and Nickel-Metal Hydride battery applications. Common classifications of ICs for hydrogen storage are \( AB_5, AB_2, AB \) and \( A_2B \); LaNi_5, TiCr_2, TiFe and Mg_2Ni being representative examples. Table 1 provides an overview of the relative attributes of these different classes of hydride forming Intermetallic compounds. This includes versatility (meaning the ability to tune the alloy to operate over a wide range of pressures and temperatures) and PCT meaning the ability to reversibly provide hydrogen gas over a narrow range of pressures (relatively flat plateau pressure) at room temperature conditions.

<table>
<thead>
<tr>
<th>Attribute</th>
<th>( AB_5 )</th>
<th>( AB_2 )</th>
<th>( AB )</th>
<th>( A_2B )</th>
<th>V-SS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Versatility</td>
<td>+</td>
<td>+</td>
<td>− / 0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>H-Capacity</td>
<td>0</td>
<td>0 / +</td>
<td>0 / +</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>PCT</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>− / +</td>
<td>+</td>
</tr>
<tr>
<td>Activation</td>
<td>+</td>
<td>0</td>
<td>− / 0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Impurity effects</td>
<td>+</td>
<td>0</td>
<td>−</td>
<td>0</td>
<td>− / ?</td>
</tr>
<tr>
<td>Cyclic stability</td>
<td>− / 0 / +</td>
<td>− / 0 / ?</td>
<td>− / 0 ?</td>
<td>0 / ?</td>
<td>?</td>
</tr>
<tr>
<td>Ease of manufacture</td>
<td>+</td>
<td>0</td>
<td>+</td>
<td>0</td>
<td>?</td>
</tr>
<tr>
<td>Pyrophoricity</td>
<td>0</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>+ / 0</td>
</tr>
<tr>
<td>Cost</td>
<td>0</td>
<td>+</td>
<td>+</td>
<td>− / 0</td>
<td></td>
</tr>
</tbody>
</table>

* Attribute key: − = problem; 0 = neutral; + = good; ? = uncertain.

Table 1. Qualitative overview of metal hydride types as to attributes.
3.3 Hydride Complexes
Hydride complexes occur when combinations of ligands, metal ions and hydrogen form molecules. Hydride complexes are differentiated by whether they contain transition metals; those that do are called TM hydride complexes, those that do not are called non-TM hydride complexes. The difference between hydride complexes and ICs is that complexes release hydrogen through a series of decomposition and recombination reactions. Some combinations of elements, for example Mg and Fe, form hydride complexes but cannot form ICs; hydrogen is required for these two elements to form a stable compound (Mg$_2$FeH$_6$).

3.4 Nano-Structured Materials
Nano-structured materials have much higher surface area to volume ratios than bulk materials, enabling adsorption to play an important role. Nano-structuring of materials also improves reaction kinetics by increasing the diffusivity, reducing the reaction distance and increasing the reaction surface area. Metal hydrides and hydride complexes can be nano-structured by a variety of different processes including sputtering, pulsed laser deposition, and mechanical milling.

Nano-structured carbons, such as nanotubes, fullerenes and graphitic sheets are examples of nano-structured materials that have been extensively studied for hydrogen storage.

3.5 Porous Materials
Porous materials are being studied for use as hydrogen storage media due to their high surface area to volume ratio and the ability of hydrogen to adsorb to these internal and external surfaces. Although adsorption via physisorption is significant for porous media, absorption may also contribute to the overall hydrogen storage capacity of advanced materials. Examples of porous materials being investigated for hydrogen storage are various forms of aerogels, clathrates and metal-organic frameworks (MOFs).

4 Summary
The intent of the preface has been to introduce terminology commonly used in hydrogen storage materials research and to provide an overview of the fundamental processes and measurement considerations. We hope that this information proves useful and ultimately contributes to progress in the research and development of new and better hydrogen storage materials.
Section 1: Introduction

This introductory section is intended to provide the reader with the basic information necessary for productive and accurate hydrogen storage measurements. Introductory topics include: 1) evaluating the experimental setup and procedures with respect to the ultimate purpose for making a set of measurements, 2) the relationships between direct and indirect (measureable vs. derived) quantities (which we will refer to as variables) used to describe hydrogen storage materials properties, and 3) an overview of common methods used to measure these hydrogen storage properties. A variety of apparatus have been developed in the past to measure sorption properties; this manuscript will focus on some instruments typically used for hydrogen storage measurements (namely volumetric and gravimetric systems).

1 Purpose of Measurements

For simplicity, hydrogen storage properties measurements can be broken down into three basic categories: storage system level performance measurements, materials development measurements, and fundamental science measurements. It is important to understand the purpose of a particular experimental investigation before making measurements because the experimental setup and procedures can vary greatly depending on the purpose of the measurements and results and conclusions can be misleading if they are not presented in the proper context. Of course, these categories overlap and support each other. Testing the design and performance of a storage system, comparing storage properties across new or improved materials, and gaining insight into the underlying phenomena of hydrogen storage require experimental apparatuses and procedures that may be significantly different. Care should be taken to match the experiment well to the purpose of the study to avoid misleading results.

1.1 Purpose: System Performance

System level measurements are concerned with the performance of materials at an operational scale. The primary difference between experimentation at the system level versus other levels is the importance of application-specific considerations. Depending on the utilization of the hydrogen storage system, any combination of properties may be most important for performance optimization. Properties such as energy density, reversibility and safety at standard operating conditions are some of the primary 'real world' considerations. For example, one consideration central to portable hydrogen storage systems like those envisioned for transportation is energy density because the system must be transported. Stationary applications, in contrast, may be more concerned with total energy than energy density, and cost.

To gain useful system performance information, enough material should be tested to provide results representative of the material’s behavior in a full-sized operating storage system. In general, this means between 10 grams and
1 kilogram of storage material. The authors believe 100 grams is a good tradeoff between the practical issues of synthesis and handling of the materials and a characteristic, uninterrupted volume of material that would be found in a large-scale system. It is recommended to take samples from different portions of larger batch to get a good average material representation.

An example of an experimental setup that may be appropriate at a system performance level but not for other purposes, is the use of flow meters to study rates of hydrogen uptake or release. This requires that the material performance has already been well characterized at a smaller scale. This is because conventional thermal mass flow meters generally have fairly narrow dynamic measurement ranges (often less than one decade). Thus, some prior knowledge of the material’s performance is needed to select flow meters with the appropriate flow range. This can be a severe limitation when a material’s hydrogen sorption/desorption rates may vary over several orders of magnitude depending on state of charge, temperature and pressure. An alternative is Coriolis flow meters, which have a robust dynamic measurement range, maintaining accuracy over several decades.

### 1.2 Purpose: Materials Development

The purpose of experimentation at the analytical level (typically tens of milligrams to a few grams) is to guide the development of advanced materials with desirable hydrogen storage properties. It is important to realize that hydrogen storage properties of a material can be strongly influenced by any number of individual sample characteristics including total chemical composition and distribution, microscopic and macroscopic structure, etc. In the realm of materials development, parametric and comparative studies are generally the best approach to producing reliable and productive studies. There is an extremely wide range of material characteristics (including such things as doping and nano-structuring) that can be modified and manipulated to produce profound changes in hydrogen storage properties. For example, the effect of elemental substitution in the LaNi\(_5\) family of intermetallic compounds (LaNi\(_{5-x}A_x\)) is demonstrated in Figure 3. In this case Ni has been substituted with tin or aluminum to form materials with decidedly different hydrogen storage properties, including changing the thermodynamic stability of the hydride that is formed.
In order to conduct instructive parametric studies, it is especially important that measurements are conducted under identical conditions (pressure, temperature, sample size, etc.) using identical measuring equipment. This minimizes the number of free variables associated with the experiment and allows for the accurate determination of the effect of a specific material property. The effect of some material properties on hydrogen storage can be quite small and easily masked by variation in testing conditions and measuring equipment if the proper precautions are not taken.

1.3 Purpose: Fundamental Science

As the title suggests, fundamental science research is concerned with identifying and understanding the physics and chemistry that govern a material’s hydrogen storage properties. This fundamental knowledge can be used in research at the materials development and system performance levels. It may help to validate models that describe hydrogen-material interactions or aid in making enlightened decisions on new directions in improving materials. Without some level of fundamental understanding, system and materials development measurements would be conducted solely through intuition or painstaking systematic studies. By using knowledge gained from fundamental science, researchers can eliminate much of the trial and error and more efficiently conduct measurements for material development. With regard to fundamental science, samples may be small (gram quantities) and indeed very small (milligram powders, liquids and thin films) as synthesis is achieved on lab-scales. As a consequence, very sensitive instruments will be needed to perform measurement on sample weighing as low as tens of milligrams. Also, fundamental science implies the
modeling of the measurements and hence, intrinsic behavior should be clearly separated from instrumental effects to match a particular model.

2 Hydrogen Storage Properties

Measurements at all three of the levels investigation described above, probe similar hydrogen storage properties of both systems and materials. The relative importance of each property may depend on the application or the purpose of investigation. The following is a very brief summary of the principal measured properties of hydrogen storage materials and systems. Each property is investigated in more detail in the individual chapters of the Recommended Practices Documents.

2.1 Property: Capacity

Capacity is the maximum steady-state hydrogen content of a storage material. Capacity can have several different definitions that reflect the application or material considerations including reversible capacity, usable capacity and excess material capacity (These terms will be defined in the coming chapter on capacity measurements and will be added to the definition sections). These definitions in turn may depend on the material’s stability, composition, temperature, pressure and number of cycles. Each of these variables has the potential to change the capacity of a material. Additionally, there is a very important distinction between concentration and capacity: capacity is a material property that does not vary at a set state (after an extended period of time). In this text, a material’s capacity will be referred to as its maximum steady-state hydrogen content and concentration as its temporal hydrogen content. For a more detailed discussion of capacity definitions and theory, please see the chapter on concentration and capacity.

2.2 Property: Kinetics

Kinetics is the relationship between hydrogen concentration and time in a storage material. Kinetics is a measure of the rate of hydrogen sorption or desorption of a material and may not be exclusively dependent on intrinsic material properties. Sample size, heat transfer effects and other parameters that are highly dependent on the experimental method can affect kinetic measurements and conclusions. Minimizing the effects of external influences on kinetics is very difficult and requires a great deal of knowledge and preparation. Thus, extreme caution should be used in ascribing measured kinetics to fundamental sorption mechanisms or an intrinsic property of a given material. Sample preparation, catalyst and additives, particle size, heat transfer capabilities, among other things can strongly influence the kinetics. As one example, degassing at high temperature can remove or diffuse oxide layer from metals and increase subsequent hydrogen absorption kinetics. For a more detailed discussion of experimental considerations for kinetics, please see the chapter on kinetics.
2.3 Property: Thermodynamics

The intrinsic thermodynamic properties of hydrogen storage materials influence a number of other parameters, most notably the hydrogen capacity based on temperature and pressure. Unfortunately, the relationship between measured temperature and pressure conditions and intrinsic thermodynamic properties may be complicated by kinetics considerations. In many of today’s materials, the temperature and pressure conditions required for hydrogen uptake and release are dictated by kinetic considerations, not necessarily the intrinsic thermodynamics of hydrogen bonding. The ability to distinguish between the two material properties is especially important because the techniques used to improve one property are often ineffective or not available for the other.

In addition, we refer in this document to thermodynamic properties with respect to thermal equilibrium (the measure equilibrium state of hydrogen concentration in the material, hydrogen pressure and temperature). Thermal equilibrium may be dictated not only by the chemical interaction thermodynamics, but other terms as well such as strain energy, dislocation production and interfacial energy. In many metal-hydrogen systems these terms lead to differences in the absorption versus desorption equilibrium pressures at a given concentration and temperature. This is generally observed as hysteresis in the Pressure Concentration Temperature (PCT) isotherms. For a more detailed discussion on the thermodynamics of hydrogen storage, please see the chapter on thermodynamics.

2.4 Property: Cycle-Life

Cycle-life testing is restricted to reversible hydrogen storage materials such as metal and complex hydrides, amides and physisorbing materials. Materials that store hydrogen irreversibly (i.e. not reversible under practical conditions), like chemical slurries of elemental hydrides (e.g. LiH, NaBH₄), cannot be cycled. Cycle-life measurements are typically performed to characterize the effect of cycling on capacity that stems from activation effects, grain growth, structural degradation or chemical degradation due to gas impurities. However, kinetics may also be impacted by cycling and some of the observed change in capacity may in fact be due to changing kinetics during cycling. For example, if measurement time intervals during a cycle-life experiment do not reflect the changing kinetic properties of the material, the measured capacity will not be representative of capacity at quasi-equilibrium. Impurities in the hydrogen gas supply may have a profound effect on the cycle-life behavior of a storage material (see section 5.3.6). For a more detailed discussion of cycle-life considerations, please see the chapter on cycle-life measurements that will follow in a later version.
3 Types of Measurement

With respect to characterizing a material's hydrogen storage performance there are two principal types of measurements, kinetics measurements and pressure-composition-temperature (PCT) measurements. Kinetics measurements can be considered the fundamental measurement of hydrogen storage because other types of measurement, including PCTs, are collections of several individual kinetics measurements. The relationship between kinetics and PCT measurements will be discussed further at the end of this chapter. Beyond these two main characterization measurements, there is a host of other measurements such as scanning temperature measurements that have value in rough screening for potential new hydrogen storage materials.

3.1 Measurement Type: Kinetics

Kinetics measurements track the rate of change of hydrogen concentration in a sample after the sample has been perturbed from quasi-equilibrium. Because hydrogen storage measurements can only approximate thermodynamic equilibrium due to experimental limitations, equilibrium will be used synonymously with quasi-equilibrium throughout the rest of this manuscript. Perturbation from equilibrium is accomplished by changing the thermodynamic state of the sample's environment, which causes the sample to sorb/desorb hydrogen in the attempt to restore equilibrium. For the gravimetric and volumetric methods, kinetics measurements are generally conducted under isothermal conditions, with changes in pressure used to perturb equilibrium. This is done partly for compatibility with PCTs and partly because of the profound effect of temperature on the kinetic character of almost all materials. The profiles and interpretations of kinetics curves and the considerations for kinetics measurements are the subject of the kinetics chapter of the Recommended Practices Document.

It is instructive to introduce a few fundamental features common to almost all kinetics measurements. The degree of perturbation from equilibrium is the driving force behind sorption/desorption in hydrogen storage materials: large perturbations cause relatively large rates of change in concentration while smaller perturbations cause relatively small rates of change. The rate of change of hydrogen concentration in a sample is largest at the beginning of a kinetics measurement, when the sample is furthest from equilibrium with its environment. The distance from equilibrium diminishes as hydrogen is sorbed by the sample, slowing the sorption rate as the sample reaches a time independent equilibrium concentration. These effects are illustrated in the sorption kinetics measurement in Figure 4. At the beginning of the experiment, the concentration changes quickly because the sample is furthest from equilibrium and with time the rate of change of concentration decreases until it becomes effectively zero.
3.2 Measurement Type: Pressure-Composition Isotherm

Pressure-composition-temperature measurements (PCTs) are the most reported hydrogen storage measurement type in academic literature (composition in this context is synonymous with the concentration of hydrogen in a sample). A PCT measurement is a collection of data points that represents the pressure, concentration and temperature of a sample in equilibrium and relates the influence of the thermodynamic variables on concentration. PCTs are also commonly referred to as PCI (Pressure-Composition Isotherms) because they are taken at isothermal conditions. This minimizes the number of free variables and allows the relationship between concentration and pressure to be presented via two-dimensional graphics. The effect of temperature on hydrogen storage properties can be determined by comparing PCT isotherms at various temperatures. Because PCTs represent a sample in equilibrium, they can also be used to determine the thermodynamic properties of a hydrogen storage material. Unfortunately, the absolute interpretation of PCT data must be kept in perspective because it is difficult to make true equilibrium measurements. This is because as the pressure approaches the true equilibrium pressure, the driving force (or potential) becomes diminishingly small and therefore, kinetics becomes increasingly slower. In other words, it would take an infinitely long time to reach equilibrium. For more information on thermodynamics as they relate to hydrogen storage properties, please review the thermodynamics chapter in the Recommended Practices Document.

It is instructive to conceptualize the relationship between PCT measurements and kinetics measurements in the volumetric method as follows: a sample is perturbed from equilibrium by a change in the pressure of the system and is allowed to reach equilibrium through the dynamic process represented by a kinetics measurement. The last data point of each kinetics measurement, which most closely represents equilibrium, provides a single point of hydrogen concentration, pressure and temperature at equilibrium. In the volumetric method, this dosing process is performed repeatedly at one temperature until
there are enough equilibrium data points are collected to construct a full PCT diagram. In this sense the PCT measurement can be thought of as a cumulative series of mini-kinetics measurements. Figure 5 demonstrates the PCT diagram of a porous material using the volumetric method by conducting a series of small dose kinetics measurements. In each kinetics measurement (separated by recharging the dosing volume which appears as discontinuities), the final pressure at the last data point represents the equilibrium pressure at a given concentration. In the volumetric method, the concentration is determined from the change in pressure with each dose (refer to the volumetric section 5.3 for details). Therefore the equilibrium pressure of the sample corresponding to each equilibrium concentration point is also known and a full PCT can be constructed from the series of kinetics measurements.

![Graph of Pressure vs. Concentration](image)

**Figure 5.** Representative measurement of a porous material. The last point in each ‘row’ of points is taken as representative of the equilibrium concentration of hydrogen in the sample at the pressure and temperature of experimentation.

This is shown in Figure 6 where the Pressure / Time data is translated to Pressure / Concentration data to form a PCT diagram. The relationship between pressure drop due to absorption (rise on desorption) and concentration is depicted in Figure 7.
Figure 6. Representative pressure / time measurement and the resulting PCT plot of hydrogen absorption to form a metal hydride. The last point of each gas sorption dose provides the concentration of hydrogen in the sample at the equilibrium pressure and temperature of each dose.

Figure 7. The relationship between pressure drop on absorption and the concentration associated with this pressure drop that is used to create each point on the equilibrium PCT diagram.
4 Hydrogen Storage Variables

Because the concentration of hydrogen in a sample is not measured directly, this important parameter necessarily relies on indirect determination from directly measured variables. Accurate measurement of direct variables is important for all hydrogen storage properties and perhaps more so for the determination of concentration than any other.

All storage properties are determined by the relationships between the variables: concentration, sample weight, temperature, pressure, cycle and time. The important distinction between direct (weight, temperature, pressure, time, cycle) and indirect (concentration) variables is made to differentiate between variables that can be measured directly from those that cannot. The last five (mass, temperature, pressure, cycle and time) are termed direct variables because they can be directly measured with traditional measuring devices such as balances, thermocouples and transducers. Concentration is considered an indirect variable because it cannot be directly measured and must be determined by correlating a direct (measurable) variable with concentration through the use of an empirical relation. Because of its dependence on measurable variables and its explicit ties to capacity, concentration will be addressed in detail in the Concentration and Capacity chapter.

The direct variables present problems in accurately collecting and interpreting data. Weight, temperature and pressure are difficult to accurately measure in all testing methods because errors can be introduced by testing equipment, sample holder design and secondary effects such as buoyancy. Errors associated with measuring the direct variables affect all property investigations and therefore the sources of these errors will be reviewed in the introductory chapter. Measurement errors specific to a particular testing method will be addressed in the associated methods section 5. In addition to this section on the direct variables, the effects of testing and variable collecting methods unique to a particular storage property (e.g. capacity, kinetics) will be addressed in the property chapters that follow.

At this point, the reader is encouraged to briefly review the section 5 on Methods of Measurement to become familiar with the nomenclature and general principles of each testing method. Material in the sections on variables (4) and also static and dynamic measurements (5.1) will refer to basic concepts associated with the measurement methods. Familiarity with the material in these sections will aid in gaining an overall understanding of the subject.

4.1 Measureable Direct Variables

Measureable or “direct” variables (weight, temperature, pressure, time, cycle) affect the intrinsic adsorption/absorption behavior of a material:
Equation 1

\[ N_a = f(P,T, \text{solid} - \text{gas system}) \]

Where \( N_a \) is the number of moles adsorbed. In addition, the measurement of direct variables is needed to determine “indirect” variables such as capacity. The following are the direct variables that are typically measured to determine the hydrogen storage properties of a material. While the means and accuracies by which these variables are measured may vary (e.g., volumetric, gravimetric, electrochemical,....) the influence on the intrinsic properties and performance of the materials should be the same.

### 4.1.1 Variable: Weight

The weight variable measured in the gravimetric method is often considered a direct measurement of hydrogen concentration; unfortunately, this is not the case. The balance in the gravimetric method (used to measure sample weight) actually measures the resultant force of a number of discrete forces: the weight of the sample, the buoyancy force caused by the displaced gas and the forces associated with mechanical disturbances and gas convection. The resultant force is often referred to as the apparent weight of the sample in order to differentiate between the measured weight and the actual weight of the sample. In order to determine the actual weight of the sample (from which concentration can be determined), it is necessary to account for and minimize the extraneous forces acting on the balance. The buoyancy force can be determined by an understanding of the skeletal density of the material while proper vibration damping can minimize mechanical forces. Forced and free convective forces are caused by pressure changes during charging/discharging and thermal gradients, respectively, and are transient in nature; patience is essential to minimize convective forces. Gravimetric measurements often use constant gas flow past the sample. In these cases, the flow drag forces must also be known and taken into account. Generally the flow drag force changes with the gas density, and therefore is a function of both temperature and pressure. Proper calibration of gas flow forces is necessary if the flow forces are significant in the particular apparatus or experimental arrangement.

In addition to the significant measurement considerations discussed above, the accuracy and precision of measuring weight depends on the type of balance used. A common weight-measuring system is an electronic strain gauge attached to a cantilever or balance (see gravimetric method section 5.4). The accuracy and precision of the strain gauge and cantilever/balance system are based on the properties of the strain gauge and the cantilever/balance material. The deflection varies with the modulus of elasticity of the material and the weight of the sample; a low modulus of elasticity leads to greater deflection and a more sensitive instrument but the instrument is more susceptible to noise due to external forces. The modulus of elasticity will vary with temperature and this must be taken into account when making measurements at different temperatures or when ramping temperatures. The sensitivity of the balance must be considered along with other trade-offs when choosing gravimetric instruments.
With respect to hydrogen storage materials, the hydrogen storage capacity is usually given on a weight basis. Therefore accurate measurement of a samples weight is critical. Both chemisorption and physisorption materials may be highly air and water reactive, therefore samples are usually prepared and weighed in an inert atmosphere such as an argon glove-box. In addition, both types of material may require significant out-gassing of residual solvents, or adsorb impurities such as water before hydrogen testing. Thus for correct hydrogen capacity measurements it is vital that the sample is weighed after degassing. This can be performed either before or after hydrogen testing and preferably in an inert gas atmosphere.

4.1.2 Variable: Pressure

The pressure variable is intimately associated with hydrogen storage research and influences several important properties including concentration correlations, capacity and the kinetics of charging and discharging. Pressure is a commonly used measurable variable in concentration correlations and is therefore of practical interest to the measurement of all hydrogen storage properties. For example, the volumetric method measures changes in equilibrium pressure, along with volume and temperature information, to determine concentration. Changes in pressure are often used to control sorption/desorption reactions during both testing and practical operation. Isothermal measurements such as PCTs, one of the most common representations of capacity, use pressure variation to drive charging and discharging during storage property characterization. Pressure variation can also be used to control sorption and desorption in functioning storage systems, depending on the application. In fuel cell (FC) applications, hydrogen supplied at initially isothermal conditions (ambient) and elevated pressure is used to charge the storage system at the fueling station. The elevated pressure causes the reversible storage material to charge with hydrogen for future consumption by the fuel cell.

In addition to its obvious effect on reaction kinetics, pressure profoundly affects capacity measurements and practical capacities. Among the many capacity definitions are two based on thermodynamic considerations, reversible and usable capacity. Reversible capacity is of concern at the materials development level and is the measure of capacity available under feasible conditions. Reversible capacity is of particular importance to chemisorbing materials. Usable capacity is the capacity defined by the thermodynamic restrictions placed on the storage system by the environment and the end application. Once again, fuel cells present an excellent example of the importance of the thermodynamic variables on practical storage. The range of temperatures and pressures accessible to charge and discharge a storage system are restricted by availability; in classic PEMFC applications, this translates to operating temperatures from ambient to (~353K) and roughly ambient to a few bar pressures (assuming no external pressurizing or regulating equipment is used). It is highly probable that maximum capacity of a given storage system does not
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occur within these limited ranges; therefore it is important to quantify usable capacity.

Electronic pressure transducers are the most common means of measuring pressure in hydrogen storage testing equipment. The accuracy limitations of transducers are generally described by two different types of error bands, those based on a percent of the actual reading (Capacitance Manometers) and those based on a percent of the full scale (Strain Gauge Transducers) (see Figure 8). For this reason, percent reading error bands are more accurate at low pressures while percent full-scale error bands may be more accurate at high pressures depending on the total error rating of the transducer. For hydrogen storage testing, particularly volumetric methods, the low-pressure range (0-15 bar) is often the most critical range for investigating storage properties. This makes the percent reading error bands the preferred error band for hydrogen storage.

![Figure 8. Depiction of the two types of error bands in pressure transducers.](image)

A simple calculation demonstrates the percent reading advantage at low pressures. Take two pressure transducers rated to 300 bar, one has a 1 % reading error band while the other has a 0.1 % full-scale error band; the ‘break even’ pressure for these two transducers is 30 bar. The percent reading transducer is more accurate for pressures below 30 bar, the range of pressures most often encountered in hydrogen storage experiments. For optimum accuracy, several percent reading transducers rated to different pressures should be used.

The sensitivity of pressure transducers requires that the pressure signal change from all other sources, \( \Delta P_0 \), is minimized. It is conceivable to potentially account for these other pressure signal changes and correct the data for their influence, but in practice it is more fruitful to minimize erroneous \( \Delta P_0 \) signals. Background
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Pressure signal change can be caused by a number of factors including transducer sensitivity, zero drift, miss-calibration and hysteresis effects.

With a near instantaneous change in pressure, there may be some error introduced into the immediate response of the pressure transducer, especially with respect to pressure measurements made by measuring the strain or deflection of a gauge’s diaphragm. There are a couple of ‘tricks’ that can help determine the true measurement signal from a pressure transducer. After the transient from gas introduction has decayed, the measurement signal becomes pseudo-differential in the sense that the current signal can be compared to the measurements that came before and will come after. The signal curve is expected to show a smooth behavior and therefore noise superimposed on the curve can be “reduced” through standard procedures such as smoothing. If the sample shows marginal sorption/desorption during the transient, the pressure vs. time curve contains both the calibration data and the sorption/desorption data. In other words, since very little absorption occurred during the transient, the transient can be ignored and the pressure reading at t = 0 is the pressure with “no absorption”. Thus the ideal sample, ideal in the sense that it is easy to measure, would have no sorption/desorption signal until after the transient but would come to equilibrium before other longer-term error signals such as temperature fluctuations have a chance to impact the data.

4.1.3 Variable: Temperature

Temperature is the second thermodynamic variable with important implications to hydrogen storage properties. It has many of the same hydrogen storage effects as pressure; it can be used to determine the binding energy of hydrogen in a sample (as in Differential Scanning Calorimetry), drive the sorption/desorption reactions in several testing methods and applications, and temperature influences a number of different definitions of capacity. To continue with the fuel cell example, temperature variation is the primary mechanism used to release hydrogen after initial charging because of the relatively limited pressures available due to FC structural considerations. Temperature affects practical capacity in much the same way as pressure. The two capacity definitions derived from thermodynamic considerations are based on temperature considerations as well as pressure considerations. Although, in hydrogen storage research, pressure and temperature are intimately linked, their measurements have their own, unique considerations.

Temperature-related measurement error is one of the most common errors associated with hydrogen storage measurements, particularly kinetics measurements. The accuracy of temperature measurements is typically limited to the accuracy of the thermocouples used and the heat transfer characteristics of the sample material and sample cell. Like pressure transducers, the accuracy of a thermocouple measurement depends on what type of thermocouple is used and the temperature regime of the measurement. Some thermocouple types have a wide temperature range but are less accurate compared to those with
limited temperature ranges. Because most hydrogen storage experiments are conducted under isothermal conditions, the authors advise the use of limited-range durable thermocouples that offer greater accuracy in measurement.

An important consideration is that thermocouples of all types generally present nearly the same voltage at room temperature. It is only at elevated temperatures that the deviation in the type (J, K, and N for example) will become apparent. As most modern temperature measuring devices offer many thermocouple options, it is important not only to validate the devices thermocouple type settings but also to validate by using separate secondary temperature measuring devices and to run calibrations on measurement devices on a regular basis.

Note that most thermocouples (if not all) are at the limit of their operating ranges under cryogenic temperatures and can have large errors when within a few degrees of 77K. The temperature is preferably measured with Pt resistance or silicon diode at around or below 77K.

Heat transfer between the sample material, sample cell and thermocouple is one of the primary sources of error in hydrogen storage measurements. Sorption and desorption reactions can be highly exothermic and endothermic and the energy loads must be transported efficiently to maintain isothermal conditions during testing. Insufficient heat transfer can lead to pockets of sample at higher temperature than the temperature read by the thermocouple, effectively invalidating the isothermal assumption and conclusions based on that assumption. Thermocouples that are not in intimate contact with the sample, or perhaps not even in contact with the sample holder, can produce very large inaccuracies with respect to the actual sample temperature. On a system level where kilograms of storage material may be used, excellent heat transfer characteristics are required to supply and dissipate the significant amounts of energy necessary to charge and discharge a sample. The importance of heat transfer considerations will be discussed on several occasions throughout the course of the Recommended Practices Document.

4.1.4 Variable: Cycle

The desire for reusable hydrogen storage systems necessitates the ability to charge and discharge (cycle) repeatedly without loss of performance. Nearly all storage properties, particularly capacity and kinetics, vary with cycling and the variation can have a profound impact on system efficacy. Cycling phenomena include activation effects and poisoning (capacity) and retardation (kinetics) due to the gettering of impurities during cycling. In general a sample should be cycled until the capacity is stable; usually 10 cycles are sufficient for that purpose.

Some important considerations for the cycle variable are primarily aimed at minimizing activation and gas stream impurity effects on measurements and are not testing method-specific. For activation effects for both capacity and kinetics, the authors advise cycling the sample at least ten times in order to measure the
intrinsic properties of the material. It is also critical to evaluate the effects of poisoning and retardation on performance, especially in metal hydrides; for practical application, storage systems will commonly be charged with hydrogen gas that contains impurities like CO₂, H₂O and NH₃. The impurities adsorb to the material, occluding catalytic sites and diffusion pathways, and can be difficult to desorb because of their high thermodynamic affinity. As cycling increases with impure hydrogen gas, the impurities build to levels that poison capacity and retard kinetics. Therefore, it is important to develop materials that can withstand the effects of gas stream impurities in order to avoid performance deterioration.

Several other phenomena occur when cycling a hydrogen storage material. In intermetallic compounds, decrepitation, self-pulverization due to stresses caused by lattice expansion upon hydriding, and disproportionation, dissociation of a compound into its fundamental components during repeated cycling (e.g. \( \text{LaNi}_5 + \text{La}_2\text{Mg}_{17} \rightarrow 3\text{La} + 5\text{Mg},\text{Ni} + 7\text{Mg} \)), cause variation in storage properties and system performance. Decrepitation and disproportionation are generally considered activation phenomena and initial cycling mitigates their effects.

Note: UHP Ultra-High purity (99.999% purity) hydrogen gas is recommended. It is also highly recommended that the source hydrogen gas is always tested for purity, especially for oxygen and water contaminants. A good solution is to have a hydrogen purifier system just before the gas enters the testing system.

### 4.1.5 Variable: Time

The effect of time on hydrogen storage measurements is manifested by the rate at which measurements are taken. However, some confusion surrounds how measurement rates affect data representation. Measurement data is collected in two distinct steps: the data acquisition hardware converts the continuous analog output from the measuring device (e.g. thermocouple, pressure transducer) to a digital value at a certain rate \( R_1 \) and the computer software samples the digital value from the data acquisition hardware at another rate \( R_2 \).

![Figure 9. Schematic of a typical hardware/software set-up and the flow of information in a hydrogen storage testing system.](image)
As Figure 9 illustrates, information is exchanged between two interfaces but not necessarily at the same rate. The sampling rate of the data acquisition hardware can be faster or slower than the sampling rate of the computer software. The difference between the rates may lead to the collection of multiple data points at the same value of the measured parameter and/or a step-wise rather than continuous change in the measured parameter. Both of these effects are most pronounced at small time steps that approach the limits of the sampling rates and are an artifact of data collection. The average results of the measured parameters are still representative of the homogeneous change in the properties being measured.

Figure 10 illustrates a number of data collection artifacts. At the beginning of the experiment, the $R_2$ sampling rate of the computer software is faster than the analog to digital conversion rate $R_1$ of the data acquisition hardware. Therefore, the computer software samples the data acquisition hardware several times before the hardware updates. This leads to several consecutive data points collected at the same pressure, an artifact that disappears as the $R_2$ sampling rate decreases (by design) below the data conversion rate $R_1$ as the experiment progress. The step-wise rather than continuous change in the data at the beginning of the experiment is caused by the difference between the sampling rates and the initial gradient in the measured parameter. Figure 10 shows the measured pressure directly after the sample is dosed with hydrogen; the pressure gradient is initially large but slowly levels off as the sample approaches equilibrium. This is reflected in the data as the pressure step size becomes less and less and the data eventually appears continuous. These kinds of data collection artifacts demonstrate the resolution limits of the instrumentation but generally do not have a significant impact on the accuracy of the measurement and the conclusions that can be drawn. However, without understanding what causes them, they may be misinterpreted as a problem with data acquisition or instrument operation when first observed.
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4.2 Accuracy and Precision of Measurements

In measuring direct variables, it is important to understand the distinction between the accuracy and precision of testing equipment and convey pertinent accuracy and precision information to the reader. Accuracy is the degree of conformity of the measured variable to the actual value while precision is the degree of reproducibility of the measurement. A measurement can be accurate and precise, one or the other, or neither. An excellent and frequently used parallel to explain the relationship between accuracy and precision is the target analogy. The distance from the marks to the center of the target is the measure of accuracy and the size of the mark cluster is the precision. Figure 11 illustrates the possible combinations of accuracy and precision in the target analogy. As a thought exercise, consider the limiting cases of perfect accuracy and perfect precision. In the case of perfect accuracy, precision must also be perfect. The reverse however, is not necessarily true: perfect precision does not ensure perfect accuracy. These cases can be easily shown in the target analogy. The limiting case of perfect accuracy necessitates that all of the marks lay one on top of another at the center of the target; obviously this ensures perfect precision because the spacing between the marks is zero. Perfectly precise marks lay one on top of another but can be located anywhere on the target, hence perfect precision does not ensure perfect accuracy.
4.2.1 Analog to Digital Conversion Error

The accuracy and precision of all direct variable measurements are influenced by analog to digital conversion effects. Traditional analog measuring devices such as thermocouples and pressure transducers output continuous signals that must be converted into discreet digital values. The analog to digital conversion process (ADC) has three principal sources or error: quantization, non-linearity and aperture error. Quantization error is caused by the finite resolution of converting a continuous analog signal to discreet digital values, non-linearity error is caused by physical imperfections in the conversion device itself (but can be mitigated by proper calibration) and aperture error arises from uncertainty in the ADC sampling rate. In addition to errors caused by ADC, measuring devices for some direct variables have unique considerations for accuracy and precision. These considerations will be reviewed in detail in the following variable sections 4.1.

4.2.2 Error Bars in Data Representation

Error bars are used in experimental science to serve a number of important functions in relating the precision of an experimental measurement to the reader. An error bar indicates the range of one standard deviation of the measured variable and can be used to visually compare the precision of two measurements, determine whether differences are statistically significant and determine the statistical fit of experimental data to a given function. It is important to include error bars in graphics in literature when reporting hydrogen storage properties to both reflect the experimental nature of hydrogen storage testing and aid in comparing the performance of materials and the precision of testing methods.

Experimental data is often a collection of dependent variable measurements at a given independent variable value. The collection of measurements is presented as their mean value and standard error, which is based on the standard deviation of the collection. Standard error of a collection of measurements is determined by dividing the standard deviation of the collection by the square root of the number of measurements in the collection. Standard error is represented in mathematical form by Equation 2 (M is a vector of measurements with m entries):
Equation 2

\[ SE = \frac{\text{std}(M)}{\sqrt{m}} \]

For example, a (24.30, 24.68, 25.73, 26.08) collection of molar specific volume (L/mol) measurements at standard temperature and pressure would be summarized as \( v = 25.2 \pm 0.38 \) L/mol and presented graphically as in Figure 12.

![Figure 12. Graphical representation of collection of data points in mean/error bar format.](image)

Error bars should be included on figures and error ranges on values whenever possible or at least stated in the experimental section. The value of these error limits can be determined from the known (or rated) errors of each measuring device (balance, pressure transducer, thermocouple, timing device….) and standard error analysis\(^{13}\).

### 4.2.3 Testing For Accuracy

A very important and common standard practice that should always be carried out before experimental measurements and on a regular and documented basis is blank runs under identical experimental conditions with a non-absorbing material (example: glass beads or low surface area, silica or stainless steel). It is important for both gravimetric and volumetric techniques that the blank sample has essentially the same mass and x-ray (bulk) density as that of the sample that will be measured. This criterion helps to emulate, to some degree, both buoyancy effects and corrections in gravimetric measurements and gas displacement volumes in making temperature corrections in volumetric measurements. The results of such “blank” measurements should show zero hydrogen absorption and desorption within the expected range of errors. This is an important step in identifying instrumental and/or procedural errors.
Ultimately, blank tests following a rigorous calibration procedure will produce a total experimental error that encompass all the uncertainties associated with the various variables, calibrations, calculations, corrections and uncertainties.

Finally, cross-checking or validation by performing both multiple measurements on the same sample and on different samples from the same batch of material should be performed not only on one instrument but, when available, on different types of instruments potentially at different institutions with different equipment operators. While it is often very difficult to perform measurements under exactly the same conditions and also difficult to ensure identical sample preparations or loading, the intrinsic hydrogen storage properties of a materials should be essentially the same. Thus, differences in capacity, kinetics, thermodynamics, etc. significantly larger than the total experimental error should be taken seriously. In this field, there are many instrumental and procedural issues that can lead to highly incorrect results including both false positives and negatives (usually the former). When in doubt, it would seem prudent to assume the worst.

4.3 Calibrations
Measurement instruments should be recalibrated on a regular basis. Most institutions have standard calibration protocols that should be followed. An important part of such protocols is a rigorous documenting of such calibrations so that if significant errors are discovered, previously recorded data can be corrected appropriately. The following are common methods for calibrating devices that measure important dependent variables.

4.3.1 Weight
It is recommended that NIST standard procedure be followed for making weight measurements. Documents detailing these procedures and calibrations can be downloaded from the NIST Weights and Measures website.

4.3.2 Volume
The standard procedure for calibrating a volume is:
1) Liquid method
2) Gravimetric method
3) Pressure method
4) Known displacement volume calibration sample


The Pressure method is commonly used in this field and requires a well calibrated pressure transducer connected to a standard volume (as measured through the Liquid or Gravimetric methods). The Pressure method is a secondary calibration technique that is generally used for convenience as no liquids are
required. It is essentially the Sievert’s method without a sample. The setup consists of a the calibration volume connected to the calibration pressure transducer with a low-displacement volume valve between the two and a rigid coupling fitting to connect the calibration system to the instrument or volume to be calibrated.

![Figure 13 Pressure Method calibration system.](image)

Gas (typically dry Nitrogen, Argon or Helium) is charged at a high pressure (0.1 to 5 bar, and at the upper 10% of the pressure transducers full scale range) into the calibration volume the valve is closed and the pressure is measured ($P_1$). Sufficient time (5-30 minutes) should be allowed for the gas to come to thermal equilibrium with the tubing and vessels before any pressure reading is taken. The pressure in the volume to be measured is then reduced (0.1 to 1 bar, and at the lower 10% of the pressure transducers full scale range) and the second pressure is measured ($P_2$). The valve is then opened gas is dosed into the unknown volume from the calibration volume. After enough time has passed for the gas to come to thermal equilibrium (5 to 10 minutes) the resulting equilibrium pressure is measured ($P_3$). Thus, the volume to be calibrated can be determined from the gas mass balance:

$$V_x = V_c \times \left( \frac{P_3}{T_3} - \frac{P_1}{T_1} \right) \div \left( \frac{P_2}{T_2} - \frac{P_3}{T_3} \right)$$

Where $V_x$ is the unknown volume and $V_c$ is the calibration volume. Here the ideal gas law is assumed to be sufficiently correct. The process should be repeated several times and should also be done in the opposite sense, where high pressure gas is dosed from the unknown volume into the calibration volume.

Ideally, the temperature of the entire calibration and unknown volume system are being held at exactly the same temperature ($T_1=T_2=T_3$). If not the temperatures should be measured carefully at the calibration volume ($T_1$) and the unknown volume ($T_2$) and an assumption that the final equilibrium temperature ($T_3$) is a weighted average of the two:

$$T_3 = \frac{(T_1 \times V_c) + (T_2 \times V_x)}{V_c + V_x}$$
Where some initial approximations for $V_x$ can be made or Equation 3 and Equation 4 can be solved uniquely for $V_x$ as a function of $V_c$, $P_1$, $P_2$, $P_3$, $T_1$ and $T_2$. Note this method of taking temperature differences into account is not very accurate because there is a temperature gradient in the gas that is not generally well represented by simply the weighted average of the two temperatures. A better approach is to take several temperature measurements along the portions of the temperature gradient and assign weighted temperature values based on the local relative volume associated with each local temperature measurement. Note that if there are valves and sections of tubing between the unknown volume to be calibrated and the calibration system, then each section must be isolated and calibrated in a step-wise manner. The displacement volume associated with a closed valve is automatically included in the calibration of the volume behind such a valve.

The known displacement volume calibration sample method is similar to the pressure calibration method except that instead of using a secondary calibrated volume as in Figure 13, a cell of unknown volume is used. A sample well known density (and when weighed, volume) is placed in the cell. The same measurements are performed as described above. Once with the sample in the cell and once without the sample in the cell. Then $V_x$ is solved for using the two measurements and the difference in $V_c$ being equal to the volume of the standard sample. An example of such a standard sample would be a large single crystal of silicon.

### 4.3.3 Pressure

Most of today’s modern pressure transducers have associated electronics that linearize the transducers output to the applied pressure. This generally only requires span (voltage, current or resistivity output versus pressure) and zero adjustments. Typically both will vary with temperature, time, and use. Spans can be determined from the comparative testing described below. The standard procedure for calibrating a mass balance is:

1) Perform a series of pressure transducer readings (5 minimum) over the transducer’s full range of pressure with simultaneous measurements of the same gas pressure against a NIST traceable (or national equivalent) standard pressure transducers.

2) Perform a series of pressure transducer readings (5 minimum) over the transducer’s full range of pressure with simultaneous measurements of the same gas pressure against a NIST traceable (or national equivalent) dead weight pressure calibrator.

3) Zeros are usually adjusted under vacuum by setting the transducer’s final output (post electronics and post software) to zero (0.000... to within appropriate significant digits).

### 4.3.4 Temperature

It is recommended that NIST standard procedure be followed for making temperature measurements and calibrations. A document detailing thermocouple
calibrations can be downloaded from the NIST Weights and Measures website.\textsuperscript{17} Checking temperature measurement accuracy against boiling (distilled) water and ice (corrected for barometric pressure) is a recommended interim test.

### 4.3.5 Time

It is recommended that NIST standard procedure be followed for making time measurements. Documents detailing these procedures and calibrations can be downloaded from the NIST Weights and Measures website.\textsuperscript{18}

### 4.3.6 Other Calibrations

A variety of other devices may be employed for measuring gas sorption or other properties of hydrogen storage materials. Most devices or instruments will come with appropriate calibration instructions. Detailed information concerning calibration theory, regulations, and procedures can be found through most scientific societies and national standards institutions.

## 5 Methods of Measurement

There are a number of measurement methods that can be used to investigate hydrogen storage materials and systems. Gravimetric and volumetric methods are the two primary methods and the most robust in terms of depth of analysis; temperature-programmed desorption, differential scanning calorimetry, and thermal gravimetric analysis are also used. All measurement methods quantify a change in a measurable property to indirectly calculate the hydrogen storage capacity of a material. It is important to keep in mind that all methods indirectly calculate hydrogen concentration and each have complications associated with indirect methods. For example, characterizing the hydrogen storage capacity of highly porous media has proven difficult using gravimetric and volumetric methods because both require an understanding of the amount of gas displaced, calculated from the skeletal density of the material, and the corresponding buoyancy (gravimetric) and volume calibration (volumetric) effects on measurements.

This section will summarize these most commonly used measurement methods and provides detailed techniques for measuring hydrogen storage properties. The section also provides a series of detailed consideration that may have a significant impact on the performance and accuracy of a measurement. These range from general sample preparation considerations to issues unique to specific types of measurements and instruments.

One clear consideration is that the measurement equipment should be constructed of materials that are both hydrogen compatible and do not have significant hydrogen permeation. Obviously metal that react with hydrogen at low pressures and temperature such as Titanium, Magnesium, Palladium and Vanadium to name a few should not be used. Common copper tubing and brass components are prone to out-gassing impurities that may have an impact measurements. 316L stainless steel is commonly used in many hydrogen
applications. Teflon and many polymer sealing materials have high hydrogen gas permeation rates and are not generally recommended for these measurement applications.

5.1 Static and Dynamic Measurements

Hydrogen storage testing can be divided into static and dynamic testing based on whether or not the sorption/desorption reaction is allowed to reach equilibrium at any point during the experiment. In volumetric and gravimetric methods for example, hydrogen gas can be introduced or removed either by aliquots (static) or through a variable flow or pressure regulator (dynamic). In thermally driven methods such as TPD and TGA, the temperature of the sample can be raised in steps (static) or continuously (dynamic).

In static testing, a sample in equilibrium at a certain temperature, pressure and composition is perturbed by a sudden change in pressure (volumetric and gravimetric methods) or temperature (TPD and TGA). The sample is allowed to reach equilibrium at some new temperature, pressure and composition before it is perturbed again. This process continues until the sample is fully sorbed or desorbed. This step-by-step, or equilibrium state-by-equilibrium state process allows static testing to be used to determine the thermodynamic properties of a material by van't Hoff diagram analysis. Static testing is the most frequently reported testing method in literature when using volumetric and gravimetric methods.

Dynamic measurements are characterized by continuous changes in pressure and temperature and do not allow for equilibrium. Although they are not ideal for investigating purely thermodynamic properties, they can provide information on the kinetic activity of a material. When hydrogen flows at a constant rate into an evacuated hydrogen storage system, the pressure in the system increases linearly until the material begins to sorb. After this point, the pressure in the system is a function of the material's ability to sorb hydrogen; if the system is to remain at constant pressure across the pressure plateau in Figure 14, the rate of hydrogen flowing into the system must be equal to the rate of hydrogen sorbed by the material.

Dynamic measurements are more useful for evaluating a material's hydrogen storage performance during application than static measurements. For example, fuel cells require hydrogen at a specified constant pressure to operate efficiently and safely. In order to supply hydrogen at constant pressure, a regulator is connected between the FC and the hydrogen storage system through which the flow rate varies. Dynamic measurements allow testing of storage materials at flow rates representative of the flow through the pressure regulator in the fuel cell/storage system and the dynamic pressure limits associated with such flow rates. These results are highly dependent on the impact of heat transfer on sorption/desorption kinetics.
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5.2 dC and dP Dosing Methods

In isothermal testing, the sample is charged and discharged by changes in hydrogen pressure in the sample holder. Hydrogen can be added step-by-step in aliquots or continuously through flow or pressure regulators. These techniques are used to provide static and dynamic methods of storage testing. Static and dynamic dosing can also be referred to as dC (differential concentration) and dP (differential pressure) dosing, respectively. In dC dosing, a specific amount of hydrogen calculated from the temperature, pressure and volume of the aliquot is dosed to the sample. In dP dosing, a flow or pressure regulator is used to increase the pressure in the sample cell a specific amount per dose. The terms dC and dP originate from the relationship between the different types of dosing and PCTs; dC dosing steps along the PCT concentration axis and dP dosing steps along the PCT pressure axis.

Volumetric instruments typically take steps in concentration (dC dosing) whereas Gravimetric instruments generally take steps in pressure (dP dosing).
Figure 15. PCT of classic metal hydride material with plateau. dC (squares) and dP (red circles) dosing provide different information in solid solution (vertical) versus hydride formation (horizontal) portions of the PCT phase diagram.

Figure 15 shows the relationship between dC and dP dosing and PCTs for a metal hydride. dC dosing (squares) enables limited investigation of the solid solution phases, represented by the regimes with greater slope at low and high concentrations, compared to dP dosing (circles). However, the reverse is true for the investigation of transitional regimes at intermediate concentrations. dC dosing can provide much greater information across plateaus in metal hydrides and the saturation regimes of physisorbing materials. A common complaint in hydrogen storage testing is incomplete or uninformative investigation of metal hydride plateau regimes due to dP testing. In some cases where the plateau slope is very small, the plateau may be missed altogether.

An advantage of the dC stepping of the volumetric method is that samples can be prepared to specific hydrogen concentrations. This may be very useful for ex-situ examination of the materials using other analytical techniques.

Figure 16. Example of PCT with missing plateau information due to inherent limitation in dP dosing.
5.3 Volumetric Method

The volumetric method of hydrogen storage measurement, also known as the manometric method, or as Sievert’s method in honor of the German chemist of the same name, uses temperature-pressure-volume correlations to determine hydrogen concentration and the storage properties of a material. A generalized volumetric system with commonly employed components is shown in Figure 17.

![Figure 17. Multipurpose Gas Sorption/Desorption Apparatus. The pressure gauge on the right is only required when using the flow controller.](image)

The apparatus consists of a gas reservoir connected to a specimen reactor. Because the volumetric method measures concentration indirectly through temperature-pressure-volume correlations, the volumes and temperatures of the reservoirs and sample holder (system) must be known in advance. A thermocouple should always be positioned in as close a contact as possible with the sample. The system volumes are carefully pre-calibrated and the reservoir and sample holder are maintained at constant (but not necessarily equal) temperatures using an external temperature controller. By fixing volume and temperature, reservoir and sample holder pressures can be measured using pressure transducers to provide isothermal pressure-concentration data.

The volume of the sample holder must be calibrated while filled with sample in order to get an accurate measurement of the free gas volume in the system. Although a generally straightforward procedure for chemisorbing media, calibration can be tricky for highly porous and nano-structured media at low temperatures due to inaccurate or incomplete skeletal density information and the possible physisorption of helium used to perform the calibration. Note however, that it is unlikely that even high-surface-area materials will have significant helium adsorption below 5 bar and at 77K, and certainly not at significantly higher temperatures. I can be verified that helium does not adsorb by measuring the sample holder volume with the sample (dead volume) as function of the helium expansion pressure at temperature. Materials that do not adsorb
helium will likely show a constant dead volume as function of pressure. Similarly, in a gravimetric measurement, such material would also exhibit a linearly decreasing mass response (buoyancy) as function of helium pressure.

Volumetric testing requires accurate measurement and control of the instrument and sample holder temperatures and the associated temperature gradients. The temperature of the instrument should be controlled in order to minimize temperature fluctuations due to external sources such as room heating, ventilation and air-conditioning. The temperature of the sample can be precisely controlled below room temperature by cryogenic cooling accompanied by PID (Proportional Integral Differential control) heating or above room temperature by PID heating and appropriate insulation.

To test a sample using the static method, the gas reservoir is filled with H₂ to a pressure P and then allowed to react with the specimen reactor by opening the top valve. When filling the gas reservoir, sufficient time should be allowed for the gas to come to thermal equilibrium with the tubing and vessels before a pressure reading is taken. An equilibrium pressure P’ is reached between the gas reservoir and the specimen reactor that, once paired with temperature, initial pressure and volume information, can be used to precisely determine the amount of hydrogen sorbed by the material. Dynamic testing allows the slow and continuous introduction of hydrogen to the specimen reactor. Hydrogen gas from the gas reservoir flows into the specimen reactor via the electronic flow controller while continuous pressure readings are taken by the pressure gauges attached to the reservoir and reactor. The pressure, temperature and volume data are analyzed to generate dynamic (non-equilibrium) pressure-composition isotherms.

One intrinsic advantage of the volumetric method is that, unlike gravimetric methods, the quantity of gas dosed to or from the sample can be small compared to the total capacity of the sample. Thus, the method allows the direct preparation of sample of known hydrogen contents. This may be useful for doing in-situ or ex-situ secondary measurements such as X-ray or neutron diffraction, NRM, IR, Raman, etc.

5.3.1 Differential Pressure Method

The differential pressure method is a modification of the standard volumetric testing method and is designed to increase the accuracy of Sievert's measurements. An example of a differential pressure method apparatus is shown in Figure 18.
The sample holder and a blank are maintained at the same temperature and are initially under the same pressure with the two valves in Figure 18 open. Immediately after dosing, the valves are closed, isolating the two volumes (sample holder, blank and associated tubing) from each other except for the differential pressure transducer connecting them. The differential pressure transducer measures the pressure change between the two volumes caused by the sorption/desorption of hydrogen by the sample. While adding significant complications, by measuring pressure with a differential pressure transducer instead of an absolute pressure transducer (gauge), the differential pressure method can take some advantage of the higher accuracy afforded to percent reading pressure transducers at low pressures (see section 4.1.2).

The most important consideration when testing with the differential pressure method is thermal equilibrium; the sample holder and the blank must be maintained at exactly the same temperature for the duration of the experiment for accurate isothermal data. This is often accomplished by placing the two volumes together in a heater or bath. Unfortunately, even small temperature variations like those caused by endothermic or exothermic reactions can lead to pressure fluctuations that can impair the accuracy of the differential pressure method. Excellent heat transfer and thermal management capabilities are essential to accurate testing using the differential pressure method.

### 5.3.2 Data Corrections

Volumetric systems determine the concentration of hydrogen in a material from changes in the concentration of hydrogen gas in the reservoir/sample holder system. The concentration of hydrogen gas in the system is determined by knowledge of the pressure, temperature and volume of the reservoir and sample holder. Pressure changes are measured on a pressure transducer inside the volumetric equipment that is at the same temperature as the instrument. Ideally, the system volumes are calibrated and the temperature is known and constant throughout so that the measured pressure is directly related to the concentration of hydrogen gas in the system. Unfortunately, temperature can vary both spatially
and temporally and can be difficult to monitor because it requires many sensors to map the true temperature gradients. A common approach is to divide the instrument into isothermal regions, but care must be taken to validate this assumption to avoid the contribution of thermal effects on kinetics (see Figure 19). For example, if a sample is held at 77 K, the pressure is the same throughout the system. However, the hydrogen gas density is greater in the fraction of the system at 77 K. The volume fraction at 77 K depends on the system design, the type of sample holder used and the skeletal density of the sample.

![Figure 19. Schematic of temperature regimes in a volumetric instrument. The volume of gas at instrument temperature and at sample holder temperature varies depending on the system design and the type of sample holder used.](image)

There are two data correction methods that can be used to account for temperature and hydrogen (gas) concentration variation in the reservoir/sample holder system; both involve calculating 'apparent' volumes that more closely represent the actual concentration of gas in the system. The apparent volume corrections can be applied to raw data after experimentation.

### 5.3.3 Temperature Correction

When the gas in a volumetric instrument is not all at one constant temperature, there will be a density gradient in the gas which must be accounted for in the measurement of hydrogen uptake or release. This is known as the “Temperature Correction” or “Apparent Volume Correction”. This typically occurs because the sample is held at a temperature which is different than the gas dosing reservoir. Below we describe two methods for compensating for this gas temperature gradient. It is important to note that the temperature and the volume of the sample cell has a large impact on this correction and should be considered in the
whole design of the system. While the corrections are generally small terms, in the present context, the volume and temperature differences can be substantial.

The method referred to here as the temperature correction method uses an empirically determined temperature correction factor to calculate the true hydrogen gas content in the system. In practice, a temperature correction factor does not affect the temperatures used in the concentration correlation equation (equation of state) but rather the volume used. It is determined from the volume fraction of the system at non-ambient temperatures and is used to define an apparent volume. In this analytical approach the “Apparent volume” of the sample holder $V_{SH,App}$ is determined at any sample temperature $T_{SH}$ using the room temperature volume calibration of the sample holder (with sample) $V_{SH}$ corrected by the volume weighted measured temperature differences between the reservoir and sample holder (see section 4.3.2). This apparent volume is then applied to the equation of state to determine the hydrogen content in the total volume of the system.

The process for applying a temperature correction is straightforward. An inert (non-sorbing) material of similar physical properties as the sample (density, size and thermal conductivity) is tested at identical conditions to the actual sample experiment. As an example, for the hydrogen uptake measurements of a high-surface area activated carbon sample, solid (non-porous) graphite pellets of the same mass could be used as the blank or control sample for determining the temperature correction factor that will be applied to the actual adsorption and desorption data. To minimize the number of free variables associated with the temperature correction factor the sorption testing on the blank sample should be performed under identical conditions as the sorption measurements on the material of interest. Using the same example, if one is investigating the hydrogen storage properties of a sample at 77 K, the inert material should be tested at 77 K with the exact same reservoir volume, tubing and sample holder. It is important the inert material have the same skeletal (gas displacement) volume as the sample, as well. Because the temperature correction factor reflects the volume fraction of the system at non-ambient temperatures, it is necessary that the volumes of gas at sample and reservoir temperatures are identical in the inert material and sample experiments. Once volumetric equivalency has been ensured, the inert sample is tested in order to establish a baseline with which to compare the performance of the sample. Theoretically, capacity should be zero at all pressures for the inert material. In practice, this first-order approximation may be sufficient for the temperature correction, however, there will be a non-zero effect of pressure on the temperature (gas density) gradient (see Figure 20).
A temperature correction factor is applied to the raw data of the inert material and varied until the associated apparent volume yields zero sorption at over the pressure range of interest. The temperature correction factor that yields zero sorption for the inert sample is then applied to the raw data of the actual sample.

Figure 21 presents an extreme example (most difficult scenario) of a low capacity, very low density material (0.1 g/ml) showing raw and temperature corrected data for a physisorption at 77 K. The temperature correction was taken from the data in Figure 20.
5.3.4 Volume Correction

A volume correction is an empirical correction for the temperature based gas density gradient based on performing an inert gas (usually helium) volume calibration measurement of the sample holder (with sample) at the testing temperature. This calibrated volume is the volume at room temperature required to contain the same amount of hydrogen gas as the volume of the sample holder at testing temperature. It provides a direct method to determine the “Apparent Volume”. This simple relationship is defined in Equation 5.

\[
 n_{SH} = \frac{PV_{SH}}{RT_{SH}} = \frac{PV_{SH,App}}{RT_R} \rightarrow V_{SH,App} = \frac{T_R}{T_{SH}} V_{SH}
\]

Where \( n_{SH} \) is moles of gas in the sample holder, \( R \) is the gas constant, \( P \) is the measured pressure, \( V_{SH} \) is the volume of the sample holder, \( V_{SH,App} \) is the “Apparent volume” of the sample holder, \( T_{SH} \) is the temperature of the sample holder, and \( T_R \) is the temperature of the reservoir (instrument). While this method is more direct and easier to perform that the Temperature Correction method described above, it may be inappropriate to use with microporous materials that exhibit significant helium adsorption at low temperatures.

To apply the empirical volume correction, one must use the apparent volume (determined at the testing temperature) along with pressure and reservoir
temperature data in the concentration standard calculation of hydrogen concentration. Reservoir temperature data must be used exclusively because the temperature variation in the system is accounted for in the apparent volume correction. Figure 22 shows the same data as that of Figure 21 with both the Temperature Correction method and the Volume Correction method applied to the data. Note that the very low density of the sample in this example (0.1 g/ml) combined with the extreme temperature difference (sample holder 77K and reservoir 302K) causes a very large correction to the data. The difference in the corrected results demonstrates the limits of accuracy of these two different correction approaches under the most extreme conditions.

Figure 22. PCT for a physisorbing material at 77 K showing the raw data, temperature corrected, and volume corrected data. This is the same raw data that is used in Figure 21.

5.3.5 Compressibility Factor

The volumetric method uses gas laws to determine the number of moles of hydrogen in the system based on pressure, temperature and volume measurements. A gas law can be written in general form as

Equation 6

\[ PV = nRTz(P,T) \]

The measurement and sources of error for pressure, temperature and volume are discussed in the variables section 4.1. The compressibility factor, z, accounts
for the non-ideality of the gas at a given temperature and pressure. For an ideal gas, the compressibility factor is unity and Equation 6 collapses into the familiar ideal gas law. The compressibility factor for hydrogen is taken from compressibility data and applied as a function of the measured pressure and temperature (Figure 23). It is recommended to determine compressibility factors using a standard reference database such as the NIST12 database for fluid properties.\(^{23}\)

![Figure 23. Compressibility factor \(z\) for hydrogen as a function of temperature and pressure.](image)

### 5.3.6 Gas Impurities

Gas impurities can be introduced into a volumetric system through the use of impure hydrogen gas and leaks during desorption experiments. In volumetric measurements the sample’s hydrogen storage capacity is determined by measuring pressure drop/increase due to sorption/desorption. Since the partial pressures of any low level impurities are a minute fraction (parts per million to perhaps parts per thousand) of the measured hydrogen pressure, the impact on the total capacity caused by impurities is essentially negligible. In the volumetric method, the impurities may still impact capacity and kinetics through surface effects such as poisoning and retardation; however, the measurement will be truly representative of the material’s performance in the presence of impurities.

Poisoning and retardation (see Capacity and Kinetics chapters) occur when gas impurities react with the sample to inhibit the total hydrogen sorbed/desorbed and/or the rates of reaction, respectively. These effects cause decreases in the measured storage capacity and kinetic activity of the sample material and may incorrectly indicate the sample material is not suitable for hydrogen storage.
applications. It is important to not eliminate a sample from consideration until gas impurity effects are mitigated. Impurities may simply cause “gas blanketing” which reduces sorption rates but does not necessarily impact capacity or desorption rates.

5.3.7 Instrument Temperature
The temperature of the instrument is an often-overlooked factor in taking accurate measurements. In most volumetric equipment, the dosing volume is inside the body of the instrument and is not in thermal equilibrium with the sample and sample holder. Therefore, knowledge of the instrument temperature and the temperature gradient between the instrument and the sample holder is necessary to accurately relate the pressure reading on the pressure transducer (inside the instrument) and the equilibrium pressure in the sample holder. Instrument temperature management is most often performed by a temperature control system inside the instrument housing.

5.3.8 Leaks
Hydrogen can be leaked to the environment (external leak) and across valves inside equipment (internal leak) during hydrogen storage testing. Both types of leaks during volumetric testing cause unexpectedly high capacities and strange kinetic behavior. During sorption testing above ambient pressures, gas leaked to the environment is assumed to be sorbed to the sample. Leaks during sub-ambient desorption also affect the capacity by increasing the reading of the amount of gas desorbed. In kinetic testing, leaking sorption/desorption profiles appear linear in time, as opposed to true sorption profiles that have a curved profile and eventually reach an equilibrium pressure. An equilibrium pressure of exactly 1 atmosphere and extraordinarily large hydrogen storage capacities are highly unlikely and should be double checked. Leaks also expose the sample and system to gas impurities that can affect property measurements and compromise samples.

It is advisable to check hydrogen storage testing systems for external and internal leaks every 20-40 experiments by running an empty sample holder. It is also good practice to check for external leaks on exposed tubing and joints using helium leak detection or a hand-held flammable gas detection instruments. These leak precautions apply to both volumetric and gravimetric testing methods. An important note is that certain types of gas fittings may have “virtual leaks” that can not be picked up by external leak checking. Therefore, conical gas fittings are not recommended.

5.3.9 Reference Point
The volumetric method measures relative changes in hydrogen concentration calculated from changes in pressure in the reservoir/sample holder system. Because only changes in hydrogen concentration can be calculated, volumetric methods require a reference point with which to calculate absolute concentrations. The most obvious and most commonly used reference point is an
uncharged sample under vacuum, which corresponds to (0 bar, 0 wt.% H₂). From this reference point, absolute concentration with respect to an indirect variable can be determined. Note that the quantity of hydrogen absorbed and desorbed in a closed set of isothermal cycles should be consistent. Thus, a second absorption would commence at a capacity that is the end point of the previous desorption.

Using mass balance of the gas by knowing the total gas volume, and measuring pressure and temperature, the exact concentration of hydrogen in a sample can be determined. In some cases, where the uptake (or release) of hydrogen from the sample is slow enough, it may be possible to adjust the concentration vs. time measurement to a concentration of zero (or fully charged) at time = 0.0 (the point at which the valve between the reservoir and sample holder was opened). With accurate volume calibration and pressure and temperature measurements the calculated concentration at time = 0.0 should likewise be zero (or fully charged).

### 5.3.10 Gas Heating and Cooling effects

An important temperature effect is the adiabatic gas expansion effect. While, hydrogen is unusual in that it has a range that exhibits reverse Joule-Thompson behavior the overriding thermal effect will be cooling of the gas on expansion and heating on compression. This effect can often be seen in the initial part of a kinetics measurement when hydrogen is dosed at high pressure from a calibrated volume to the sample held at low pressure or vacuum. There is a small temperature drop in the gas that is seen as a pressure drop and consequently what appears to be initial sorption. The gas expansion effect impacts desorption kinetics measurements in a similar manner: as the hydrogen desorbs from the sample and expands into the vacated volume, the temperature and pressure spike can be mistaken as increased desorption. These transient effects usually last only a few seconds or less depending on the thermal mass and internal surface area of the gas handling system and sample holder (see Figure 24). However, with large quantities of gas and large pressure differentials, it may take the gas, vessels and tubing several tens of minutes to return to thermal equilibrium. This effect disappears as the gas comes into thermal equilibrium with its surroundings. One should be careful to not misinterpret this temperature relaxation behavior as gas sorption by the sample. This is one good reason why it may be quite important to have a temperature sensor in close proximity if not intimate contact with the sample. To be able to determine when transient temperature related effects have subsided, one should ideally monitor the gases return to return to isothermal conditions.
5.3.11 Sample Size

During single-dose measurements, the largest practical calibrated volume available limits the amount of material that can be tested. At a more sophisticated level an automatic volumetric doser with multiple volumes can be used to perform such kinetic measurements on wide range of sample sizes.

Small samples (less than 100 mg) exhibit small pressure changes due to sorption/desorption. It is therefore important to minimize the background pressure noise in order to measure the small pressure changes associated with small sample sorption/desorption. Also, small samples require that the instrument must be capable of high sensitivity both in the pressure sensor itself and the volume size. In particular, volumes should be match in size relative to the sample volume.
5.3.12 Skeletal Density
The low skeletal density of highly-porous materials can cause difficulties in accurately determining hydrogen storage properties. This is primarily due to two factors: first, knowledge of the amount of gas displaced by the sample is required to determine the free volume in the sample holder. This can be difficult to quantify for samples of low skeletal density that have the potential to adsorb helium, the gas commonly used to calibrate volume in a volumetric system. Helium adsorption during volume calibration may lead to unexpectedly large volume calibrations, an effect that becomes especially pronounced at cryogenic temperatures. Second, low skeletal density makes it difficult to test a sufficient mass of sample for an accurate measurement. Insufficient sample mass leads to small pressure changes and, therefore, less accuracy due to pressure transducer sensitivity. It is important to have enough sample mass that pressure changes are significantly above the level of resolution of the pressure transducers.

When the skeletal density of the sample is very low and difficult to determine, the two factors discussed above lead to opposite conclusions regarding the amount of sample necessary for accurate testing. On the one hand, it is desirable to minimize the error associated with the free volume in the sample holder by using a small sample volume compared to the overall volume of the sample holder. This suggests using as small a sample as possible. Recent literature from Gray et al. supports this view and finds that the volume calibration error associated with skeletal density is proportional to the square of the volume of the sample. On the other hand, pressure transducer sensitivity suggests a larger sample will lead to more accurate measurements due to the larger pressure changes associated with sorption and desorption. These opposing conclusions engender a tradeoff between small and large sample sizes in this special case; one possible solution is to test both small and large amounts of material and compare the results.

5.3.13 Calibration Volume Changes
The effects of miss-calibration of the reference volume and a possible volume change with pressure can contribute error through the volume term; the latter effect is only considered significant in the sensor head, the valves with their accompanying diaphragm or bellows components and changes in the sample volume itself. Pressure-induced volume changes in the tubing, calibration volumes, and gas component are generally considered negligible.

5.3.14 Volume Dilatation of Sample
Some hydrogen storage materials, especially metal hydrides, undergo volume dilatation and contraction during hydrogen sorption and desorption. The volume of a system (sample, sample holder and appropriate tubing) is typically calibrated using helium before or after experiments are conducted when the sample is in the uncharged state. Because volumetric methods require knowledge of the volume of a system in order to calculate concentration, changes in the volume of the sample during testing are not accounted for in the calculations and have the
potential to cause errors in measurement. Volume dilatation and contraction of a sample is often negligible when compared to the overall volume of the system. However, dilatation effects should be considered when testing large quantities of materials that exhibit large volume expansion coefficients.

5.4 Gravimetric Method

The gravimetric method of measurement uses weight changes measured on a balance to determine concentration and the storage properties of a material. A schematic of a simple gravimetric system is presented in Figure 25. Note additionally, that gravimetric systems are also often configured as flow-through systems, in which case there is also a gas exit port attached to the chamber through a pressure control device.

![Figure 25. Schematic of counterbalanced gravimetric method system for hydrogen storage testing.](image)

Before gravimetric testing, the weight of the anhydride sample is measured and the sample is placed on one end of a symmetric microbalance in a sample holder. In more advanced gravimetric systems, an inert tare with the same weight and comparable density to the unhydried sample is placed on the other end of the microbalance to provide a counterbalance. The tare is designed to minimize the effects of buoyancy caused by the hydrogen gas displaced by the sample volume and must be inert in a hydrogen atmosphere.

After the chamber containing the gravimetric equipment is evacuated, hydrogen from an external hydrogen source enters the chamber and is generally increase in incremental pressure steps as it is sorbed by the sample in the sample holder. The microbalance is typically equipped with an electronic circuit that measures the strain on the balance material that is directly related to the change in weight of the sample. This information combined with pressure and temperature readings provided by a pressure gauge connected to the vacuum chamber and a thermocouple located next to the sample holder can be used to construct various forms of hydrogen storage properties.

Measurements can generally be made under isobaric conditions with no loss of sensitivity and modification of the thermodynamic driving force or altering the
kinetics by significant pressure changes. This can be accomplished even in static experiments by providing makeup gas during absorption or bleeding the evolved gas during desorption.

5.5 Thermal Gravimetric Analysis Method

We make a distinction here between the Gravimetric instruments described above and Thermal Gravimetric Analysis (TGA) in that Gravimetric instruments generally operate under isothermal conditions with a controlled over pressure of gas. Whereas, TGA’s typically operate under vacuum or low-pressure flowing gas conditions and ramping temperatures. Some equipment, however, may be setup to operate in all of these modes. TGA is a thermal analysis technique often used in conjunction with DSC to determine the hydrogen storage properties of a sample. The first documented use of TGA was in the study of the efflorescence of hydrated salts in 1912 but it is now traditionally used in the quantitative investigation of decomposition reactions. TGA is experimentally similar to DSC but instead of measuring the heat flow as function of temperature, TGA measures sample weight as a function of time or temperature. This is accomplished by placing the sample in an environment that is heated or cooled at a controlled rate and measuring the weight change.

The equipment necessary to perform TGA consists of an accurate balance, a programmable furnace, a reaction chamber and a data collection system. As the temperature inside the furnace and reaction chamber changes, the balance measures the variation in weight due to various chemical reactions including dehydriding. TGA is affected by many of the same issues as the gravimetric testing method such as buoyancy and mechanical disturbances. The buoyancy force exerted on the sample by the displaced fluid varies with temperature and must be taken into account either during the experiment, e.g. the tare technique in the gravimetric method, or during data analysis; mechanical disturbances must be minimized through leveling and anti-vibration supports. Another complication TGA shares with the gravimetric method is how to determine the sample temperature. Using a thermocouple to directly measure the temperature affects the weight reading of the sample. Consequently, care must be taken to place the thermocouple in such a way that it accurately reads the sample temperature but does not affect weight readings.

Scanned temperature experiments (TGA) are often done using flowing inert gas, such as argon. Although useful for comparing different materials (catalyzed vs. uncatalyzed, for example), desorption occurs into an essentially H$_2$-free environment, and thus it does not reflect the actual conditions encountered in a real storage system application, where there is always H$_2$ gas present at pressures at or exceeding 1 bar. Interpretation of desorption temperatures thus must be done carefully. For this reason TGA may be used as a screening tool, but not as a technique to rely on for realistic system desorption temperatures. When one really wants to know practical desorption temperatures, the
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experiment must be conducted by flowing H₂ gas into the TGA at the particular pressure of interest (say, 1-8 bar).

5.5.1 Buoyancy
The weight of the fluid displaced by the sample exerts an upward force on the sample holder that can affect the measured weight change appreciably. The degree of the buoyancy effect is proportional to the volume displaced by the sample and the density of the surrounding gas. The upward force due to buoyancy on a symmetric microbalance like that in Figure 25 is

\[
F_b = g \rho \Delta V = g \rho \left( \frac{m_s - m_t}{\rho_s - \rho_t} \right)
\]

where \( \rho \) is the density of the hydrogen gas, \( m_s \) and \( \rho_s \) are the mass and density of the sample and \( m_t \) and \( \rho_t \) are the mass and density of the tare which may include terms representing the mass and density of the sample holder (bucket, pan) and the hang-down wires. At elevated pressures \( \rho \) is not only a function of pressure and temperature but must also include corrections for non-ideal gas behavior.

5.5.2 Gas Impurities
Gas impurity effects are generally much more severe in gravimetric methods than volumetric methods because samples are much smaller and the total amount of gas surrounding the samples is many orders of magnitude larger than in the volumetric case. This means that even at low impurity levels the relative exposure of the sample to impurities may be orders of magnitude higher. In addition, the weight increase due to contaminants can not be distinguished from increase due to hydrogen sorption and therefore gas impurities have the potential to greatly affect measurements; in the volumetric system, the partial pressure of contaminants is typically negligible compared to hydrogen and therefore doesn’t affect measurements to the same degree. This makes it important to reach relatively high levels of vacuum in order to take accurate measurements.

In contrast to the volumetric method, the effects of gas impurities tend to lead to false positives, measurements that indicate a sample is a better hydrogen storage material than it really is. This is because weight increase due to sorption of impure species is mistaken for hydrogen sorption, although in reality the sorption of hydrogen is decreasing (poisoning). Once again, it is necessary to minimize gas impurity effects during testing in order to gain an understanding of a material's true hydrogen storage properties. When a gravimetric system is used in a flow-through mode, the sample is continuously exposure to fresh source gas, thus impurities in the source gas must be minimized. Use of high purity gas, non-permeable and non-contaminating tubing such as stainless steel, and, if necessary, passing the source gas through water and oxygen scrubbers will mitigate the effect of impurities.
The effects of impurities is increased at low temperatures where a (porous) sample can act as “getter” for molecular species condensing at temperature above the particular cryogenic temperature used. This can lead to large errors in gravimetric measurements. There has been considerable concern in the case of hydrogen storage on carbon nanostructures where important errors can be attributed to impurities in gases, vacuum quality and gas purity.

Because gravimetric measurements are much more sensitive to co-generation of other gases (evolved mass, not evolved moles) they often send up warning flags (like when the weight loss exceeds the total hydrogen content) even when the evolved gas is not being monitored.

5.5.3 Heat Transfer
Heat transfer effects are critically important to hydrogen storage measurements and are resolved from temperature data collected by thermocouples. Ideally, the thermocouple measuring the temperature of the sample material and the sample are in perfect thermal contact, allowing the thermocouple to best measure the temperature of the sample. In gravimetric systems, this is not the case. A thermocouple cannot be attached directly to the sample because it will affect the sample’s measured weight and is therefore placed near but not touching the sample. This makes it difficult to measure the true temperature of the sample and resolve heat transfer effects. In cases where the thermal gradients in the vicinity of the sample holder are large enough that the temperature reported by the thermocouple does not represent the true temperature at the sample holder, calibration with known temperature standards is necessary. System design that minimizes thermal gradients near the sample is highly desirable.

5.5.4 Leaks
The effects of leaks on the gravimetric method vary widely depending on the type of experiment. One advantage of the gravimetric method is that leaks that occur during sorption testing have little effect on the measurement because the gas lost to the environment is not included in the indirect calculation of concentration like in the volumetric method. However, the flow of gas in the instrument caused by a leak can affect the forces measured by the cantilever. Leaks during gravimetric desorption testing can affect the sample material as well as the measurement. Sub-atmospheric pressures sometimes used for desorption promote gas contamination and can lead to poisoning, retardation and other issues associated with gas impurities. This is a serious consideration when evacuating the sample holder or degassing a sample at elevated temperatures under vacuum.

5.5.5 Sample Transfer
Some hydrogen storage materials are sensitive to common gases and vapors like those found in the atmosphere. Physisorbing materials readily adsorb atmospheric gases and vapors and must be baked out under vacuum for a sufficient time (3-12 hours) before hydrogen storage testing. Other materials,
such as magnesium-nickel metal hydrides, oxidize rapidly and can not be exposed to oxygen. Both cases necessitate ‘airless’ sample transfer between storage containers and the hydrogen storage instrument. Airless sample transfer is difficult in gravimetric systems because the sample must be loaded onto a microbalance, a process that often exposes the sample to atmosphere. Some gravimetric and volumetric systems offer airless sample transfer devices but the process is typically awkward and while special designs make this manageable, typical systems are prone to mistakes that can compromise a sensitive sample. One approach is to incorporate the entire instrument inside an inert gas glovebox.

5.5.6 Volume Dilatation
Volume dilatation during sorption affects weight measurements in gravimetric systems by changing the buoyancy forces acting on the microbalance. As the volume of the sample increases, the amount of hydrogen displaced by the sample also increases. This increases the buoyancy force acting on the volume that in turn affects the resultant force measure by the microbalance. This is a particularly challenging consideration because taring the microbalance with an inert material, a common technique used to minimize buoyancy effects, will not account for buoyancy effects due to volume dilatation. If necessary, these can be taken into account by post-data acquisition processing provided a measurement or reasonable estimate of the sample density in the sorbed state is available.

5.5.7 Thermal Gradients and Gas Flow Forces
Thermal gradients in the gas phase inside the chamber cause natural convection currents that can affect the microbalance, leading to noisy weight measurements. Once perturbed, the microbalance can take a long time to stabilize; therefore it is important to heat and cool the system slowly to minimize thermal gradients. Careful system design, including controlled flow channels and baffles to prevent establishing large convective cells, can help mitigate against thermal gas flow noise. In flow-through systems, gas flow drag forces on the sample holder and hang-down wires must be taken into account. These require either running a second, null experiment under identical conditions in the absence of the sample as a baseline for comparison of the sample run, or else calibration of the flow forces under various conditions of flow velocity (which generally changes with the kind of gas, temperature, and pressure).

5.6 Temperature-Programmed Desorption Method
Temperature-Programmed Desorption (TPD) refers to a wide range of experimental methods that rely on temperature variation and generally include mass spectroscopy to investigate and quantify desorption reactions. The technique can be used for both reversible and irreversible processes, with the latter referred to as Temperature-Programmed Reaction Spectroscopy (TPRS).
TPD measurements may be as simple as measuring pressure rise in a volumetric instrument or weight loss in a gravimetric instrument while ramping the sample temperature. However, TPD generally refers to spectroscopic desorption measurements. The basic set-up for TPD techniques is illustrated in Figure 26. The sample is loaded into the experimental apparatus (a temperature-programmed heater contained in a vacuum chamber) and charged with hydrogen until fully loaded. Note that hydrogen loading may also be performed in a separate apparatus prior to putting the sample in the TPD instrument. After the remaining gas has been drawn off, the computer-controlled heater slowly raises the temperature of the sample. This releases hydrogen that is evacuated to vacuum. A mass spectrometer connected to the evacuation line analyzes the relative composition of the desorbed gas and quantifies the amount of hydrogen desorbed by the sample. As with other methods (gravimetric and volumetric) that employ simultaneous mass spectrometry analysis, TPD systems have an advantage, in that it can distinguish between hydrogen and other constituents in the evacuated stream.

Concentration, temperature and time data is obtained through TPD measurements and can be used to determine capacity, kinetic and thermodynamic properties of a material. A common data representation format for TPD experiments is shown in Figure 27.
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The area under the peak at \(~475\text{K}\) in Figure 27 is proportional to the amount originally sorbed to the sample. In the case of full charging, it represents the capacity of the material. Kinetic information is obtained from the contour of the spectroscopic peak and knowledge of the relationship between temperature and time based on the computer program. Lastly, the temperature associated with the peak is related to the reaction enthalpy of hydrogen-substrate desorption.

Unfortunately, TPD measurements can only be done at vacuum due to limitations in analytical equipment, effectively limiting the amount of thermodynamic information that can be collected. Quantitative analysis requires accurate calibration of the mass spectrometer against known flow rates. It is also important to understand that TPD or any dynamic type measurement gives results that are a convolution of both thermodynamic and kinetic properties of the hydrogen storage material. This may lead to misinterpretation of the results. For example, does a decrease in desorption temperature of a material that is modified through the addition of a dopant mean that the thermodynamics of the host material has been altered? Or, is the dopant acting as a catalyst, or a thermal conductor, or modified the materials morphology such that kinetics are improved, increasing desorption rates at lower temperatures? For the development of new (reversible) materials it is important to have information on both the dynamic and equilibrium hydrogen sorption behavior of the storage material.

5.7 Differential Scanning Calorimetry Method

Differential Scanning Calorimetry (DSC) is a thermal analysis technique used to investigate the thermodynamic properties of a material by measuring the energy necessary to maintain a sample material and an inert reference material at the same temperature over a range of temperatures. The relative heat flow to the sample material as a function of temperature can be used to determine thermodynamic properties such as specific heat and enthalpy.

Temperature variation in DSC is controlled by a computer and is typically linear in order to simplify calculation, although nonlinear temperature variation can be used as well. The specific heat at constant pressure $C_p$ of a sample material (on a per mass unit basis) as a function of temperature is determined by the equation

$$c_p(T) = \left(\frac{\partial h}{\partial T}\right)_p = \frac{q(T)}{dT}$$

where $T$ is temperature, $h$ is energy, $q$ is heat flow and $t$ is time. The heat flow can be taken from the data and the temporal temperature variation is based on the computer program controlling the experiment. In this way, the specific heat as a function temperature can be determined. Furthermore, the enthalpies of reactions can be determined from the measurement. For example, the enthalpy of crystallization is calculated by integrating the specific heat capacity determined from Equation 9 between the temperatures at which the heat flow varies ($T_{c,1}$ and $T_{c,2}$).
Equation 9

\[
\Delta H_{\text{crys}} = \int_{T_{c,1}}^{T_{c,2}} C_p(T) dT
\]

The two most commonly used methods for conducting DSC measurements are power-compensation DSC and heat-flux DSC. In the power-compensation method, the sample and reference material are placed in independent, identical furnaces. The furnaces are maintained at the same temperature over a variety of temperatures by varying the power input. The power input and temperature data are used to construct the DSC diagram. The indirect and direct variables in power-compensation DSC are flipped in heat-flux DSC. The sample and the reference material are placed in one furnace and exposed to the same heat flux. The variation in temperature between the sample and reference is used to determine the relationship between heat flux and temperature.

In the context of hydrogen storage, Differential Scanning Calorimetry is primarily used for desorption testing because DSC equipment is not typically designed to handle the high pressures required for sorption with some exceptions. The advantage of DSC over other methods is that other thermal events such as melting or crystal structure changes may be observed. A significant limitation for testing hydrogen storage using DSC is that there is no way to determine the amount of hydrogen desorbed by a sample, only the total enthalpy of a given reaction. For instance, DSC alone would be unable to distinguish between a materials that desorbs 0.1 mol H\textsubscript{2} with an enthalpy of reaction of 30 kJ/mol H\textsubscript{2} from one that desorbs 1 mol H\textsubscript{2} with an enthalpy of reaction of 3 kJ/mol H\textsubscript{2}. It is important to have an understanding of the enthalpy of reaction per mol hydrogen in order to compare thermodynamics across materials. This vital consideration for hydrogen storage should not be overlooked but can be remedied by coupling DSC with quantitative measurements of hydrogen uptake and release.

5.8 Important Consideration: Gas Composition

All testing methods (with the possible exception of TPD spectroscopy) rely on the assumption that the change in the direct variable used for correlation during desorption is due to hydrogen gas alone. Unfortunately, this assumption is not always valid. During desorption, it is possible to evolve gases other than hydrogen by chemical reaction of the sample material itself. These evolved gases can affect pressure and weight measurements and can contaminate experimental data.

Unless the gas evolution properties of the material are well established, for many materials it may be highly incorrect to assume that the change in the [directly measured] correlation variable, such as weight or equilibrium pressure, is due entirely to hydrogen. In particular, this can produce seriously misleading results in making hydrogen storage measurements during desorption. In addition, impurities (water, oxygen...) in the hydrogen gas stream used to charge a sample may be gettered out of the hydrogen, causing a significant weight change that may be misinterpreted as hydrogen absorption. In terms of hydrogen
absorption capacity measurement this contaminated gas problem is much more significant for gravimetric measurements than volumetric measurements because of the ratio of the quantity of gas to sample size and the fact that sample weight change is being equated to hydrogen content. In volumetric measurements the pressure change due to sorption of ppm level impurities would be minimal, if even measureable. Whenever possible we recommend using a secondary technique, such as mass spectrometry, to validate weight loss or gain measurements in gravimetric systems. Volumetric measurements should likewise be tested for evolution of gases other than H\textsubscript{2} which can lead to capacity loss or possibly decreased kinetics with cycling (e.g. nitrogen loss in amides) or detrimentally affect fuel cell catalysts (evolution of ammonia from amides).

One way to account for non-hydrogen gases is to use a mass spectrometer to determine the composition of the desorbed gas. The mass spectrometer is connected to the outlet line of the sample holder and tests a representative sample of the evolved gas, outputting the relative composition of the individual species via their partial pressures. One caveat when using mass spectrometers is the production of a virtual H\textsubscript{2}O signal. Water and air are often present in the background spectrum, especially in the absence of a complete bake-out of the experimental equipment. However, depending on the state of the mass spectrometer being used, the filament itself can be partially oxidized and when testing H\textsubscript{2} for impurities, the H\textsubscript{2} can reduce the filament to create water. Although the mass spectrometer registers the water, it may not necessarily be coming from the material.

In some alanate-amide mixtures, the gas evolved from a sample during desorption is not entirely hydrogen. Trace amounts of ammonia can be found as well, as shown in Figure 28.

![Residual gas analysis of alanate-amide sample discussed in the Kinetics section.](image)

Figure 28. Residual gas analysis of alanate-amide sample discussed in the Kinetics section.
Water is a common background contaminant, but is also found as a contaminant of hydrogen storage materials, generally due to adsorption from air due to improper sample handling prior to testing. The presence of ammonia is much more interesting in this case. Under elevated temperature, the alanate-amide mixture reacted to measureable amounts of ammonia as well as hydrogen, although the exact mechanism for the reaction is unclear. This type of unexpected gas evolution points to the critical need to apply other analysis to validate the gas composition assumption inherent in the testing methods. Simultaneous measurement of gas composition and (inferred) concentration is highly preferred until it can be assured that co-evolution of other species does not occur from the particular material.

6 Summary
It is our intent that this overview of fundamental processes and measurement considerations will aid research and development of new and better hydrogen storage materials. In particular we hope to have clarified some of the recommended practices and caveats in performing high-quality experiments to measure the hydrogen storage properties of advanced materials.
Section 2: Kinetics

Measurement of hydrogen sorption and desorption kinetics is important for practical hydrogen storage system performance, advanced materials development and fundamental research on the mechanisms of hydrogen uptake and release.

In this chapter we will provide an overview of the purpose of kinetics measurements, introduce some modeling of kinetic mechanisms, discuss important considerations that affect kinetics measurements, and present some methods to improve kinetics.

The proper characterization of hydrogen storage properties is one of the most crucial and time-consuming steps on the road to the discovery and development of advanced and practical materials. Because of this, it is important that measurements, including kinetics measurements, are informative, reliable and consistent with accepted standards. The intent of this chapter is to describe some common practices accepted in the field of hydrogen storage and to highlight important considerations both positive and negative that can have a strong impact on kinetics measurements.

1 Recommended Reading

This document is not meant to be a thorough review of the leading work in this field, but rather an overview and series of instructive examples of important issues concerning the measurement of the kinetic properties of hydrogen storage materials. The following resources are a good place to find more detailed information on the key topics.

1.1 Kinetics
The book “Basic Chemical Kinetics” by Eyring, Lin and Lin.\(^{27}\)
The book “Chemical Kinetics and Reaction Mechanisms” by Espenson.\(^{28}\)
The book “Chemical Kinetics and Catalysis.” By, Richard I. Masel\(^{29}\)
The book “Chemical Kinetics.” by J. E.Nicholas\(^{30}\)

1.2 Hydrogen Storage
The book “Hydrogen Storage Materials (Materials Science Forum)” edited by Barnes.\(^{31}\)
The review article “Hydrogen-storage materials for mobile applications” by Schlapbach and Züttel.\(^{32}\)

1.3 Hydrides
The book “Hydrides” by Wiberg and Amberger.\(^{33}\)
Section 2: Kinetics

1.4 Metal Hydrides
The book "Metal Hydrides" edited by Muller, Blackledge, and Libowitz.\textsuperscript{34}
The book "Transition Metal Hydrides" edited by Dedieu.\textsuperscript{35}
The review article "A panoramic overview of hydrogen storage alloys from a gas reaction point of view" by Sandrock.\textsuperscript{5,10}

1.5 Physisorption Storage
The review article "Review of hydrogen storage by adsorption in carbon nanotubes" by Darkrim, Malbrunot and Tartaglia.\textsuperscript{36}
The journal article "Hydrogen Storage in Microporous Metal-Organic Frameworks" by Rosi et al.\textsuperscript{37}

2 Introduction and Definitions
As an introduction, we will briefly review definitions in chemical kinetics in general and then discuss kinetics as it pertains to hydrogen storage materials.

2.1 Chemical Kinetics
The following description of kinetics involves excerpts from general textbooks on the topic. This is intended only as a general introduction to the field. For a more detailed discussion, please see the references at the end of this chapter.

"Kinetics deals with the speed or rate at which reactions proceed and the rate of reaction is defined simply as the rate of change of concentration for reactants and products… The rate of a chemical reaction depends on several factors in addition to reactant concentration. In some cases the rate is influenced by the products, substances such as catalysts, or even chemically inert species. Of greater general importance is the fact that in almost all cases the rate varies with temperature, often very considerably."\textsuperscript{30}

The following factors affect reaction rates:

2.1.1 Nature of the Reactants
"Depending upon what substances are reacting, the reaction rate varies. Acid reactions, the formation of salts, and ion exchange are fast reactions. When covalent bond formation takes place between the molecules and when large molecules are formed, the reactions tend to be very slow."\textsuperscript{38}

2.1.2 Heterogeneous vs. Homogeneous Reactions
The physical state (i.e. solid, liquid, gas) of the reactants is also important. If the state of all of the reactants is the same, they will mix and the result will be a homogeneous reaction. If the reactants are of different phases or for some reason cannot mix, then the reaction is
heterogeneous and limited only to the interface between the reactants. Heterogeneous reaction rates therefore scale with surface area.\textsuperscript{29}

### 2.1.3 Concentration

“Concentration plays an important role in reactions. According to the collision theory of chemical reactions, this is because molecules must collide in order to react together. As the concentration of the reactants increases, the frequency of the molecules colliding increases, striking each other more often by being in closer contact at any given point in time. Imagine two reactants in a closed container. All the molecules contained within are colliding constantly. By increasing the amount of one or more of the reactants you cause these collisions to happen more often, increasing the reaction rate.” An equation that describes the relationship between the rate of reaction and the concentration of all species in the reactor is called a rate equation and is presented in detail in section 3.2.2.

### 2.1.4 Temperature

“Temperature usually has a major effect on the speed of a reaction. Molecules at a higher temperature have more thermal energy. When reactants in a chemical reaction are heated, the more energetic atoms or molecules have a greater probability of colliding with one another. Thus, more collisions occur at a higher temperature, yielding more product in a chemical reaction. More importantly however, is the fact that at higher temperatures molecules have more vibrational energy; that is, atoms are vibrating much more violently, so raising the temperature not only increases the number of collisions but also collisions that can result in rearrangement of atoms within the reactant molecules.” Temperature has a profound effect on reaction rates, most often described by the Arrhenius equation. This is presented in detail in section 3.2.1.

### 2.1.5 Catalysts

A catalyst is a substance the increases the rate of a reaction without modifying the overall standard Gibbs energy change in the reaction… this definition is equivalent to the statement that the catalyst does not enter into the overall reaction; it is both a reactant and a product of the reaction.\textsuperscript{1} A catalyst increases the rate reaction by providing a different reaction mechanism to occur with a lower activation energy (Figure 29).

“Ostwald defined a catalyst as ‘a substance one adds to a chemical reaction to speed up the reaction without the catalyst undergoing a chemical change itself.’ …this definition is not quite accurate. Catalysts do undergo chemical changes during the course of reaction. It is just that the changes are reversible, so that the catalyst is not consumed as the reaction proceeds. Example of catalysts include the acids in your stomach that you use to break down food, and the enzymes that people
put in detergents to make the detergent work better. Most chemical processes use catalysts at some stage in the production process. …One of the key features of catalysts is that they change the form of the rate equation. In the gas phase, rates of reactions are often proportional to the reactant concentrations to some simple powers. However, catalytic reactions follow much more complex rate equations. It is common for the rate of a catalytic reaction to be constant or even go down as the concentration of one of the reactants increases. This is quite different from gas-phase reactions, where rates generally increase with increasing reactant pressure.”

Figure 29. Generic potential energy diagram showing the effect of a catalyst in an hypothetical exothermic chemical reaction. The presence of the catalyst opens a different reaction pathway (shown in red) with a lower activation energy. The final result and the overall thermodynamics are the same.

2.1.6 Equilibrium
The concept of equilibrium is an important link between thermodynamics and chemical kinetics. Equilibrium does not determine the reaction rate of a system, but rather is defined by the statement “at equilibrium, the rate of any forward chemical reaction… must equal the rate of the reverse chemical reaction." This is most easily represented by the concept of the equilibrium constant.

For example, for the reaction $A + B \leftrightarrow C + D$, the equilibrium constant is defined as
Where $k_1$ is the rate of the forward reaction, $k_2$ is the rate of the reverse reaction, and brackets representation concentration. Therefore, if $k_1 = k_2$, the product of the concentration of the products will equal the product of the concentration of the reactants at equilibrium.

“In chemical kinetics, reactions that simultaneously and independently proceed in two directions – the forward and reverse ones – but at different rates, are called opposing, two-way, or reversible. It is characteristic of reversible reactions that in a certain time after they begin the rates of the forward and the reverse reactions become equal and a state of chemical equilibrium sets in.

All chemical reactions are reversible, but in definite conditions some of them can proceed only in one direction up to the practically complete disappearance of the reactants. Such reactions are called irreversible. Usually reactions are irreversible in which at least one product is removed from the sphere of the reaction (when reactions proceed in solutions a product precipitates or evolves in the form of a gas), or which are attended by a great positive heat. When considering ionic reactions, a reaction is practically irreversible if its products include a very poorly soluble or poorly dissociated substance.

The concept of reversibility of a reaction here does not coincide with that of thermodynamic reversibility. A reaction reversible in the kinetic sense may be irreversible in the thermodynamic one. For a reaction to be called reversible in the thermodynamic sense, the rate of the forward process should differ from that of the reverse process by an infinitely small amount, and, consequently, the process as a whole should proceed infinitely slowly.”

### 2.1.7 Free Energy

The concept of Gibbs free energy change, $\Delta G$, is another link between kinetics and thermodynamics. It is “the indicator of spontaneity of a reaction or physical change at constant T and P. If there is a net decrease of useful energy, $\Delta G$ is negative and the process is spontaneous.” It is defined as

$$\Delta G = \Delta H - T\Delta S$$
Where $\Delta H$ is the change in enthalpy, $T$ is the temperature and $\Delta S$ is the change in entropy. The heat of formation, $\Delta H_f$, and the entropy of formation, $\Delta S_f$, are roughly independent of temperature, and therefore can be used to determine the spontaneity of the reaction at various temperatures by inserting them into Equation 11.\(^{40}\) It is important to note that the Gibbs free energy only determines whether the reaction will occur, but says nothing about the speed of reaction. For that we must rely on kinetics.

### 2.2 Kinetics in Hydrogen Storage

For hydrogen storage, kinetics is generally taken to mean the dynamic rates at which hydrogen sorption and desorption from a storage material occur in time. A primary distinction between capacity and kinetics is that capacity measurements are theoretically taken at thermodynamic equilibrium, independent of the time required to reach equilibrium, while kinetics investigates how the material approaches equilibrium and what influences this approach. The availability of hydrogen in a storage material is dependent on the kinetics of the material under the system operating conditions. While a material might demonstrate high hydrogen storage capacity, the true amount of hydrogen practically available may be significantly less depending on the material's intrinsic kinetics. A number of different intrinsic properties of a storage material may control kinetics including surface interactions, transport phenomena, hydrogen-substrate storage mechanisms and phase change.\(^{41}\) External factors such as temperature and pressure also have a profound effect on hydrogen sorption and desorption kinetics.

In its most simple form, a kinetics experiment provides a useful measure of the rate of hydrogen uptake or release from a storage material. Unfortunately, comparing kinetics across materials and experimental setups can be complicated. For example, one way to compare kinetics is to consider average rates. A common practice is to define the average kinetic rate as the time to reach 95% of the full capacity. However, as is demonstrated in Figure 30, it is possible to derive the same average sorption rate for materials that, in fact, exhibit very different kinetic character. Thus, it is important not only to compare average rates but also to compare the shape of the kinetic curves. Factors that can influence the experimental rates are discussed in the section on experimental and analysis considerations (Section 4).
Figure 30. Hypothetical concentration versus time curves showing 3 different kinetic behaviors that have the same average rate at $t_{95\%}$.

2.3 Impact of Heat Transfer

Kinetics measurements are conducted to quantify the kinetic performance of hydrogen storage materials and identify the intrinsic mechanisms controlling hydrogen uptake and release. Identification of the kinetic mechanisms, most specifically the rate-controlling mechanism, is instrumental in developing materials with improved kinetic properties. It is not easy to perform measurements to accurately determine the intrinsic rate-controlling mechanism however. Early in hydrogen storage research, intrinsic material properties like surface effects, mass transport and storage mechanisms were generally assumed to be the rate-controlling mechanisms. In reality, heat transfer and other thermal effects dominate kinetic rates for nearly all reactions in hydrogen storage systems and measurements.\footnote{42}

In taking kinetics measurements, heat transfer is the most important effect for which researchers must account. The temperature of sorption/desorption is the most influential variable in hydrogen storage kinetics for both chemisorbing and physisorbing materials. It is imperative that measurements are taken under isothermal conditions in order to minimize the effects of heat transfer and identify the intrinsic kinetic mechanisms. Isothermal measurements are difficult in systems with poor heat transfer and fast intrinsic kinetics because the heat generated or taken up during hydrogen-substrate interactions can cause local temperature excursions that profoundly affect kinetic rates. All too often, kinetic rates reported in literature are in fact heat transfer rates because the sorption/desorption process is limited by an experiment’s ability to supply or dissipate energy. Heat transfer-limited kinetics measurements occur primarily because of improper equipment design and errors in experimental protocol. The effect of heat transfer on kinetics measurements is underscored by Dantzer in
1997: with regard to kinetic measurements on hydrogen sorption in LaNi$_5$, before 1983-1985, attempts to interpret data in terms of a microscopic [kinetics] model are questionable since no proof is provided on the isothermal conditions of the experiments. The effects of heat transfer are addressed often to reinforce their importance to the field. Further discussion on steps that can be taken to minimize the effects of heat transfer can be found in the experimental and analysis considerations section 4 of this chapter.

3 Kinetic Theory
This section is designed to familiarize the reader with knowledge of the fundamental mechanisms of hydrogen storage kinetics. It will also briefly review the process used to identify rate-limiting mechanisms through mathematical modeling.

Kinetic theory is a complicated and heavily researched field and a complete review of its content is beyond the scope of this work as an introductory document on hydrogen storage properties and testing. Accordingly, the kinetic theory section 3 is restricted to a brief overview. The authors will use references extensively in order to both summarize information and provide the reader with avenues for further inquiry into the subject of kinetic theory.

3.1 Fundamental Mechanisms of Kinetics
In order to improve the kinetic properties and performance of potential hydrogen storage materials, it is important to gain a fundamental understanding of the underlying rate-controlling mechanisms. The sorption and desorption of hydrogen by a material involves a number of important steps that occur in series and in parallel. For different sorption and desorption processes as well as for different hydrogen storage materials, the mechanisms controlling these steps differ. In general, there are three broad categories of kinetic mechanisms: surface interactions, mass (hydrogen, potentially the host elements) and energy transport (heat transfer), and binding mechanisms. Surface interactions, mass and energy transport and binding mechanisms are considered intrinsic processes. Mass and energy transport also depend on a number of extrinsic considerations including the particle size, packing density, the shape of the sample and sample holder, the thermal contact between the sample and sample holder, and the mobility and pressure of the gas. Any one mechanism may be the slowest, rate-controlling mechanism, although heat transfer is often the rate-controlling mechanism in practical application. Moreover, for reversible storage materials, the rate-controlling mechanisms may not necessarily be the same for sorption and desorption. It is generally difficult to differentiate experimentally between sorption and desorption mechanisms without careful experimental control.

Sorption and desorption in physisorbing materials is a two-step process consisting of mass and energy transport and surface interactions/binding mechanisms. Hydrogen is only transported through the void volume of the material and physisorption storage occurs at the surface of the skeletal material
with minimal effect on the structure of the material. In this sense, surface interactions/binding mechanisms can be considered equivalent phenomena in physisorbing material and will be addressed in the physisorption binding mechanisms section below. In chemisorbing media, surface interactions and bulk diffusion present distinct steps in the process of hydrogen storage and one or both may play the key role in hydrogen storage kinetics. The following section on surface interactions will discuss the various surface interactions that affect hydrogen storage kinetics in chemisorbing media.

### 3.1.1 Binding Mechanisms

The mechanisms that store hydrogen differ for physisorbing and chemisorbing media. The relative strength of the bonds greatly influences the kinetics and thermodynamics of the hydrogen storage material.

#### 3.1.1.1 Physisorption

For physisorption, molecular hydrogen bonds to the surface via van der Waals forces and remains in molecular form, $H_2$. Storage through physisorption is primarily limited to porous and nano-structured materials, where high surface area-to-volume ratios allow the physisorption mechanism to contribute significantly to storage capacity. In these types of materials, the external surface dictates the accessibility of the internal pore structure and the internal surface area dictates the degree of physisorption. Bonding can occur through dipole-induced dipole and induced dipole-induced dipole interactions (Figure 31), with heats of adsorption (physisorption) ranging from 4-6 kJ/mol depending on the nature of interaction.

![Figure 31](image)

**Figure 31.** 1) Dipole-induced dipole van der Waals interactions. 2) Induced dipole-induced dipole van der Waals interactions.

In Figure 33 below, the heat of adsorption is shown on the $H_2 + M$ potential energy curve as $E_P$. Van der Waals forces dominate kinetic forces at low temperatures because molecules have relatively low kinetic energy under these conditions, allowing weak intermolecular forces to bond hydrogen to a substrate.

It should be mentioned that the energy of physisorption is enhanced in small pores owing to the overlap of attractive fields from opposite walls. Therefore, there is considerable interest in the development of porous materials with pore of nanometric dimensions. The extent of adsorption depends on the surface area and the microporous volume. The characterization of samples in terms of surface area is subject to controversy, especially for microporous materials. On these materials, the surface measured will depend on the size of the probe molecules, the model use (BET or Langmuir), the extent of micropores filling etc.
3.1.1.2 Chemisorption
The dissociation of hydrogen and the subsequent surface chemisorption (bond formation) and/or the formation of one or more bulk hydrides with a hydrogen storage material forms the basis of chemisorption hydrogen storage. Depending on the type of hydride bond formed, hydrides may be classified as ionic, metallic or covalent hydrides. A thorough review of the chemistry of hydrides can be found in the book “Hydrides” by Wiberg and Amberger and in particular on metal hydrides in the book “Metal Hydrides” edited by Muller, Blackledge, and Libowitz. The important point is that the kinetics of hydrogen uptake and release may involve many complex steps including adsorption, dissociation, surface diffusion, bulk hydrogen diffusion, hydride formation, reactant diffusion, phase separation, etc.

3.1.2 Mass Transport
The kinetics of hydrogen storage is strongly influenced by the mobility of hydrogen and potentially the storage material itself and the transport of energy through the storage media (material and gas). The mobility of hydrogen, reactants, and energy within the storage media involve a wide range of transport properties. For simplicity we have divided the discussion of transport mechanisms into physisorbing and chemisorbing storage media.

3.1.2.1 Physisorption
For porous materials, molecular hydrogen easily passes through macroscopic boundaries and is transported through the network of pores to the internal surfaces of the material. Transport through the internal networks is governed by porous media-specific transport laws. Darcy’s Law is a well-known momentum transport constitutive equation in porous media that relates the velocity with respect to the porous media \( \nu \) to the viscosity of the fluid (hydrogen in this case) \( \mu \), the diagonal permeability tensor \( K \), and the piezometric pressure, \( P \).\[ \] \[ \text{Equation 12} \]

\[
\langle \nu \rangle = -\frac{1}{\mu} [K \cdot \nabla \langle P \rangle]
\]

Transport of fluids in porous media is the subject of extensive study and the reader is encouraged to review several books on the subject.\[ \], \[ \]

3.1.2.2 Chemisorption
For chemisorption in metal hydrides, hydrogen transport in the bulk occurs via atomic diffusion through the interstitial sites of the host metal structure. The dynamics of interstitial motion cover a wide range of time scales, from \( 10^{14} \) Hz for vibrational motion to \( 10^{10} \) Hz for collective long-range diffusion. The mechanisms controlling local diffusion in chemisorbing materials are highly temperature dependent. Values measured for long-range diffusion can vary over 12 orders of magnitude for different samples at different temperatures.\[ \] In the high-temperature region of practical interest to chemisorption hydrogen storage research, a classic barrier-hopping model can be used to describe the diffusion
of hydrogen through a metal hydride. The temperature dependence of the diffusion rate can be expressed by the relation

\[ D = D_0 e^{\frac{-E_A}{k_B T}} \]

where \( D \) is the number of successful jumps per unit time, \( D_0 \) is the number of vibrations or attempts per unit time, \( k_B \) is the Boltzmann constant and \( E_A \) is the activation energy. At lower temperatures, quantum tunneling effects of hydrogen atoms become dominant.\(^{48}\)

### 3.1.3 The Impact of Surface Interactions on Kinetics

As described above, hydrogen can be stored on the surface (adsorption) and in the bulk (absorption) of chemisorbing materials. For superficial chemisorption, hydrogen is weakly bound to the host’s surface. For bulk chemisorption, only atomic hydrogen can pass through the surface layers and be transported into and out of the bulk. Therefore, the surfaces of chemisorbing materials must split hydrogen gas into atomic hydrogen for storage. This is accomplished by the dissociation of \( \text{H}_2 \) via the transfer of electrons to the sample material. Hydrogen dissociation on the surface of a metal hydride is shown in Figure 32. Generally, hydrogen exists as a diatomic molecule before interacting with the surface of the material (left side of the figure), dissociates at the surface into two atomic hydrogen atoms, and diffuses into the bulk of the material (small molecules in the lattice). Doping with metal catalysts is a common technique used to increase the efficiency of hydrogen dissociation at the surface of chemisorbing materials. A considerable number of investigations at the materials development and fundamental mechanism levels seek to maximize the ability of a material to dissociate hydrogen through doping and surface texturing.\(^{49,50}\)

![Figure 32. Schematic representation of the interaction of hydrogen with a clean crystalline surface.](image-url)
The interaction of hydrogen with the surface of a chemisorbing material is complicated by the fact that the surface or interfacial properties of the material may be substantially different than those of the bulk. The crystal structure, electronic, magnetic and dynamic properties, as well as the chemical composition of a material can be dramatically altered near the surface due to the discontinuity of the surface. By definition then, the surface layer consists of the atomic region that shows different properties from those of the bulk. For a clean single crystal, the surface layer is generally no thicker than the first few monolayers. Oxidized, contaminated, or multiphase materials may have a surface layer as thick as 10 nm. From the crystal structure point of view, atoms in the surface layer may relax or reconstruct. Surface relaxation may involve changes in the lattice dimensions and inter-atomic distances in the first few layers. Reconstruction consists of a rearrangement of atoms in the top atomic layers and often results in a loss or change of symmetry. Electronic states are also altered in the surface layer.

Details on surface contamination and the effect on kinetics is discussed in section 3.1.3.2 below.

### 3.1.3.1 Surface Energetics

Surface energetics governs the dissociative chemisorption of hydrogen in the surface layer. The potential energy of the interaction of gaseous hydrogen with a surface can be depicted using a simple two-dimensional set of potential energy curves (Figure 33). (Complex hydrides do not fit into this simple picture, since the absorption and desorption of hydrogen involve structural phase transitions.) Away from the surface the two curves are separated by the heat of dissociation, $E_D = 218 \text{ kJ/molH}$, which represents the amount of energy required to split H$_2$ into two hydrogen molecules. The interaction of gaseous hydrogen with the metal to form a stable solid solution or intermetallic is described by following the H$_2$ + M potential from its minimum ($E_P$) to its intersection with the 2H + M potential in the surface layer. If the potential energy curves intersect above the zero energy level, positive activation energy $E_A$ is required for hydrogen dissociation; if the curves intersect below zero, dissociation is energetically favorable ($E_{NA} < 0$) and occurs spontaneously. After the curves intersect, dissociation occurs and the potential continues along the 2H + M curve as atomic hydrogen is transported into the bulk. The potential energy of hydrogen in the material reaches a deep minimum on the 2H + M potential (the heat of chemisorption, $E_C \approx 50\text{kJ/molH}$) close to the surface/bulk interface, which dictates that only atomic hydrogen transports through the bulk of metal hydrides.
For chemisorption, some surface properties that can limit the overall hydrogen sorption rates (such as $E_C$ and the sticking probability) become concentration dependent as the surface coverage of atomic hydrogen increases. The coverage, in turn, depends on the mobility of atomic hydrogen on the surface and into the bulk. Perhaps most importantly, diffusion from the very stable superficial chemisorption sites ($E_C$) into the bulk absorption sites ($E_S$) may be a rate-limiting step and is usually closely linked to surface structure. Thus, kinetics may be influenced by a feedback loop between concentration and surface interactions.

### 3.1.3.2 Surface Contamination

So far, the interaction of hydrogen with metals has been described by the ideal case of a clean elemental metal surface. Clearly, this model does not describe the real surface environment found on most samples. Elemental metals generally have surfaces that have been passivated by oxygen, thus preventing the dissociation reaction and the diffusion of atomic hydrogen into the bulk (Figure 34).
Intermetallic compounds, on the other hand, often show highly reactive surfaces even after exposure to oxygen or other contaminants (H₂O, N₂, CO₂, etc.). This is due to reorganization in the chemical composition of the surface to minimize the surface energy. This form of surface segregation is an intrinsic property of alloys.

Schlapbach et al. showed that surface segregation can be caused by selective oxidation of certain elements in the alloy. In the case of LaNi₅, they found that the selective oxidation of La prevents the surface from becoming passivated. Using depth profiling by combining photoelectron spectroscopy and sputtering, they demonstrated that there was a surface enrichment of La, as well as the precipitation of metallic Ni in the subsurface. This suggests that the decomposed...
surface layer is permeable to molecular hydrogen that may dissociate in the subsurface on active Ni precipitates and on the un-oxidized host alloy (Figure 35). Surface segregation at room temperature has also been observed in many other hydride-forming intermetallic compounds such as ErFe, ZrMn$_2$, TiMn$_2$, CeRu$_2$, and Th$_7$Fe$_3$, and at elevated temperatures for the hydrogen storage compounds TiFe and Mg$_2$Ni.

An active surface does not necessarily guarantee rapid kinetics. TiFe provides a good example. This compound requires activation at high pressures and temperatures to absorb hydrogen. Yet, the detection of HD molecular species created through H$_2$–D$_2$ exchange interactions at the surface showed that this compound actually has an active surface even at room temperature. It is thought that the difficult activation is due to bulk diffusion properties.$^{54}$

In practice, there are different ways to overcome the kinetic barriers of oxides and other contaminants on the surface (activating the material or sample). By heating under vacuum oxygen may diffuse from the surface into the bulk, essentially cleaning the surface. By heating under an inert gas flow unstable surface layers may be removed. By heating under hydrogen the oxides may be reduce and removed. Any one of these processes may cause surface segregation and the formation of a mixed surface layer consisting of nano-scale oxide and catalytic metal particles (such as La-oxides and Ni clusters on the surface of LaNi$_5$ as described above).

### 3.1.3.3 Surface Hydriding

Hydride formation that occurs at the surface of the material can hinder the further transport of hydrogen into the bulk.$^{55,56}$ This phenomenon is depicted in Figure 36. When hydrogen dissociates at the superficial layer, it must be transported away from the surface through the first several monolayers of material. If the hydrogen transport properties of the material are poor, the hydrogen and substrate will react to form hydride at the surface. The hydride acts as a barrier to further hydrogen transport into the bulk and can significantly affect kinetic performance. The sorption rate eventually becomes limited by the diffusion of hydrogen through this hydride layer.
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Figure 36. A schematic representation of the formation of a surface hydride which creates a diffusion barrier blocking the transport of atomic hydrogen into the bulk.

3.1.4 Heat Transfer

Because of the exothermic nature of most hydrogen storage materials, heat transfer often plays the most critical role in the kinetics of hydrogen sorption and desorption. This topic was brought out early in this chapter because of the importance of heat transfer on the study of sorption kinetics. Consideration of heat transfer on experimental measurements are presented in detail in section 4.9. At this point it is useful to discuss heat transfer within the context of the materials themselves.

Heat transfer in porous media occurs through the internal pore network and the material itself. Heat transfer performance is generally dictated by the pore size distribution relative to the mean free path of the hydrogen molecule at various temperatures and the structure (e.g. characteristic dimension, linkage) of the skeletal material.\(^{57, 58}\)

At first glance, heat transfer in chemisorbing materials would seem to be governed by thermal conduction through the storage material itself. However, with few exceptions, chemisorption materials are often powders with limited particle-to-particle contact. Thus, heat transfer is generally governed by many of the same principles as porous media. For example, the large lattice expansion that occurs in hydriding metals and intermetallic compounds creates enough internal stress within the crystal lattice and grain boundaries that bulk samples will turn to powders on hydriding. With cycling these powders will break down into smaller and smaller particles. This is referred to as decrepitation. This process itself will change the thermal conductivity of the storage material. Thus, the
repeated process of hydrogen sorption and release can have a strong influence on the kinetic behavior of the storage material system with cycling. Heat transfer in other chemisorption materials such as alanates and amides as well as nanomaterials of any kind may also be affected by mass transport of the reactants and grain growth.

Over the years, a great amount of effort has been put into the design and development of materials and devices optimized for controlling material mobility and heat transfer in hydrogen storage systems.59

### 3.2 Kinetics Modeling

Mathematical models are used to provide insight into the rate controlling mechanisms of a material’s hydrogen uptake and release. The model that best fits experimental observation is typically assumed to be the rate-limiting mechanism.

The kinetics modeling section will be structured around models for metal hydrides to illustrate the process of determining sorption mechanisms. Kinetics in metal hydrides can be mathematically modeled by the general relationship

\[
dC = K(T)F(C,P)
\]

where \(K(T)\) is the temperature-dependent rate constant and \(F(C,P)\) is a function dependent on the hydrogen concentration in the material \(C\) and gas pressure \(P\). A brief introduction will be given on the rate constant \(K(T)\) and how it is determined followed by a more in-depth discussion on modeling the hydrogen concentration function.

#### 3.2.1 Temperature Dependence

When kinetics measurements are made at different temperatures, it is generally found that gas sorption and desorption rates are highly dependent on temperature. This is indicated in the temperature dependence of the rate constant \(K\) in Equation 15. In 1889, Arrhenius recognized that most chemical reactions occur at rates that increase exponentially with temperature. This empirical observation is valid in hydrides and can be conveniently written as

\[
K(T) = Ae^{-\frac{E_A}{k_BT}}
\]

where \(A\) is the frequency factor, \(E_A\) is the activation energy, \(k_B\) is the Boltzmann constant and \(T\) is the temperature. Note however, that the Arrhenius equation is actually only an approximation and, while it is a good approximation over the temperature range of interest for hydrogen storage, it works best in for temperature of about 50-100 K where the activation energy may be considered temperature independent.29
From Equation 15 we find that the activation energy \( E_A \) is given by

\[
E_A = -k_B T \ln(K)
\]

The activation energy of a reaction can be determined by measuring the rate constant \( K(T) \) at several different temperatures and then plotting \( \ln(K) \) versus \( 1/T \). This is known as an Arrhenius plot.

An example of such an Arrhenius plot is shown in Figure 37. This plot presents data for the same \( \text{La}_2\text{Mg}_{17} \) sample given in the example in the following section. The rate constants were determined from a series of absorption measurements at different temperatures using the same pressure-dependent curve fitting approach as in Figure 39. A line was then fit to the Arrhenius data in Figure 37. The activation energy for the hydride formation reaction can be obtained from the slope of this \( \ln(K) \) versus \( 1/T \) plot. The activation energy calculated from these measurements is 63.7 kJ/mole.

Figure 37. Arrhenius plot of rate coefficients \( K \) versus \( 1/T \), for hydrogen absorption by \( \text{La}_2\text{Mg}_{17} \).
3.2.2 Concentration Dependence

The rate of adsorption and desorption is dependent on the availability of hydrogen in the sample. The hydrogen concentration function is used to model this availability and is often normalized to allow different kinetics measurements to be compared on the same plot. The functional dependence of concentration is modeled non-dimensionally as

Equation 17

\[ F(\alpha) = \kappa \left( t / t_{1/2} \right) \]

where the reacted fraction \( \alpha \) is the concentration \( C \) normalized by the final concentration \( C_0 \), \( \kappa \) is an empirically-determined rate constant, and \( t \) is the time normalized by \( t_{1/2} \), the time at half concentration \( (C_0/2) \).60

With great care, experimental data can be compared with theoretically derived models for different bulk-process rate limiting mechanisms (diffusion or reaction propagation). This type of mechanistic analysis was described early on by Sharp et al. This is presented below as a series of models that have been derived for different hydrogen transport mechanisms and geometries. The nine models presented, divided by mechanistic category, are compared to experimental data in Figure 38. In the absence of heat transfer and/or surface interaction effects, such an analysis can be used to determine the rate-limiting concentration driven mechanism of a material. The subscripts of the equations below denote the kinetics profile they describe in Figure 38.

Please note that it is extremely difficult to make measurements that are sufficiently isothermal and have little-to-no surface effect considerations. Such an analysis should be restricted to experimental configurations designed specifically for this type of analysis (see section 4.9) and well defined high-purity samples and gases.

3.2.2.1 Diffusion Models

Diffusion models can be used to describe one-, two-, and three-dimensional diffusion processes. A simple one-dimensional diffusion process with a constant diffusion coefficient can be described by

Equation 18

\[ D_1(\alpha) = \alpha^2 = .2500(t / t_{1/2}) \]

Two-dimensional diffusion into a cylindrical body gives

Equation 19

\[ D_2(\alpha) = (1 - \alpha)\ln(1 - \alpha) + \alpha = .0426(t / t_{1/2}) \]

Diffusion from the surface towards the center of a sphere is61

Equation 20

\[ D_3(\alpha) = \left( 1 - (1 - \alpha)^{1/3} \right)^2 = .0426(t / t_{1/2}) \]
Another well-known model for a spherical body is\textsuperscript{62}

\begin{equation} \label{eq:21}
D_4(\alpha) = \left(1 - \frac{2\alpha}{3}\right) - (1 - \alpha)^{\frac{2}{3}} = 0.0367 \left(t / t_{1/2}\right)
\end{equation}

\subsection{3.2.2.2 Reaction Models}

Kinetics may be limited by the chemical reaction rate i.e. the rate of transition from solid solution to intermetallic. This mechanism can be modeled by

\begin{equation} \label{eq:22}
\frac{d\alpha}{dt} = \kappa (1 - \alpha)^n
\end{equation}

where \(n\) is the order of the reaction. In some cases, solid-state reactions appear to follow first order kinetics

\begin{equation} \label{eq:23}
R_4(\alpha) = \ln(1 - \alpha) = -0.693 \ln \left(t / t_{1/2}\right)
\end{equation}

If kinetics are controlled by a constant velocity propagation of the phase transition boundary, then useful relationships can be derived from Equation 22. The case of a cylinder, \(n = 1/2\), gives

\begin{equation} \label{eq:24}
R_6(\alpha) = 1 - (1 - \alpha)^{\frac{1}{2}} = 0.2929 \left(t / t_{1/2}\right)
\end{equation}

For a sphere of radius \(r\) in which the reaction proceeds from the surface towards the center with \(n = 2/3\),

\begin{equation} \label{eq:25}
R_7(\alpha) = 1 - (1 - \alpha)^{\frac{1}{3}} = 0.2063 \left(t / t_{1/2}\right)
\end{equation}

The above models of phase boundary propagation assume a virtually instantaneous and homogeneous nucleation of the product phase. In such a case the surface of a material is quickly covered by a hydride layer and the kinetics are described by the propagation of the phase boundary towards the center of the material. However, this is generally not how a hydride is formed. Typically the hydride phase nucleates at discrete locations in the bulk and grows outward. Hydride domains come into contact with each other as they grow and growth continues along the remaining reactant-product boundaries, eventually consuming the entire material. This process of nucleation and growth can be described by the following equations derived by Avrami and Erofe’ev\textsuperscript{63 64}

\begin{equation} \label{eq:26}
A_8(\alpha) = (\ln(1 - \alpha))^{\frac{1}{2}} = 0.8362 \left(t / t_{1/2}\right)
\end{equation}

\begin{equation} \label{eq:27}
A_9(\alpha) = (\ln(1 - \alpha))^{\frac{1}{3}} = 0.8850 \left(t / t_{1/2}\right)
\end{equation}
The following is an example presented to show how one could potentially determine the rate-controlling mechanism. The nine models described above are plotted along with the experimental hydrogen absorption data of a metal hydride sample in Figure 38. The models are shown in Figure 38 on a plot of $\alpha$ versus $t/t_{1/2}$. At low concentrations, it is clear that diffusion (Equation 18 through Equation 21, 1-4 in Figure 38) is the rate-controlling mechanism, although it is difficult to distinguish between the different diffusion models. However, this distinction becomes more obvious at higher hydrogen concentrations. The experimental data best fits curve 3, which describes diffusion from the surface to the center of a sphere. Therefore, absent heat transfer and/or surface interaction effects, this diffusion process is the mechanism that would best describe the rate limiting mechanism of hydrogen sorption in the metal hydride in question. Once again, it is important to remember that it is very difficult to accurately determine the intrinsic mechanisms of hydrogen sorption/desorption due to thermal effects. Heat transfer effects dominate kinetics for most measurements. Therefore, conclusions drawn from modeling must be kept in perspective when analyzing sorption/desorption mechanisms.

![Figure 38. Various solid-state kinetics models. 1) One-dimensional diffusion, 2) diffusion in a cylinder, 3+4) diffusion in a sphere, 5) 1st-order phase transformation, 6) constant velocity phase-boundary propagation in a cylinder, 7) constant velocity phase-boundary propagation in a sphere, 8+9) nucleation and growth.](image)

3.2.3 Pressure Dependence
The volumetric method commonly used for measuring kinetics is not done at a constant pressure. Therefore, the pressure term in Equation 14 must be included to account for the effect that the changing potential energy of the surrounding
gas will have on sorption kinetics. The following set of equations were proposed by Wang and Suda\textsuperscript{65} for the sorption kinetics of the hydride phase transition:

\textbf{Equation 28} \quad \frac{dC}{dt} = K_h F_h (P, C)

\textbf{Equation 29} \quad K_h = A e^{\frac{-E_h}{R T}}, F_h (P, C) = \left( \frac{P}{P_{eh}} \right)^a \left[ 1 - \left( \frac{P_f}{P} \right)^a \left( \frac{C}{C_f} \right)^b \right]

\textbf{Equation 30} \quad \frac{dC}{dt} = K_d F_d (P, C)

\textbf{Equation 31} \quad K_d = A e^{\frac{-E_d}{R T}}, F_d (P, C) = \left( \frac{P}{P_{ed}} \right)^b \left[ 1 - \left( \frac{P_f}{P} \right)^a \left( \frac{C_f}{C} \right)^b \right]

\(K_h\) and \(K_d\) are the absorption and desorption rate constants, \(P_{eh}\) and \(P_{ed}\) are the equilibrium plateau pressures, \(P_f\) and \(C_f\) are the final pressure and concentration, and \(a\) and \(b\) are the orders of the relationships between the sorption rates and pressure and concentration, respectively. Corrections due to pressure change effects on rates can be overcome by simply adding pressure regulation or using relative volumes such that sorption results in measurable pressure changes but not significant with respect to reaction rates (quasi-isobaric).

As an example of relationships given in Equation 28 through Equation 31, experimental measurements of the hydrogen absorption kinetics of the intermetallic compound La\textsubscript{2}Mg\textsubscript{17} Figure 39 at 300°C are presented in Figure 39. The most linear behavior was found for reaction orders \(a = 2\) and \(b = 1\). This relationship proves to be valid for up to 95% of the absorbed capacity. The slow non-linear residual absorption at the bottom of the plot is most likely due to reestablishing thermal equilibrium after initial fluctuations in temperature caused by the exothermic hydriding reaction. The rate constant \(K_h\) from Equation 29 was determined from the slope of a linear fit of the upper part of the data.
4 Experimental and Analysis Considerations

Measured sorption and desorption rates are dependent on experimental and data analysis considerations as well as intrinsic material properties. As a consequence, experimental conditions and procedures that vary from system to system and from researcher to researcher can have a tremendous effect on measurements and the conclusions drawn from them. This makes comparisons between measurements with experimental equipment, different experimental conditions, different sample shapes, sizes, morphology and composition difficult if not impossible. Considerations that can have a significant effect on kinetics measurements will be discussed in this section along with suggestions to minimize their impact.

This section will rely heavily on examples from literature to support and illustrate the effects of various kinetics considerations. For more detailed information on specific considerations, refer to the reference of the cited example.

4.1 The Limitations of Kinetics Measurements

While a great deal can be learned from a good set of sorption and desorption kinetics measurements, from a materials discovery perspective, it is critical that the underlying hydrogen sorption process that is occurring is well understood.
Unfortunately, kinetics measurements themselves often give very little direct information about these processes. In many ways kinetics plots look similar to each other. For that reason, basic sorption and desorption measurements should be complemented by a reliable set of supporting measurements. These may include characterization by PCT isotherm, TPD, X-ray and neutron diffraction, NMR, FTIR and Raman measurements among others.

To give an example, the measurement of hydrogen uptake or release (kinetics) may appear quite similar for extremely dissimilar chemical processes. Two examples are presented to show how desorption measurements from two different types of hydrogen storage materials can look quite similar (no distinguishing features other than being measured at different temperatures and showing differing overall rates) even though the mechanisms of hydrogen uptake and release are very different.

In the first example, the room temperature release of hydrogen from a classic AB₅ (LaNi₅-based) intermetallic hydride is shown in the kinetics measurement of Figure 40.

![Figure 40. Kinetics measurement of LaNi₅ Intermetallic compound with a single phase transition. The experiment was conducted at room temperature. Units in H per formula unit released.](image_url)
Figure 40 was made by measuring the increase in pressure using a calibrated volumetric instrument as a sample of approximately 5 g desorbed hydrogen. For the most part, this material exhibits the classic kinetics profile of a single phase transition from the intermetallic hydride compound to a metal alloy with some hydrogen in solid solution (commonly referred to as beta and alpha phases, respectively). An intermediate crystal structure transition is known to occur (gamma phase), but for current illustrative purposes the kinetics can be considered as a single chemical reaction. Atomic hydrogen is released from its interstitial binding site in the host metal’s lattice, diffuses to the surface and combines with another hydrogen to form gaseous hydrogen, which gives rise to the measured increase in pressure.

The capacity change as a function of time represents the material’s total hydrogen content change (in H per LaNi$_5$) during the desorption transition from LaNi$_5$H$_6$ to LaNi$_5$. This does not mean that the concentration of hydrogen in the hydride is changing but rather that the sample’s total composition of the two phases is changing with time. The sample’s total composition at any one time is:

\[ (1 - X(t))\text{LaNi}_5\text{H}_6 + X(t)\text{LaNi}_5 \]

with boundary conditions $X(t=0) = 0$ and $X(t=\infty) = 1$.

This transition from solid solution $\alpha$-phase through the $\gamma$-hydride phase to the $\beta$-hydride phase in LaNi$_5$ is best observed by performing a series of equilibrium PCT measurements (Figure 41). Absorptions are shown for increasing temperatures in purple to red and desorptions at the same set of temperatures in blue. The flat portion (plateau region) of each measurement is present because of the equilibrium coexistence of the $\alpha$-phase and $\beta$-phases (or $\gamma$-phase as the case may be). On moving from left to right along an absorption plateau the measured sample is being transformed, gas aliquot by gas aliquot, from the $\alpha$-phase intermetallic alloy to the $\beta$-phase hydride. The reverse is true going from right to left across the desorption plateau.
Hydrogen sorption through a single gas-solid phase transition such as the one above represents the ideal case for studying the mechanisms of hydrogen sorption kinetics. However, today’s advanced hydrogen storage materials are becoming increasingly complex. They can consist of materials that undergo multiple hydriding reactions, complete structural decompositions and potentially the long-range transport of reactants other than hydrogen. And yet, these complex processes may present absorption and desorption kinetics that are quite similar at first glance to a classic hydride.

Our second example shows the measurement of the third cycle of gas desorption from titanium-doped alanate-amide mixture with a starting composition of

$$LiH + Al + 2LiNH_2 + Mg + TiF_3$$

It should be noted that, unlike the intermetallic hydride example above, this is not an isothermal measurement but rather a ramp and soak-style TPD measurement. However, within the isothermal portion of the measurement at $t > 0.4$ hours, the desorption resembles the classic kinetics curve of Figure 40 to a large extent.
Without doing a more detailed PCT isotherm, it may be incorrectly assumed by looking only at the kinetics measurement that the desorption involves a single reaction. By performing an equilibrium desorption PCT measurement of the same sample (Figure 43) in the fourth cycle makes it clear that the continuous evolution of gas observed in the kinetics measurement of Figure 42 is the result of a complex series of chemical reactions instead of the single phase-transition as in classic hydrides.

Each of the three distinct plateaus corresponds to a different chemical transition within the material. This complex system has not been optimized or fully characterized by any means, but from what is known of its complex hydride and amide precursors, these reactions undoubtedly involve chemical decomposition and re-formation reactions. For hydride and amide products to form, this necessitates the long-range transport, not only of hydrogen, but also of other reactants including one or more species of Li, Al, and N. Later studies that coupled a high-pressure mass spectrometer to these same experiments showed the presence of ammonia in the evolved gas, adding additional complexity to the gas-solid system. 8
4.2 Discrete Kinetics Measurements

In the opening paragraph of Kinetics, it is noted that the difference between kinetics and capacity is that kinetics are inherently dynamic while capacity measurements are taken at thermal equilibrium. However, in principal, each data point on a PCT isotherm is the equilibrium concentration after the gas/material system has been perturbed from equilibrium. In fact, a volumetric PCT measurement consists of a series of small doses which, when measured with respect to time, individually represent measurements of kinetic properties. More precisely, with each dose (or aliquot of gas), a specified over-pressure (or under-pressure) is applied to the sample and the system is allowed to reach equilibrium through a dynamic process. The last data point of each kinetics measurement, which is assumed to represent equilibrium in composition, pressure and temperature, provides a single point on the PCT curve. This process is performed repeatedly until there are enough equilibrium data points collected to construct a full PCT diagram.

In a volumetric measurement where the quantity of gas is controlled (and thus the reactions are controlled), each dose is performed within a narrow band of concentration and pressure. In this sense the time-resolved data of a PCT measurement can be thought of as a series of mini-kinetics measurements that can separate different portions of the overall sorption process. This is demonstrated using the same PCT isotherm measurement of the second example above.
Figure 44 presents the pressure time plot of what was used to create the PCT isotherm of Figure 43.

![Figure 44. Pressure versus time plot of data used to create the PCT diagram of alanate-amide mixture in Figure 43.](image)

By controlling the dosing pressure so that only one reaction occurs at a time, it is possible to determine relative reaction rates of each independent reaction. Indeed with the same applied pressure differential of 2 bar, the desorption rates for the three observed reactions from the highest to the lowest pressure plateaus are 0.29, 0.10 and 0.15 wt.%/hr. Note that such data is not only useful for determining the kinetics of individual reactions, but it also demonstrates that due to poor kinetics the final PCT isotherm was derived from relatively non-equilibrium conditions.

These examples serve not only to demonstrate the wide range of complexity encountered in the development of advanced storage materials, but also point to the importance of utilizing multiple methods of characterization in evaluating what may at first appear to be a simple sorption process. Given limited time and resources, the ability to perform different measurements simultaneously in a single experiment becomes a real advantage in understanding complex systems.

### 4.3 Matching Experimental Setup to the Measurement Purpose

Perhaps more than any other type of measurement, the manner in which kinetics measurements are performed will greatly depend on the purpose of the
measurement. The purpose of measurements is the deciding factor in determining the measurement method to be used, the design of the experiment or a series of experiments, and ultimately the size of the sample and the design of the containment. This section will provide some guidelines to help the reader understand the purpose of kinetics measurements at the different levels of research and highlight considerations associated with measurements at each level.

4.3.1 Experiments for System Performance
The kinetic rates of a hydrogen storage system are determined by a number of factors in addition to the intrinsic kinetic properties of the storage material. Some of the factors that affect kinetic rates at the system level include heat transfer between the storage material, material housing and environment, the operating over- or under-pressure relative to thermodynamic equilibrium and the packing density of the storage material. Heat transfer plays an important role in the kinetic performance of storage systems because some materials require the delivery and dissipation of significant amounts of energy during charging and discharging. Dedrick presented an example of the extreme energy releases characteristic of some metal hydrides upon charging: hydriding 5 kg of sodium alanate storage material in 2.5 min releases an average of 668 kW. The total heat released is enough to boil 64 gallons of water.\textsuperscript{67} The ability to handle sorption/desorption energy loads such as those for sodium alanate necessitates the use of sophisticated thermal management systems such as internal plates, fins and tubing in system performance measurements. System-level considerations like thermal management devices must be included in the design and evaluation of the kinetic performance of a storage system.

For a system level understanding, this may mean, for example, designing experiments to be able to examine material from different sections of a test bed after performing kinetics measurements under a different set of charging or discharge conditions. In such a case, being able to access the material multiple times would be important. Or perhaps it is the heat produced and heat transfer that is the focus of interest. Then one may want to instrument a test bed with multiple temperature sensors at different position in the material and then to run a series of kinetics measurements under different test conditions.

To gain useful system performance information, enough material should be tested to provide results representative of the material’s behavior in a full-sized operating storage system. In general, this means between 10 grams and 1 kilogram of storage material. 100 grams is probably a good tradeoff between the practical issues of synthesis and handling of the materials and a characteristic, uninterrupted volume of material that would be found in a large-scale system. For kinetics measurements it will be important to be able to measure hydrogen flow rates into or out of the test bed. If the material performance is already fairly well characterized, it may be possible to use flow meters to study sorption/desorption rates at a system level. However, with the exception of Coriolis-based flow
measurement devices that maintain accuracy over two decades, conventional flow meters generally have fairly narrow dynamic measurement ranges (often less than one decade). This is a severe limitation when sorption/desorption rates may vary over several orders of magnitude depending on state of charge, temperature and pressure.

Mass change measurements on a system scale are also possible, but complicated by the connections between the system on a balance and the need for hydrogen transfer and potential heating or cooling. System scale kinetics measurements are most easily performed volumetrically using a normal regulator for sorption and a backflow regulator for desorption to maintain a constant or simulated pressure profile on the storage system. In its most simple form the amount of material is limited by the largest practical calibrated volume for a single desorption dose.

One example of scale up measurements were the experiments performed by Gary Sandrock on Ti-doped alanates. Hydrogen absorption/desorption studies were performed on a simple high-pressure cylindrical vessel shown in Figure 45 designed to simulate the heat transfer and gas impedance conditions of a larger bed. This 316 SS reactor has a 4.6cm OD, 0.38cm wall thickness, about 11.4cm internal length and was pressure rated to 3000 psia (204 atm) at 350°C. It was loaded with about 100 g of bicatalyzed NaAlH₄ (78 g of NaAlH₄ and 22 g of Ti(OBu)₄ + Zr(OPr)₄). The reactor had internal thermocouples but no internal heat exchange structure. The reactor was heated within an air furnace. Absorption kinetics and capacity were measured volumetrically and desorption measured with flow meters backed up by a wet test meter (a device to measure gas flow by counting the revolutions of a shaft upon which water-sealed, gas-carrying cups of fixed capacity are mounted).

Figure 45. Scale up (100 g) reactor with end cap removed in Ar-glovebox to show catalyzed alanate.
The higher than expected initial charging kinetics, combined with limited heat transfer, results in exothermic temperature excursions, as is common with hydrides. An example is shown in Figure 46 for the 4th hydrogen absorption. The fully dehydrided bed was heated to 155°C and then charged with hydrogen at 172 atm. Within one minute, the exothermic hydriding reaction resulted in an internal temperature of 234°C. This is essentially the van’t Hoff temperature for NaAlH₄ at this applied pressure. However, the melting point of NaAlH₄ is only 182°C; thus, any NaAlH₄ formed during the first 0.5 hr of Figure 46 would do so directly into the liquid phase. As one can see in the figure, a thermal arrest occurs at 182°C due to solidification during cooling (about t = 0.6-0.8 hr). This is precisely as expected and shows that liquid NaAlH₄ was formed during the exothermic temperature excursion associated with the rapid initial charge.

![Figure 46. Exothermic temperature excursion during scale up bed charge half-cycle 4A \((P_i = 172 \text{ atm, } T_i = 155^\circ\text{C})\).](image)

It is natural to ask if such melting could be detrimental to the subsequent performance of the alanate bed. From subsequent absorption and desorption measurements, the answer seems to be “no” or at least “not much”. In fact that series of scale up experiments pointed to a possible benefit of partial melting. When the reactor was opened between cycles 4 and 5 (photograph shown in Figure 16) the material in the bed was found to be sintered into a porous, solid mass. Such a structure may have distinct advantages for actual applications. In particular, such a sintered structure should reduce particulate migration, increase packing densities, allow expansion, and provide a constant internal gas impedance and enhanced safety.
Later, a more advanced test bed (Figure 47) was developed which had thermocouples placed at regular radial spacings and at different depths within the bed to examine heat transfer properties of the material during charge and discharge measurements.

Figure 47. Image of scaled up Alanate test bed with series of internal and external thermocouples.\textsuperscript{70,71}

4.3.2 Experiments for Materials Development

Kinetics measurements for materials development must focus on the intrinsic kinetics of the storage material in order to effectively compare materials. This can be difficult for materials with highly exothermic or endothermic reactions and good intrinsic kinetics because heat transfer often masks the intrinsic kinetic character of the materials. Intrinsic kinetics testing emphasizes testing at isothermal conditions and conducting measurements under identical conditions (pressure, temperature, sample size etc.) using identical measuring equipment in order to minimize the effects of heat transfer. The importance of testing kinetics under identical conditions using identical measuring equipment and setup is illustrated in the many dozens of papers before 1982 reporting absolute absorption and desorption rates for LaNi\textsubscript{5}. Reported kinetics rates at half capacity varied between < .01 min\textsuperscript{-1} to 35 min\textsuperscript{-1} at 5 atm and 298 K.\textsuperscript{72} The measurements in the review were performed with different quantities of material on completely different equipment without special considerations for heat transfer. Therefore conclusions drawn from these measurements concerning intrinsic kinetics are questionable at best. There are a few special instances when non-isothermal kinetics measurements are informative, particularly when applied to developing functional hydrogen storage applications.
Section 2: Kinetics

For research focused on materials development, a good approach is to run a series of comparative measurements on different samples while ensuring that all other material properties (sample size, packing density, purity...) and measuring conditions remain as consistent as possible. The experimental setup, sample size and measurement technique should be centered on simplicity and the ability to make comparative measurements as efficiently as possible.

Heat transfer and measurement of the true sample temperature at low gas pressures, in addition to buoyancy corrections at high gas pressures, make kinetics measurements by gravimetric methods difficult if not impractical. These complications are due to the limitation that the sample in a gravimetric analysis system cannot make physical contact with a thermal mass for heat exchange. This can be accommodated by reducing sample size or, in the case of desorption, by using a carrier gas in flow through mode to maintain the sample temperature. Volumetric kinetics measurements on samples ranging in size from a few milligrams to a couple of grams are quite straightforward. A major consideration is that, in particular for materials with elevated enthalpies of formation (> 10 KJ/mol H₂) and good intrinsic kinetics, sorption/desorption rates may be limited more by heat transfer in the measuring equipment than by physical or chemical sorption mechanisms. The flexibility in sample holder design and sample compositions (addition of thermal ballasts) can overcome this problem. With this in mind, it is still very relevant to measure sorption rates under imperfect (non-isothermal) conditions in order to obtain relative hydrogen uptake and release rates and to develop modified materials with improved kinetic performance. However, it is exceptionally important that materials are measured under essentially the same conditions (pressure, temperature, sample size, perhaps sample packing density) using identical measuring equipment and setup to be able to make quantitative and even qualitative comparisons between materials. Comparison of the intrinsic kinetic properties under the aforementioned instrumental limitations can be further qualified by considering or measuring, for example, differences in the isobaric heat capacity and thermal conductivity among the materials under investigation.

A classic example of such a successful comparative analysis with the goal of developing optimized hydrogen storage materials was the systematic study of increasing levels of TiCl₃ additives in NaAlH₄. By performing these measurements under identical conditions a clear trend in kinetic performance was observed ultimately leading to key findings in understanding the effect of Ti additives on alanates. Another example of comparative measurements for improved performance of storage materials was through the investigations of many different types of alanate additives by Anton et al.
4.3.3 Experiments for Fundamental Studies

Fundamental mechanism studies look to identify and understand intrinsic kinetic phenomena. Like those for materials development, fundamental investigations require isothermal conditions in order to eliminate the effects of heat transfer. Investigation of hydrogen diffusion mechanisms, the effects of catalysts on hydrogen dissociation, and the nature of hydrogen/sample material interaction are examples of research at the fundamental level. If the purpose of a kinetics measurement is to study intrinsic mechanisms, it is absolutely necessary to design the experimental measurement equipment to achieve essentially isothermal conditions.

For these basic science experiments, simplicity (reducing the number of free experimental parameters) is key. While it may not be possible to determine rate-limiting mechanisms without specially designed sample cells, sorption and desorption kinetics measurements on standard equipment and laboratory scale samples (e.g. 1 gram) can provide not only relative measurements for materials optimization but also some fundamental insight into sorption processes. Gary Sandrock’s work on Ti doping in sodium alanates provides a good example of how detailed kinetics measurements and the determination of activation energies have been employed for improved understanding and development of better hydrogen storage materials.\(^7\) Hydrogen desorption rates were measured at several temperatures for the decomposition of NaAlH\(_4\). For convenience, these rates were determined using a linear fit to the initial portion of each desorption kinetics measurement. The desorption rates for both undoped and 4 mol.\% TiCl\(_3\)-doped NaAlH\(_4\) are plotted on a log scale versus 1/T in Figure 49. This is essentially an Arrhenius plot. The effect that Ti doping has on the desorption behavior of NaAlH\(_4\) is apparent from these measurements.
Figure 49. Log desorption rates versus inverse temperature plotted for NaAlH₄ with no TiCl₃ and 4 mol% added TiCl₃ [Original plot modified for simplicity, Na₃AlH₆ removed from figure].

The activation energies calculated from this plot as well as other Ti doping levels are presented in Figure 50. This series of kinetics measurements demonstrate that adding only small amounts of TiCl₃ to the alanates significantly reduces the activation energy for hydrogen desorption. It is interesting to note from Figure 50 that increasing the level of TiCl₃ in NaAlH₄ shows little change in the activation energy. Thus, the chemistry of the Ti-enhanced kinetics is unchanged with Ti concentration. Further improvement in the desorption rates by increased TiCl₃ doping is observed through an increase in the pre-exponential factor A of Equation 15. This indicates that performance is improved by the increased distribution and access to titanium in the material.
If the purpose of the kinetics measurements are indeed to study fundamental mechanisms then it is absolutely necessary to design the experimental measurement equipment to achieve essentially isothermal conditions and at an even higher level of precision, isobaric measurements. This was in fact done with painstaking detail by Goodell et al. for hydrogen absorption and desorption in LaNi$_5$. Their thermal ballast technique is described in detail in the Thermal Effects section 4.9.1 of experimental considerations.

4.4 Efficient Testing

Evolution of hydrogen storage material discovery, development and characterization is directly dependent on the ability to conduct measurements with speed and efficiency. Kinetic activity is often the limiting factor in hydrogen testing. For example, the time required to reach equilibrium in performing PCT measurements is directly dependent on the kinetics of the sample material. Any technique that reduces sample characterization time will help in improving the pace of material discovery and development.

An excellent example of efficient testing used by Gary Sandrock drastically reduces the time required to determine the activation energy of a sample with slow kinetics. To develop an Arrhenius diagram necessary to determine the activation energy of a chemical reaction, several experiments must be performed at different temperatures to determine the relationship between the rate constant $K$ and temperature. Instead of performing a series of isothermal experiments, Sandrock’s method changes the temperature of the sample in steps, as shown in Figure 51.
Figure 51. Example of efficient testing in determining the activation energy of a sample with slow intrinsic kinetics. The figure itself is a kinetics measurement used to relate the rate constant to temperature.

Several rate measurements can be performed using this method in the same amount of time as a single isothermal measurement. It is important to remember that the temperature is stepped instead of ramped because isothermal conditions are still required to perform kinetics tests. For such type of measurements to be valid the sample holder and heater design should allow temperatures to be changed and establish a steady on a time scale that is short compared to the kinetic rates. This technique can only be used with samples that have slow enough intrinsic kinetics that the sample temperature can be increased in several steps before the sample completely desorbs. In addition, rates should be measured when the sample is below about 60% conversion to avoid the impact of changes in reactant concentration on the rate measurements.

### 4.4.1 Kinetics and Capacity

With respect to kinetics measurements it is important to review how capacity is reported. In particular, it may be more useful to examine relative sorption rates of materials based on what we term here as “active capacity” in contrast to “material capacity”.

Today, hydrogen storage capacities are typically presented in units of weight percent (wt.%), where wt% is defined as:

\[
\text{wt.\%} = \frac{\text{mass}_{H_2}}{\text{mass}_{\text{sample}} + \text{mass}_{H_2}} * 100\%
\]
This makes sense from a practical perspective, because it is a measure of how much hydrogen can actually be delivered from a storage material. And, in this vein, it is important to recognize that ultimately the practical hydrogen storage capacity must be on a systems level that includes the material, containment vessel, heat transfer system, controls and balance of plant. However, from the perspective of a materials development level, it is equally important to be able to quantify capacity in terms of the active material only. In particular, where this becomes important is in the evaluation of improvements of kinetic performance. Since rates are typically being measured in units of wt.% per time, to make a comparative analysis of the effect on kinetics of different catalysts or additives, rates should be compared on an active wt.% capacity basis and not on material wt.% capacity. Thus, for this purpose, we may define

\[
\text{Equation 34} \quad \text{Material wt.\%} = \frac{\text{mass}_{H_2}}{\text{mass}_{\text{sample}} + \text{mass}_{H_2}} \times 100\%
\]

\[
\text{Equation 35} \quad \text{Active wt.\%} = \frac{\text{mass}_{H_2}}{\text{mass}_{\text{active sample}} + \text{mass}_{H_2}} \times 100\%
\]

The mass of the active sample only includes the mass of material that participates in hydrogen storage and the mass of the sample is the total weight of the sample including both the active and inactive material in the sample. Another possible way to perform such a comparative analysis is to use rates based on reacted fraction per time rather than absolute capacity values. As will be seen in the next section, for comparative kinetics it can be very important to present capacity in terms of reacted fraction or active wt.% rather than true wt.%.

### 4.4.2 Active Capacity

A specific example is presented here to demonstrate the difference between “material capacity” and “active capacity” and to aid in understanding the impact of this differentiation in evaluating relative kinetics of a series of different samples.

, NaAlH<sub>4</sub> doped with TiCl<sub>3</sub> will react during the milling preparation process to form NaCl according to:  

\[
\text{Equation 36} \quad \text{NaAlH}_4 + x\text{TiCl}_3 \rightarrow x\text{Ti} + 3x\text{NaCl} + 3x\text{Al} + (1 - 3x)\text{NaAlH}_4 + 3x\text{Al} + 6xH_2
\]

This means that only

\[
\text{Equation 37} \quad \frac{54(1 - 3x)}{54 + 154.5x}
\]
of the sample, representing the fraction of active sodium alanate to the total sample, is active for hydrogen uptake and release. Without taking this into account, improvements in alanates by doping with 6 mol% vs. 2 mol% TiCl$_3$ would be underestimated by up to 30% (Figure 52). In this case, this has little impact on the conclusions with respect to the level of dopant since the correction actually improves the rates with increasing levels of doping. However, another example may serve to demonstrate how this effect could impact such an evaluation.

![Figure 52](image)

**Figure 52.** Arrhenius plot comparing desorption rates of NaAlH$_4$ Generation II, III-D, and III-E alanates (rates: II on a material weight basis, III-D and III-E on a NaH + Al weight basis). Red arrows indicate change in data points on converting from a material to an active weight basis.\(^{74}\)

To investigate the effects of active capacity versus material capacity on kinetics measurements, a sample was tested using only TiH$_2$ instead of TiCl$_3$. In this case, a mixture of NaH, Al, and TiH$_2$ in a 1:1:1 composition was mechanically milled and tested for hydrogen absorption and desorption. From an applications perspective, TiH$_2$ as a precursor should overcome the problem of capacity loss associated with the formation to Na-halides when using Ti-halide precursors.
This composition did absorb hydrogen (130°C and 82 bar H₂) to form NaAlH₄. In this case, the inactive component (TiH₂) makes up almost half of the samples mass. Thus rates would be off by a factor of 2 if capacity were measured on wt.% basis using the sample’s total weight. However, by plotting rates on an active material mass basis (NaH + Al), the desorption rates are nearly identical to those of the indirectly doped material and the 2 mol% TiCl₃ doped alanates (Figure 52). Had the total weight of the sample been used, an incorrect conclusion may have been drawn about Ti-hydride vs. Ti-halide additives. With optimization, kinetics may be achieved without a severe degradation in hydrogen storage capacity. This example of kinetic analysis based on active versus material weight capacity serves to demonstrate that even small details in how the analysis is performed can have a big impact on the conclusions and ultimately on decisions concerning the direction of materials research and development.

Note that it may be better to report results for this type of analysis in terms of reacted fraction. Capacities reported as wt% of the active material can be misleading if not fully and carefully explained, because the capacity reduction that accompanies additives and catalysts is suppressed.

4.5 Activation Effects

Activation effects are important in kinetics measurements because the kinetic character of a reversible storage material can change drastically during the first five to ten cycles. The change in kinetic character is most significant early in cycling and gradually approaches the intrinsic kinetic character of the material. In application, a hydrogen storage material will typically cycle hundreds, if not thousands, of times so it is important to characterize the intrinsic kinetics in order to evaluate a material’s suitability for end-use. It is also important to run samples of new materials through several absorption/desorption cycles to determine their true intrinsic kinetic character. While the activation process is time-consuming, it is better than missing a viable storage material because its kinetic properties appear uninteresting after the first few cycles.

Pre-reacted sodium alanate is an excellent example of kinetic activation effects. Figure 53 is kinetic data taken from a cycling experiment on sodium alanate and clearly shows the drastic impact activation effects can have on kinetic character.

Each desorption measurement is taken in a fixed time interval so the increasing hydrogen desorption indicates that the kinetics of the sample are improving. The rate of hydrogen evolution increases markedly the first five cycles and gradually approaches the intrinsic kinetic rate from cycle 5-13. The class of pre-reacted sodium alanates would have been completely disregarded if judgment were passed based only on the kinetic character and capacity of the first few cycles.
Figure 53. The kinetic activation effects of pre-reacted sodium alanate. The marked increase in desorbed hydrogen indicates increasing kinetic rates.

Digging a little deeper into the kinetic character of sodium alanate, it is interesting to note that there are two distinct rates in each desorption cycle in Figure 53. Each represents a separate step in the two-step equilibrium desorption reaction.

Equation 38

\[
NaAlH_4 \rightarrow \frac{1}{3} Na_3AlH_6 + \frac{2}{3} Al + H_2 \rightarrow NaH + Al + \frac{3}{2} H_2
\]

The difference in the kinetic rates of the two reactions is probably due to the difference in the thermodynamic stability of NaAlH_4 and Na_3AlH_6. NaAlH_4 is less stable than Na_3AlH_6 and forms at a higher equilibrium pressure (Figure 54). In these desorption measurements, the under-pressure is below both equilibrium plateau pressures, allowing the decomposition of both alanate phases. However, the pressure difference driving the desorption reaction from NaAlH_4 to Na_3AlH_6 (DP_d (1)) is greater than for Na_3AlH_6 than for NaH (DP_d (2)), leading to higher kinetic rates for NaAlH_4 than Na_3AlH_6. It is important to note that the situation is exactly the opposite during absorption measurements. The lower plateau pressure of the NaH to Na_3AlH_6 transition would provide a greater driving pressure differential (DP_a (2)) compared to the Na_3AlH_6 to NaAlH_4 transition (DP_a (1)). Thus, in absorption, higher kinetic rates would be observed for the reverse NaH + Al to Na_3AlH_6 step in the reaction of Equation 38.
Figure 54. The reaction follows the equilibrium pressure (darkest line) from bottom left to top right for absorption and top right to bottom left for desorption. The horizontal section of the equilibrium pressure line represent phase transitions (reactions), the vertical lines represent the alanate compounds as indicated.

At first glance it appears that only the first reaction in Equation 38 is affected by activation because nearly all of the increased desorption capacity occurs during the first section of each desorption profile. As the material is cycled, the NaAlH₄ to Na₃AlH₆ reaction becomes more active while the Na₃AlH₆ to NaH reaction remains constant. This is easily rationalized with the aid of Figure 54. Before the first hydriding of the material, the sample is in its as prepared NaH + Al state. After the first charging which is incomplete (a1), the sample is dehydrided (d1) but can not completely desorb all of the hydrogen taken up in the first absorption due to the poor desorption kinetics of the second Na₃AlH₆ to NaH step and the fixed absorption/desorption times used for each cycle. This creates a permanent offset in total hydrogen concentration. During subsequent cycles (a2-d2 and a3-d3), only the NaAlH₄ to Na₃AlH₆ reaction improves (activates) because the NaH to Na₃AlH₆ to has already been taken to completion in the first cycle. It is important to remember that Figure 53 presents desorption rates only and not both sorption and desorption rates.
4.6 Gas Impurities

4.6.1 Retardation

Retardation in kinetics occurs when impurities such as CO₂, H₂O and NH₃ in the H₂ gas stream react with storage material surfaces, leading to decreased kinetic activity. The gas stream impurities occlude catalytic sites and diffusion pathways and can be difficult to desorb from the storage material because of their high thermodynamic activity. These effects influence the mechanisms discussed in the theory section 3.1. The primary retardation considerations for physisorbing materials like porous media are decreased surface area due to physisorbed impurities and the occlusion of internal pore networks; blocked catalytic sites and diffusion pathways affect chemisorbing materials like metal hydrides. Unlike poisoning, retardation does not significantly affect ultimate hydrogen storage capacity: the sorption curve of a material experiencing retardation asymptotically approaches the original hydrogen storage capacity of the material, albeit at a slower rate.

Sandrock et al. presents an excellent example of the retardation effects of NH₃ on LaNi₅ during cycling.⁷⁵ Although Figure 56 only shows the beginning of the sorption curve, each arc eventually approaches the ultimate hydrogen sorption capacity dictated by cycle 0. The effects of retardation are often reversed by flushing and subsequent use of high-purity H₂.
4.6.2 Gas Impurity Effects on Kinetic Performance

An important consideration is whether fresh or recycled gas is used during cycling kinetics measurements. For recycled gas measurements, the temperature of the sample is increased and decreased with each cycle with the sample exposed to the same gas in a fixed gas volume. Thus, the total amount of impurities in the hydrogen gas/system remains the same. When fresh gas is used for every cycle, gas impurities may have a much larger impact on the retardation of the material’s kinetics. This is because the gas impurities are strongly bound to the surface of the material and are not released upon desorption. With cycling, the concentration of impurities on the sample increases, degrading the material’s storage performance. In this sense the storage material is simply acting as an impurity getter. The advantage with gettering storage materials is that the hydrogen delivered to the application (e.g. fuel cell) will be much higher purity than that provided by the gas supplier. The disadvantage, of course, is that the performance of the storage system may be significantly affected.

As a side note, kinetics measurements performed on gravimetric systems in flowing gas mode, impurities in the gas stream that react with the sample will be observed as a slow linear increase in weight rather that the curved behavior expected for a true kinetics curve (much like leaks in a volumetric system discussed below). Such behavior should be taken as an indicator that impurities are present in the hydrogen gas stream.

4.7 Leaks

Leaks cause kinetic sorption/desorption profiles to appear linear in time, as opposed to true sorption profiles that exhibit curvature. An example of a leak is demonstrated in Figure 57, where the expected capacity of the sample is 1.39
wt.%. Sorption steadily increased with a linear behavior until a fitting was tightened about two hours into the experiment. With respect to new materials, one should weigh on the side of skepticism and review the setup and measurements if the results appear too good to be true.

**Figure 57.** Example of a leak on kinetics measurement. A little after an hour into the experiment, the leak was eliminated.

### 4.8 Pressure Effects

Observed kinetic rates are highly dependent on the pressure differential between the gas reservoir and the sample reactor that drives the sorption/desorption reaction. The theoretical description of the effect of pressure on kinetics was presented in section 3.2.3. and the experimental results can be measured as shown in the example of Figure 59 in section 4.8.1 below.

From an experimental perspective, in order to investigate the intrinsic kinetic properties of a material, it is important to reduce the degree of variation in pressure differential during testing. If the system pressure changes significantly during a sorption/desorption reaction, the changing pressure differential will affect the observed kinetic rates and mask the intrinsic kinetic properties of a material. It is, however, important to balance loss in sensitivity in the measurement due to the smaller total pressure change with the ultimate impact that a changing pressure may have on the kinetic behavior. In the volumetric method, this is done by appropriate selection of the volume reservoir so that the system pressure change is relatively sufficient for accurate measurements but not large compared to the driving pressure differential during reaction.
4.8.1 Reservoir Selection

In volumetric measurements, it is important that the dosing volume, sample holder volume and dosing pressures are selected to match not only the size of the sample but also the type of measurement. One simple way to estimate an appropriate selection is to calculate an assumed amount of gas that the sample will absorb and divide this by the number of doses that are needed to collect a reasonable set of data. For an absorption kinetics measurement the dosing volume should be large enough to be able to dose enough gas to completely charge the sample with the final pressure being greater than any plateau pressure and still result in enough of a pressure drop to make an accurate measurement. A typical value would be 110% of the capacity of the sample at the resulting pressure. For a desorption kinetics measurement the same is true with the sample being able to nearly completely desorb all of its gas to the reservoir volume. A typical value would be 95% of the capacity of the sample at the resulting pressure. For PCT isotherm measurements the volume of the reservoir and dosing pressures essentially determine how many data points will be collected on an isotherm. A good rule of thumb is that the size of the reservoir and pressure step should be selected to completely charge or discharge the sample in about 30 doses. This of course is dependent on the size of the sample, its capacity and charging pressures.

As a particular example, the effect of pressure on kinetics is illustrated in Figure 58 for a 1 g sample of LaNi\(_5\). When using a volume reservoir that is too small (10 mL, dashed line), there is not enough hydrogen to hydride the entire sample to LaNi\(_5\)H\(_6\). Therefore the pressure in the system must be the plateau pressure of LaNi\(_5\). The amount of hydrogen in the reservoir is

\[
\text{Equation 39} \quad n = \frac{PV}{RT} = \frac{5\text{bar} \times .01L}{.083\text{bar} \times \text{L} \times .083\text{mol} \times \text{K} \times 298K} = .002\text{molH}_2
\]

A small portion of the hydrogen remains as a gas in the system to satisfy the thermodynamic plateau pressure.

\[
\text{Equation 40} \quad n = \frac{PV}{RT} = \frac{1\text{bar} \times .01L}{.083\text{bar} \times \text{L} \times .083\text{mol} \times \text{K} \times 298K} = .0004\text{molH}_2
\]

The remaining hydrogen, .0016 mol H\(_2\), enters the LaNi\(_5\) sample to create .234 g of LaNi\(_5\)H\(_6\). Because there is not enough hydrogen in the 10 mL reservoir, the system pressure drop is from 5 bar to 1 bar and is equal to the driving pressure differential; such a situation would greatly affect the observed kinetics.
In contrast, a 400 mL reservoir (solid line) under the same conditions has enough hydrogen to hydride the entire sample. The amount of hydrogen in the 400 mL reservoir is

\[ n = \frac{PV}{RT} = \frac{5 \text{bar} \times 0.083 \text{bar} \times 0.083 \text{L} \times 298 \text{K}}{0.083 \text{mol} \times 298 \text{K}} = 0.08 \text{mol} \text{H}_2 \]

After complete hydriding, there is 0.073 mol H\(_2\) that remains in gas phase. The pressure associated with 0.073 mol H\(_2\) is

\[ P = \frac{nRT}{V} = \frac{0.073 \text{mol} \times 0.083 \text{bar} \times 0.083 \text{L} \times 298 \text{K}}{0.083 \text{mol} \times 298 \text{K}} = 4.5 \text{bar} \]

In contrast, a large reservoir under the same conditions has enough hydrogen to hydride the entire sample. After complete hydriding, there is enough H\(_2\) remaining in the gas phase to maintain a pressure of 4.5 bar. The change in system pressure during hydriding is about 0.5 bar (solid line), much less than the 4 bar pressure differential driving the sorption reaction. Data from the large reservoir experiment would have measured sorption to full capacity. The larger reservoir also provides the quasi-isobaric sorption kinetics of the LaNi\(_5\) sample as the rates are not impacted by a significant change in the driving which is logarithmic with pressure differential.

**Figure 58.** Diagram of the pressure in sorption/desorption system during hydriding reaction. The dashed line represents a test using a reservoir that is too small and the solid red line represents a test using a sufficiently large reservoir.
Another example is presented below to show the important effect of the driving pressure on kinetics. A series of absorption and desorption measurements were performed on NaAlH₄ by Weifang Luo who then fit the data with an empirical rate model. In this model, the absorption and desorption rates were dependent not only on temperature but also on the driving pressure (ΔP is the difference between the equilibrium pressure at a given temperature and the applied pressure). The model was validated by comparing predicted desorption curves with measured curves for changing pressures and temperatures. Once validated, the model was then used to predict absorption and desorption rates over a wide range of pressure and temperatures. Absorption rates for the formation of NaAlH₄ from Na₃AlH₆, Al and H₂ are presented in Figure 59. These results have important consequences for how such materials would be used in a hydrogen storage application and a first look may be counter intuitive. That is because the plateau pressures increase with temperature for a given applied pressure and the driving pressure differential decreases. Therefore, it is possible that even though mobility increases with temperature, there are practical conditions where it is better to operate at a lower temperature to have faster hydrogen charging of the storage material. This can be seen if one compares the absorption rate at 120°C versus 140°C at a charging pressure of 50 bar in Figure 59.

![Figure 59. Calculated formation rates of NaAlH₄ at several temperatures and applied pressures](image-url)
4.9 Thermal Effects

Arguably the most significant factor in accurate kinetics measurements is the interplay of thermal effects in the hydrogen storage material, material holder and measurement system. The importance of thermal effects has been discussed on several occasions in this recommended practices manuscript and will be reiterated. Isothermal or controlled scanning temperature conditions are required to take meaningful fundamental kinetics measurements, in particular when attempting to identify rate-limiting mechanisms. Unfortunately, large temperature variations can occur locally due to the exo- and endothermic reactions and inadequate material thermal conductivity.

4.9.1 Thermal Ballast

Several techniques have been developed to ensure isothermal conditions and minimize thermal effects. Goodell and Rudman were two of the first to obtain approximately isothermal conditions in the kinetics testing of hydrogen storage materials. Their method is simple: ‘thermal ballast’ is added to the hydride to increase the heat capacity of the sample and dampen the thermal gradients in the system. Ballast material is carefully paired with the hydride being tested to ensure porosity, non-reactivity and minimal contamination. The effect of ballast on the temperature excursions upon hydriding is defined by

\[
\Delta T = \left( -\frac{\Delta H}{6.2R} \right) \ln\left(1 + (1-b)x_p\right)
\]

with \(\Delta H\) the heat of reaction, \(b\) the atomic fraction of ballast and \(x_p\) the fractional hydride composition of the sample. Equation 43 is presented graphically in Figure 60. To give an example of the significant thermal effects at work in kinetics studies, 98% of a LaNi\(_5\)–ballast sample must be thermal ballast in order to limit the temperature change during reaction to within 10K of ambient temperature.
4.9.2 Heat Transfer-Maximizing Cell

The second method used to eliminate heat transfer effects is to employ a specially designed testing cell to maximize heat transfer to the sample. Maximum heat transfer is accomplished by using a reaction bed that is as thin as possible along with a thermocouple in the bed itself. Supper et al. designed a specialty cell to eliminate heat transfer effects and study the influence of other parameters on kinetics. In the Supper cell, a thin disk of sample material is sandwiched between a porous metal disk that allows for gas transfer and an integrated water-copper heat pipe that acts as an isothermal heat source or sink. The heat pipe is regulated using an external water loop.
The Supper cell in Figure 61 was used to investigate the effects of temperature and sample thickness on kinetics in LaNi$_{4.7}$Al$_{3}$. For the sample with 1 mm bed thickness, increasing the temperature of the sample enhanced the kinetics of the material in agreement with Arrhenius’ law.

![Graph showing concentration over time](image)

Figure 62. Measurements taken with the Supper Cell on LaNi$_{4.7}$Al$_{3}$ to investigate the effects of temperature and sample thickness on kinetics. Sample bed thickness of 1 mm.

### 4.9.3 Effect of Sample Thickness

The effect of increasing sample thickness on the kinetics at various temperatures in the example above is worth noting. As the thickness of the sample bed is increased, the relation between temperature and kinetics is inversed; reaction rate decreases with elevating temperature, as seen in Figure 63. Supper et al. provide an excellent explanation: the increased thickness of the bed increases the temperature in the bed as the reaction proceeds because the heat of reaction cannot be removed effectively. The van’t Hoff thermodynamic relation dictates that the increase in temperature of the sample requires a commensurate increase in the equilibrium pressure in the system. The increased equilibrium pressure reduces the driving pressure differential at higher temperatures because the experiments were all performed at the same distance from equilibrium at the initial sample temperature. This decrease in the true pressure differential because of non-isothermal conditions in turn decreased the rate of reaction. This example of the effect of sample thickness on kinetics shows the complexity of heat transfer effects in drawing conclusions from experimental observation.
Figure 63. Measurements taken with the Supper Cell on LaNi$_{4.7}$Al$_{0.3}$ to investigate the effects of temperature and sample thickness on kinetics. Sample bed thickness of 6 mm.
5 Approaches to Improve Kinetics

The kinetic behavior of a material is controlled by mechanisms that are determined by specific material properties. Consequently, it should be possible to improve kinetics by altering these material properties.

5.1 Alloying

By alloying hydrogen storage materials with catalysts and/or other foreign substances, it is possible to improve kinetics in a variety of ways. In chemisorbing materials, catalysts aid in the dissociation of hydrogen at the gas/material interface and allow for more rapid diffusion of atomic hydrogen into the bulk. A stylized view of how this may affect the rate-controlling mechanisms is presented in Figure 64. Although oxygen (white) may passivate one metal (dark gray), the introduction of a second element (light gray) may provide active sites for dissociation, and may also create diffusion paths into the bulk. Metal hydrides can also be blended together to manipulate the thermodynamic and kinetic properties of the base material. The addition of catalysts in physisorbing media may enable the material to utilize the spillover effect (see associated paragraph in the preface). This effect is most pronounced in carbon-based physisorbing media including graphitic sheets and carbon nano-tubes.

Figure 64. The presence of a second element, either in solution or as a compound, can improve kinetics by providing active sites for dissociation and creating a diffusion path through the lattice.11
5.2 Grain Boundaries
Intimate contact between two different phases may improve kinetics by enhanced diffusion through inter-phase grain boundaries (Figure 65). Intimate contact between separated phases can be produced by mechanically milling a mixture of phases together, by vapor deposition of one phase onto the others, or by the solid-state disproportionation reaction of a compound upon hydriding.

![Diagram of intimate contact between two different phases with hydrogen diffusion](image)

**Figure 65.** Intimate contact between two different phases may allow atomic hydrogen to diffuse through an active phase and cross the phase boundary into the passivated phase.

5.3 Decrepitation
Kinetics may be enhanced by crack formation due to lattice expansion during hydrogen sorption. In metal hydrides, these cracks bring clean metal surfaces into contact with hydrogen. The oxygen-free surface may promote the hydrogen dissociation reaction and diffusion into the bulk. This process may also be enhanced in multi-phase systems by cracks that form along inter-phase boundaries because of the different amount of lattice expansion in separate hydride-forming as well as non-hydride phases. Such a situation is pictured in Figure 66.
5.4 Nano-Structuring

Kinetics may be inhibited by diffusion through the lattice of chemisorbing media or through the internal pore network of high-surface area physisorbing materials. In both cases, nano-structuring can increase the diffusivity of hydrogen. In chemisorbing media, nano-structuring increases the surface area-to-volume ratio, which allows for greater access to the bulk and shorter diffusion paths (Figure 67). Diffusion pathways through the internal pore network of physisorbing media are often tortuous and can become occluded. Nano-structuring can offer more direct diffusion pathways and greater access to internal surfaces.
5.5 Example of Improvements

An example of enhanced sorption kinetics by altering material properties is the observed rapid absorption and desorption kinetics of Mg-hydride formation in multi-phase composite materials based on Mg, Mg$_2$Ni and La. A combination of the above concepts was ascribed to improved kinetics measured in these composite materials. The composites were produced by mechanically milling La$_2$Mg$_{17}$ together with LaNi$_5$. In the process, fine particles of LaNi$_5$ were encompassed by the softer La$_2$Mg$_{17}$ to create an intimate contact between the two phases (Figure 68). In a second step the composites underwent a phase and morphology transformation during the rapid absorption and desorption of hydrogen at 300°C. This resulted in the disproportionation of the La$_2$Mg$_{17}$ and LaNi$_5$ phases. Morphological changes occurred as the La coalesced into fine particles and the Ni combined with Mg in a solid-state reaction to form larger grains of Mg$_2$Ni. During desorption cycles the remaining Mg likely diffused and sublimated to form a matrix around the other phases. The final product was a composite of fine particles (<1 micron) composed of Mg$_2$Ni covered with small grains of La-hydride all held together and coated with Mg (Figure 69).
The kinetics of these composite materials proved to be far superior to those of the individual Mg and Mg$_2$Ni phases. A study of the relative kinetics of each phase in the composite revealed that the probable mechanisms for the enhanced kinetics were enhanced catalytic dissociation of hydrogen on the Mg$_2$Ni and La phases, rapid diffusion along inter-phase boundaries and a reduction of the MgH$_2$ diffusion barrier due to the extremely large surface area of the Mg phase.
6 Distinguishing Between Kinetics and Thermodynamics

In pursuit of the ultimate hydrogen storage material it is important to separate kinetic effects from the thermodynamic properties of a material. This is because a material may be kinetically hindered, requiring high temperatures and pressures to achieve reasonable hydrogen sorption/desorption rates, and these high operating conditions are regularly mistaken as inherent thermodynamic properties of the sample. All too often, a material is hastily excluded from hydrogen storage consideration on the basis of its initially observed inability to sorb/desorb hydrogen at moderate temperatures and pressures. In reality the equilibrium thermodynamics may be perfectly adequate for a useful hydrogen storage system but poor kinetics require testing outside a practical range of temperatures and pressures. Figure 70 shows what is considered the range of standard operating conditions for a Proton Exchange Membrane (PEM) fuel cell (shown in grey). As an extreme example, consider a hypothetical material that can thermodynamically operate at room temperature, but which is kinetically hindered such that much higher temperatures would be required to overcome the kinetic barriers. Because the equilibrium pressure increases with increasing temperature Figure 70, it is possible that the kinetic barrier could suppress hydrogenation over the entire range of experimentally accessible pressures, i.e., by the time a temperature is reached where the kinetics allow reaction, the required pressure is too high to be achieved.

![Figure 70](image-url)

**Figure 70.** Equilibrium thermodynamics of some commonly researched hydrogen storage materials. The boxed area represents the range of practical operating conditions for a PEM Fuel Cell.
A classic example of the relationship between kinetics and thermodynamics is investigated in the work by Bogdanovic and Schwickardi on the effect of Ti-additives on the hydrogen sorption properties of alanates. Prior to their work, alanates were not seriously considered as hydrogen storage materials because of the extreme pressures and temperatures required for hydrogen absorption. Bogdanovic et al. systematically doped sodium alanate with varying levels of titanium compounds and found that this significantly improved kinetics and by consequence reversibility. This opened the doorway to what followed as an intensive worldwide study of the alanate family as practical hydrogen storage materials. Until the Bogdanovic work, little was done to determine the thermodynamics of the sodium-alanate system because it was nearly impossible to reach thermodynamic equilibrium due to slow intrinsic kinetics of the pure alanate system. Only after the discovery of greatly enhanced kinetics in samples doped with titanium precursors was it even feasible to make equilibrium PCT measurements, and even these achieve only partial equilibrium conditions at lower pressures and temperatures due to kinetic limitations.
Summary
It is our intent that this overview of fundamental processes and measurement considerations will aid research and development of new and better hydrogen storage materials. In particular we hope to have clarified some of the recommended practices and caveats in performing high-quality experiments to measure the hydrogen sorption and desorption kinetics properties of advanced materials.

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