Hydriding of titanium: Recent trends and perspectives in advanced characterization and multiscale modeling

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A B S T R A C T

Titanium (Ti) and its alloys are attractive for a wide variety of structural and functional applications owing to excellent specific strength, toughness and stiffness, and corrosion resistance. However, if exposed to hydrogen sources, these alloys are susceptible to hydride formation in the form of TiHx (0 < x ≤ 2), leading to crack initiation and mechanical failure due to lattice deformation and stress accumulation. The kinetics of the hydriding process depends on several factors, including the critical saturation threshold for hydrogen within Ti, the specific interaction of hydrogen with protective surface oxide, the rates of mass transport, and the kinetics of nucleation and phase transformation. Unfortunately, key knowledge gaps and challenges remain regarding the details of these coupled processes, which take place across vast ranges of time and length scales and are often difficult to probe directly. This work reviews recent advances in multiscale characterization and modeling efforts in Ti hydriding. We identify unanswered questions and key challenges, propose new perspectives on how to solve these remaining issues, and close knowledge gaps by discussing and demonstrating specific opportunities for integrating advanced characterization and multiscale modeling to elucidate chemistry and composition, microstructure phenomena, and macroscale performance and testing.

1. General overview of hydriding of Ti

1.1. Introduction to Ti

Titanium (Ti) and Ti-based alloys possess superior combined physical and mechanical properties in addition to good corrosion resistance, which make them outstanding materials for a variety of structural and functional applications [1–8]. For instance, the high specific strength (strength-to-weight ratio) and corrosion resistance make Ti a widely used material in aerospace (such as airframe parts and jet engine blades), automotive, and energy applications (such as petrochemical and nuclear-power-generation components) [3,5,6]. Because of their excellent biocompatibility in vivo, Ti alloys have also become increasingly attractive for biomedical applications [2,9].

Ti exists in two allotropic crystalline forms: the high-temperature β phase with a body-centered cubic (bcc) structure and the low-temperature α phase with a hexagonal close-packed (hcp) structure [5,8,10]. Consequently, Ti alloys can be categorized using a nomenclature based on the predominant phases within their microstructure at room temperature. Alloys consisting of mainly of the α phase are referred to as α-alloys (or near-α alloys if a small amount of β phase is present). Conversely, β-dominated Ti alloys are called β-alloys, whereas those featuring a mixture of α and β phases are called α/β alloys.

Alloying elements in Ti alloys usually act as α or β stabilizers, which are required to achieve the desired mechanical properties, including tensile strength, creep, and fracture toughness, as well as resistance to degradation modes such as fatigue crack propagation, stress-corrosion cracking, and oxidation [1,3,11,12]. Further optimization of
mechanical properties can be achieved through thermal and mechanical treatments to engineer microstructures with desired sizes, shapes, spatial distributions, and interface coherency states of α precipitates within the β matrix. Although the microstructure is primarily composed of two simple phases, many different phase transformation pathways from β to α exist, providing ample opportunities for engineering and optimizing the microstructure towards specific applications. A commonly employed Ti metal is the unalloyed commercially pure (CP) Ti family, which is typically classified by the amount of impurities such as oxygen and iron \[13\]. A representative Ti alloy of particular interest is Ti-6Al-4 V, which is considered the “workhorse” among α/β Ti-alloys \[14,15\].

1.2. Hydriding in Ti

Despite the general corrosion resistance, there is a major concern associated with hydrogen or hydride phases residing in the Ti alloy matrix. Historically, this has led to issues with environmentally assisted cracking, including hydrogen embrittlement (HE) and hydride-induced cracking (HIC) \[5,16\]. Specifically, titanium hydride in the form of TiH\(_x\) (0 < x ≤ 2) often precipitates at the matrix surface and in the bulk material in application environments featuring H\(_2\) gas, atomic H, or H\(^+\) \[17–21\], which can eventually induce crack initiation and mechanical failure as the lattice deforms and stress accumulates \[22,23\]. Hydride formation can also be induced by an electrochemical process, which involves oxidation of Ti and reduction of H\(^+\) (proton) or atomic H \[17–20,24,25\]. Externally applied tensile loads and residual stresses promote ingress of hydrogen into the Ti matrix, eventually inducing embrittlement, with cracks preferentially propagating along phase boundaries or directly across matrix grains \[7\]. Hydrides can also play a role in the corrosion susceptibility of Ti and Ti alloys, either acting as a protective barrier on the matrix surface inhibiting corrosion attack \[17,26\] or assisting crevice corrosion and stress-induced corrosion cracking at local microstructural defects such as inclusions and phase interfaces \[7,27\].

In the past few decades, there have been significant efforts to better understand the physicochemical phenomena underlying Ti hydrogenation, including H adsorption, transport, and hydride phase formation, as well as their impact on the mechanical, microstructural, and corrosion properties of base materials \[5,17–20,23–35\]. Broadly speaking, these
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Recent research advances in Ti hydriding phenomena have been driven by increasingly mature and emerging experimental characterization tools (see Fig. 1) including, but not limited to: in operando X-ray diffraction (XRD) and imaging [36,37], atom probe tomography (APT) [33,34], in situ atomic force microscopy/scanning Kelvin probe (AFM/ SKP) [38,39 40–46], in situ scanning and/or transmission electron microscopy (S/TEM) [24,35], and cryo-controlled characterizations [34]. Meanwhile, the fundamental understanding of hydriding has been catalyzed by progress in modeling efforts from atomistic to continuum scale with significantly improved computational capabilities (Fig. 1): density functional theory (DFT) calculations [32,33,47], molecular dynamics (MD) [18,48], Monte Carlo (MC) [48,49], phase-fielding modeling (PFM) [8,15,50], and integrated models [21,51–53] that transcend idealized representations. However, a number of knowledge gaps and challenges, some of which are listed in Section 1 (Table 1) and will be discussed in detail in Section 2, remain at different length scales. In our view, tighter integration of experimental and modeling research, either through a validation feedback cycle or direct comparison, is sorely needed to form a comprehensive story of Ti hydriding phenomena and to resolve these remaining challenges.

This work is intended to briefly review the recent advances in characterization and modeling efforts for investigating a variety of fundamental chemical, physical, and materials processes associated with Ti hydriding, as well as how these phenomena are coupled. In particular, we focus on how such experimental and modeling efforts have been applied to shed light on key identified challenges. Finally, we propose new perspectives on how advanced characterization and multiscale modeling could be more effectively integrated to bridge remaining knowledge gaps, as illustrated schematically in Fig. 1.

2. Recent advances and key challenges

2.1. Chemistry and composition

2.1.1. Hydrogen uptake

There are three methods reported in the literature for charging H into the Ti matrix, namely, electrochemical charging, gaseous charging, and H ion irradiation. The first two are the most frequently used. Here the mechanisms of these methods are briefly introduced, and their advantages and disadvantages are discussed.

2.1.1.1. Electrochemical charging. In the electrochemical charging method, TiH₂ formation can occur in Ti metals/alkyls via corrosion processes in acidic or deaerated aqueous environments. In these processes, H adsorption is readily accessible from cathodic discharge of H⁻—i.e., cathodic galvanostatic or potentiostatic polarization or by interfacing Ti with a more galvanic metal [7,17,18,26,27,54–58]. Further penetration through the outer oxide film into the alloy matrix involves coupled processes of additional hydrogen absorption and redox chemistry, which are generalized in equation (1):

\[
\text{TiO}_2 + x\text{H}^+ + xe^{-} \rightarrow (\text{Ti}^{IV})_{1-x} (\text{Ti}^{III})_x \text{O}_{2-2x} (\text{OH})_x \quad (1) .
\]
Experimental observations demonstrate that H absorption through intact TiO₂ films into Ti becomes efficient at a cathodic potential range around ~0.6 to ~1 V_SCE [18,56]. The resulting TiH₂ composition varies depending on values of the cathodic charging current densities and/or charging potential [17]. Notably, defects can aid hydrogen permeation through the oxide film, for example via the weak points associated with structural defects including oxygen vacancies, grain boundaries and triple junctions in polycrystalline oxides, and lower-density regions associated with amorphous oxides or porosity; in such cases, the required cathodic potential for penetration can be less negative, e.g., ~−0.35 V_SCE [18,28]. External stress also introduces defects in the oxide film and matrix that eventually lead to less negative cathodic potentials for H charging [7,57]. Accordingly, Ti hydrides formed on Ti metal surfaces have been shown to be unevenly distributed, and preferentially located at intermetallics and grain boundaries [58]. As another strategy to avoid the oxide-induced impedance, Abul-Hamid and Latanision sometimes high H₂ partial pressure, gaseous hydrogenation can penetrate the surface oxide rapidly and crack the surface. The higher temperatures for gaseous charging can also lead to microstructural evolution of the matrix due to an annealing effect, which is a concurrent process with hydrogenation. Ion implantation adds the factor of irradiation damage to the hydrogenation layer and matrix by introducing a significant amount of microstructural and point defects, including vacancies and cation interstitials. The hydriding layer in this case is also usually limited by a short implantation depth range of nanometers to a few micrometers.

For all three charging modes, the exposure times and resulting hydrogen concentration gradients can also introduce additional complexities. In particular, the nature of the surface oxides (e.g., thickness and composition) changes as a function of charging time, so the uptake behavior is rarely uniform. Moreover, the time required for complete hydrogen penetration and dissipation of intrinsic hydrogen concentration gradients in Ti depends on the specimen size and can be limited by the continual accumulation of near-surface hydrogen and hydrides. Such accumulation can impede diffusion of hydrogen and promote additional steep concentration gradients between sample surface and center.

For the reasons stated above, a direct comparison between the three methodologies is not straightforward. Accordingly, development of models that represent these three methods should take in to account the different physical and chemical phenomena that govern hydrogen uptake in each case. In addition, the following efforts towards more systematic and reliable comparisons across different charging methods are suggested: 1) use a readily accessible pseudo reference material, such as a single crystal, that can be used by different researchers to minimize the effects of individual materials properties on charging behavior; 2) explicitly specify the charging method and strictly control and report charging parameters to ensure repeatability and reproducibility; 3) develop proper analysis protocols that can eventually be published as “standard or gold” rules for certain standard alloys or pseudo reference materials, such as a standard protocol for TDS measurement.

### Table 2

Representative hydrogen diffusion coefficients in Ti oxide, Ti matrix, and Ti grain boundaries based on experimental measurements.

<table>
<thead>
<tr>
<th>Materials or features</th>
<th>Diffusion coefficient, cm² s⁻¹</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single crystal rutile</td>
<td>7.5 × 10⁻¹⁰</td>
<td>[76]</td>
</tr>
<tr>
<td></td>
<td>10⁻¹¹ - 10⁻¹³</td>
<td>[85]</td>
</tr>
<tr>
<td>Polycrystalline rutile thin film</td>
<td>2.9 × 10⁻¹⁰</td>
<td>[77]</td>
</tr>
<tr>
<td>Polycrystalline anatase thin film</td>
<td>4 × 10⁻¹⁵</td>
<td>[80]</td>
</tr>
<tr>
<td>Amorphous TiO₂ thin film</td>
<td>1 × 10⁻¹⁵</td>
<td>[80]</td>
</tr>
<tr>
<td></td>
<td>10⁻¹⁸ - 10⁻¹⁷</td>
<td>[28]</td>
</tr>
<tr>
<td>a-Ti matrix</td>
<td>2.6 × 10⁻¹⁰</td>
<td>[86]</td>
</tr>
<tr>
<td>β-Ti matrix</td>
<td>– 5 × 10⁻⁹</td>
<td>[87]</td>
</tr>
<tr>
<td>Grain boundary in pure Ti matrix</td>
<td>9.1 × 10⁻⁸</td>
<td>[59]</td>
</tr>
</tbody>
</table>

2.1.1.4. Comparison and discussion. Although electrochemical charging, gaseous charging, and ion implantation for hydrogen share some similarities (e.g., hydrogen adsorption, penetration through the oxide film, hydride formation, as well as coupling of these processes to defects) [72,73], datasets collected using different methods should not necessarily be interpreted in the same way. This is mainly because the driving forces and associated mechanisms differ across the three methods: electrochemical potential/current for electrochemical charging, temperature and pressure for gaseous charging, and irradiative damage for ion implantation.

Electrochemical charging usually develops a much higher hydrogen partial pressure (e.g., on the order of 100 atm or higher at the metal-liquid interface [72]), which leads to the amount of absorbed hydrogen in the Ti surface region being significantly higher. However, unlike gaseous charging, the amount of hydrogen electrochemically charged into the matrix is limited by the TSS of hydrogen in that matrix [73], which means hydrogen diffusion and hydriding kinetics in the subsurface are usually limited by the ambient charging temperature (≤100 °C). Another difference is that electrochemical charging often induces a highly hydroxylated surface due to the abundance of water, leading to fundamentally different surface processes. In addition, because of its much higher temperature range (480–650 °C) and sometimes high H₂ partial pressure, gaseous hydrogenation can penetrate the surface oxide rapidly and crack the surface. The higher temperatures for gaseous charging can also lead to microstructural evolution of the matrix due to an annealing effect, which is a concurrent process with hydrogenation. Ion implantation adds the factor of irradiation damage to the hydrogenation layer and matrix by introducing a significant amount of microstructural and point defects, including vacancies and cation interstitials. The hydriding layer in this case is also usually limited by a short implantation depth range of nanometers to a few micrometers.
2.1.2. Hydrogen transport through oxide surface layers

The surface Ti oxide film can exist in either a crystalline (typically anatase and/or rutile) or a noncrystalline form, or else as a mixture of the two. This surface oxide layer acts as a robust barrier layer that protects the Ti matrix from H ingress [74,75], significantly slowing down the hydrogen diffusion rate. As summarized in Table 2, an apparent hydrogen diffusion coefficient was reported in the range of 10^-18-10^-15 cm^2 s^-1 in amorphous TiO_2 thin films [28,76] and around 2.9 × 10^-20 in rutile thin films [77]. In bulk TiO_2, a H diffusion coefficient was estimated around 5.3 × 10^-16 cm^2 s^-1 at ambient temperature, based on experimental measurements on single-crystal rutile [76]; a similar diffusion coefficient of 2.84 × 10^-16 cm^2 s^-1 at 373 K was found for oxides covering pure α-Ti metal [78].

The hydrogen barrier efficacy of Ti oxides strongly depends on the oxide structure and composition [55,79]. Structural defects such as vacancies, grain boundaries, and triple junctions at the Ti surface, as well as chemical heterogeneities such as cathodic intermetallic compounds (e.g., Ti-Pd) can significantly impact hydrogen transport [13]. In general, faster H diffusion is expected in amorphous or polycrystalline TiO_2 than in single-crystal TiO_2 considering the structural and chemical defects in the former. However, the existing data is not always consistent with this notion. For example, a recent study reported a much higher apparent H diffusion coefficient of 10^-15 cm^2 s^-1 that was similar in both amorphous and amorphous/anatase mixed oxide films [80]. This discrepancy is likely due to porosity in their column structure contributing to the fast diffusion, which renders the amorphous and partially crystalline samples indistinguishable.

Moreover, the H-charging process itself can lead to concurrent compositional, structural, and electronic changes in the surface oxides, which in turn can influence H transport and trapping in the oxides and the underlying matrix metal as the reaction proceeds. Hannula et al. [81] recently indicated that atomic H treatment of deposited TiO_2 (at 300 °C in UHV chamber) created an electronically “leaky” and photoelectrochemically unstable film due to the formation of O vacancies and TiO_2 dissociation. This “leaky” phenomenon, which signals a change in electronic structure, should be heeded during routine hydrogen charging of the Ti oxide surface and subsequent analysis.

In parallel with the experimental work, computational studies based on first-principles DFT calculations have been carried out to investigate the mechanism of hydrogen transport in Ti oxides. These studies have largely focused on crystalline systems with specific orientations; nevertheless, several important findings have been reported. Specifically, H migration from surface to subsurface sites in anatase has a lower kinetic barrier than that of H2 desorption, which indicates that H likely preferentially diffuses to the subsurface and further transports into the bulk [82,83]. In addition, subsurface oxygen vacancies can favorably accommodate H [82,83], and preexisting atomic H in the oxides enhances H diffusion into the subsurface while simultaneously preventing the escape of H [84].

Collectively, these studies not only provide a detailed understanding of hydrogen transport in the oxide, but also suggest a possible strategy to enhance the hydrogen storage capability of the material. Nevertheless, given that H diffusion data are inconsistent, and that the specific connection to structural and compositional complexities in TiO_2 are not well established, much remains to be understood regarding the mechanism and degree to which the Ti oxide film can act as a hydrogen barrier. Approaches to precisely predict hydrogen transport through the oxide and its potential interaction with specific heterogeneities within the oxide are particularly needed.

2.1.3. Hydrogen transport and trapping in Ti

Following surface oxide permeation, H can be transported, redistributed, and/or trapped within the Ti matrix. H transport in Ti usually occurs via three pathways: short-circuit transport via lattice defects, dislocation movement, and diffusion along grain boundaries. It was reported that for the pure Ti matrix, the effective diffusion coefficient of H along grain boundaries is around 9.1 × 10^5 cm^2 s^-1 [59] (Table 2) - significantly higher than the H transport within the grain lattice (e.g., 2.6 × 10^15 cm^2 s^-1 in Cu α-Ti grains [86]).

Compared to the Ti matrix, the diffusion kinetics in the hydride layer are reported to be about two orders of magnitude slower than in the Ti matrix, around 4 × 10^12 cm^2 s^-1 at room temperature [28,88]. This is in general agreement with extrapolations from first principles [89]. Once the hydride forms near the Ti surface due to continuous H adsorption, further H ingress into the metal would therefore be significantly inhibited. In addition, the H diffusion coefficient in the hydride further decreases with increasing content of H in Ti hydride [87].

H transport can also be inhibited by irreversible trapping, for instance at various types of matrix defects (e.g., dislocations, precipitates, and grain boundaries). In principle, the concentration of trapped H, as well as the associated binding energies at trapping sites, can be deduced through relevant calculations and analysis of adsorption/desorption profiles [90,91]. Section 2.2 provides a more complete discussion of defect interactions with H. In addition to the specific connection between these defects and hydrogen trapping in Ti metal and Ti-based alloys [91], insights may be gleaned from analogous studies on various steels. For instance, several trapping energies (0.14, 0.11, 0.27 and 0.54 eV) corresponding to different peaks in the thermal desorption spectra of H have been reported for martensitic steel. These values of trapping energy are comparable to those reported for other tempered martensite (0.15 and 0.68 eV) [92], pure iron (0.17, 0.35, 0.60 eV) [93,94], and low-carbon steel (0.13, 0.24 and 0.50 eV) [95] samples. The consistency among these independent reports suggests that H is preferentially trapped in the same types of trapping sites with comparable trapping energies. These trapping energies from low to high are associated respectively with interstitial sites (energy of ~0.16 eV) [93], reversible trapping sites (~0.29 eV) related to elastic field near-edge dislocations [90,94] or to semi-coherent precipitates [90,96,97], and irreversible trapping sites (~0.59 eV) related to dislocation cores [98].

2.2. Microstructure phenomena

2.2.1. H-dislocation interaction

When H concentrates around dislocations in metals, it impacts the interactions between these dislocations and other dislocations/defects and can eventually lead to local plasticity, with impacts similar to macroscopic brittle fracture. Whereas the nature of hydrogen-dislocation interactions has been extensively studied in BCC metals, there have been far fewer relevant studies in HCP metals such as Ti. It is often supposed that hydrogen affects dislocation mobility via the kink-pair nucleation and migration mechanism [99–102]. Low concentrations of hydrogen (e.g., 0.1 atom ppm) enhance dislocation mobility by promoting kink-pair nucleation whereas high concentrations of hydrogen (e.g., 10 atom ppm) decrease dislocation mobility by impeding kink-pair migration [102]. Note that mobility of screw dislocations is much lower than edge dislocations, so plasticity is majorly limited by the former in bcc metals [103]; however, the relevant reports of hydrogen-dislocation interaction in HCP metals are very limited [99–102]. The impact of hydrogen on dislocation mobility has also been discussed in the context of the hydrogen-enhanced local plasticity (HELP) mechanism on local hydrogen-dislocation interactions [48]. Within HELP, the movement of hydrogen clouds around dislocations follows the movement of the dislocations during plastic deformation, and HE is largely determined by the H-dislocation interactions via the population of free electrons in the hydrogen clouds [51].

To reveal these linkages, advances have consistently been made both experimentally and theoretically on the quantification of H concentration and dislocation interactions in various HE-sensitive metals, including Ti alloys [29,30,32,48,49,89,104–120]. From the experimental side, attempts to link local H-dislocation interactions to microscopic embrittlement in Ti could benefit from more detailed studies of H in other metal alloys using advanced techniques. For example, the series...
of works on Ni and steel HE reported by Barnoush et al. [105–107,111,118] using the in situ electrochemical nanoindentation (ECNI) method falls within the category of nanometer-to-micro-scale examination of both H-defect interactions and mechanical responses. Quantitative in situ TEM nanomechanical characterization could likewise provide key insights, as utilized by Zhu et al. [110] in a pioneering study on local H-dislocation interactions at free metal nanowire surfaces, which reported hydrogen-induced suppression of dislocation nucleation, contrary to the well-reported hydrogen-induced promotion of such nucleation in many bulk metals. Note these most other investigations of H-defect interactions (e.g., with vacancies, dislocations, grain boundaries, or intermetallics) have relied on macroscopic testing [113,114,117,121,122], which normally combines effects of multiple defects with different length scales. The challenge remains to separate the intrinsic impact of dislocations from other defects, such as twinning nucleation and stacking faults, on hydriding behavior.

Ab initio simulations based upon DFT can provide a reliable computational tool to quantitatively evaluate the energetics of defect interactions during the Ti hydriding processes [123,124]. However, complex defects such as dislocation networks or clusters, as well as stacking faults and interfacial/phase boundaries, feature large strain fields that are often difficult to address with DFT alone. Classical molecular dynamics simulations are one alternative, but it is difficult to validate reliable interatomic potentials for defect interactions in the Ti-H system [89], which affects predictions of hydrogenation phenomena. For example, the inconsistency between classical molecular dynamics and ab initio simulations for H diffusion in Ti hydride was found to most likely be due to the Ti-H interatomic potential [112]. Beyond this, the materials in service are often under external load, which concentrates stresses that can facilitate H transport and lead to local buildup of H, promoting dislocation nucleation. Such mechanisms are common in metals and add to the difficulty with modeling these systems, typically requiring integration of models at different length scales.

Overall, despite advances in understanding, robust quantitative descriptions of localized H at or near dislocations and their specific linkage to the microscopic/macroscopic brittleness of the Ti matrix are still lacking. Associated questions remaining to be answered are: what is the local H content near dislocations? What is the strength of the H-dislocation interaction and how can the dislocation interaction be deconvoluted from that of other defects? How do the interactions impact local hydride formation and plasticity and further link to macroscopic mechanical response, including brittleness? Closer feedback between multiscale modeling and experimental investigations, which provide complementary benefits and liabilities, may provide a more suitable approach to isolate specific H-dislocation interactions and trace their impacts across scales in hydride-forming materials.

2.2.2. H-grain/phase boundary interaction

In addition to its interaction with dislocations, H can also impact Ti properties by interacting with grain boundaries. A few studies have been carried out at atomic level on the hydrogen impact on the mobility of twin boundaries (TB) in Ti, e.g., (112 T) [125]. These studies confirm that the (112 T) TB glides easily and would readily encounter the mobile H atoms [126]. Note also that the energy barrier for TB shifting increases gradually with increasing hydrogen concentration. Unfortunately, such studies remain limited, and information regarding the interaction of H with other grain boundary orientations is sorely needed.

Literature reports also indicate that grain boundaries and phase boundaries in Ti can act as preferential nucleation sites for growth of hydrides [127,128]. There have been additional reports exhibiting a high fraction of intragranular hydrides found in matrices even with relatively large grains [24,129]. For example, the nucleation and growth of TiH₃ hydride in Ti64 initially occurred in the β phase because of the high diffusivity and contiguity of H in this phase [30,31]. In such instances, the process is accompanied by local stress accumulation and H confinement near the alpha/beta boundary, which eventually leads to hydride formation preferentially along the boundary. Such studies highlight the complex interaction between atomic disorder, stress, and composition at phase and grain boundaries, which remains difficult to unravel with most current approaches.

2.2.3. H-impurities/inclusions interaction

There are several major metallic impurities in CP Ti, such as Fe, Pd, and Ni, that may influence the hydrogenation of Ti because they are H sensitive. Fe is the most common impurity and present in many grades of CP Ti [130]. Pd is usually present in multi-grades of CP Ti metals such as Grades 7, 16, 18, and 24 [131], and Ni is usually found in CP Ti metal Grade 12 [131]. These elements usually dissolve in the solid solution of Ti, either in the α-phase, the β-phase, or at the α/β interface. Their solubility tends to be higher in the β-phase than in the α-phase due to their affinity and stability in the BCC structure. In fact, these metallic impurities prefer to incorporate substitutionally as stabilizers in β-Ti regardless of their atomic size or chemistry [132]. However, when the impurity content exceeds their solubility in the Ti solid solution, they tend to precipitate to form Ti-R (R = Fe, Pd, or Ni) intermetallic compounds such as TiFe and Ti₅Fe [133], TiPd and Ti₃Pd [134,138], and Ti₂Ni [13,139,140], further complicating the hydriding mechanism. For example, Ti-Fe particles can precipitate even if the Fe impurity concentration is as low as ~0.03 wt% [130].

Extensive research regarding the influence of the elemental impurities (Fe [27,58,59,141–143], Pd [144,145], and Ni [13,139]) on hydrogen adsorption/absorption, hydrogen transport (diffusion and penetration), and hydriding has been carried out. However, the findings are often inconclusive or contradictory. Some studies reported that enriched elemental impurities in Ti solid solutions provide an entry point for hydrogen to the underlying Ti metal and promote hydrogen adsorption/absorption and hydrogen transport [139,144,146]. However, other studies determined that hydrogen incorporation and transport efficiency decreased with the rise in elemental impurity content in Ti [141,145]. For example, Cotton [146] reported that diffusion rate was proportional to the concentration of elemental Fe, and that hydriding was promoted by elemental Fe impurities. However, Covington [141] demonstrated that less hydrogen was absorbed as the Fe content increased and indicated that low levels of Fe in Ti do not make it more susceptible to hydriding.

The impact of Ti-R intermetallic compounds on hydriding phenomena have also been investigated. Ti-Fe particles can impede hydrogen absorption and diffusion because hydride formation in/around Ti-Fe particles usually causes less lattice distortion than in Ti solid solutions [147]. Among these lines, Yan et al. [27] found that the hydrogen penetration depth into the matrix decreased with the presence of Ti-Fe particles. Liu et al. [17] reported preferential H aggregation at Fe-containing particles along grain boundaries in CP Ti-2, which might be due to either enhanced H diffusion or preferential formation of hydrides. Grade 16 CP Ti was shown to be more resistant to hydride formation and HIC than Grade 2 CP Ti and 12104–105 Ti due to the presence of Pd-Ti particles in Grade 16 and the high solubility of H in these Ti-Pd intermetallic compounds, which act as a H sink [154,155]. Authors of these works further stated that the Ti-Pd compounds themselves do not provide an entrance for H absorption but rather catalyze proton reduction and hydrogen evolution, which apparently contradicts with the high solubility of H in Ti-Pd particles and with Pd content promoting Ti adhesion [134–140]. Hydrogen adhesion can also be promoted by the β-phase of Ti as it is usually rich in impurities and can be considered as an impurity-containing intermetallic compound. The β-phase usually distributes discontinuously along grain boundaries in CP Ti and can form a continuous network in (α + β) Ti that acts as a fast diffusion pathway for H [13,139].

The enrichment and distribution of Pd, Ni, and Fe can clearly have a significant influence on the hydrogen adsorption, penetration, and hydriding behavior of Ti. However, the conflicting nature of some of the
literature reports, as well as a lack of understanding of key mechanistic details, prompt additional investigation. Critical information is missing to thoroughly understand the role of these elemental or particle impurities in hydrogen adsorption, transport, and hydride formation. Research efforts to reveal the decoupled effects on each of these processes is particularly important not only for Ti, but also for other hydrogen-sensitive metals, e.g., Zr and Al [148-150]. Note that non-metallic impurities e.g., C, O, and N are not discussed here, but it is helpful to realize that these impurities can also compete with H adsorption within Ti matrix [151], particularly at defects that also function as H sinks.

2.2.4. Hydride nucleation and growth

Hydride formation and the resulting composition are also strongly dependent on microstructural features. In general, three types of Ti hydrides with different compositions and crystal structures are consistently reported: face-centered tetragonal (FCT) ε-hydride (TiH₂, c/a < 1); stable face-centered cubic (FCC) δ-hydride (TiH₂, 1.5 < x < 1.99); and metastable FCT γ-hydride (TiH, c/a > 1) [24,152,153]. The lowest-stoichiometry hydride was reported to be TiH₀.₅±₀.₂ with a lattice parameter a of 4.40 Å, formed in α-Ti [19,154]. At lower concentrations, H remains in α-Ti with no hydride precipitate. The δ-phase hydride has been the most frequently observed Ti hydride. Hydrides grown under galvanostatic conditions, regardless of variation in charging durations or charging current densities, primarily consisted of TiH₁.₅ (δ-hydride) with a minor fraction of TiH₁.₂ and TiH₂ [177]. The preferential formation of δ-hydride has been attributed to the relatively small misfit strain at the δ-hydrate/matrix interface [155], close-packed substrate textures for both δ-hydride and α-matrix [156,157], preferential co-existence of solid solutions of both the δ-hydride and α-matrix phases [127,158], and preferred orientation of δ-hydrate (200) along Ti (002) [34,127]. Formation of plate-shaped or lath δ-hydrate was also confirmed in α/β Ti-55 alloys [159].

It has been reported that hydride nucleation in Ti matrix usually follows four different orientations, as tabulated in Table 3 [24,152,153]. Among these, two of the most frequent orientations (ORs) are OR1 and OR2 for the α-Ti/δ-hydride transition [160]. Details of ORs between hydrides and the Ti matrix have been documented elsewhere [24,152,153]. Nevertheless, the specific impacts of evolving strain, dislocation density, composition, interfacial coherency, and similar factors on hydride nucleation probability and preferred ORs have eluded detailed investigation.

Experimentally, one challenge in addressing key unknowns regarding hydride nucleation is the sensitivity of conclusions to sample preparation. Hydride formation can readily occur even from sample grinding and polishing, which complicates interpretation. In particular, Ti hydride nucleation and phase transformation are commonly identified in the process of thin sample fabrication using focused ion beams (FIB) and in subsequent TEM analysis [34,35]. For example, the conventional FIB-liftout process often induces δ- and/or γ-hydrides [35,161]. It has been further demonstrated that FIB-induced Ti hydrides do not precipitate within grains but rather preferentially form along α-phase grain boundaries and along α/β phase boundaries [33]. The temperature of sample preparation also plays a critical role, as no hydride formation was observed in cryo-plasma-FIB-prepared samples [34]. On the other hand, thermally produced FCC Ti-X phases are closely related to contamination by oxygen in high-purity Ti thin samples based on in situ TEM heating studies at ~ 600 °C [35,162], likely with analogous effects for hydride formation. Zr has a similar concern of contamination during sample preparation [163,164]. Sample handling is likewise critical because exposure to air can result in H contamination in the form of moisture [165]. To minimize H (and also O) contamination, Breen et al. [166] performed all transfers and APT analysis under vacuum and cryogenic conditions. Such cryogenic conditions served two purposes: (1) to slow out-diffusion of H from the sample; and (2) to reduce background H by removing water vapor, which is often a major component of residual gas in vacuum chambers.

2.3. Mechanical performance & testing

2.3.1. Mechanical properties of Ti hydrides

The mechanical behavior of hydrides is complex: the deformation and brittleness of hydrides are determined by the hydride distribution, morphology, size, and orientation, as well as environmental factors such as temperature and applied stress [22,47,167-174]. Acquiring a thorough understanding of these intrinsic mechanical properties and their dependencies on materials features and environmental conditions is critical to discern potential impacts on HE and HIC tendency. Although some hydride mechanical property data have been documented in literature [32,47,172,174], reports are still very limited, particularly from detailed experiments [167,174]. Some existing data regarding the mechanical properties of hydrides, including fracture toughness (K), elastic modulus (E), bulk modulus (B), and shear modulus (G), are summarized in Table 4.

The fracture toughness of hydrides, for example, has been studied only for the δ-phase (TiH₁.₅) using bending experiments, and it was reported to be around 2.2 MPa*m⁰.₅ [167]. This is a common value for ceramics but significantly below the value of 50-70 MPa*m¹.₂ reported for the metallic matrix α-Ti [168]. The fracture toughness of the hydrides was calculated to be 0.7 and 0.4 MPa*m¹.₂, respectively, for the δ-phase (TiH₁.₅) and ε-phase (TiH₂) based on ab-initio calculations and Griffith-Irwin theory [32]. Note that the discrepancy between experimental observations and DFT calculations can be explained in part by temperature effects, given that the ground-state DFT calculations are performed at 0 K, whereas low temperature usually decreases toughness [169].

The difference in fracture toughness between the δ-phase (TiH₁.₅) and ε-phase (TiH₂) correlates with the Young’s modulus, which is about two times larger for the δ-phase (125 MPa*m¹.₂ by experiment [174] and 151 MPa*m¹.₂ by DFT [47]) than for the ε-phase (60.1 MPa*m¹.₂ by DFT [47]). Note only one existing experimental dataset [174] that includes Young’s modulus (125 MPa*m¹.₂), bulk modulus (125 MPa*m¹.₂), and shear modulus (43 MPa*m¹.₂) has been reported for δ-hydride to date, and no experimental values for the other Ti-hydrides exist, highlighting the data scarcity of mechanical properties of Ti-hydrides.

2.3.2. Mechanical/stress response to hydride formation

Hydrogen incorporation into the Ti matrix causes lattice deformation, and further formation of Ti hydrides locally destroys the original matrix lattice. These structural changes are likely to contribute to the accumulation of volumetric and interfacial stresses, which can eventually lead to HE and/or HIC. Such effects have been observed in other hydriding-susceptible metals.

For example, H incorporation in Zircaloy-4 was found to induce significant strain in the δ lattice, leading to a transformation from the FCC δ-hydrate to the HCP γ-hydrate (Zr₂H) [165]. Similarly, H charging in steel led to austenite phase peak splitting and the development of heterogeneous tensile strains on the basis of synchrotron grazing-incidence X-ray diffraction (GIXRD) studies [186]. Nevertheless, the specific connection between stress and hydride formation in the case of Ti derivatives is not entirely clear. For instance, J. Wen et al. [128], found that electrochemical H-charging in the β-21S Ti alloy over time did not lead to hydride nucleation even though a proportional increase

<table>
<thead>
<tr>
<th>Orientation relationship</th>
<th>Interface plane</th>
</tr>
</thead>
<tbody>
<tr>
<td>OR1 (0001)₀ // (1 1 0)₀, c&lt;1 (0 0 1)₀ // (0 1 1)₀</td>
<td>(1 0 1)₀ // (1 1 0)₀</td>
</tr>
<tr>
<td>OR2 (0001)₁ // (0 0 1)₁, c&lt;1 (1 1 0)₁ // (0 1 1)₁</td>
<td>(0 0 1)₁ // (1 0 1)₁</td>
</tr>
<tr>
<td>OR3 (1 1 1)₂ // (1 1 1)₂, c&lt;1 (1 1 0)₂ // (0 1 1)₂</td>
<td>(1 0 1)₂ // (1 1 0)₂</td>
</tr>
<tr>
<td>OR4 (0 0 1)₃ // (0 0 1)₃, c&lt;1 (1 1 0)₃ // (0 1 1)₃</td>
<td>(1 0 1)₃ // (1 1 0)₃</td>
</tr>
</tbody>
</table>
example, an increase in the average corrosion potential and decreased surface hydride formation [17,26,58,188]. The increased corrosion potential is an indication of decreased overpotential for the cathodic hydrogen evolution reaction (HER) reaction and corresponds to the Ti potential. These findings suggest that grains with orientations that can more efficiently accommodate the strain associated with both OR1 and OR2 types (in Table 3) of hydride phase provide useful measurements to compare to computational modeling work. However, a general consensus regarding a single mechanism, including hydrogen-enhanced localized plasticity (HELP) [122,196–198], adsorption-induced dislocation emission (AIDE) [199,200], hydrogen-enhanced decohesion (HEDE) [201–203], and hydrogen-enhanced strain-induced vacancies (HESIV) [204–208], have been proposed in the past based on microscopic observations and modeling work. However, a general consensus regarding a single dominant mechanism by which hydrogen causes embrittlement has not been reached. More detailed discussions of HE mechanisms have been well documented elsewhere [91,109,122,196–200,204–208].

2.3.4. Hydrogen embrittlement without hydride formation

A significant amount of work has been dedicated to the mechanism of HE-induced failure in hydrogen-sensitive metals. Several possible HE mechanisms, including hydrogen-enhanced localized plasticity (HELP) [122,196–198], adsorption-induced dislocation emission (AIDE) [199,200], hydrogen-enhanced decohesion (HEDE) [201–203], and hydrogen-enhanced strain-induced vacancies (HESIV) [204–208], have been proposed in the past based on microscopic observations and modeling work. However, a general consensus regarding a single dominant mechanism by which hydrogen causes embrittlement has not been reached. More detailed discussions of HE mechanisms have been well documented elsewhere [91,109,122,196–200,204–208].

2.3.5. Hydride impact on localized stress corrosion cracking

Ti is sensitive to localized corrosion in an aqueous environment of low pH, high temperature, and/or high concentration of halides e.g., 0.5 wt% HCl or 1 wt% H₂SO₄ at 100 °C [209,210]. External mechanical loading or internal residual stress at localized corrosion sites further adds to the sensitivity of Ti to stress-assisted localized corrosion and cracking (SCC). Studying SCC under external stress loads is also practically relevant, as Ti alloys used as structural materials in a service environment must usually withstand external mechanical loading. Some recent progress and remaining issues regarding the mechanism of SCC under loading or at the crack tip as it pertains to hydride formation, hydrogen enrichment, or mixed oxygen/hydrogen enrichment are briefly discussed below.

Recent work has provided evidence that hydride formation itself can induce cracks, leading to mechanical failure in Ti alloys. Crack deflection and failure was observed in fatigue crack growth testing performed on Ti-10 V-2Fe-3Al alloys in high-pressure (8.3 MPa) H₂ gas [211]. The analysis revealed hydride-induced fatigue cracking, as TiH₂ was the dominant residual phase on the cracking surface. Furthermore, hydrogenation of the Ti β-phase was found to induce lattice deformation, introducing a localized stress field to the α/β boundary region that facilitated hydriding [31]. In this process, nanoscale cracks nucleated in the hydrides and propagated along the interface of the hydride with the β-phase, eventually leading to transgranular cracking across β grains. The large mismatch between the cracked β grains and the α grains also

in the lattice parameter of BCC β-Ti was observed.

We note that it can be difficult to experimentally reveal local mechanical responses to hydrogenation, in part due to the high mobility of H and the length scales associated with the strain. An efficient methodology to bridge this gap involves combining in situ nano-to-micron scale chemical and mechanical analysis with in situ hydrogen charging, which can eliminate the impact of hydrogen diffusion and loss during tests while simultaneously providing fundamental insights into local mechanical responses to hydrogenation with high resolution.

Meanwhile, these nano-to-micron scale experimental apparatus can provide useful measurements to compare to computational investigations at the atomic scale. For example, anisotropic strains associated with both OR1 and OR2 types (in Table 3) of hydride phase transformations have been modeled to reveal the orientation dependence [24]. The results show that for OR1-type δ-hydrides, the maximum and minimum transformation strains are achieved for directions along the {10T2.6}α and {0001}α plane normals, respectively. In contrast, for OR2-type hydrides, the maximum (of 21.8%) and minimum (of 17.6%) strains are achieved for directions along the {10T0}β and {0001}β plane normals, respectively. These findings suggest that grains with orientations that can more efficiently accommodate the strain associated with Ti-hydride transformation may also be more favorable for hydride formation. Further studies along these lines are recommended to reveal the full magnitude of the volumetric and interfacial stress effects on hydride formation and vice versa.

2.3.3. Hydride impact on general corrosion resistance

Similar to the functionality of the surface oxide film, the hydriding process creates a layer on Ti metal that ennobles the surface and is believed to provide some degree of corrosion inhibition [17,26,187]. For example, an increase in the average corrosion potential and decreased hydrogen discharge kinetics were reported for Ti metal and alloys upon surface hydride formation [17,26,58,188]. The increased corrosion potential is an indication of decreased overpotential for the cathodic hydrogen evolution reaction (HER) reaction and corresponds to the Ti potential. Ti was also found to be in the active dissolution state during tests while simultaneously providing fundamental insights into local mechanical responses to hydrogenation with high resolution.

Table 4

Summary of experimental (Expt) and DFT calculated lattice parameters (in angstroms) and mechanical properties (in MPa*m¹/²) of Ti hydrides.

<table>
<thead>
<tr>
<th>Hydride</th>
<th>Expt/DFT</th>
<th>a</th>
<th>c</th>
<th>E</th>
<th>B</th>
<th>G</th>
<th>K</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>TiH</td>
<td>Expt</td>
<td>4.21</td>
<td>4.59</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[158,175]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.168</td>
<td>4.584</td>
<td>183</td>
<td>126</td>
<td>73</td>
<td></td>
<td>[176]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.164</td>
<td>4.581</td>
<td>85.5</td>
<td>129</td>
<td>31.7</td>
<td></td>
<td>[177]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.171</td>
<td>4.581</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[47]</td>
</tr>
<tr>
<td>TiH₁.₅</td>
<td>Expt</td>
<td>4.40</td>
<td>4.40</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[157]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.360</td>
<td>4.404</td>
<td>125</td>
<td>125</td>
<td>43</td>
<td>2.2</td>
<td>[167,178]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.372</td>
<td>4.372</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[177]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.355</td>
<td>4.394</td>
<td>151</td>
<td>137</td>
<td>57.5</td>
<td>0.72</td>
<td>[32,47]</td>
</tr>
<tr>
<td>TiH₂</td>
<td>Expt</td>
<td>4.53</td>
<td>4.28</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[157]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.53</td>
<td>4.28</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[180]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.513</td>
<td>4.179</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[181]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.554</td>
<td>4.210</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[182]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.532</td>
<td>4.187</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[176]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.534</td>
<td>4.209</td>
<td>137</td>
<td>17.6</td>
<td></td>
<td></td>
<td>[183]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.517</td>
<td>4.201</td>
<td>121</td>
<td>139</td>
<td>45</td>
<td></td>
<td>[177]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.486</td>
<td>4.352</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[184]</td>
</tr>
<tr>
<td></td>
<td>DFT</td>
<td>4.527</td>
<td>4.199</td>
<td>60.1</td>
<td>141</td>
<td>21.1</td>
<td>0.43</td>
<td>[32,47]</td>
</tr>
</tbody>
</table>
ultimately resulted in transgranular cracking across the α grains. Adsorbed hydrogen can also chemically react with carbon at grain boundaries and produce methane gas, which can in turn lead to void formation and mechanical failure [212].

Nevertheless, hydride formation-aided SCC at a crack tip has been questioned in certain types of Ti alloys (e.g., Ti-4Al [30] and Ti-8Al-1Mo-1V [213–216]) due to two aspects: the slow hydride formation kinetics and its orientation. On one hand, the onset of hydride nucleation was not observed in Ti-4Al until about 14 min after gaseous H₂ charging under a pressure of 16 kPa [215], which is indeed significantly slower than the crack propagation rate of around 10⁻³ m/s in an aqueous environment [214]. On the other hand, intergranular hydride formation was observed along phase (or grain) interfaces in Ti-811 in an aqueous NaCl solution around neutral pH; however, SCC fracture produced hydride-free intragranular facets across grains [215,216], which is contrary to the finding of hydride-induced transgranular-cracking across grains in Ti64 [31]. Adding to this, S. Cao et al. [213,214] found that absorbed H at the crack tip accelerated dislocation emission, enhanced the dislocation density, and introduced severe plastic deformation ahead of the crack tip that produced additional dislocations and eventually led to cracking. These studies make it apparent that SCC and HE at and ahead of a crack tip are highly dynamic processes and at very localized sites involving microstructure evolution, passive film formation/breakdown, chemical/electrochemical reactions, and dynamic environmental conditions.

Recently, S. Joseph et al. [217] suggested that a synergistic effect of O and H solute atoms in the solid solution is responsible for cracking in Ti-6Al-2Sn-4Zr-6Mo in an aqueous (D₂O and H₂O) environment. This proposed scenario, in which O also plays a vital role, challenged the decades-old assumption that cracking of Ti alloys is related only to H (as interpreted by the HELP mechanism and the AIDE mechanism). This work inspires a new perspective for cracking studies that transition from a focus only on H towards the ingress of O—a critical insight for designing corrosion-resistant Ti-based materials.

It should also be noted that at this stage, none of the above mechanisms can be completely ruled out in favor of the others. Studies of the proposed mechanisms are at different stages of maturity and not always on the same materials or under the same conditions. As such, it is difficult to interpret characteristics beyond each individual experiment. Instead, it is necessary to carry out in-depth and highly systematic analysis to identify and validate missing characteristics, as well as to determine the conditions under which each of the representative mechanisms might manifest. Advances in both characterization capabilities and integrated multiphysics models could provide significant benefit for elucidating the different stages of cracking, including localized corrosion processes, crack nucleation, and crack propagation.

3. Opportunities and perspectives

Charting a course towards improving the performance of Ti metals and alloys in the presence of hydrogen relies on improved understanding of hydrogenation phenomena at the atomic to microstructural scales [34,35]. Achieving this understanding is an ongoing challenge, in large part because of the difficulty of making observations at the necessary length scales and deconvoluting causation associated with particular materials features. Integration of experimental approaches with modeling offers a unique opportunity to address these concerns [21,51–53,218–221]. Some general experiment-modeling integration strategies to probe relevant phenomena across multiple scales are presented in Fig. 2. Governing thermodynamic and kinetic processes associated with Ti hydriding are described in terms of composition and chemistry (atomistic scale), microstructure and mass transport (microstructural scale), and bulk hydriding and corrosion performance (macroscale). In each case, some physical quantities that can be both measured and modeled are identified, along with corresponding classes of techniques that can be utilized to achieve the handshaking between experiments and simulations: chemical spectroscopy, diffusion kinetics, phase growth/dissolution, and overall hydriding rate. The schematic in Fig. 2 underscores the need to combine multiscale models, multimodal materials characterization, and testing in order to achieve a holistic view.

**Fig. 2.** Strategies for integration of advanced characterization and modeling to tackle challenges listed in Table 1 in Ti hydriding, with representative examples from the authors’ work probing surface chemistry, mass transport, and hydrogen mass uptake. Top from left to right: Raman spectra for rutile and anatase TiO₂ native oxides at ambient conditions, dark field STEM imaging of hydrided α-Ti, and potential evolution during the galvanostatic H-charging on an α-Ti surface in deaerated NaCl solution. Bottom from left to right: DFT modeling of H atom in an amorphous TiO₂ native oxide, phase field modeling (PFM) of hydride formation in α-Ti, and integrated modeling of H uptake kinetics in TiO₂ native oxides with different phase and grain boundary composition approximations.
of Ti hydriding. Recent progress, opportunities, and perspectives in both experimental characterization and modeling will be discussed in this section, along with additional specific examples of proposed integration strategies.

### 3.1. Opportunities and perspectives in characterization methods

Although challenging, it is critical to develop and implement experimental characterization methods with high spatial resolution and dynamic capabilities. Table 5 summarizes the commonly used characterization methods, their capabilities, and the challenges listed in Table 1 that can potentially be addressed using these methods. Shaded techniques (rows 2–6) are destructive, whereas unshaded techniques are non-destructive.

#### Table 5
Characterization methods, their capabilities, and the related challenges in Table 1 that can potentially be addressed using these methods. Shaded techniques (rows 2–6) are destructive, whereas unshaded techniques are non-destructive.

<table>
<thead>
<tr>
<th>Methods</th>
<th>Length scale</th>
<th>Information</th>
<th>H/hydride associated defects</th>
<th>Associated challenges*</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEM-based (ex/in situ)</td>
<td>Sub-atomic to tens of μm</td>
<td>Atom position, lattice parameter, oxidation state, phase, microstructure, orientation, composition</td>
<td>Point defects, dislocations, boundaries, precipitates, nanocracks</td>
<td>1, 2, 3, 4, 5, 6, 7</td>
</tr>
<tr>
<td>SEM/EBSD-based (ex/in situ)</td>
<td>nm to mm</td>
<td>Phase, microstructure, orientation, composition</td>
<td>Dislocations, boundaries, microcracks, precipitates</td>
<td>2, 3, 4, 5, 6, 7</td>
</tr>
<tr>
<td>NanoSIMS (conventional/flash-frozen)</td>
<td>50 nm to μm on lateral, nm in depth</td>
<td>H content, phase (qualitative), depth profile</td>
<td>Boundaries, microcracks, fracture surface, precipitates</td>
<td>1, 2, 3, 4, 5, 6, 7</td>
</tr>
<tr>
<td>APT (conventional/flash-frozen)</td>
<td>Atomic to tens of nm</td>
<td>H content, composition (quantitative)</td>
<td>Substitutional solutes, dislocations, boundaries, precipitates</td>
<td>1, 2, 3, 4, 5, 6, 7</td>
</tr>
<tr>
<td>TDS (ex situ)</td>
<td>Bulk measurement, ppm to sub-ppm</td>
<td>H content in bulk &amp; defects, trap energies in defects (derived)</td>
<td>Point defects, dislocations, boundaries, precipitates</td>
<td>1, 2, 3, 5</td>
</tr>
<tr>
<td>SKPFM (ex situ)</td>
<td>nm to tens of μm</td>
<td>Topography, surface potential, H content</td>
<td>Boundaries, precipitates, microcracks</td>
<td>1, 2, 3, 5, 6, 7</td>
</tr>
<tr>
<td>ERAD</td>
<td>Tens of μm on lateral, nm in depth</td>
<td>H content (quantitative), depth profile</td>
<td>Charged surface</td>
<td>1, 4, 6</td>
</tr>
<tr>
<td>Neutron diffraction/scattering (ex/in situ)</td>
<td>nm spatial resolution</td>
<td>Atom position, lattice parameter, phase, orientation, strain</td>
<td>Point defects, Lattice displacement, precipitates, microcracks, charged bulk</td>
<td>1, 4, 5, 6</td>
</tr>
<tr>
<td>Neutron imaging (ex situ)</td>
<td>Tens of μm spatial resolution, 10 ppm mass resolution</td>
<td>H content (quantitative in 2D and 3D)</td>
<td>Precipitates, microcracks, macrocracks, charged bulk</td>
<td>1, 4, 5, 6</td>
</tr>
<tr>
<td>NRA</td>
<td>nm in depth</td>
<td>H content, depth profile</td>
<td>Charged surface</td>
<td>1, 4, 6</td>
</tr>
<tr>
<td>X-Ray diffraction/(imaging) (ex/in situ/operando)</td>
<td>Tens of nm to tens of μm on lateral, nm in depth</td>
<td>Lattice parameter, phase, strain, orientation, strain</td>
<td>Point defects, Lattice displacement, precipitates, microcracks, charged surface</td>
<td>1, 2, 3, 5, 6, 7</td>
</tr>
<tr>
<td>XAS (ex/in situ/operando)</td>
<td>nm in depth</td>
<td>Local coordination, oxidation state, bond distances, phase</td>
<td>Electronic and point defects</td>
<td>1, 2, 3, 5, 6, 7</td>
</tr>
</tbody>
</table>

*Indicates the most closely relevant challenges that can be routinely studied but does not exclude the possibility of studying other challenges.
example, leading methods for sample characterization include tomography and mass spectrometry for precise compositional analysis, and in situ or operando techniques like electron microscopy for structure and phase identification during the hydriding processes. Non-destructive, high-resolution experimental tools based on X-ray techniques and atomic force microscopy-based methods are likewise useful to collect information during materials degradation and have the advantage of doing so with minimal perturbation or impact of external factors, especially when these tools are employed in operando or in situ to monitor topography, phase, and electronic state evolution. In the case of hydriding, additional challenges arise from the difficulty of directly probing the tiny atom. Techniques based on neutron probes are particularly useful in this regard; additional hydrogen-sensitive characterization techniques are also documented in reference [222].

3.1.1. Development of high-resolution, localized hydrogen/hydride detection methods

3.1.1.1. Electron microscopy. Environmental electron microscopy, including environmental SEM and TEM [223,224], particularly when integrated with in situ nanomechanical testing, provides a powerful tool to investigate hydriding of metal alloys. The mechanical behavior of a hydrogenated matrix can be directly observed [197], and nanomechanical testing can be carried out on samples at the nano-to-micrometer scale e.g., nanopillars [225,226], nanowires [110], and nanocantilevers [107]. As an example of how this could be done, Xie et al. utilized quantitative environmental TEM nanomechanical testing [225] to report that dislocations in nanopillars of Al metal can be locked with hydrogen exposure and reactivated under cyclic loading without hydrogen exposure. Using a similar method, Yin et al. [110] observed that hydrogen suppresses the nucleation and movement of dislocations on the surface of Ag nanowires, rather than the usually reported hydrogen-enhanced dislocation nucleation and movement. Electron microscopy has also been applied to study hydriding of Ti-based metals. Q. Wang et al. [24] evaluated the orientation relationship between hydrides and the Ti metal matrix in CP titanium using an interrupted in situ electron backscatter diffraction (EBSD) setup. Kim et al. [30] developed an apparatus that combines electrochemical hydrogen charging in an aqueous environment with a high vacuum-based SEM/EBSD capability to study the dynamic hydrogenation process of Ti64. The emerging electron channeling contrast imaging (ECCI) approach [227,228] has also shown to be very useful in revealing the correlation between subnanometer-scale H-dislocation interactions and micrometer-scale heterogeneous phenomena and mechanical response by imaging dislocations in larger-area grains.

However, it must be realized that it is challenging to directly observe hydrides based on contrast of collected images using electron microscopes; instead, indirect approaches are generally required. TEM usually needs to operate in electron diffraction mode to capture characteristic diffraction patterns specific to each type of hydride. SEM typically identifies hydrides through equipped EBSD by collecting phase maps or Kikuchi patterns and comparing them to known crystallographic phase databases. Interestingly, Kooi et al. [229] very recently demonstrated unprecedented imaging of H atoms and hydrides in Ti by developing an integrated differential phase contrast method in STEM mode. Some other limitations of electron methods include limited field of view and sample size, as well as reduced vacuum (gaseous environment) and occasional radiolysis of water (in humid or liquid cell environments), which can reduce resolution due surface contamination [223,224]. The sample preparation process for TEM (e.g., through electro-polishing or FIB) also tends to introduce H contamination and facilitates preliminary hydride formation if not controlled properly.

3.1.1.2. Secondary ion mass spectrometry (SIMS). SIMS techniques, including Time-of-flight (ToF) SIMS and NanoSIMS, have been used to analyze the surface and depth distribution of hydrogen, hydride, or hydroxide, and to correlate with microscopy (including 3D) of surfaces at microstructurally relevant length scales in Ti alloys [17,27,29,133,220,230–240]. The advantages of SIMS are that it can analyze all elements (including H), distinguish isotopes, achieve extremely low detection thresholds (in ng/g or ppb range), and collect images of lateral chemical distributions (in certain instances) [236–239]. However, accurate SIMS analysis requires that the reference material be very similar to the studied material, given that sputter yields are highly dependent on chemical composition [239,241,242]. Therefore, SIMS is more often used for qualitative evaluation and visualization of hydrogen distribution and usually needs to be combined with other hydrogen-sensitive techniques for more reliable quantitative analysis. As an example, Y. Duan et al. [232] studied the influence of Ti-Mo interdiffusion on the distribution profile of deuterium (D) across TiDx/Mo films using the SIMS method, as well as a combined Rutherford backscattering (He3RBS) and D(3He,p)He nuclear reaction analysis (NRA) approach. It turned out that the SIMS-measured D concentration is remarkably smaller than the concentration obtained via RBS-NRA (this technique is discussed further below). Beam damage effects is another concern of SIMS, which compromises the analysis accuracy of localized hydrogen/deuterium signals [243].

3.1.1.3. Atom probe tomography (APT). APT utilizes a strong electric voltage to activate individual atoms of elements emitted from a needle-shaped sample, then chemically maps the 3D distribution of those elements [244]. APT is the only currently known technique with atomic spatial resolution that can precisely map hydrogen distribution and quantify its content in a material [33,34,161,165,166,217,245–247]. Independent researchers have further demonstrated that APT tips prepared by cryo-FIB can significantly lower hydrogen contamination compared with tips prepared under non-cryogenic conditions [34,166]. Cryogenic environment control throughout the sample tip preparation processes using, for example, cryo-FIB combined with a cryo-transferring chamber can further eliminate pre-existing H attached to samples [34,166]. These advances in technique development make it possible to properly leverage the unique advantages of APT towards analysis of hydrogen in Ti and Ti hydrides.

However, it is important to note that the APT tip can still be contaminated, usually in the range of 0.5–1.5 at%, by the residual H in the APT instrument chamber [35]. Hydrogen charging in a deuterium environment represents an alternative strategy that has been verified by others [165,166,246,247], although the noise from residual H in the collected spectra still perturbs the quantitative analysis to some degree. [165,166,246,247]. If other isotopes or used, it must be considered whether deuterium or tritium would exhibit the same segregation, phase, and distribution behavior compared to hydrogen, which is many cases is an unanswered question.

3.1.1.4. Flash freezing techniques. Beyond eliminating contaminations through cryo-environment control, an even more promising and exciting development involves the use of flash-freezing techniques, which have been successfully applied both to APT [248–252] and SIMS [253–255] at different length scales at solid–liquid interfaces. Flash freezing can “lock” the instantaneous elemental and ionic information (e.g., hydrogen, water, protons) at the regions of interest for performing “quasi in situ” analysis. For example, flash-frozen APT work [250] has been performed for 3D nanoscale analysis of composition and structure at the solid-water interface in corroded glass. These characterization advances should be transferable to hydriding studies of Ti, especially for trapping of H and local pH gradients in phase/grain boundaries, crevices, and matrix/inclusion interfaces. Depending on the size of the features of interest, APT, or SIMS, or a combination of the two could then be adopted for analysis and data acquisition.
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3.1.2. Development of non-destructive and in situ or operando methods

3.1.2.1. Scanning Kelvin probe force microscopy (SKPFM). SKPFM is a promising method to non-destructively detect H with high spatial resolution. It allows detection through a change in surface potential caused by H incorporation into the surface oxide and the matrix following hydrogen charging of the metal [42,256–263]. Direct quantification of H is made possible by applying a thin Pd layer, which effectively controls for the impact of oxide heterogeneity on hydrogen adsorption, thereby enabling the detection of limiting concentrations as low as 0.01 ppm and at nm resolution [40–46]. The measured surface potential via the Pd layer follows a logarithmic correlation with the charged H content [39,42]. It would be interesting to carry out feasibility studies of hydrogen adsorption quantification and hydriding phenomena in Ti alloys by coupling in situ SKPFM with an electrochemical charging process.

3.1.2.2. Elastic recoil detection analysis (ERDA). As a non-destructive and highly accurate quantification method, ERDA [264] can be used to measure both the lateral distribution at a resolution of tens of micrometers and the depth profile at a resolution of nanometers for light elements such as H and C. A beam of projectile ions that are usually heavier than the light elements to be probed collides elastically with the nuclei of the light element targets, ejecting them from the matrix material. ERDA has been successfully used to study H distribution in Ti64 by measuring the volume density [265] and depth profile [151,266]. Note that despite its usefulness in depth profiling, ERDA has relatively low lateral resolution and accumulative signals. As such, it should be classified as a surface-averaged measurement tool.

3.1.2.3. Thermal desorption spectroscopy (TDS). TDS is a reliable tool to collect information pertaining to H trapped in metals and alloys. The lattice H concentration, the reversibly incorporated H concentration and the irreversibly trapped H can all be estimated through desorption spectra. TDS also enables analysis of trapped energies at different types of matrix defects (e.g., vacancies, interstitial sites, dislocations, precipitates, and grain boundaries) [90]. Note that the accuracy of such analysis is typically around 0.1 eV for the binding energy. Uncertainties can be ascribed to surface defects, inaccuracies of temperature monitoring devices, and microstructural thermal instability. In interpreting TDS data in terms of H binding energies, one must remember that the temperature of H release implicitly contains information regarding both thermodynamic and kinetic factors, which can be difficult to disentangle if significant kinetic limitations are active.

3.1.2.4. Neutron diffraction and imaging. Hydrogen adsorption, transport and hydriding phenomena in the bulk materials can be studied non-destructively via neutron-based techniques such as neutron diffraction, scattering, vibrational spectroscopy, and imaging. For example, in situ neutron diffraction was utilized to study the decomposition of Ti hydride powder during constant heating in Ar environment [267]. In situ and operando neutron imaging was used to investigate the structural stability and the spatiotemporal H distribution in bulk (up to 40 mm in diameter) metal hydride composites (hydrided Hydralloy) [268]. In a similar vein, SCC phenomena associated with low H content (10–300 ppm) in Zr alloys were studied using neutron imaging with a mass resolution of ~5 wt ppm and a spatial resolution of ~25 μm [269].

3.1.2.5. X-ray techniques. XRD as a non-destructive and readily accessible method that is particularly useful for studying surface/sub-surface hydriding phenomena relating to hydride nucleation, lattice degradation, phase transformation, and cracking or fatigue. In addition to bulk phase identification, strain effects correlating to H content can be probed. For example, the evolution of the lattice parameter of δ-TiHx, which depends on the H concentration in pure Ti samples, was determined by XRD measurements in the Bragg-Brentano symmetric geometry using Co Kα radiation [19]. Liu et al. used the technique to estimate different hydride compositions, identifying a primary composition of TiH1.5 (5 hydride), with a small fraction of TiH1.7 and TiH2 in electrochemically charged pure Ti [17].

In situ or operando XRD has also been shown to be very powerful to collect the dynamic information of hydriding phenomena in Ti, Zr, and Pd [128,185,186,211,221,270–274]. In particular, 3D X-ray diffractive imaging (XRDI) is a new in situ/operando X-ray imaging technique developed to resolve 3D distributions of strain and phase transformations of solid materials exposed to reactive environments [36,275–277]. For example, Ulvestad et al. [118] used 3D XRDI to monitor the strain and phase transformation evolution during hydriding of Pd nanocubes and corroborate strain distributions with H concentration distributions. Hahn et al. [221] revealed the dynamics and hysteresis mechanism of H-Pd interactions during intercalation and deintercalation based on a multimodal in situ XRD and absorption spectroscopy (XAS) apparatus.

XAS has also been demonstrated as a probe of phase composition within Ti hydrides [278]. Although less direct than XRD for phase identification, XAS has the intrinsic advantages of sensitivity to both surface and bulk regions depending on the collection mode (electron and fluorescence yield, respectively), as well as the ability to probe hydrogen effects on electronic structure in non-crystalline local environments or prior to hydride phase formation. The combination of X-ray-based multimodal diffraction, spectroscopy, and microscopy could open a new characterization frontier to understand various metal hydriding phenomena.

3.1.2.6. Nuclear reaction analysis (NRA). Through the resonant reaction H(12N, α)12C [279] or non-resonant reaction D(3He,p)4He [232], nuclear reaction analysis (NRA) is capable of quantitatively depth profiling and non-destructively measuring H (or D) content distribution at solid surfaces, subsurfaces, and interfaces. The method utilizes a 12N or 3He ion beam to detect the H or D isotope in a depth range of up to 1–2 μm. The near-surface depth resolution is a few nanometers for surface-normal ion incidence and can be further improved to 1 nm by applying a grazing-angle beam. NRA can measure hydrogen with a surface concentration of ~10^13 cm^-2 (~1% of a typical atomic monolayer density) and volume concentration at a sensitivity of ~10^18 cm^-3 (~100 at. ppm). NRA has been successfully demonstrated in some hydriding studies including: quantitative measurement of surface and bulk H concentration in the near-surface region of a H-treated Pd single crystal [280]; determination of H depth and density near the interfaces of thin SiO2 films and Si(100) substrates [280]; and depth profiling of D, Ti, and Mo in metal hydride films (TiD2/Mo) [232].

3.2. Opportunities and perspectives in multiscale modeling

In close concert with experimental probes, multiscale, multiphysics models have a key role to play in providing foundational understanding of Ti hydriding. Here we focus on predictive models that range from atomistic to mesoscopic to macroscopic, which are emerging as a powerful tool for generating “synthetic data” that can be treated on equal footing with measurements for unbiased comparison. Table 6 lists several classes of methods that can provide much-needed insight, whereas Fig. 2 shows how these models can provide tie-ins with experiments (Table 5) to more comprehensively address the challenges listed in Table 1.

In general, the methods fall into two categories: atomistic/discrete methods and continuum methods. The first category includes first-principles and quantum chemical approaches (most notably, DFT), as well as DFT-based molecular dynamics, which provide high-accuracy energies, forces, charge states, and atomic positions, as well as thermodynamics for phase diagram prediction, but are limited to very small
or ideal systems for short times. Other atomistic methods include force-field based molecular dynamics (MD) for simulating local diffusion, segregation, and trapping, in addition to coarse-grained approaches such as kinetic Monte Carlo (KMC) for transcending much larger ranges of time scales. On the continuum side, more complex models that incorporate microstructural features and nonequilibrium conditions can be incorporated, but at the expense of physical accuracy. Among the continuum methods, phase-field modeling (PFM) provides a highly flexible framework for analyzing phase transformation, microstructure evolution, and chemomechanical coupling under different environments. This approach can be well integrated with atomistically derived quantities, as detailed in the sections below. Finite-element models (FEM) lack the flexibility, resolution, and microstructure awareness of the phase-field approach, but are well suited to examine solid mechanics and fracture effects. Finally, integrated kinetic models can be built around parameterized nucleation-growth and diffusion equations to simulate the full hydriding process; unfortunately, these models typically forego details of microstructure and defect interactions and are therefore less useful for addressing the gaps in Table 1. In addition, they are typically empirically fitted and therefore provide minimal predictive capability beyond the specific conditions and materials for which they were parameterized. Nevertheless, we include them here because they can retain some predictive power when coupled with finer-scale modeling approaches to obtain needed parameters. The following sections detail three key needs for progressing modeling methods to better address hydriding issues in Ti and other metals: multiphysics integration, multiscale integration, and incorporation of beyond-ideal factors.

### 3.2.1. Multiphysics integration

One of the major challenges in modeling metal hydride formation and associated microstructures is the involvement of concurrent chemical, physical, and materials processes, as well as their complicated (and often unknown) coupling. For instance, to comprehensively account for nucleation and growth of hydride phases in realistic polycrystalline Ti and its alloys, the model should incorporate surface reactions; surface, grain, and boundary diffusion, thermal transport, grain boundary H segregation, crystalllographic structural transformations, and mechanical interactions involving large volume changes. Of the available approaches, perhaps the most versatile in this regard is the phase-field method [281–283], which has provided a general framework for integrating these multiphysics phenomena for modeling phase transformations and associated microstructure evolution in a wide variety of structural applications [284], energy applications [285], and functional applications [286]. PFM is based on a diffuse-interface description and can track the evolution of a diffuse grain, phase, or particle boundary according to underlying governing physical equations. As such, it can easily be extended to include different physicochemical phenomena, provided the necessary parameters and functional relationships can be properly defined.

For modeling hydride formation in metals using PFM, α-Zr (hcp) has often been employed as a model system [287,288]. The fundamental
Recently, Heo et al. established a comprehensive phase-field model for simulating hydride formation with general metal/hydride interfacial coherency in a generic polycrystalline Zr system [50]. Because hydride formation in α-Ti exhibits very similar features and characteristics to Zr, this same framework can directly be applied to model Ti hydride formation. Preliminary simulations along these lines are reported in Section 3.3, wherein we have parameterized the model to simulate reaction-induced hydride formation in polycrystalline pure α-Ti using available materials databases (e.g., thermodynamic free energies [300], diffusivities [89,301], elastic modulus (in Table 4), etc.). A more detailed description of this work is forthcoming.

Despite the power of PFM and related continuum approaches to integrate multiphysics factors within a single framework, the underlying governing relationships must be known or readily derivable. Because defects, compositional heterogeneity, microstructure, and other factors discussed in Section 2 play an important role in determining local interaction with H, determining such relationships in practice can be extraordinarily difficult. Moreover, if nonequilibrium hydriding kinetics are to be properly incorporated, then the effects of the environment must likewise be considered either implicitly or explicitly. In our view, this is best accomplished by incorporating a combination of experimental inputs and atomistic modeling inputs to extract important parameters and functional relationships. Examples of needed quantities for accurate PFM include phase free energies, detailed microstructures, diffusivities, interfacial and surface energies, and elastic constants.

3.2.2. Scale integration

Another challenge closely related to the need for multiphysics model integration is the intrinsically multiscale nature of hydrogen-material interactions. In general, each of the methods listed in Table 6 can be integrated with other modeling methods to extend scales of space, time, or both. For example, KMC methods are well suited to extend atomistic simulation times to experimentally relevant scales by incorporating DFT- or MD-derived kinetic barriers, provided events can be discretized (e.g., diffusion or trapping) and mapped onto a regular lattice. However, it is far more challenging—yet ultimately more impactful—to integrate discrete atomistic approaches and continuum approaches. This combination can leverage the superior accuracy and capability of atomistic approaches to account for local features such as point defects, while also appealing to the microstructure awareness and rapid nonequilibrium simulation capabilities inherent to many of the continuum approaches.

Along these lines, H-dislocation interactions and hydriding phenomena were recently explored via multiscale approaches in which thermodynamic and analytical models were combined to achieve both atomic-level resolution and continuum-level computational efficiency [48,49,108]. These approaches allow for determination of the concentration of hydrogen at the atomic scale upon perturbation of thermodynamic variables, including temperature and hydrogen chemical potential. In general, the thermodynamic parameters are linked to, and controlled by, continuum-scale models (e.g., FEM or PFM).

Another example is the integrated thermodynamic and analytical model developed by Leyson et al. [49]. In this model, input parameters (e.g., H-H interaction energies) were derived from coupled embedded atom force-field and Monte Carlo simulations. The integrated model assessed the dependence of hydride size on environmental temperature and bulk hydrogen concentration and also successfully predicted both nanometer-scale hydride nucleation and the activation of hydrogen-enhanced local plasticity. A sharp transition from a non-hydride-forming domain to a hydride-forming domain was identified, for which the transition point was characterized by a critical hydrogen chemical potential for hydride nucleation. The predicted results were also consistent with nanoindentation tests carried out on H-charged materials.

Similarly, Zhao et al. [108] developed a continuum model, the so-called hydrogen-informed expanding cavity model, that was calibrated by the large-scale MD simulations. This model incorporates the collective interactions between hydrogen and dislocations and their impact on softening of HE metals and alloys. The simulation results show that the activation energy for dislocation nucleation decreases with the concentrated local H, leading to decrease in indentation force and hardness. Note that this behavior is consistent with the hydrogen-promoted dislocation migration and multiplication along with the matrix softening reported by many studies as discussed above.

The PFM examples discussed in the previous section provide additional opportunities to improve multiscale integration. DFT, MD, and KMC simulations can provide transport data, whereas DFT and force-field models can provide needed interfacial energy and solid mechanics inputs. Some specific examples of how this can be done are provided in Section 3; nevertheless, additional progress is needed to incorporate the needed diversity and complexity into the lower-length scale simulations. A particular priority involves modeling static and dynamic properties of disordered and heterogeneous systems including grain and phase boundaries, as well as amorphous surface oxides that offer unique configurational and chemical challenges at the interface with the Ti metal.

3.2.3. Beyond ideal models

To correctly capture physical, chemical, and materials behavior in simulations of hydriding phenomena, it is important to incorporate models capable of transcending idealized crystalline structures and purely thermodynamic considerations. Critical frontiers in computation include the development of models with more sophisticated and coupled physics, more realistic microstructures, more complex atomic configurations of interfaces and boundary regions, and better approximations of real operating conditions.

This need was recognized in a recent paper outlining how “beyond-ideal” factors could be incorporated at both the atomistic and continuum scales to improve hydriding models being developed by within the Hydrogen Materials—Advanced Research Consortium (HyMARC) [51]. In that study, four key beyond-ideal factors were identified and shown to influence performance predictions in hydriding reactions: (1) surface anharmonic dynamics; (2) interface and surface energy penalties; (3) mechanical stresses induced by constrained volume expansion; and (4) the presence of the native surface oxide. Although the focus did not include Ti, the identified beyond-ideal factors echo many of the themes introduced in Table 1 and Section 2. An additional priority is the development and application of methods capable of evaluating the
properties of Ti surface oxides, which are notoriously configurationally complex. Graph neural network-based approaches [302] are emerging as a promising approach to quantify and efficiently explore this complexity and are currently being applied by our team to investigate H in TiO$_2$ films.

Another challenge involves accounting for atomistic defect interactions within continuum models that are otherwise incapable of achieving the necessary fidelity. One example of such an approach that has achieved widespread use is the point defect model (PDM). The PDM has been applied to predict the growth or dissolution of oxides and hydrides in hydride-forming metals, typically by integrating experiments and numerical models [52,53]. For instance, Ai et al. [52,53] employed the PDM to analyze H-associated degradation mechanisms in pure Zr in hydrogenated pressurized water reactor primary coolants.

Fig. 3. Modeling and experimental integration strategies for studying hydrogen transport and trapping in surface Ti oxides, with specific examples from the authors’ work. Computational approaches on left side, top to bottom: Relative probability of H binding energy (in eV with respect to H$_2$) at diverse local sites in amorphous TiO$_2$ (α-TiO$_2$) with respect to oxygen coordination number (2-fold [2F], 3-fold [3F], 4-fold [4F]); KMC setup for molecular modeling of H hopping in α-TiO$_2$ based on a graph neural network description (inset shows Ti (blue), O (red), and Ti-O bonds (green) within the network); mesoscopic continuum modeling of hydrogen diffusion through a complex microstructure in polycrystalline TiO$_2$ (insets: input microstructure and corresponding prediction of local variation in H composition). Experimental approaches in right side, top to bottom: NMR measurement of O coordination number distribution in $^{17}$O-labeled α-TiO$_2$, APT analysis of as-deposited (top level) and polycrystalline (bottom level) Ti oxide film showing compositional variations (scale bar: 50 nm) indicative of interfaces and local heterogeneities; measured H permeation through a 400 nm thick as-deposited Ti oxide film on Ti metal using a Devanathan cell as a function of charging time in an aqueous environment (charging current: 200 nA/cm$^2$, discharge potential: 0.2 V SCE, electrolyte: deaerated 0.6 M NaCl).
They found that the barrier layer comprises hydrogen-deficient zirconium hydride (ZrH_{2-x}), while the outer layer comprises porous, stoichiometric ZrO_{2} that forms via hydrolysis of ZrH_{2-x} at the interface between the layers. This analysis indicates that corrosion resistance depends on the physical features of the porous oxide layer, including its porosity and thickness. Furthermore, hydride platelets were found to precipitate below the metal/barrier layer interface, with their number density decreasing with distance from that interface [53]. This behavior verifies that hydride platelets form due to atomic H that diffuses through the metal phase from the interface with the barrier layer. Importantly, this mechanistic understanding provides practical guidance on slowing hydriding by engineering the porosity and/or thickness of the outer layer for enhanced outer-layer resistance.

Considering the mechanistic similarity between Ti hydriding and Zr hydriding, the PDM could likewise be applied to predict key aspects of hydriding and hydrogen-induced corrosion of Ti and its alloys. For electrochemical charging, the PDM could be optimized to describe the partial anodic process and incorporate cathodic hydrogen evolution. The optimized model could then be used to compute the passive dissolution rate, allowing for reliable estimation of the time to perforation of the porous oxides layer. Once perforated, hydriding of the base Ti alloy would occur. The thickness of the barrier layer becomes larger as the potential is made more negative and can serve as a reliable measure of the extent of hydriding, which can in turn be predicted by the PDM based on in situ EIS measurements along with optimized model parameters [52,303]. Suggested details of how PDM could be applied to studying Ti hydride growth process are provided step-by-step in the supplementary information.

### 3.3. Examples of experiment-model integration

In this section, we provide representative examples of our current efforts to more closely integrate experiments and models to investigate some of the key thermodynamic and kinetic mechanisms associated with Ti hydriding. These examples, which are outlined alongside preliminary representative results in the subsequent sections, demonstrate the feasibility of the integrated approach for analyzing coupled chemical, physical, and materials processes that constitute hydrogenation of Ti. In particular, we focus on probes of three key governing phenomena: (1) H transport through the protective Ti surface oxide; (2) micromechanical response of surface oxide to hydrogenation of underlying Ti; and (3) the beginnings of hydride formation in polycrystalline Ti.

#### 3.3.1. H transport through Ti surface oxide

Fig. 3 illustrates how experiments and models can be integrated to probe the effects of local binding characteristics and microstructural features on hydrogen transport in stoichiometric and nonstoichiometric titanium surface oxides (TiO_{x}) at multiple length scales. As discussed in Section 2.1, this oxide layer forms at the Ti metal surface in service, and understanding its interaction with H is critical for elucidating the key processes in early-stage hydrogenation of Ti and its alloys. The TiO_{x} oxide layer incorporates a variety of chemical and structural complexities, including atomic structural disorder, non-stoichiometry, and structural defects such as grain boundaries.

To isolate physicochemical factors that determine the impacts of the structural complexities on hydrogen transport at the atomic level, quantum–mechanical computational approaches based on DFT can be employed. Such approaches are uniquely capable of interrogating the relationship between relevant local atomic structures of the oxide and the associated H binding characteristics. Ab initio molecular dynamics simulations can be used to generate both ordered and highly disordered atomic structures of TiO_{x}, which can then be analyzed to identify classes of unique environments for hydrogen binding, parameterized by structural factors such as the local oxygen coordination number (Fig. 3, top left). Particularly for atomically disordered regions, it is critical to validate that the models are correctly capturing the available local environments. For this purpose, we can use DFT to compute spectral “fingerprints”, which can be compared directly to measurements. One example of is NMR, which can probe the relative expression of local O coordination environments throughout the sample (Fig. 3, top right). In addition to validating the possible local environments in the oxide, such spectroscopic measurements can be used to “tune” the model towards a calibrated result that better expresses the correct fraction of local environments found in the material.

Having validated the unique local environments, we can proceed to compute hydrogen hopping events among them. Coarse-grained featureization using graph neural network-based approaches aid in rapidly analyzing the role of diffusion network topology within the complex oxide (Fig. 3, center left). This analysis informs kinetic Monte Carlo simulations, by which hydrogen hopping kinetics in TiO_{x} are directly extracted. The network models can be informed and validated by precise elemental distribution and composition analysis of TiO_{x} using APT, which provides a compositional and microstructural guide to inform factors such as stoichiometry (Fig. 3, center right).

Finally, the measured microstructural features are reproduced in digital representations using PFM, and the microstructure-level effective diffusivity of hydrogen through TiO_{x} is computed in the presence of different microstructural features (e.g., grain size, grain boundaries, crystallinity fraction, and compositional variation). This can be done using a microstructure-aware mass transport modeling approach, such as the one our team recently reported for complex oxides [304] (Fig. 3, bottom left). This mesoscopic modeling approach incorporates input diffusion parameters from atomistic simulations and digital representations of realistic oxide microstructures. Fig. 3 (bottom left) shows an example of computed effective diffusivities of hydrogen in polycrystalline TiO_{x} (see figure inset for the corresponding input microstructure) at different temperatures. The resulting H transport prediction can then be compared to independent measurements of H permeation. At mesoscopic (e.g., grains) and macroscopic (e.g., entire film) levels, hydrogen diffusion through Ti oxide thin film can be evaluated experimentally by gaseous permeation or electrochemical permeation. Fig. 3 (bottom right) includes an example of electrochemical permeation of hydrogen under controlled environment through a Ti oxide thin film. The onset of discharge on the anodic side of the cell indicates the permeation of hydrogen through the film and metal foil. Diffusion coefficients can be estimated from such measurements and compared with mesoscopic effective diffusivity calculations with identical microstructural features and chemical composition (We caution that microcracks in the oxide films can affect these measurements; however, such effects can be partially compensated by comparing charging and discharging currents.). Having validated the model, we can then proceed to systematically quantify the impacts of microstructural features such as grain size, grain morphology, and grain boundary types on hydrogen diffusion properties in TiO_{x}.

#### 3.3.2. Micromechanical response to H incorporation

Fig. 4 shows how the relationship between microstructure and mechanical response of the surface oxide to hydrogenation-induced strain can be assessed by combining experiments and simulations. As discussed in Section 2.3, hydrogenation of Ti metal involves significant volume expansion, which exerts mechanical strain within the oxide layer on the metal surface. The associated lattice distortion in the oxide can affect hydrogen transport kinetics or cause physical damage (e.g., cracking) to the surface oxide layer, potentially accelerating hydrogen permeation through the layer [305]. When analyzing the mechanical response of the oxide layer, additional potential complexity may arise from possible phase transformations or hydrogen reactions activated by the exerted strain [305].

The relevant phase and reaction behavior in TiO_{x}, as well as their sensitivity to microstructural features, can be probed experimentally via controlled experiments employing TiO_{2} thin films with varied microstructures. Fig. 4 (top left) shows examples of crystalline TiO_{2} surface...
films produced with grain sizes ranging from ~ 5 nm to ~ 200 nm by ion beam sputter deposition method and subsequent thermal treatment. Electron microscopy is used to determine grain structure, whereas XRD is used to identify phases. Fig. 4 (top right) demonstrates that the anatase phase is the only crystalline phase during thermal treatment at an intermediate temperature (700 °C), whereas both the anatase and rutile phases are present at 1100 °C. The impact of mechanical loading on the phase and hydrogen reaction behavior is gauged by monitoring phase evolution and strain of TiO_2 in situ under pressure loading in an H_2 environment in a diamond anvil cell (DAC) coupled with Raman...
spectroscopy. In this case, the strain is visible, although no evidence of hydriding is detected in the Raman spectroscopy measurement up to 30 GPa at ambient temperature.

A companion computational effort involves the design of a microstructure-aware micromechanics modeling approach [306] to analyze the mechanical response of polycrystalline TiO$_2$ to the tensile strain caused by the volume expansion associated with hydriding of the underlying Ti metal. First, a microstructural representation is produced (Fig. 4, bottom left) using PFM or direct tomographic reconstruction to match the experimental grain structure. Next, the stress distribution within this digital microstructure is computed based on local elastic moduli of the individual grains, as parameterized from DFT calculations. Fig. 4 (bottom right) shows how von Mises stress distributions qualitatively differ for two average grain sizes. Based on these computed distributions, we can identify mechanical “hotspots” featuring extreme local stresses that might initiate cracks or induce phase transformations.

By further combining with experimental mechanical testing and microstructural probes, the models can in principle be calibrated to quantify the relationship between oxide microstructural features and observed micromechanical and phase behavior.

Note that by combining models with experiments, we can directly probe local stress variation within the oxide film under hydriding conditions, as well as isolate the micromechanical effects within the film from those of the underlying metal substrate. By comparison, direct measurement of stress distributions within an isolated, standalone thin oxide film without a supportive substrate would be extremely challenging experimentally. As an alternative, TiO$_2$ thin films could instead be deposited onto a carefully selected substrate, or else well-controlled bulk TiO$_2$ could be used. The latter solution would also enable use of bulk experimental probes such as XRD, permitting monitoring of average lattice deformation and strain accumulation during hydrogen charging and transport for direct comparisons with model predictions as a function of microstructure.

### 3.3.3. Hydride phase formation

Fig. 5 shows how hydride phase formation at grain boundaries and related microstructural features within Ti metal can be probed at three different scales by combining simulations and experiments. Here, experimental and modeling approaches are synergistically combined to examine multiscale mechanisms incorporating coupled physicochemical processes and metal microstructural effects that determine the hydrgenation kinetics and the associated hydride phase microstructure evolution. Experimental and modeling efforts benefit one another by providing input parameters, measurement or simulation conditions, and cross-validation.

The modeling effort begins with atomistic DFT and MD simulations of grain boundary structures to compute hydrogen incorporation and segregation energetics (Fig. 5, upper left). The simulations are performed in concert with experimental characterization of α-Ti/hydride interfaces and grain structures using electron microscopy at lattice resolution (Fig. 5, lower left).

Next, we adapt a mesoscopic PFM approach to simulate hydride formation and corresponding microstructure using the model formulation by Heo et al. [50] (Fig. 5, top middle). The PFM simulations account for multiple multiphysics factors governing hydriding of polycrystalline Ti, including the thermodynamic driving force [300], hydrogen diffusion rates [301], crystallographic structural changes [283], microstructure evolution [282], hydrogen-grain boundary/surface interactions [307,308], and hydrogen reactions at the surface. The atomistic simulations provide key parameter inputs for these quantities. To analyze and quantify the realistic metal microstructural impacts on the hydride formation tendency and kinetics, it is further necessary to include experimentally relevant microstructural features and reliable energetics related to experimentally probable grain boundaries. As such, we also analyze the spatial distribution of hydrogen (H or D) and hydride in Ti microstructures based on imaging data (Fig. 5, bottom middle), which are then compared with simulated hydride phase microstructures. For instance, the orientation relationship between the growing hydride phase and the α-Ti metal phase can be captured using EBSD, providing information comparable to PFM-modeled α-Ti/hydride mixed microstructures.

Finally, a fully integrated model based on PFM allows us to simulate the overall H absorption kinetics within the metal microstructure under controlled conditions (Fig. 5, top right). The validity of this integrated model is verified by comparing the PFM results with experimental measurements performed at the macroscale (Fig. 5, bottom right). For instance, the experimental hydrogen uptake curves as a function of H-charging time and/or temperature under controlled hydrogen pressure can be collected via a pressure-composition-temperature (PCT) apparatus. The model-predicted and PCT-measured hydrogen uptake behavior represent macroscopic characteristics of hydriding resulting from collective effects of the involved multiphysics factors, allowing for direct comparison. The model can then be used to probe various “what if” scenarios in which materials or environmental parameters are altered. Various physical ingredients can also be turned off to explicitly isolate their effects, as in the example of elastic micromechanical interactions given in Fig. 5 (top right). In addition, by linking performance to H-metal interactions across all three scales (atomistic, microscopic, and macroscopic), we can in principle distinguish reversible (e.g., H in hydride phases) and irreversible (e.g., H in trapping sites) H contents, with implications for developing practical microstructural or compositional engineering guidelines for improving performance or degradation resistance.

### 3.4. Summary

Recent advances in multiscale, multiphysics simulation and modeling, as well as ongoing improvements in the resolution and sophistication of experimental probes, continue to provide new opportunities to address lingering unknowns in Ti hydriding. However, in our view, it is the integration of atomistic and continuum simulations with multimodal experimental characterization that offers the highest value. Indeed, such integration has already begun to elucidate contributing factors in several key underlying phenomena. The demonstrated examples for hydrogen transport in surface oxide, micromechanical responses of surface oxide, and hydride formation in polycrystalline Ti illustrate how improved experiment-theory integration might be achieved. Nevertheless, these examples represent only a few of the many emerging integrated approaches for examining atomistic and microstructural impacts that could be applied in the near future to understand thermodynamics and kinetics of Ti hydriding. Additional extensions should be prioritized to account for more complex hydrogen-related phenomena, including hydriding-induced cracking and hydrogen-dislocation interactions, which form key bases for analyzing mechanical failure mechanisms due to hydrogen exposure.

Opportunities are also available for improvements in the models to better facilitate direct comparisons with experiments. For instance, as first-principles simulations of complex surfaces become more routine, the feedback cycle between spectroscopic predictions and measurements can be expected to play a much larger role in elucidating the effects of surface chemistry and other materials features on hydriding. By using model systems that represent features of realistic systems under working conditions, first-principles methods can also be employed to illustrate the specific relationship between surface chemistry and transport at the interface, and to identify the most impactful materials or environmental factors. Likewise, mesoscopic “microstructure-aware” models can be used to integrate atomistically derived parameters with inhomogeneous microstructural features that match experimental realities, allowing far more representative investigations of hydrogen-materials interactions under more realistic conditions. Capturing such nuances is likely to prove increasingly important as our understanding of Ti hydriding improves.
On the experimental side, a variety of useful approaches have been demonstrated to measure composition, phase evolution, and mass transport at surfaces and interfaces during Ti hydriding. Such quantities already provide a compelling roadmap for direct integration with theoretical calculations. Nevertheless, the development of complementary experiments capable of validating theoretically predicted phenomena at all relevant length scales remains challenging: single experiments usually result in a qualitative or quantitative measurement that at best corresponds to a single value within a theoretical model. Such experiments offer little help to validate theory at multiple length scales simultaneously—key to extracting reliable connections between atomistic, microstructural, and macroscale behavior—and are therefore subject to sample-to-sample variation. One way to address this problem is to prioritize development in situ and in operando methodologies capable of taking multiple measurements during the hydrogen uptake process. Ideally, the measurement modalities would be capable of quantifying multiple physical changes that span length scales (e.g., resistivity, optical properties, mass, and dilatometry, followed by imaging and destructive techniques). Having access to these simultaneous probes would provide significant value for existing efforts to integrate experiments and theory towards a deeper understanding of materials changes during hydriding.

In conclusion, we emphasize that because combined theory–experiment approaches can uniquely and holistically investigate surface morphology, chemical composition, and hydrogen transport, future investments along these lines will be crucial. Such investments, properly realized, can guide forthcoming strategies for slowing hydriding of Ti alloys in corrosion-resistant structural applications. On the opposite side of this coin, we suggest that the obtained scientific understanding could likewise be applied to accelerate hydriding Ti-based materials for emerging functional applications, including hydrogen and thermal storage.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.

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