Recent progress in microstructural hydrogen mapping in steels: quantification, kinetic analysis, and multi-scale characterisation

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ABSTRACT

This paper gives an overview of recent progress in microstructure-specific hydrogen mapping techniques. The challenging nature of mapping hydrogen with high spatial resolution, i.e. at the scale of finest microstructural features, led to the development of various methodologies: thermal desorption spectroscopy, silver decoration, the hydrogen microprint technique, secondary ion mass spectroscopy, atom probe tomography, neutron radiography, and the scanning Kelvin probe. These techniques have different characteristics regarding spatial and temporal resolution associated with microstructure-sensitive hydrogen detection. Employing these techniques in a site-specific manner together with other microstructure probing methods enables multi-scale, quantitative, three-dimensional, high spatial, and kinetic resolution hydrogen mapping, depending on the specific multi-probe approaches used. Here, we present a brief overview of the specific characteristics of each method and the progress resulting from their combined application to the field of hydrogen embrittlement.

INTRODUCTION

Microstructure-specific hydrogen mapping has been recognised as the most important challenge on the pathway towards a better understanding of the nature of hydrogen embrittlement in metallic alloys, specifically in steels. Local presence of hydrogen is challenging to map due to its low mass, huge embrittling effect already at low concentrations, and high diffusivity in presence of compositional, thermal and mechanical gradients. For instance, the average hydrogen content in steels is of the order of $10^{-1}$ to 100 mass ppm, and some of the hydrogen is diffusible even at room temperature – an effect further enhanced in the presence of stress localisation effects. These complications render conventional chemical and spatially sensitive mapping methods such as energy dispersive X-ray spectroscopy or wavelength dispersive X-ray spectroscopy not readily applicable in the context of hydrogen embrittlement.

To enable both, spatially resolving and microstructure-sensitive hydrogen mapping, various approaches have been used, e.g. autoradiography [1–3], scanning photoelectrochemical microscopy [4–6], scanning electrochemical microscopy [7,8], secondary ion mass spectroscopy (SIMS) [9–12], silver reduction and decoration [13–17], the hydrogen microprint technique (HMT) [18–21], scanning Kelvin probe force microscopy (SKPFM) [22–26], atom probe tomography (APT) [27–30], prompt γ activation analysis [31], proton–proton (pp) scattering [32–34], and neutron radiography [35–39]. Additionally, a quantitative method for determining the hydrogen content at specific microstructural constituents with a characteristic trapping depth such as dislocations is available in terms of thermal desorption spectrometry (TDS) [40–42]. Although this method does not provide spatially resolving mapping, hydrogen desorption rate profiles detected by TDS are microstructure-sensitive.

Initial efforts were devoted to achieving spatially resolved hydrogen mapping to correlate hydrogen segregation with specific microstructural characteristics, defect sites and fracture surface features. Since hydrogen-assisted cracking has been demonstrated to occur along internal interfaces such as phase and grain boundaries, the detection of localised hydrogen at such microstructural sites provided an explanation for the occurrence of hydrogen embrittlement.
even at very low average hydrogen content [17,43].

With the recent advances in highly defect-specific microstructure probing techniques, the request for corresponding progress also in highly spatially resolving hydrogen mapping has gained momentum. Regarding the spatial resolution necessary for correlating local hydrogen accumulation with specific phenomena such as interface decohesion, cracking, void formation and localised plasticity effects, ‘multi-scale’ hydrogen probing concepts have become important [11]. This means that the transport and trapping of hydrogen has to be conducted over several microstructural scales in the same material. An atomic-scale example is the future challenge of mapping hydrogen trapping phenomena for instance at point or line defects e.g. occurring in the form of vacancy decoration or hydride formation at dislocations. Larger scale effects such as void and crack formation at hetero-interfaces, grain boundaries or triple interface junctions ranging over sub-μm to μm scales represent more collective decoration phenomena. At mesoscopic strain-localisation zones such as shear bands, hydrogen accumulation extends over a few tens to hundreds of μm. Regarding yet larger scales, hydrogen also segregates at the tips of stress concentration sources such as cracks and notches or surface reaction zones between the material surface and its environment which can even extend over sub-mm to mm regimes. Consequently, a corresponding demand for improved mechanism-sensitive, multi-probe and multi-scale hydrogen mapping protocols enabling micro- to mesoscopic hydrogen mapping has grown in recent years.

In addition to enhancing spatial resolution, quantification of local hydrogen content is another important goal of hydrogen mapping. A phenomenological relationship between the average hydrogen content and fracture strength has been well established [44–46]. Yet, the capability of elucidating the underlying damage mechanisms behind this relationship requires experimental means that allow us to relate the hydrogen to its effects on specific microstructure features [47,48]. A crucial reason for the pronounced microstructure dependence of hydrogen embrittlement is the difference in the degree of hydrogen trapping and localisation. For predicting hydrogen embrittlement susceptibility in a comprehensive fashion, it is indispensable to reveal the quantitative relationships between specific microstructure features and the associated local hydrogen content. However, only a few prior works succeeded so far in applying such spatially highly resolved hydrogen mapping routines for quantifying the local hydrogen content in a microstructure-sensitive fashion. Therefore, an increasing number of studies has been recently devoted to the quantification of the local hydrogen content at microstructural resolution [28,49,50].

Another equivalently essential recent challenge lies in microstructure-sensitive hydrogen mapping with kinetic resolution. Kinetic hydrogen tracking is essential since the associated embrittlement phenomena are all characterised by pronounced strain rate [46,51,52] and hydrogen exposure time dependences [53–55]. The latter dependence is known as hydrogen-induced delayed fracture [53,54,56]. It can be principally understood in terms of the permanent diffusion and redistribution of hydrogen inside materials with their wide range of potential trapping sites such as vacancies, dislocations, precipitates, grain boundaries, phase interfaces, triple points, micro- and nanocracks, voids and other defects that bind hydrogen and/or accelerate its transport, Figure 1. Yet, to understand the specific underlying mechanisms that lead not only to the trapping of hydrogen but actually to time-dependent fracture, kinetic analysis of local hydrogen is also required.

![Diagram showing various types of lattice defects and hydrogen trapping](image)

**Figure 1.** Hydrogen can get trapped and transported at various types of lattice defects such as vacancies and certain solutes, dislocations, grain boundaries, phase interfaces, triple points, precipitates, micro- and nanocracks, surface steps, voids and also inside the surface oxide layers. The main challenge in identifying the governing mechanisms behind hydrogen embrittlement lies in mapping hydrogen trapping and kinetics with microstructural sensitivity.
Furthermore, three-dimensional characterisation of hydrogen distribution is another important challenge, since the hydrogen segregation behaviour depends also on stress triaxiality, distance from inner and outer surfaces, and microstructure features that are hidden underneath a specific target location. In addition, effects associated with hydrogen diffusion to a probed surface from microstructures that are hidden underneath a specific target location may lead to misleading conclusions when only mapped in two-dimensions. These examples underscore the wide variety in methods, scales and goals associated with the mapping of hydrogen developed in recent years. Hence, in this paper, we review the recent progress made on microstructure-specific hydrogen detection methods enabling quantitative, three-dimensional, high spatial resolution, multi-scale, and kinetically resolved hydrogen mapping.

**Quantification of local hydrogen content**

The global hydrogen content, yet, energetically resolved as pertaining to specific types of lattice defects through their associated characteristic trapping energy, can be quantified through TDS. More specifically, TDS measurements conducted at different heating rates enable the determination of the activation energy of de-trapping from each type of lattice defect by analysing the resulting hydrogen desorption rate profiles \[40–42\]. Accordingly, the cumulative hydrogen content associated with each of the TDS peaks that is characterised by a specific activation energy of the underlying trapping mechanism gives the local hydrogen content at each of these lattice defect traps. This approach depends on the assumption that the hydrogen trapping energies and the associated hydrogen escape kinetics do not overlap \[60\]. Specifically, when the activation energy of hydrogen diffusion in the matrix is high, a corresponding trapping energy analysis cannot be properly applied. An important problem is for instance the analysis of hydrogen trapped at very weakly bonded trap sites in BCC steels which are characterised by an intrinsically high lattice diffusivity of hydrogen. The weakly trapped hydrogen can be desorbed even under cryogenic temperature conditions as schematically shown in Figure 2(a).

![Figure 2](image)

**Figure 2.** (a) Schematic of a hydrogen desorption profile together with indication of the corresponding trap sites. (b) Example of an experimentally obtained cryogenic TDS profile in two different Fe-C alloys with different carbon contents \[61\]. Reproduced with permission from Mater. Sci. Tech. (MS&T) 2011, 1279 (2011). Copyright 2011, Wiley-TMS. The region emphasised in terms of the red box shows an example where a characteristic hydrogen trapping peak observe in an alloy with 6 mass ppm C has been shifted to much higher temperatures in an alloy with 50 mass ppm C. By initiating the TDS measurement at cryogenic temperatures, the full configuration of the TDS profile is obtained without losing the otherwise non-measurable, weakly trapped and diffusible hydrogen in the analysis. Accordingly, accurate and trap-specific activation energies and the corresponding trapped hydrogen content are acquired by such C-TDS experiments. In addition, C-TDS measurements are capable of uncovering new types of hydrogen desorption behaviour such as the recently discovered carbon effect on hydrogen desorption discussed below \[61\]. An example of the C-TDS profile is shown in Figure 2(b). This result indicates that an earlier undiscovered hydrogen desorption peak exists in the cryogenic temperature region as outlined by the red broken lines. Moreover, carbon addition of only 0.005 wt-\% shifted the position of the main peak from ambient temperature to 0°C. These features could not have been revealed when subjecting hydrogen-decorated samples to conventional TDS analysis.

A next step in the microstructure-sensitive probing of hydrogen lies in using SIMS maps. A schematic of
the SIMS technique is illustrated in Figure 3. Spatially resolved SIMS that can detect the ion intensity of hydrogen or respectively deuterium has been studied as a quantitative mapping methodology for hydrogen that is trapped inside a specimen, as opposed to hydrogen that diffuses out towards the surface. Important aspects that may affect and even reduce the sensitivity of hydrogen detection by SIMS are as follows, in addition to other more minor experimental factors that were discussed in previous papers [10,63].

(i) Hydrogen emission from moisture and adsorbed hydrocarbons deposited on the sample surface which alter the results.
(ii) Background hydrogen in the atmosphere of the SIMS chamber since hydrogen exists even under excellent ultra high vacuum (UHV) conditions.
(iii) Diffusion and desorption of hydrogen during the measurement in steels.
(iv) Effects associated with sputter direction, crystallographic texture and surface topology (also referred to as matrix effects).

Further, yet minor factors affecting SIMS results are specimen edge effects and an influence of friction heat during mechanical polishing [10]. Several measures have been suggested to reduce errors arising from these effects: The hydrogen emission from sources on the sample surface can be reduced by sputtering silicon onto an area surrounding the observation region [10]. The use of high-purity sputtered silicon inhibits hydrogen emission. Reducing the background hydrogen existing in practically every SIMS chamber can be realised by a cold trap method using liquid nitrogen [10]. Effects associated with surface roughness, sputter direction and texture can be reduced by carefully mapping these features and probing their specific influence on the results. Suited counter-measures include excellent surface preparation and representative selection of crystallographic orientations. As a result, a quantitative relationship between hydrogen content and hydrogen ion intensity was obtained successfully in a type 304 austenitic steel as shown in Figure 4(a). Moreover, SIMS enables spatially resolved hydrogen mapping along the depth direction from the specimen surface as shown in Figure 4(b) owing to its through thickness serial sectioning enabled by continuing sputtering. Such a depth profile was indeed obtained on a type 304 austenitic steel with Al and Fe–Al surface coating layers that prevented hydrogen uptake into the base material region of interest. Here, it was clearly demonstrated that hydrogen segregates to the interface of the Al/Fe–Al layers. In addition, hydrogen in a ferritic steel could be detected by SIMS measurements at 83 K [64]. This successful example of a cryogenic SIMS measurement will be most likely further exploited and enhanced in the future to enable quantification of the hydrogen content in materials where it has high diffusivity such as in ferritic steels.

Recently, the scanning Kelvin probe (SKP) approach has received attention as an alternative approach to determine local hydrogen content quantitatively owing to its potential to resolve hydrogen both with high spatial microstructural resolution on the micro- and even submicroscale as well as with temporal resolution, i.e. it also reveals the permeation and release kinetics.
Figure 4. An example of SIMS analysis: hydrogen mapping from the Al-based layer-coated surface of a type 304 austenitic stainless steel. (a) Quantitative relationship between ion intensity and hydrogen content. TDA: thermal desorption analysis [65]. cps: count per second. $I_{\text{HN}}$: net intensity of hydrogen. (b) Depth profile of ion intensity [49]. Hydrogen segregates at the Al/Al-Fe interface. ‘Reproduced with permission from Int. J. Hydrogen Energy, 38, 10152 (2013) and 40, 10336 (2015). Copyright 2013 & 2015, Elsevier’.

Figure 5. A schematic illustration of a setup for scanning Kelvin probe (SKP) mapping [22]. KP: Kelvin probe. $U_{\text{ext}}$: applied external bias. $I_{\text{ac}}$: ac-current resulting from harmonic modulation of the capacitance. $\Delta\psi_{\text{KP sample}}$: difference between Volta potentials of the KP and the sample. ‘Reproduced with permission from Electrochem. Acta, 53, 290 (2007). Copyright 2007, Elsevier’.

at high spatial resolution [24,50,66]. Figure 5 shows a schematic illustration of a SKP analysis. Under certain conditions, such as when a palladium detection layer is used [24,66] (see also Section ‘Multi-scale and high spatial resolution hydrogen mapping’), the probed surface potential of a sample can be calibrated to provide direct correlation with the local hydrogen concentration. In such as case SKP mapping is capable of revealing the hydrogen distribution on sample surfaces. More specifically, the surface potential decreases with increasing hydrogen content [66]. Also the potential mapping directly on the passive film of steels or aluminium provides information about hydrogen distribution inside the material. For instance, the hydrogen distribution in the vicinity of a pit formed in an MgCl$_2$ droplet in a UNS S46500 precipitation-hardened stainless steel could be resolved with high spatial resolution as shown in Figure 6. Since the potential around the pit is locally low as shown in Figure 6(b), this result indicates that the MgCl$_2$ droplet enabled formation of hydrogen during the pitting.

Multiscale and high spatial resolution hydrogen mapping

A wide range of spatial resolution regimes has become accessible through the use of various types of hydrogen mapping techniques as shown in Table 1. To realise even nm-scale microstructural hydrogen mapping, atomic force microscopy-based SKPFM has been applied instead of SKP [11,23,25,67]. Since SKPFM is not equipped with a reference electrode and switching to a suitable reference is difficult to achieve during scanning a given area of the investigated sample, the values obtained are not calibrated. This prevents exact hydrogen quantification, as it is possible when conducting SKP on Pd coated samples (see [24,66]). However, the potential contrast within the scanned area and the relative changes in that contrast can provide valuable information about the locally evading hydrogen. In a recent study involving SKPFM, a pronounced improvement in hydrogen detection was accomplished by coating the specimen surface with a thin 100-nm palladium layer serving as a buffer and storage material as schematically shown in Figure 7(a–c). The two main aspects that are currently under investigation concerning the use of the atomic force microscopy-based SKPFM in conjunction with the use of a palladium coating are as follows:
Figure 6. Diffusible hydrogen distribution at a pit visualised by SKP [50]. (a) The schematic shows a droplet pre-exposure on a surface with the location of the cross-section cut depicted in (b). (b) SKP potential map obtained on a UNS S46500 precipitation-hardened stainless steel. The SKP measurement was carried out on a cross-sectional area of the specimen containing the pit as shown in (a).


(i) The chemical potential of hydrogen in palladium is much lower than that in steel. Therefore, the palladium stores the hydrogen that diffuses out of the steel microstructure and serves, hence, as a detection or respectively buffer layer. This effect still allows visualisation of the location of the hydrogen accumulation even after larger time delays, as lateral diffusion of hydrogen inside the Pd layer is inhibited as consequence of its nano-crystalline fcc structure [24].

(ii) The microstructure-dependent formation of locally different oxide films on the surface of the sample under inspection complicates establishing the relationship between the surface potential detected by atomic force microscopy SKPFM and the resulting hydrogen concentration values. This was up to now a critical problem for corresponding Kelvin probe hydrogen probing techniques but the use of palladium coatings has resolved this problem as its buffer effect collects the locally evading hydrogen independent of the nature of the underlying oxide, reliably accumulating it over a certain time constant [24].

As a result, the use of SKPFM enables measurement of hydrogen down to a scale below 0.01 atomic ppm at a spatial resolution of several tens of nanometres [25]. Figure 8(a) shows an example of a SKPFM measurement in an austenite/$\alpha’$-martensite dual-phase microstructure. Here, low-potential regions appear in the $\alpha’$-martensite. Moreover, additional low-potential regions are observed at the martensite–martensite grain boundaries as indicated by arrows. Hence, hydrogen localises on the martensite–martensite grain boundaries. Correspondingly, hydrogen-assisted damage was observed at the martensite–martensite grain boundaries [17].

The silver reduction and decoration technique schematically shown in Figure 9 is helpful as a more qualitative hydrogen detection method to supplement the quantitative SKPFM results for the following reasons: Silver decoration is a technique that utilises a hydrogen-driven reduction reaction of silver ions from a potassium silver cyanide aqueous solution on the specimen surface, which easily enables multi-scale observation of hydrogen distribution from microstructure-interfacial segregation to sub-mm-scale hydrogen partitioning. This approach is particularly helpful when aiming at first identifying preferred hydrogen storage positions in the microstructure which then can be subjected to higher locally resolving or more quantitative probing methods. Figure 8(b) shows an optical micrograph of the same steel microstructure as in Figure 8(a). In the optical micrograph, silver particles appear as black (in this case, diffusible hydrogen localises inside the $\alpha’$-martensite), visualising hydrogen partitioning. Further magnification of the image using scanning electron microscopy indicates hydrogen segregation on martensite/martensite grain boundaries as indicated by the yellow arrows in Figure 8(c). In such cases the silver decoration technique can be coupled with SKPFM, since both methods allow to detect hydrogen that evades from the surface with high microstructural sensitivity. Regarding kinetic resolution, it should be noted that the silver decoration technique is capable of detecting diffusible hydrogen also at high effusion rates already at early stages immediately after hydrogen charging. In contrast, the SKPFM is preferentially suited for studying lower effusion rates at later stages, as it takes some time before the first measurement can be started. Therefore, a combined use of silver decoration and SKPFM enables ‘multi-time and multi-scale’ hydrogen mapping with complementary information regarding effusion kinetics.

The maximum attainable spatial and chemical resolution for resolving the position of hydrogen can be achieved by APT (Figure 10). This method combines
Table 1. Summary of specifications of currently used hydrogen mapping methods.

<table>
<thead>
<tr>
<th>Hydrogen Mapping Method</th>
<th>Spatial Resolution</th>
<th>Time Resolution</th>
<th>Viewfield</th>
<th>Main Advantage</th>
<th>Main Challenge</th>
</tr>
</thead>
<tbody>
<tr>
<td>TDS</td>
<td>None (the spatial resolution is the specimen size)</td>
<td>None (the measurement time required for probing a specimen)</td>
<td>Entire specimen</td>
<td>Diffusible and non-diffusible hydrogen contents are determined separately and quantitatively. Furthermore, microstructure-specific local hydrogen content can be determined through identification of the trap-energy of a specific site.</td>
<td>A part of hydrogen starts to be desorbed at a cryogenic temperature, thus activation energy at weak trap sites cannot be identified by conventional TDS that starts from room temperature. Cryogenic TDS (C-TDS) resolved trap energies even at weak trap sites.</td>
</tr>
<tr>
<td>Ag reduction and decoration</td>
<td>Sub-micrometer when used in conjunction with a SEM</td>
<td>Seconds with an optical microscope</td>
<td>mm² scale</td>
<td>Adaptive with various microscopy techniques, enabling multi-scale mapping</td>
<td>This classical method has already been established well. As an extra advancement, microstructure-specific kinetic analysis was realized with this method during in situ hydrogen permeation.</td>
</tr>
<tr>
<td>HMT</td>
<td>Sub-micrometer when used in conjunction with a SEM</td>
<td>None</td>
<td>mm² scale</td>
<td>Adaptive with various microscopy techniques, enabling multi-scale mapping</td>
<td>This classical method has already been established well. As an extra advancement, this method was utilized to visualize effects of stress and deformation on segregation.</td>
</tr>
<tr>
<td>SKP</td>
<td>The best spatial resolution is a few tens of micrometers, but usually it is about 100 μm</td>
<td>Seconds, but first measurement about 1 h after H-charging</td>
<td>up to several cm²</td>
<td>Quantitative and kinetically resolved hydrogen mapping with a micrometer-scale resolution</td>
<td>Kelvin probe potential measured on the oxide covered surface is difficult to quantify. Through use of a palladium film and by controlling the atmospheric conditions such as humidity, highly accurate hydrogen quantification is possible and very high sensitivity can be achieved.</td>
</tr>
<tr>
<td>SKPFM</td>
<td>Several tens of nanometres</td>
<td>seconds, but first measurement about 1 h after H-charging</td>
<td>Several thousands of μm²</td>
<td>Microstructure-specific hydrogen mapping together with kinetic analysis</td>
<td>Since in SKPFM calibration of the measured potential values is challenging, full quantification of hydrogen is difficult to achieve. However, the interpretation of potential contrast and its evolution contains valuable information. Also here the use of a palladium enhances the sensitivity to hydrogen and enables reliable mapping of hydrogen activity.</td>
</tr>
<tr>
<td>SIMS</td>
<td>A few micrometers in an x–y plane spatial resolution. For depth (z) direction, the resolution is a few tens of nanometres in semiconductors</td>
<td>None</td>
<td>500 μm × 500 μm in an x–y plane view field</td>
<td>Hydrogen trapped within a specimen can be detected, which enables three-dimensional hydrogen mapping</td>
<td>Detection of hydrogen in a ferritic steel was difficult, because of the high diffusivity and low content of hydrogen. This was achieved by cryogenic SIMS in combination with silicon sputtering and a cold trap.</td>
</tr>
<tr>
<td>APT</td>
<td>Best case: lattice parameter resolution near crystallographic poles in pole direction; Worst case: several nm in plane resolution near crystallographic poles</td>
<td>None</td>
<td>Up to one billion atoms</td>
<td>Highest spatial resolution among all hydrogen-mapping techniques; can resolve nano-hydrides and positions of single hydrogen atoms e.g. at specific traps; can be operated in correlative mode to reveal both, structure and composition at the same location</td>
<td>All UHV system contain hydrogen, hence experiments are better done with D in order to differentiate between environmental H in the chamber and deuterium coming from the material; H atoms are mobile when not in traps; this can lead to undesired H-H recombination or H mobility.</td>
</tr>
<tr>
<td>Neutron radiography</td>
<td>Approximately 20–30 μm</td>
<td>~ 28 s and ~ 1 h for two- and three-dimensional analyses, respectively</td>
<td>Entire specimen (e.g. 45 mm height × 10 width × 5 mm thickness)</td>
<td>Non-destructive three-dimensional hydrogen mapping. The two dimensional analysis enables kinetic characterization of the hydrogen distribution.</td>
<td>Further improvements in spatial and temporal resolution have been attempted. Future developments of better detectors and neutron sources with higher intensity will lead to better results.</td>
</tr>
</tbody>
</table>
well-controlled field evaporation and ionisation of surface atoms layer-wise from a tip-shaped specimen in combination with time of flight mass-to-charge spectroscopy and a position-sensitive detector. Field evaporation is triggered by exposing the positively biased tip-shaped specimen to a combination of a permanent high electrical field and either additional ultra-fast voltage pulses for local field enhancement or additional ultra-fast laser pulses for local temperature increase. The specimen apex has typically a radius of curvature of 60–100 nm. The time of flight serves for chemical analysis though mass-to-charge spectroscopy and the impact regions on the position-sensitive multi-channel detector serve for the determination of the original position of the field-evaporated ions within the sample. A three-dimensional elemental map can thus be reconstructed from the collected data, applying an inverse projection of the detector coordinates to the tip surface. Depth-scaling is done taking into account the sequence and volumes of the detected ions. Typical dimensions of the analysed volume can comprise up to one billion atoms on current state of the art instruments [68–70].

As an example, Figure 11(a) shows deuterium segregation at a vanadium carbide (VC) precipitate observed by APT. Here, to distinguish hydrogen in the specimen from background hydrogen, deuterium was introduced instead of hydrogen. The deuterium distribution at VC precipitates is stable during the measurement at 50–70 K, since VC precipitates act as hydrogen trap sites [72]. The atomistic visualisation is helpful as a direct evidence of hydrogen traps at a specific microstructural feature. Since hydrogen is present even in ultrahigh vacuum environments deuterium charging is typically more suited for revealing specific microstructure features and decoration sites, Figure 11(b).

Figure 7. The specimen is (a) first mechanically polished and (b) subsequently hydrogen-charged. (c) The scanning Kelvin probe force microscopy (SKPFM) is carried out after the palladium coating [25]. ‘Reproduced with permission from J. Electrochem. Soc., 160, C643 (2015). Copyright 2015, The Electrochemical Society’.

Figure 8. Multi-scale, multi-kinetics and multi-probe visualisation of interfacial hydrogen segregation through an optimised silver reduction and decoration method coupled with SKPFM in a Fe–32Ni–0.2C austenite/α′-martensite dual-phase alloy (wt-%) covered by a palladium protective surface layer [17]. (a) Scanning Kelvin probe force micrograph taken after an exposure time of 5 h in nitrogen gas atmosphere. (b) Optical and (c) scanning electron micrographs. The arrows in Figs. 8a and 8c, respectively, indicate the matching observation of interfacial segregation of hydrogen. ‘Reproduced with permission from Mater. Corros., DOI: 10.1002/maco.201609104, Copyright 2016 WILEY-VCH Verlag GmbH & Co. KGaA’.

Figure 9. Mechanically polished specimens for the silver reduction and decoration method are immersed into a chemical solution containing silver ions [11]. In a redox reaction the silver ions are then locally replaced by evading hydrogen atoms so that Ag is deposited, thereby marking the regions of enhanced hydrogen concentration. ‘Reproduced with permission from Acta Mater., 109, 69 (2016). Copyright 2016, Elsevier’.

\[ \text{Ag}^+ + \text{H}_2 \rightarrow \text{Ag}^0 + \text{H}^+ \]
Figure 10. APT enables three-dimensional elemental mapping with near atomic resolution [71]. By applying either ultra-fast high voltage or laser pulses to a positively biased tip-shaped specimen, surface atoms are layer-wise field-evaporated. They get ionised and accelerated towards a position-sensitive detector that records their time of flight (for chemical analysis) and impact positions (for determination of the original position within the sample). A three-dimensional elemental map can be reconstructed from the collected data, applying an inverse projection of the detector coordinates to the tip surface. Depth-scaling is done taking into account the sequence and volumes of detected atoms [68–70]. ‘Reproduced with permission from Ultramicroscopy, 111, 1619–1624 (2011). Copyright 2011, Elsevier’.

Figure 11. (a) APT revealing the deuterium distribution at a VC precipitate [28]. (b) Comparison of deuterium and hydrogen distributions in Ag, revealing that deuterium is more indicative when aiming to reveal microstructure-related effects [29]. Hydrogen is ubiquitous in any ultrahigh vacuum system, hence, APT experiments are better conducted using deuterium charging. ‘Reproduced with permission from Scripta Mater., 67, 215 (2012). Copyright 2012, Elsevier and Int. J. Hydrogen Energy, 39, 12225 (2014). Copyright 2014, Elsevier’.

Three-dimensional hydrogen mapping

From a viewpoint of tomographic hydrogen mapping at microstructure scales, APT provides the most precise access to imaging hydrogen in three dimensions at highest, namely near-atomic spatial resolution as mentioned above. In addition, conducting SIMS mappings in conjunction with through thickness serial sectioning enabled by continuing material removal via sputtering is expected to also map hydrogen in three dimensions at a wider field of view than APT.
These techniques are both powerful and promising as destructive methods for the three-dimensional microstructure-specific hydrogen mapping.

As non-destructive approaches to the microstructure-specific three-dimensional mapping of hydrogen, the proton–proton-scattering method in conjunction with split field magnetic detectors and the related neutron radiography have been developed. In particular, the neutron radiography has been applied successfully to the detection of hydrogen in iron [38]. Figure 12 shows a schematic image of the setup used for the tomographic analysis of hydrogen via neutron radiography. Neutrons interact with protons, i.e. with the cores of hydrogen. Since the degree of the interaction between the incoming neutrons and the protons is higher than that between neutron and isotopes of iron, a clear contrast between hydrogen-enriched and hydrogen-depleted regions appears on the neutron intensity map. Through a scintillator, the neutron intensity map can be visualised as an optical image. The images are taken along different orientations for obtaining a corresponding reconstructed tomographic characterisation. This is achieved by rotating the sample stepwise through an angular range of 360°. This method enables a full scale three-dimensional mapping of the hydrogen distribution in a specimen including the tomographic imaging of hydrogen-related cracks and voids. An example for the case of hydrogen in ARMCO iron is demonstrated in Figure 13. Figure 13(a) exhibits the specimen surface with blisters after electrochemical hydrogen-charging. The corresponding cracks are observed in Figure 13(b). The three-dimensionally mapping reveals that hydrogen segregates at blisters on the specimen surface.

**Kinetically resolved hydrogen mapping**

The temporal resolution column given in Table 1 provides a comparison among the different hydrogen mapping methods in terms of their respective capability to reveal hydrogen release kinetics. Table 1 shows that a substantial advantage of the SKP and SKPFM approaches lies in their accessibility of kinetic release data. Figure 14 shows the surface potential change as a function of time in an austenitic twinning-induced plasticity steel [25]. The specimen surface was coated with palladium. The low-potential regions appear in the form of a platelike morphology, here visible as dark regions. The platelike low-potential regions correspond to the locations of deformation twins. In particular, a clear contrast change was observed in the outlined region that contains fairly thick deformation twins. The contrast in the outlined region is not distinct at first (Figure 14(a)); but upon continuing probing the changing contrast reveals increasingly clearly the preferred hydrogen release from the deformation twins (Figure 14(b)). The released hydrogen is reshuffling from the specimen interior into the palladium layer where it is accumulated and detected upon release.

![Figure 12](image12.png)

*Figure 12.* A schematic showing the setup used for tomographic analysis by neutron radiography [38]. ‘Reproduced with permission from Acta Mater., 78, 14 (2014). Copyright 2014, Elsevier’.

![Figure 13](image13.png)

*Figure 13.* A three-dimensional hydrogen map of a hydrogen-charged ARMCO iron sample taken by neutron radiography [38]. (a) Surface region with blisters. (b) Crack distribution inside the sample. (c) Hydrogen distribution. ‘Reproduced with permission from Acta Mater., 78, 14 (2014). Copyright 2014, Elsevier’.
Figure 14. Time-dependent variation of the hydrogen distribution analysed by SKPFM in a Fe–18Mn–1.2C twinning-induced-plasticity steel (wt-%) covered with a palladium buffer layer [25]. Surface potential images taken at exposure times of (a) 6, (b) 67, (c) 300 h, respectively. (d) Line profiles of the detected surface potential corresponding to the white lines in (a–c). The black arrows indicate hydrogen segregation at deformation twins. ‘Reproduced with permission from J. Electrochem. Soc., 160, C643 (2015). Copyright 2015, The Electrochemical Society’.

Figure 15. Results of the in situ optical microscopy observations of Ag deposition during the hydrogen permeation testing at (a) 0, (b) 55, (c) 90 min in an annealed pure iron specimen [73]. ‘Reproduced with permission from Scripta Mater., 129, 49 (2017). Copyright 2017, Elsevier’.

After prolonged probing time (300 h), the contrast disappears again because of the gradually completed hydrogen desorption (Figure 14(c)), i.e. the source that causes the potential change that is being exploited as signal indicative of the presence of hydrogen becomes exhausted. The sequential change in the local hydrogen release signal with time is shown in the form of variations of the corresponding line profiles in Figure 14(d). This result indicates that deformation twins play an important role in hydrogen segregation, and the hydrogen atoms at deformation twins are diffusible at ambient temperature. This kinetic approach offered by the SKPFM has also been successfully applied to revealing also other hydrogen trap sites such as at inclusions [24] and second phases [23,26] in steels.

Kinetic analysis of the microstructural hydrogen distribution has been also attempted through different other approaches in recent years. For instance, the exposure-time dependence of the mapped hydrogen distribution was analysed through an in situ silver decoration technique [73]. Figure 15 shows the evolution of the silver decoration with time during hydrogen permeation in an annealed pure iron specimen. In this experiment, the top surface of the specimen was immersed in a potassium silver cyanide aqueous solution. Hydrogen was introduced by electrochemical charging on the bottom surface of the specimen at ambient temperature. A hydrogen-driven redox reaction occurs when hydrogen permeates through the specimen. An optical lens with a long focal length was used for the optical microscope to enable in situ observations of the specimen surface without contacting the potassium silver cyanide aqueous solution. Silver deposition occurred preferentially at grain boundaries, and the occupation fraction of silver particles on the grain boundaries increased with time. A time-dependent variation of the evading hydrogen flux along the grain boundaries was revealed through this methodology. When coupled with electron backscatter diffraction (EBSD) analysis, this experiment also revealed that the hydrogen flux at low-angle grain boundaries is lower than that at high-angle grain boundaries.

In addition to these effects, the influence of residual elastic stresses has recently been suggested to explain the strain rate dependence associated with the steel’s hydrogen embrittlement susceptibility. Characterisation of the effects of stresses on hydrogen mapping has been carried out through HMT probing [74,75]. The basic procedure of the HMT is demonstrated in Figure 16. In a recent work, a slow strain rate test
was carried out in a specimen in conjunction with an HMT emulsion containing AgBr. Figure 17 shows a set of results obtained from a HMT experiment conducted on a martensitic steel. A HMT result without any mechanical loading is shown in Figure 17(a) for reference. This image demonstrates that no specific hydrogen segregation occurs when the specimen is not mechanically stressed. In contrast, as shown in Figure 17(b), the HMT result obtained after the slow strain rate test that was conducted until fracture at a strain rate of \(8.3 \times 10^{-6} \text{s}^{-1}\) demonstrates that hydrogen segregates along grain boundaries. More specifically, when coupled with an EBSD analysis as shown in Figure 17(c), preferential hydrogen segregation occurs along the prior austenite grain boundaries, but not on packet boundaries or any other of the characteristic martensite boundaries such as lath interfaces (please see references for the microstructure features of lath martensite [76,77]). In fact, fracture at the here imposed low strain rate occurred already when the sample was loaded only into the elastic regime, and such hydrogen segregation was not observed when the specimen was fractured at a high strain rate (\(8.3 \times 10^{-1} \text{s}^{-1}\)). Accordingly, the intergranular fracture was observed when deforming at a low strain rate, and the brittle features disappeared when the sample was deformed at a high strain rate. These observations indicate the essential role of mapping both, stress and time in concert with a microstructure-sensitive and spatially resolving signal associated with hydrogen segregation as well as with the resulting microstructural hydrogen embrittlement features.

Detection of dynamic hydrogen desorption associated with martensitic transformation

In the previous sections, we introduced recent hydrogen mapping experiments demonstrating hydrogen localisation behaviour under static or quasi-static conditions. Another crucial factor affecting the hydrogen distribution is the martensitic transformation itself. Since hydrogen solubility and diffusivity both strongly depend on crystal structure, the dynamic change in crystal structure occurring upon martensitic transformations must necessarily affect the hydrogen redistribution in metastable steels. With these considerations in mind, a C-TDS process accompanied by cooling was utilised to detect the dynamic hydrogen desorption during martensitic transformation [78]. Figure 18 shows the hydrogen desorption profile observed in a hydrogen-charged Fe–31 wt-%Ni metastable austenitic

The alloy during cooling and subsequent heating. Initially, the alloy was in the fully austenitic phase state. Then, the austenite started to transform into $\alpha'$-martensite when reaching the transformation start temperature $M_s$ upon cooling. Accordingly, hydrogen desorption occurred at the $M_s$ temperature as shown in Figure 18. This observation indicates that hydrogen desorption occurs when $\alpha'$-martensite is induced even at cryogenic temperatures. Factors causing the hydrogen desorption associated with martensitic transformations are considered to be (1) hydrogen-decorated motion of misfit dislocations that are associated with the transformation and (2) a change in local diffusivity because of the crystallographic transition. Since the heating process without the transformation also displays similar hydrogen desorption behaviour, the major reason for the transformation-induced hydrogen desorption was attributed to a change in local diffusivity caused by the transition from a close-packed structure (FCC) to a non-close-packed structure (BCC). The same experiment was also performed for an $\epsilon$-martensitic transformation which has practically the same packing density as the FCC phase. In the case of $\epsilon$-martensitic transformation, the peak intensity of the hydrogen desorption rate was relatively small compared to the case of the $\alpha'$-martensitic transformation, but it was still readily observed [78]. This technique will be helpful as an important tool to enable a more quantitative understanding of hydrogen embrittlement susceptibility of transformation-induced-plasticity (TRIP) steels.

Summary and outlook

Here we reviewed recently developed and matured methods for the microstructure-sensitive mapping of hydrogen. More specifically, we discussed TDS, silver reduction and decoration, the HMT, SIMS, APT, neutron radiography, and the scanning Kelvin probe methods (SKP and SKPFM). The specifications of the respective methods were summarised in Figure 19 and Table 1.

We have shown results that demonstrate how suited combinations of these methods can help to satisfy recent demands regarding better multi-scale, multi-probe and multi-kinetics hydrogen mapping. For instance, a combination of silver decoration and SKPFM enables multi-scale and kinetically resolved hydrogen mapping. A combination of SKP and SKPFM enables quantitative mapping with high spatial resolution of several tens of nanometres. SIMS coupled with TDS allows for quantitative mapping of local hydrogen in a specimen with microstructural resolution with particular focus on the roles of interfaces and second phases. Possibly, APT coupled with SIMS and neutron radiography will enable multi-scale three-dimensional hydrogen mapping. However, for future lines of research, there are still important challenges remaining: these are the quantification of SIMS data in ferritic steels, visual detection of plastic-deformation-induced hydrogen desorption, atomistic or near atomistic observation of diffusible and weakly trapped hydrogen and improved direct correlations between hydrogen and single lattice-defects and their change upon mechanical loading. Addressing, improving and finally solving these challenges opens a pathway towards elucidating hydrogen segregation-related mechanisms of hydrogen embrittlement in steels.

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