Hydrogen blistering under extreme radiation conditions

Maciej Sznajder, Ulrich Geppert and Miroslaw R. Dudek

Metallic surfaces, exposed to a proton flux, start to degrade by molecular hydrogen blisters. These are created by recombination of protons with metal electrons. Continued irradiation progresses blistering, which is undesired for many technical applications. In this work, the effect of the proton flux magnitude on the degradation of native metal oxide layers and its consequences for blister formation has been examined. To study this phenomenon, we performed proton irradiation experiments of aluminium surfaces. The proton kinetic energy was chosen so that all recombined hydrogen is trapped within the metal structure. As a result, we discovered that intense proton irradiation increases the permeability of aluminium oxide layers for hydrogen atoms, thereby counteracting blister formation. These findings may improve the understanding of the hydrogen blistering process, are valid for all metals kept under terrestrial ambient conditions, and important for the design of proton irradiation tests.

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INTRODUCTION

High vacuum and intense corpuscular radiation are among those environmental conditions that can change irreversibly the metal properties on a short timescale. Similar conditions are met in nuclear fusion reactor technologies. There, the so-called plasma facing materials are exposed to intense deuterium ion fluxes where the ion energy, of a few tens of eV, is released just below the metal surface. These metals, e.g., tungsten, suffer then from the formation of tiny blisters (few tens of μm in diameter) filled with molecular deuterium gas. Another field which opens new challenges to material science is space technology. There, materials have to withstand rapid temperature changes, intense corpuscular and electromagnetic radiation as well as impacts caused by space debris and micro-meteorites. Metals play a key role in space technology since they are used in many applications, e.g., for building primary structures of spacecrafts, as surface mirrors for space telescopes, as thin coatings deposited on thin elastic films (e.g., polyimide) used for insulation purposes to reflect light and propel solar sails. For space applications the blistering process causes e.g., a change of thermo-optical properties of metallic surfaces which can affect thermal properties of satellite interiors. In case of solar sailing, when the sail surface is populated by blisters, the effective momentum transfer from the solar photons is reduced by a drop of sail surface reflectivity. As a consequence, the propulsion abilities of the sail are reduced by the growing population of the blisters.

In irradiation tests, the proton flux magnitude can be set freely from low to high values. The selected value is determined by the requirement to shorten laboratory time as much as possible to reach the number of incident protons which corresponds to the physical situation under study. The main motivation is to reduce the costs for maintenance of the facilities. However, such an approach has serious implications. An irradiation test should reveal the true nature of the changes in the material properties after the accumulation of the required proton number. This number is usually an equivalent of days, months or years during which the material is exposed to the respective environment. Certainly, the materials will change their physical properties in dependence on the applied proton energy and flux. The kinetic energy of the protons is released as heat in so-called thermal spikes. A larger proton flux and/or a larger proton kinetic energy will increase the number of thermal spikes. This, in turn, will augment the hydrogen diffusivity because the irradiation “drilled” holes into the test material structure. More holes enhance the mobility of hydrogen that can easier escape. This process can be named “radiation induced diffusion”.

A standard of the European Cooperation for Space Standardisation stated that metals, e.g., aluminium, copper, nickel, titanium, and stainless steels “do not suffer from space-environment conditions” also that “radiation at the level existing in space does not modify the properties of metals”. However, it has been shown that solar wind protons, while bombarding metallic surfaces, recombine with free electrons to hydrogen atoms within the metal structure. As a consequence surface blisters filled with molecular hydrogen gas are formed.

In 2014, a new standard was published which states that metals can be embrittled by hydrogen. Hydrogen embrittlement corresponds to four corrosion mechanisms: formation of a hydride phase, enhanced local plasticity, grain boundary weakening, and blistering.

Blistering is a fundamental process of metallic surface erosion. When taking place on metallic surfaces caused by exposure to a proton flux, the recombination processes of the protons with metal electrons to neutral hydrogen atoms are precondition for the blistering phenomenon. Hydrogen blisters occur as a consequence of supersaturation of hydrogen within the target material structure. The blisters change the physical properties of the exposed metals and increase their erosion rate.

The growth of blisters depends strongly on environmental conditions, i.e., proton energy, integrated proton flux (hereafter called fluence), and sample temperature. However, the efficiency of the growth is also influenced by the number of

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The dependence of blister growth mechanism on the crystallographic orientation has been studied experimentally. It has been proven that for aluminium the blisters preferably form on \{111\} instead on \{100\} and \{110\} planes.\textsuperscript{15}

**RESULTS AND DISCUSSION**

There are three possibilities of implanting hydrogen into the test material, i.e., the gas plasma method, the electrochemical method, and the chemical method. Sample exposure to gas ions avoids the well-known problem of Al(OH)$_3$ layer formation after performing the electrochemical or chemical charging of a test sample.\textsuperscript{18} Hence, we have used the gas plasma method to implant protons to the aluminium structure. Ten foil samples have been exposed to the proton flux. Irradiation time $t_{irr}$, accumulated fluence $F$, as well as its systematic errors are given in Table 1.

The test material is a metallised Upilex-S polyimide foil covered on both sides with 100 nm vacuum deposited aluminium (VDA) layers provided by the UBE Company. Since the foil was stored under room conditions (relative humidity 55 ± 5%, temperature 22 ± 2 °C) its surface is covered with a native AlO\textsubscript{x} layer, approx. 5 nm thick.\textsuperscript{19}

Irradiation tests have been performed by use of a linear proton accelerator at the German Aerospace Center in Bremen. The accelerator is part of the so-called Complex Irradiation Facility (CIF) established to investigate the changes of physical material properties, caused by electromagnetic and corpuscular radiation. The facility has been built in Ultra High Vacuum (UHV) technology and without rubber sealings to avoid self-contamination. It is equipped with electron and proton linear accelerators, where both can generate independently particles in the energy range from 1 to 100 keV with a maximum current of 100 µA. Three electromagnetic sources are also available, i.e., an argon VUV source, a deuterium and a xenon lamp, also called as a solar simulator. When working simultaneously they cover a wide wavelength range from 40 nm to 2150 nm. The facility is also equipped with a mass-spectrometer which can analyse the particle spectrum coming out of a test object generated during an irradiation test. The facility is presented in Fig. 1. All beams of the radiation sources meet at the irradiation chamber where the test sample is placed.

The protons are generated by ionisation of the hydrogen gas stored in a bottle and placed within a secure deck close to the accelerator. The hydrogen is guided through a thermo-mechanical gas inlet valve with remote control to the ion source. The ionisation takes place within a glass bulb with a radio frequency, which is capacitively coupled to the bulb. The source output is optimised by control of the source gas pressure and oscillator loading. The plasma is then pushed out of the bulb to the acceleration section by a proper value of the so-called extraction voltage. Protons are accelerated in an appropriate tube by a high voltage, which corresponds to the required energy. The accelerator tube is manufactured as a metal to ceramic brazed assembly with no organic compounds. After the acceleration to beam line vacuum tubes which end within the irradiation chamber where a sample material under study is placed.

The irradiation procedure together with discussion of the error bars are given in the Supplementary Notes.

The outcome of the irradiation tests was unexpected. Based on previous experimental studies\textsuperscript{2} an increase of the average blister radius has been expected for larger proton fluences. However, it turned out that on samples which received the largest number of protons appear the smallest size and number of blisters.

| Sample | $t_{irr}$ [s] | $F$ [$10^{17}$ [p·cm$^{-2}$]] | $\delta F$ [$10^{17}$ [p·cm$^{-2}$]] | $<R>$ [µm] | $\delta R$ [µm] |
|--------|----------------|----------------|----------------|-------------|-------------|
| S1     | 0.62           | 1.40           | ± 0.003       | 0.112       | ± 0.063     |
| S2     | 1.39           | 2.97           | ± 0.007       | 0.136       | ± 0.096     |
| S3     | 2.84           | 6.10           | ± 0.015       | 0.154       | ± 0.098     |
| S4     | 3.41           | 7.74           | ± 0.019       | 0.126       | ± 0.073     |
| S5     | 5.52           | 11.86          | ± 0.029       | 0.132       | ± 0.067     |
| S6     | 5.95           | 13.51          | ± 0.033       | 0.090       | ± 0.046     |
| S7     | 9.74           | 20.93          | ± 0.051       | 0.095       | ± 0.044     |
| S8     | 9.97           | 22.66          | ± 0.055       | 0.099       | ± 0.052     |
| S9*    | 9.41           | 12.81          | ± 0.092       | –           | –           |
| S10**  | 3.92           | 12.77          | ± 0.019       | –           | –           |

*Sample S9* was exposed to ~1.5 times smaller proton fluence than the samples S1–S8 while the sample S10* was exposed to ~1.6 times larger proton flux than the samples S1–S8.

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The test material is a metallised film, i.e., 7.5 µm Upilex-S polyimide foil covered on both sides with 100 nm vacuum deposited aluminium (VDA) layers provided by the UBE Company. Since the foil was stored under room conditions (relative humidity 55 ± 5%, temperature 22 ± 2 °C) its surface is covered with a native AlO\textsubscript{x} layer, approx. 5 nm thick.\textsuperscript{19}

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In Fig. 2, the selected Atomic Force Microscope (AFM) images are shown which represent eight samples denoted as S1–S8, respectively. Based on this figure, one may note that samples from S1 to S5 are populated with the highest number of blisters, while in the case of samples denoted as S6, S7, and S8 a considerable decrease of both the number and size of blisters can be observed.

Figure 3 shows magnified sample areas which in turn allow to see blisters in more detail. In particular, there are two samples which deserve special attention, i.e., samples denoted as S2 and S8. The sample S2 in Fig. 3b has on its surface a region where the aluminium layer was mechanically removed before irradiation process and, therefore, no blisters have been formed on its surface. This area is marked by the white rectangle. Sample S8 (in Fig. 3h) indicates two special features. The first of them...
corresponds to a long exposure of the sample to the proton flux which results with AlOx surface delamination (white circle) and the second effect which is associated with the crack of a bubble cup (shown within a small white-bordered window). The influence of the proton fluence on the blister growth is discussed in the following paragraphs.

The blister population in S1–8, was studied by help of digital processing of the corresponding AFM images shown in the Fig. 2 (see Methods). In this approach, the images were converted into a grid composed of black and white pixels and the Hoshen–Kopelman clustering algorithm was used to discriminate blisters and to calculate their average radius $<R>$.

The outcome of the studies of S1–8, suggests that the evolution of blisters can be classified into three stages (marked with different colours, see Fig. 4b). They are all determined by the degradation level of the native AlOx-layer. Figure 4a shows the relation between blister numbers and their radii. Special attention was given to find out how the average blister radius evolves with increasing irradiation time and how it affects the number of blisters. This is visualised in Fig. 4c. The radii and their error bars are given in the last two columns of Table 1. The parameter $\delta R$ denotes standard deviation which is a measure of how spread out the values of blister radii are.

The dependence of the average blister radius $<R>$ on exposition time of the aluminium foil to proton flux is presented in Fig. 4b. It suggests that blistering is a discontinuous process. Initially, excess hydrogen agglomerate at the interface between aluminium and the protective oxide layer; stage marked with green colour. The hydrogen is also collected in different kinds of defects which are natural traps for diffusing interstitial atomic hydrogen. The increase of internal hydrogen gas pressure is the reason for the observed blister growth. The appearance of a large excess of hydrogen at the AlOx/Al interface may lead to Al surface reconstruction through the faceting growth mechanism. This growth stage of hydrogen blisters (marked with yellow colour) is interrupted by detachment of the AlOx layer from the internal aluminium layer. In this situation, large pieces of the oxide layer can be removed. The third stage of the blistering process seen in Fig. 4c (marked with red colour) takes place when the oxide layer is mostly degraded. In the former work of the authors, the applied proton flux was weaker than here. Then, the evolution of hydrogen blisters is a continuous process and their average radius $<R>$ increases in accordance to a particular power law.

The progressing degradation of the AlOx layer can explain the decreasing blister number while the samples receive larger proton fluences (Fig. 4c). The hydrogen permeability increases while the sample surface is scarred with a larger number of cracks. Therefore, a smaller amount of hydrogen within the aluminium structure can fuel the blisters and, consequently, a smaller number of blisters appear on the surface.

The dependency of the AlOx ageing on the proton flux has been examined too. Three samples, i.e., S9*, S6, and S10* have been exposed to three different proton fluxes: $1.36 \times 10^{12}$, $2.27 \times 10^{12}$, and $3.26 \times 10^{12}$ p$^+$/cm$^2$/s, respectively. Here, a Field Emission Microscope (FEM) has been used to analyse the samples. All three samples received comparable proton fluences, see Table 1.

Sample S9* has been exposed to the lowest proton flux, see Figs. 5a, b. Here, small dark areas can be spotted. These are places where the AlOx layer has been delaminated from the aluminium substrate (marked with blue circles). Sample S6 received an intermediate proton flux. Here, the presence of small delamination centres but also large surface areas delaminated from the aluminium substrate are seen (marked with green circles, Figs. 5, d). One can also recognise structures which appear as crack-lines formed at the AlOx surface (marked with red dashed lines). They are thin dark intersecting lines, covering the sample area exposed to protons.

Sample S10* received the highest proton flux (Figs. 5e, f). Clearly, there the AlOx layer is cracked. Figure 5e shows in particular a blister cup scared with the crack-lines, while at Fig. 5f a
large crack located at the base of a blister is seen. Note that the delamination of the AlOx layer (marked with white circle, Fig. 3h), as well as cracking of the blisters (seen in white rectangular window, Fig. 3h) have been observed for the sample S8 as well. It proofs the progressing material degradation and its influence on the blistering process.

The considered proton fluxes and energies can be compared with proton fluxes present in space. The chosen kinetic energy of
the protons (2.5 keV) was intended to deposit the recombined hydrogen completely within the aluminium. In the close vicinity of the Earth, i.e., up to altitudes of the geostationary orbit, the recorded proton fluxes start from energies of approx. 50 keV. However, for the interplanetary space the available data start from energy of few eVs.

In the interplanetary space a test object is subjected to a wide energy spectrum of the protons, from eVs up to hundreds of eVs. 

Fig. 3  Surface morphology of the specimens for eight different proton fluences. a. Sample S1. b. Sample S2. c. Sample S3. d. Sample S4. e. Sample S5. f. Sample S6. g. Sample S7. h. Sample S8. Black scale bars, 1 μm. White scale bar, 0.3 μm. Photographs were taken with AFM.
MeVs. The 100 nm aluminium would collect protons up to energy of 8 keV. Protons with higher energies travel through the aluminium and may stop within the polyimide substrate of the foil. They are of no interest, since they do not recombine to hydrogen, and therefore, they provide no contribution to the blister formation. A typical proton flux in the solar wind at 1 astronomical unit (AU) distance from the Sun and energy lower than 10 keV is \(-3 \times 10^8\) p\(^+\)cm\(^{-2}\)s\(^{-1}\). The flux can be estimated using e.g., the Advanced Composition Explorer (ACE) database. Taking the data from the lowest recorded energy to an energy of 8 keV in steps of 1 eV the cumulative flux is \(-2 \times 10^{12}\) p\(^+\)cm\(^{-2}\)s\(^{-1}\). It is comparable to the flux to which the samples S1–8 has been exposed. Sample S9* received an about 1.5 times smaller flux while that of sample S10* is nearly 1.6 times larger.

The experiments show clearly that the proton flux magnitude dictates the blister evolution when a native AlOx layer is formed on aluminium. It cracks and delaminates when the sample surface is bombarded by a flux of 2.27 \times 10^{12} p^+cm^-2s^-1 or higher. For lower fluxes the native AlOx is delaminated from its aluminium substrate as well, however on relatively small areas (Figs. 5a, b). The evolution of blisters can be classified into three stages. First, the blisters grow most dynamically and have dimensions of up to 0.3 \(\mu\)m in diameter before the AlOx starts to delaminate or crack. Second, the AlOx degrades and loses its ability to hold hydrogen within the aluminium structure. This results in blister diameters of about 0.25 \(\mu\)m. Third, if the aluminium is bombarded furthermore incessantly by protons, the process of AlOx degradation continues. Then, the typical blister diameter is about 0.2 \(\mu\)m.

In the context of space applications, proton fluxes received by the here shown samples represent their state at 1 AU from the Sun. After only a few days under space conditions the aluminium foil would be covered by hydrogen blisters as shown in the samples S1–8. A small increase of the flux by a factor of \(-1.6\) causes a significant change in the aluminium surface morphology (sample S10*). Hence, the radiation driven blistering phenomenon is very sensitive to process parameters.

These results are important for the planning of proton irradiation tests, especially for the well-suited selection of the flux magnitude. A careful choice of the applied proton flux is necessary. Too high fluxes will lead to an acceleration of material degradation that does not reflect the true nature of ageing processes as take place in the environment under study.

**METHODS**

In this work, two microscopy methods were used to visualise the blisters, an Atomic Force Microscope (AFM–Bioscope Catalyst by Bruker) and a Field Emission Microscope (FEM–FE-REM, HITACHI 4004 with EDX-System EDAX Genesis 5). In order to study in more detail the blister population, the blisters were counted and their radius was measured. This has been achieved by digital processing of the AFM images. They show different blistering stages at the aluminium foil surface. In order to approximate the average blister size the image data were converted first into ppm format (portable pixmap format) providing a map of N\(\times\)M colour pixels (defined by a RGB triplet of integers, each of them ranging from 0 to 255). They were next converted into black and white pixels—the pixel colour was replaced by black if the RGB sum was less than a threshold value of 459 and the white colour otherwise. In this study, the Hoshen–Kopelman algorithm\(^{20}\) for labelling clusters on a grid composed of black and white pixels was used to estimate the average radius \(<R>\) of blisters. Clusters representing blisters were defined by the collection of these black pixels which were in the nearest-neighbour relationship on the bitmap grid under consideration.

**Data availability**

The data presented and discussed in this study is available from the authors upon reasonable request.
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AUTHOR CONTRIBUTIONS

M.S. conducted and performed the proton irradiation experiments. M.D. processed the AFM images with the Hoshen–Kopelman algorithm. All authors contributed equally both with discussion of the results and with writing the paper.

ADDITIONAL INFORMATION

Supplementary information accompanies the paper on the npj Materials Degradation website (https://doi.org/10.1038/s41529-017-0024-z).

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Fig. 5  Morphology of samples exposed to three different proton fluxes: 1.36 × 10^{12} (top row), 2.27 × 10^{12} (middle row), and 3.26 × 10^{12} p cm^{-2} s^{-1} (bottom row). Specimens receive similar proton fluence. First row, sample S9* a and b, small number of AlOx delamination areas are present on the sample surface exposed to protons. Second row, sample S6 c and d, crack lines as well as delamination centres cover the irradiated area. Third row, sample S10* e and f, crack lines dominate the exposed areas. Black line: scale bars, 1 μm. Grey line: scale bars, 0.7 μm. White line: scale bars, 0.5 μm. Orange line: scale bar, 0.2 μm. Photographs were taken with FEM.
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