Suppressed hydrogen embrittlement of highstrength Al alloys by Mn-rich intermetallic compound particles

Xu, Yuantao Department of Mechanical Engineering, Kyushu University

Toda, Hiroyuki Department of Mechanical Engineering, Kyushu University

Shimizu, Kazuyuki Department of Physical Science and Materials Engineering, Iwate University

Wang, Yafei Department of Mechanical Engineering, Kyushu University

他

https://hdl.handle.net/2324/6792843

出版情報:Acta materialia. 236, pp.118110-, 2022-09-01. Elsevier バージョン: 権利関係:

| 1 | Suppressed hydrogen embrittlement of high-strength Al |
|----|--|
| 2 | alloys by Mn-rich intermetallic compound particles |
| 3 | Yuantao Xu ^{a,b*} , Hiroyuki Toda ^{a*} , Kazuyuki Shimizu ^c , Yafei Wang ^a , Baptiste Gault |
| 4 | ^{d,e} , Wei Li ^{b*} , Kyosuke Hirayama ^f , Hiro Fujihara ^a , Xuejun Jin ^b , Akihisa Takeuchi ^g , |
| 5 | Masayuki Uesugi ^g |
| 6 | ^a Department of Mechanical Engineering, Kyushu University, Fukuoka 819-0395, Japan |
| 7 | ^b Shanghai Key Laboratory of Materials Laser Processing and Modification, Shanghai Jiao Tong |
| 8 | University, Shanghai 200240, PR China |
| 9 | ^c Department of Physical Science and Materials Engineering, Iwate University, Morioka 020-8551, |
| 10 | Japan |
| 11 | ^d Max-Planck-Institut für Eisenforschung, Max-Planck-Straße 1, 40237 Düsseldorf, Germany |
| 12 | ^e Department of Materials, Royal School of Mines, Imperial College, Prince Consort Road, |
| 13 | London SW7 2BP, United Kingdom |
| 14 | ^f Department of Materials Science and Engineering, Kyoto University, Kyoto 606-8501, Japan |
| 15 | ^g Japan Synchrotron Radiation Research Institute, Hyogo 679-5198, Japan |
| 16 | *Corresponding author. Tel.: +81-928023246; E-mail: xu.yuantao.158@m.kyushu-u.ac.jp |
| 17 | (Yuantao Xu); toda@mech.kyushu-u.ac.jp (Hiroyuki Toda); <u>weilee@sjtu.edu.cn</u> (Wei Li) |
| 18 | Abstract |
| 19 | The pursuit of strong and ductile Al alloys with superior resistance to hydrogen |
| 20 | embrittlement (HE) is practically significant across the aerospace and transportation |
| 21 | industries among others. Unfortunately, effective ways to progress on the strength-HE |
| 22 | trade-off for Al-alloys remain elusive. A strategy of suppressing HE by introducing |

| 23 | intermetallic compound (IMC) particles to achieve hydrogen redistribution in various |
|----|--|
| 24 | trapping sites was proposed. Here, we systematically induce the precipitation of a |
| 25 | constant volume fraction of intermetallic compound (IMC) particles by adding one of |
| 26 | 14 elements in a ternary Al-Zn-Mg high-strength alloy. We show a strong correlation |
| 27 | between hydrogen trapping energies of the IMC obtained from ab initio calculations |
| 28 | with the resistance to HE. Mn-rich $Al_{11}Mn_3Zn_2$ particles exhibit the highest hydrogen |
| 29 | trapping energy (0.859 eV/atom), leading to a decrease by approximately 5 orders of |
| 30 | magnitude in the hydrogen occupancy in η_2 (MgZn_2) phase interfaces and grain |
| 31 | boundaries, where HE cracks initiate. The Mn-addition did not deteriorate the |
| 32 | ductility and most Al ₁₁ Mn ₃ Zn ₂ particles remained intact during plastic deformation |
| 33 | which was revealed by in-situ 3D X-ray tomography. Hydrogen-induced strain |
| 34 | localization at η_2 phase interfaces and grain boundaries were inhibited due to strong |
| 35 | hydrogen trapping capacity of Al ₁₁ Mn ₃ Zn ₂ , hence preventing HE cracks initiation. |
| 36 | Our approach effectively suppresses hydrogen-induced cracks without sacrificing the |
| 37 | ductility, and our strategy can help the design roadmap of HE-tolerant high-strength |
| 38 | metallic alloys. |

Keywords: Hydrogen embrittlement; Hydrogen trapping; Intermetallic compound
particles; Aluminum alloys; Ductility

41 **1. Introduction**

42 Degradation of the mechanical properties of structural materials due to hydrogen 43 intake has become the main concern for key components in many industries, such as 44 nuclear energy [1-4], aerospace [5,6], oil and gas [7]. Hydrogen causes cracking [4,5],

45 hydride precipitation [1-3], embrittlement [8], and other harmful effects [9], thereby 46 limiting the service life of parts, increasing operating costs, and creating potentially 47 unsafe working conditions. Al-Zn-Mg alloys have attracted significant research 48 interest for structural applications in aerospace, transportation, and many other 49 industries owing to their high strength and low density [10,11]. However, their low 50 resistance to stress corrosion cracking and hydrogen embrittlement (HE) severely 51 affects the service life and applications [12-14].

Environmentally assisted HE in high-strength 7xxx series aluminum alloys leads 52 53 to low ductility and catastrophic failure in structural applications [11,15,16]. The 54 dissolved hydrogen atoms can easily penetrate surface oxide layer and diffuse into various trapping sites (e.g., dislocations, vacancies, grain boundaries, and precipitates) 55 56 in the Al alloys [17]. Hydrogen not only reduces the ductility but can also change the fracture mode from ductile to brittle, macroscopically featuring as intergranular or 57 transgranular quasi-cleavage fractures [18-20]. Intergranular fracture (IGF) is 58 primarily associated with the hydrogen segregated at the grain boundary that weakens 59 the interface bonding strength (hydrogen-enhanced decohesion (HEDE) [20]). The 60 mechanisms of quasi-cleavage fracture (QCF) remain controversial due to the lack of 61 direct experimental evidence, for which the HEDE mechanism, hydrogen-enhanced 62 localized plasticity (HELP) [21] mechanism, and hydrogen-accelerated spontaneous 63 microcracking at precipitate interfaces have been proposed [22]. 64

The effects of various trapping sites such as micro pores, grain boundaries,
 dislocations, vacancies, precipitates, and IMC particles on HE mechanism of Al alloys

were widely investigated [13,17,22-24]. A high density of micro pores can store 67 hydrogen and contribute to ductile fracture [23]; however, a small amount of 68 dissolved hydrogen can cause brittle fracture [24]. Controlling the distribution of 69 dissolved hydrogen between grain boundaries, dislocations, and vacancies in 70 Al-Zn-Mg alloys was proven ineffective in suppressing HE [17]. The precipitate 71 72 interfaces (Al-MgZn₂) was reported to be a more preferable hydrogen trap site 73 compared to other defects in terms of binding energy and hydrogen partitioning, which leads to spontaneous hydrogen-accumulated QCF [22]. The design concept of 74 IMC particles primarily focused on their role in grain refinement in Al alloys [25-27], 75 whereas little information was reported regarding their role in the HE suppression of 76 Al alloys. A recent study showed that the Al₇Cu₂Fe particles are effective hydrogen 77 78 traps to reduce the trapped hydrogen at the precipitate interfaces and confirmed to be able to suppress QCF; however, the improvement of the ductility was not satisfactory 79 due to the brittle nature of Al₇Cu₂Fe particles. Therefore, achieving a superior HE 80 resistance without sacrificing the ductility remains a challenge in Al alloys and 81 82 exploring suitable IMC particles that can realize the balance between hydrogen trapping ability and brittle nature seems a potential route to effectively control HE. 83

Here, we propose a hydrogen redistribution approach by introducing IMC particles as strong hydrogen trapping site, which significantly reduces trapped hydrogen at precipitate interfaces and grain boundaries and suppresses QCF and IGF originating from hydrogen segregation. First-principles calculations are employed to calculate the binding energies between hydrogen and various IMC particles formed by

adding a fourth alloying element to Al-Zn-Mg alloys. The effect of IMC particles on the hydrogen partitioning behaviors among various trap sites during plastic deformation are investigated by 3D X-ray technique and first-principles calculations.
In addition, the role of IMC particles in initiation and propagation of hydrogen-induced QCF and IGF are discussed, and the mechanism of how IMC particles can act as strong hydrogen traps for resisting HE is clarified.

95 2. Materials and methods

To investigate the effects of IMC particles on the hydrogen partitioning and 96 97 embrittlement behaviors, various IMC particles were formed by adding a fourth 98 element, such as 1.00 Nb, 0.94 Mo, 1.00 V, 0.15 Ca, 0.09 Y, 0.15 Sr, 0.10 Ni, 1.19 Zr, 0.15 Fe, 0.26 Cr, 1.02 Ti, 0.19 Co, 0.94 Sc, or 1.91 Mn to the Al-10.0 Zn-1.06 Mg 99 100 (wt.%) Al alloys. The element content was designed to obtain IMC particles with a volume fraction of approximately 0.28%. The detailed thermomechanical treatment 101 process is presented in Figure S1. The corresponding alloys were prepared by 102 homogenization at 500 °C for 24 h after casting and then hot rolling at 450 °C with a 103 104 rolling reduction of 50%. A solution-aging process was performed as follows: first, a solution treatment was conducted at 500 °C for 5 h. Subsequently, an over-aging 105 106 treatment was performed at 120 °C for 40 h and 180 °C for 7 h. To investigate the influence of the pre-charged hydrogen content on the embrittlement behavior, 107 specimens with different hydrogen contents were prepared using LH and HH 108 treatments. The specimens with HH content were charged using electrical discharge 109 110 machining (EDM) cutting in distilled water, while the specimens with LH content

were prepared by EDM cutting in oil. The pre-charged hydrogen contents were approximately 6.97 mass ppm and 1.40 mass ppm for the specimens after HH and LH treatments, respectively, the total hydrogen content was analyzed by the vacuum fusion method, which are similar to literature values [13]. The hydrogen concentration in the Mn-added Al alloys after HH and LH treatments were also measured by thermal desorption apparatus (TDA) method, the corresponding desorption curves were shown in Figure S2.

The gauge length of in-situ tensile specimens was 0.7 mm, and these specimens 118 had cross-sectional dimensions of 0.6 mm \times 0.6 mm. The deformation and fracture 119 120 behaviors were observed using high-resolution X-ray tomography during in-situ tensile tests at BL20XU of Spring-8 in Japan; the detailed experimental setup is 121 shown in Figure S3. In-situ tensile tests (strain rate $\sim 3 \times 10^{-3} \text{ s}^{-1}$) were conducted using 122 a DEBEN CT 500 machine at room temperature. 3D images were taken using 123 monochromatic 20 keV X-rays; the detailed methods used for tomography 124 observation and in-situ tensile tests are reported in the literature [13]. 125

Transmission electron microscopy (TEM; JEOL JEM 2100F/ARM 200F) was used to characterize the detailed microstructure of the precipitates, IMC particles, and Al matrix. Scanning electron microscope (SEM) (TESCAN MIRA3) equipped with an electron back-scatter diffraction (EBSD) detector was performed to analyze strain distribution and phase features by transmission Kikuchi diffraction (TKD) technique. Thin foils were prepared for TEM and EBSD observations by electro-polishing using Struers (Tenupol-5) equipment at -30 °C and 15 V; the electrolyte was 7% perchloric acid and 93% ethanol. Atom probe tomography (APT) measurements were performed using a local electrode atomic probe (LEAP 4000 HR) to obtain the spatial distribution of elements and the number density and size of nanoparticles. APT data was obtained at 50 K under ultrahigh vacuum (4.5×10^{-11} Torr), a pulse fraction of 0.2, and a repetition rate of 200 kHz. The APT data was reconstructed using IVASTM 3.6.2 software.

Hydrogen trapping at MgZn₂ precipitates interface and various intermetallic 139 compound particles was calculated using first-principles calculations. The 140 first-principles calculations were conducted within the DFT framework using the 141 142 Vienna ab initio simulation package (VASP 5.2) with the Perdew-Burke-Ernzerhof generalized gradient approximation exchange-correlation density functional. The 143 144 Monkhorst-Pack algorithm was selected for the Brillouin-zone k-point samplings. A plane-wave energy cutoff of 360 eV was used with a first-order Methfessel-Paxton 145 smearing scheme, employing a smearing parameter of 0.2 eV. The total energy 146 converged to 10^{-6} eV in all the calculations. The relaxed atomic configurations were 147 obtained by the conjugate gradient method in which the search terminated when the 148 forces on all atoms were below 0.01 eV/Ang. The crystal structure parameters and 149 150 sampling k-point mesh of various particles, and the details of calculation process can be found in the Supplementary Materials 1.1. 151

The hydrogen trap sites in this study were grain boundaries, dislocations, pores, vacancies, precipitate interfaces, and IMC particles. To quantitatively analyze these trap site densities, the area of the grain boundaries, dislocation densities, pore

morphology, vacancy concentrations, area of precipitate interfaces, and volume
fractions of IMC particles were obtained using a combination of experimental
techniques and mathematical models, which are detailed in the Supplementary
Materials 1.2.

159 **3. Results and discussion**

160 3.1 Trapping energies of IMC particles to hydrogen and their HE resistances

Various IMC particles were formed by adding a fourth element such as Nb, Mo, 161 V, Ca, Y, Sr, Ni, Zr, Fe, Cr, Ti, Co, Sc, or Mn to Al-10.0Zn-1.06Mg alloys. 162 First-principles density functional theory (DFT) was employed to calculate the 163 trapping energies of various IMC particles to hydrogen, and the corresponding results 164 are shown in Figure 1. The hydrogen trapping energies of various defect structures, 165 such as edge/screw dislocations [28], grain boundaries [29], vacancies [30], 166 precipitates [22,31], and micro pores [32] are also summarized. The maximum 167 hydrogen trapping energy in the interior of Al₁₁Mn₃Zn₂ was 0.859 eV/atom, much 168 higher than that in other IMC particles and defect structures, which indicates that it 169 can be used as a strong hydrogen trapping site for absorbing a large amount of 170 hydrogen at interstitial sites. 171

The HE resistances of samples with and without IMC particles were evaluated using in-situ tensile tests. In-situ tensile stress–displacement curves for the standard-state, Sc-added, and Mn-added Al alloys after low-hydrogen (LH) and high-hydrogen (HH) treatments are shown in Figure 2(a). The mechanical properties of the standard-state and Sc-added Al alloys were severely affected by the hydrogen

content, where the elongation were dramatically reduced after the HH treatment, 177 compared to the LH treatment. However, in the Mn-added alloys, the elongation 178 179 remained almost unchanged after the HH treatment, compared to the LH treatment, which means that the Mn-added Al alloys overcomes hydrogen-induced ductility loss. 180 181 Detailed fractography analysis revealed that the fractional areas of IGF and QCF were 182 50.9% and 46.0% in the standard-state Al alloys and 9.3% and 11.5% for the 183 Mn-added Al alloys after the HH treatment, with 81.7% and 75.0% reduction, respectively. This confirms that the Mn-added Al alloys significantly improves HE 184 185 resistance (Figure 2(b)). The effect of a fourth element such as Nb, Mo, V, Ca, Y, Sr, Ni, Zr, Fe, Cr, Ti, Co, Sc, or Mn on the HE fracture area is shown in Figure 2(c); the 186 corresponding fracture morphology is presented in Figure 3 and Figure S4. The 187 relationships between hydrogen trapping energies of IMC particles and fractional 188 areas of QCF are summarized in Figure 2(d). Fractional areas of QCF were 189 significantly reduced with the increase of trapping energies of different IMC particles 190 in the Al alloys treated by HH. Compared with the addition of the other elements, the 191 192 Mn-added Al alloys exhibited the best HE resistance owing to the highest trapping energy between the Al₁₁Mn₃Zn₂ particles and hydrogen. 193

HE occurred in the 7 xxx series Al alloy (Al-10.1Zn-1.2Mg) with hydrogen content of 0.41 mass ppm [12], and also appeared in LH-treated standard-state Al alloys (1.40 mass ppm) in this work (Figure 3). It was reported that the introduction of Mn-rich particles by adding Mn to Al-Zn-Mg alloys under hydrogen-free conditions results in a slight decrease in ductility [33,34]. However, the introduced Mn-rich

particles in LH-treated Mn-added Al alloys contributes to a significant increase in ductility compared with LH-treated standard-state Al alloys, which reconfirms that strong hydrogen trapping capacity of $Al_{11}Mn_3Zn_2$ particles.

202 *3.2 Hydrogen partitioning behavior*

203 The microstructures of the standard-state and Mn-added Al-Zn-Mg alloys were investigated (Figure 4(a-b)). The standard-state alloys primarily consisted of MgZn₂ 204 and Al matrix, while the Mn-added Al alloys was primarily comprised of Al₁₁Mn₃Zn₂, 205 MgZn₂, and Al matrix (Figure 4(a-b), Figure S5, and Figure 5), which were identified 206 207 using high-resolution transmission electron microscopy (HRTEM) and atom probe tomography (APT) analysis. MgZn₂ has a hexagonal close-packed (hcp) structure, and 208 its edge and side surfaces of the plate are coherent and semi-coherent along the 209 [112]_{Al} zone axis, respectively. The orientation relationship between MgZn₂ and the 210 matrix is $\left[\overline{112}\right]_{Al}//\left[2\overline{110}\right]_{MeZn_2}$. Al₁₁Mn₃Zn₂ has an orthorhombic structure; the 211 orientation relationship between it and the matrix is $\begin{bmatrix} 011 \end{bmatrix}_{Al} / / \begin{bmatrix} 2\overline{2}1 \end{bmatrix}_{Al_1,Mn,Zn}$. The 212 number density of n₂ (MgZn₂) precipitates in standard-state and Mn-added Al alloys 213 are 1.13×10^{23} and 0.78×10^{23} m⁻³, and their equivalent spherical particle radius are 6.5 214 and 7.4 nm, respectively, which were determined via APT (Figure 5). The average 215 grain sizes were measured to be 124 and 75 µm for the standard-state and Mn-added 216 Al alloys, respectively (Figure 6). 3D perspective views of IMC particles and micro 217 pores obtained by high-resolution X-ray tomography are shown in Figure 4(c). These 218 parameters were used in the calculation of hydrogen occupancy of the corresponding 219 220 trap site.

The hydrogen partitioning behavior in the standard-state and Mn-added Al alloys are shown in Figure 4(d). The hydrogen occupancy of a trap site depends on both the trap density and trap binding energy, which were derived from experimental results (Figure 4(a)-4(c)) and first-principles calculations. See the Supplementary Materials 1.2 for detailed calculation procedures of hydrogen occupancy of each hydrogen trap site.

It was found that hydrogen atoms were mainly trapped at micro pores, η_2 227 precipitates, and vacancies in the standard-state alloys; however, the Al₁₁Mn₃Zn₂ 228 229 particles are the strongest hydrogen trap, which reduced the hydrogen occupancy at 230 other trap sites such as micro pores, η_2 precipitates, vacancies, and grain boundaries. In particular, the hydrogen occupancy of η_2 interfaces and grain boundaries were 231 decreased by approximately 5 orders of magnitude (from 1.15×10^{-1} to 6.20×10^{-7} and 232 from 4.42×10^{-1} to 4.60×10^{-6} atomH/site, respectively), which contributes to the 233 suppression of hydrogen-induced QCF and IGF (Figure 2). 234

3.3 Damage behavior of Al₁₁Mn₃Zn₂ particles and hydrogen partitioning behavior
during deformation

Figure 7(a)-7(h) show the damage behavior of Al₁₁Mn₃Zn₂ particles in Mn-added Al alloys after the HH treatment during in-situ tensile tests. 3D images around the crack tip were obtained using high-resolution X-ray tomography. The observation regions ahead of the hydrogen-induced quasi-cleavage crack tip and near the crack ridge were approximately 5.7×10^2 and 4.3×10^3 µm³, respectively, as shown in Figure 7(i). Particle debonding and fracture at different steps were classified by analyzing the voids and their neighboring particles: debonding was confirmed if only one particle
was found adjacent to one void; however, particle fracture was identified if two
particles were connected to one void.

246 The damage ratios, R_d , are defined as follows:

248

247
$$R_d(fractured) = n_F / n_{total} \times 100\%$$
(1)

$$R_d(debonded) = n_D / n_{total} \times 100\%$$
⁽²⁾

249
$$R_d = R_d(fractured) + R_d(debonded)$$
(3)

250 where n_F and n_D are the numbers of fractured and debonded particles, 251 respectively; n_{total} is the total number of IMC particles for each condition.

At a macroscopic tensile strain of 5.7%, the damage ratio of IMC particles near 252 the crack ridge was approximately 7% (2/28). With the further increase of tensile 253 254 strain to 9%, the damage ratio slightly increased to approximately 20% (2/10). Therefore, most of these IMC particles remained intact during plastic deformation, 255 implying that these particles, with strong hydrogen trapping capacity, did not 256 deteriorate the ductility due to good damage resistance. It is thereby expected that the 257 IMC particles can effectively suppress hydrogen-induced cracks without sacrificing 258 the ductility. This is consistent with the significantly increased fracture strain of the 259 particles-containing Al alloys compared with the particles-free Al alloys (Figure 2(a)). 260 Figure 7(i) and 7(j) show the hydrogen partitioning behavior during deformation 261

in Mn-added Al alloys after the HH treatment. At the in-situ tensile strain of 3.6%, the
volume fractions of IMC particles at the regions ahead of the hydrogen-induced
quasi-cleavage crack tip and near the crack ridge were 1.54% and 0.26%, respectively.

Hydrogen partitioning calculations revealed that the hydrogen occupancy of the η_2 265 interfaces at the region ahead of the crack tip was approximately 8 orders of 266 magnitude lower than that near the crack ridge owing to the higher density of IMC 267 particles. This means that regions with a low density of IMC particles are prone to 268 269 crack nucleation and propagation (Figure 7(i)). Al₁₁Mn₃Zn₂ particles were the main hydrogen trap, and the trapped hydrogen content could reach 2.0×10^{25} atomH·m⁻³ 270 (Figure S6), which was much higher than other tapping sites. The hydrogen 271 occupancy of Al11Mn₃Zn₂ particles remained almost unchanged before and after 272 273 deformation (3.6% strain). Moreover, they exhibited good damage resistance even at high tensile strain (up to 9%). This confirms that the $Al_{11}Mn_3Zn_2$ particles are 274 beneficial for improving the HE resistance without harming the ductility during 275 276 deformation.

277 *3.4 Suppression of hydrogen-induced strain localization at grain boundaries*

Micro-deformation mechanisms of Al11Mn3Zn2 particles under HH and LH 278 conditions were revealed by transmission Kikuchi diffraction (TKD) technique and 279 TEM (Figure 8). Al₁₁Mn₃Zn₂ particles did not deform during tensile failure under HH 280 281 and LH conditions, strain concentration around Al₁₁Mn₃Zn₂ particles mainly resulted from dislocations piled-up. The strain distributions around Al₁₁Mn₃Zn₂ and in Al 282 matrix remained almost the same when changing treatment conditions from LH to HH, 283 indicating that hydrogen-induced strain localization caused by the interaction of 284 hydrogen and dislocations was suppressed [13,35-37], which is mainly attributed to 285 strong hydrogen trapping capacity of Al₁₁Mn₃Zn₂. 286

The microstructures near the fracture surfaces of Mn-added and standard-state Al 287 alloys treated by HH after tensile failure are shown in Figure 9. The 288 hydrogen-induced cracks mainly propagated along the grain interiors in the Mn-added 289 Al alloys (Figure 9(a)), while they mainly propagated along the grain boundaries in 290 the standard-state Al alloys (Figure 9(e)). Hydrogen-induced cracks preferably 291 292 nucleated at IMC-poor regions, and IMC-rich regions could effectively blunt and suppress hydrogen-induced cracks (Figure S7), which is consistent with the result at 293 3.6% tensile strain (Figures 7(i) and 7(j)). The piled-up dislocations at grain 294 boundaries were significantly reduced during tensile failure after adding Mn in 295 standard-state Al alloys (Figure 9(b)-(d) and 9(f)-(h)), supporting that the trapped 296 hydrogen at grain boundaries were remarkably decreased owing to the introduction of 297 298 $Al_{11}Mn_3Zn_2$.

299 3.5 Suppression of hydrogen-induced strain localization at η_2 phase interfaces

As shown in Figure 10, strain maps were plotted by geometric phase analysis 300 (GPA) [38], inverse FFT patterns of HRTEM images were used to characterize 301 302 dislocations around η_2 phase interfaces. 10 HRTEM images (each image contains 2 precipitates) were collected for the HH-treated Mn-added Al alloys and HH-treated 303 standard-state Al alloys, respectively, and 20 precipitates were counted and analyzed 304 for each condition. Strain concentration around η_2 phase interfaces was not very 305 obvious in HH-treated Mn-added Al alloys subjected to tensile failure, however, the 306 strain values were very high for almost the entire η_2 phase interface in HH-treated 307 standard-state Al alloys subjected to tensile failure (Figure 10(c)-(f) and 10(j)-(m)). 308

309 Moreover, more significant dislocations pile up occurred near the η_2 phase interfaces in HH-treated standard-state Al alloys, compared with that in HH-treated Mn-added 310 Al alloys (Figure 10(g) and 10(n)). Based on the analysis results of 20 precipitates 311 under each condition, it can be confirmed that the strain localization at n_2 phase 312 interfaces was effectively suppressed during tensile failure after adding Mn in 313 314 standard-state Al alloys, which indicates that the trapped hydrogen at η_2 phase interfaces were largely removed by Al₁₁Mn₃Zn₂. These results demonstrate that IMC 315 particles inhibit hydrogen-induced strain localization η_2 phase interfaces by 316 consuming most of the hydrogen, hence preventing HE cracks initiation. 317

318 The QCF mechanism induced by hydrogen in structural materials, such as ferritic, martensitic, and austenitic steels, and nickel, can be explained by the HELP model 319 [39], where hydrogen-affected dislocations contribute to structural failure. However, a 320 QCF mechanism assisted by hydrogen in an Al-Mg-Zn alloy was reported [22]; 321 hydrogen atoms tended to be trapped at the interfaces rather than at grain boundaries 322 and dislocations owing to the higher binding energy with hydrogen, which induced 323 hydrogen-accelerated spontaneous microcracking at aluminum-precipitate interfaces. 324 In this work, a similar fracture mechanism caused by precipitate interfaces decohesion 325 326 can also be used to explain the QCF. The hydrogen occupancy of the η_2 coherent interface decreased from 8.4×10^{-10} in the unloaded state to 8.7×10^{-11} atomH/site at 327 3.6% in-situ tensile strain (the corresponding hydrogen concentration is 5.3×10^{-10} 328 atomH/nm²; the hydrogen trap interval for a coherent n precipitate interface is 6.1 329 atomH/nm², assuming that the hydrogen occupancy is equal to one [17,31]). The 330

hydrogen occupancy of the η_2 semi-coherent interface decreased from 1.84×10^{-5} 331 atomH/site in the unloaded state to 1.9×10^{-6} atomH/site (1.2×10^{-5} atomH/nm²). The 332 interfacial decohesion of η_2 precipitates initiates when the trapped hydrogen density 333 reaches 18.9 atomH/nm² [13,17], which is approximately 3.6×10^{10} and 1.6×10^{6} times 334 335 higher than the hydrogen concentration at the coherent and semi-coherent interfaces of η_2 precipitates in the Mn-added Al alloys, respectively. Moreover, it was observed 336 that hydrogen-induced strain localization at η_2 phase interfaces was suppressed after 337 adding Mn in standard-state Al alloys (Figure 10). As a result, the fractional areas of 338 QCF originating from the hydrogen-induced decohesion of precipitate interfaces was 339 remarkably reduced (Figure 2(b)). 340

Hydrogen trapped at grain boundaries decreases the cohesion strength of the 341 342 boundaries (HEDE model) and results in IGF, which is a common HE mechanism in structural materials. The critical hydrogen concentration at the grain boundaries for 343 the initiation of intergranular cracks is approximately 2.8×10^{22} atomH·m⁻³ in 344 345 Al-Zn-Mg-Cu alloys [30]. The hydrogen content trapped at the grain boundaries in Mn-added Al alloys in the unloaded and 3.6% tensile strain states were 4.04×10^{12} and 346 4.19×10^{11} atomH·m⁻³, respectively, which are much lower than the critical value 347 required for hydrogen-induced intergranular cracking. In addition, it was found that 348 hydrogen-induced strain localization at grain boundaries was inhibited after adding 349 Mn in standard-state Al alloys (Figure 9). Therefore, this is the main reason for the 350 significant reduction in the fractional areas of IGF (Figure 2(b)). 351

352

Hydrogen-induced strain localization at grain boundaries and η_2 phase interfaces

were suppressed due to strong hydrogen trapping capacity of Al₁₁Mn₃Zn₂, which 353 could remove hydrogen from the grain boundaries and η_2 phase interfaces. Therefore, 354 the Al alloys with Mn-rich particles exhibited the superior HE resistance by 355 eliminating 75.0% and 81.7% of hydrogen-induced QCF and IGF, respectively, and 356 357 suppressing hydrogen-induced ductility loss compared to the standard-state Al alloys. Moreover, in terms of HE sensitivity $(El_{loss} = (El_{LH} - El_{HH})/El_{LH} \times 100\%$, El_{LH} and El_{HH} 358 are fracture elongations of LH and HH-treated specimens, respectively), the Al alloys 359 with Mn-rich particles showed lack of HE sensitivity, which indicates that most of the 360 361 hydrogen is occupied by Mn-rich particles and does not deteriorate the mechanical properties. 362

IMC particles are often used to inhibit recrystallization during solution heat treatment; however, they are detrimental to the mechanical properties of structural materials, especially the ductility [40]. Although IMC particles such as Al_7Cu_2Fe have been proven to improve the HE resistance due to hydrogen trapping effect, this still cannot solve the problem of ductility deterioration [13].

In this work, we proposed a strategy of suppressing HE by introducing IMC particles to achieve hydrogen redistribution in various trapping sites, rather than eliminating the hydrogen. IMC particles with HH trapping energies were designed using first-principles calculations; Mn-rich $Al_{11}Mn_3Zn_2$ particles exhibit the highest hydrogen trapping energy (0.859 eV/atom), leading to a decrease by approximately 5 orders of magnitude in the hydrogen occupancy in η_2 (MgZn₂) phase interfaces and grain boundaries, where HE cracks initiate. Most $Al_{11}Mn_3Zn_2$ particles remained intact during plastic deformation and exhibited good damage resistance. Hydrogen-induced strain localization at grain boundaries and η_2 phase interfaces were suppressed due to strong hydrogen trapping capacity of Al₁₁Mn₃Zn₂, hence preventing HE cracks initiation. Superior HE resistance without sacrificing the ductility was achieved through this hydrogen redistribution approach, which can also be applied to a variety of high-strength structural materials containing IMC particles, such as iron, titanium, and zirconium-based alloys.

4. Conclusions

We systematically induce the precipitation of a constant volume fraction of intermetallic compound (IMC) particles by adding one of 14 elements in a ternary Al-Zn-Mg high-strength alloy, the effect of IMC particles on HE resistances and HE mechanisms were investigated. The main conclusions are as follows:

(1) A strong correlation between hydrogen trapping energies of the IMC obtained 387 from ab initio calculations with the resistance to HE was confirmed. The Al₁₁Mn₃Zn₂ 388 particles had a trapping energy value (0.859 eV/atom) that was much higher than that 389 of all known hydrogen trap sites, which caused the hydrogen occupancy at other trap 390 391 sites such as micro pores, η_2 precipitates, vacancies, and grain boundaries to be significantly reduced. In particular, the hydrogen occupancy of η_2 interfaces and grain 392 boundaries were decreased by approximately 5 orders of magnitude compared to the 393 standard-state Al alloys (from 1.15×10^{-1} to 6.20×10^{-7} and from 4.42×10^{-1} to 4.60×10^{-6} 394 atomH/site, respectively). 395

396

(2) Hydrogen occupancy and damage ratio of Al₁₁Mn₃Zn₂ particles changed

| 397 | slig | htly when the tensile strain reached 3.6% and 9% after HH treatment (6.97 ppmw | | | |
|-----|--|--|--|--|--|
| 398 | Н), | H), respectively, suggesting that they did not deteriorate the ductility. In addition, | | | |
| 399 | hyd | hydrogen-induced strain localization at grain boundaries and η_2 phase interfaces were | | | |
| 400 | supj | suppressed due to strong hydrogen trapping capacity of Al ₁₁ Mn ₃ Zn ₂ , hence preventing | | | |
| 401 | HE | HE cracks initiation. Therefore, the Al alloys with Mn-rich particles exhibited the | | | |
| 402 | supe | superior HE resistance by eliminating 75.0% and 81.7% of hydrogen-induced QCF | | | |
| 403 | and IGF, respectively, and suppressing hydrogen-induced ductility loss compared to | | | | |
| 404 | the standard-state Al alloys. | | | | |
| 405 | References | | | | |
| 406 | [1] | A. T. Motta, et al., Hydrogen in zirconium alloys: A review, J. Nucl. Mater. 518 (2019) | | | |
| 407 | | 440-460. | | | |
| 408 | [2] | S. Sobieszczyk, Hydrogen-enhanced degradation and oxide effects in zirconium alloys for | | | |
| 409 | | nuclear applications, Int. J. Hydrogen Energy 6 (2011) 8619-8629. | | | |
| 410 | [3] | S. Suman, M. K. Khan, M. Pathak, R. N. Singh & J. K. Chakravartty, Hydrogen in Zircaloy: | | | |
| 411 | | Mechanism and its impacts, Int. J. Hydrogen Energy 40 (2015) 5976-5994. | | | |
| 412 | [4] | M. Shahzad, H. Qureshi, H. Waqas & N. Hussain, A case study of environmental assisted | | | |
| 413 | | cracking in a low alloy steel under simulated environment of pressurized water reactor, | | | |
| 414 | | Mater. Des. 32 (2011) 5154-5158. | | | |
| 415 | [5] | M. P. Brady, et al., Rapid diffusion and nanosegregation of hydrogen in magnesium alloys | | | |
| 416 | | from exposure to water, ACS Appl. Mater. Interfaces. 9 (2017) 38125-38134. | | | |
| 417 | [6] | A. S. H. Kabir, et al., Effect of post-weld heat treatment on microstructure, hardness, and | | | |
| 418 | | tensile properties of laser-welded Ti-6Al-4V, Metall. Mater. Trans. A 43 (2012) 4171-4184. | | | |
| 419 | [7] | Z. Tarzimoghadam, et al., Multi-scale and spatially resolved hydrogen mapping in a Ni-Nb | | | |
| 420 | | model alloy reveals the role of the d phase in hydrogen embrittlement of alloy 718, Acta | | | |
| 421 | | Mater. 109 (2016) 69-81. | | | |
| 422 | [8] | G. Mcmahon, B. D. Miller & M. G. Burke, High resolution NanoSIMS imaging of | | | |
| 423 | | deuterium distributions in 316 stainless steel specimens after fatigue testing in high pressure | | | |

- 424 deuterium environment, Npj Mater. Degrad. 2 (2018) 1-6.
- 425 [9] I. I. Chernov, M. S. Staltsov & B. A. Kalin, Some problems of hydrogen in reactor structural
 426 materials: A review, Inorg. Mater. Appl. Res. 8 (2017) 643-650.
- 427 [10] A. Heinz, A. Haszler, C. Keidel, et al., Recent development in aluminium alloys for
 428 aerospace applications, Mater. Sci. Eng. A 280 (2000) 102-107.
- 429 [11] J. C. Williams, Jr. E. A Starke, Progress in structural materials for aerospace systems. Acta
 430 Mater. 51 (2003) 5775-5799.
- [12] K. Shimizu, H. Toda, K. Uesugi, A. Takeuchi, Local deformation and fracture behavior of
 high-strength aluminum alloys under hydrogen influence, Metall. Mater. Trans. A 51 (2020)
 1-19.
- 434 [13] H. Su, H. Toda, K. Shimizu, et al., Assessment of hydrogen embrittlement via image-based
 435 techniques in Al-Zn-Mg-Cu aluminum alloys, Acta Mater. 176 (2019) 96-108.
- 436 [14] H. Toda, T. Hidaka, M. Kobayashi, et al., Growth behavior of hydrogen micropores in
 437 aluminum alloys during high-temperature exposure, Acta Mater. 57 (2009) 2277-2290.
- 438 [15] S. K. Dwivedi, M. Vishwakarma, Hydrogen embrittlement in different materials: a review,
 439 Int. J. Hydrogen Energy 43 (2018) 21603-21616.
- 440 [16] J. T. Burns, S. Kim, R. P. Gangloff, Effect of corrosion severity on fatigue evolution in
 441 Al-Zn-Mg-Cu, Corros. Sci. 52 (2010) 498-508.
- 442 [17] K. Shimizu, H. Toda, H. Fujihara, et al., Hydrogen partitioning behavior and related
 443 hydrogen embrittlement in Al-Zn-Mg alloys, Eng. Fract. Mech. 216 (2019) 106503.
- 444 [18] M. L. Martin, B. P. Somerday, R. O. Ritchie, et al., Hydrogen-induced intergranular failure
 445 in nickel revisited, Acta Mater. 60 (2012) 2739-2745.
- 446 [19] Z. Zhang, G. Obasi, R. Morana, et al., Hydrogen assisted crack initiation and propagation in
 447 a nickel-based superalloy, Acta Mater. 113 (2016) 272-283.
- W. W. Gerberich, R. A. Oriani, M. J. Lji, et al., The necessity of both plasticity and
 brittleness in the fracture thresholds of iron, Philosophical Magazine A 63 (1991) 363-376.
- 450 [21] H. K. Birnbaum, P. Sofronis, Hydrogen-enhanced localized plasticity-a mechanism for
 451 hydrogen-related fracture, Mater. Sci. Eng. A 176 (1994) 191-202.
- 452 [22] T. Tsuru, K. Shimizu, M. Yamaguchi, et al., Hydrogen-accelerated spontaneous
 453 microcracking in high-strength aluminium alloys, Scientific Reports 10 (2020) 1-8.

- 454 [23] H. Toda, H. Oogo, K. Uesugi, M. Kobayashi, Roles of pre-existing hydrogen micropores on
 455 ductile fracture, Mater. Trans. 50 (2009) 2285-2290.
- 456 [24] M. S. Bhuiyan, et al., Influences of hydrogen on deformation and fracture behaviors of high
 457 Zn 7XXX aluminium alloys, Int. J. Fract. 200 (2016) 13-29.
- [25] I. Nikulin, A. Kipelova, S. Malopheyev, et al., Effect of second phase particles on grain
 refinement during equal-channel angular pressing of an Al-Mg-Mn alloy, Acta Mater. 60
 (2012) 487-497.
- 461 [26] R. Hu, T. Ogura, H. Tezuka, et al., Dispersoid formation and recrystallization behavior in an
 462 Al-Mg-Si-Mn alloy, J. Mater. Sci. Technol. 26 (2010) 237-243.
- 463 [27] Y. B. Kim, Y. H. Chung, K. K. Cho, et al., Recrystallization behavior of
 464 Al-2Li-3Mg-1Cu-0.12Zr-XMn (X= 0, 0.5) alloy, Scr. Mater. 36 (1997).
- 465 [28] M. Itakura, M. Yamguchi, K. Ebihara, K. Matsuda, H. Toda, First-principles calculations of
 466 hydrogen trapping energy at screw and edge dislocations of aluminum [In preparation].
- 467 [29] T. Tsuru, M. Yamaguchi, K. Ebihara, et al., First-principles study of hydrogen segregation at
 468 the MgZn2 precipitate in Al-Mg-Zn alloys, Comput. Mater. Sci. 148 (2018) 301-306.
- [30] L. Ismer, M. S. Park, A. Janotti, et al., Interactions between hydrogen impurities and
 vacancies in Mg and Al: A comparative analysis based on density functional theory, Phys.
 Rev. B 81 (2010) 139902.
- T. Tsuru, M. Yamaguchi, K. Ebihara, et al., First-principles study of hydrogen segregation at
 the MgZn2 precipitate in Al-Mg-Zn alloys, Comput. Mater. Sci. 148 (2018) 301-306.
- 474 [32] M. Yamaguchi, T. Tsuru, K. Ebihara, et al., Surface energy reduction by dissociative
 475 hydrogen adsorption on inner surface of pore in aluminum, J. Jpn. Inst. Light Met. 68 (2018)
 476 588-595.
- 477 [33] D. S. Park, S. W. Nam, Effects of manganese dispersoid on the mechanical properties in
 478 Al-Zn-Mg alloys, J. Mater. Sci. 30 (1995) 1313-1320.
- 479 [34] D. H. Lee, J. H. Park, S. W. Nam, Enhancement of mechanical properties of Al-Mg-Si alloys
 480 by means of manganese dispersoids, Mater. Sci. Tech. 15 (1999) 450-455.
- 481 [35] Y. Liang, P. Sofronis, N. Aravas, On the effect of hydrogen on plastic instabilities in metals,
 482 Acta Mater. 51(2003) 2717-2730.
- 483 [36] M. Dadfarnia, A. Nagao, S. Wang, et al., Recent advances on hydrogen embrittlement of

- 484 structural materials, Int. J. Fracture 196 (2015) 223-243.
- [37] I. M. Robertson, P. Sofronis, A. Nagao, et al., Hydrogen embrittlement understood, Metall.
 Mater. Trans. A 46 (2015) 2323-2341.
- 487 [38] M. J. Hÿtch, E. Snoeck, R. Kilaas, Quantitative measurement of displacement and strain
 488 fields from HREM micrographs, Ultramicroscopy 74 (1998) 131-146.
- [39] M. L. Martin, M. Dadfarnia, A. Nagao, et al., Enumeration of the hydrogen-enhanced
 localized plasticity mechanism for hydrogen embrittlement in structural materials, Acta
 Mater. 165 (2019) 734-750.
- 492 [40] Q. G. Wang, C. H. Caceres, J. R. Griffiths, Damage by eutectic particle cracking in
 493 aluminum casting alloys A356/357, Metall. Mater. Trans. A 34 (2003) 2901-2912.
- 494 Acknowledgments
- 495 This work was supported by JST, CREST Grant Number JPMJCR 1995 Japan and JSPS
- 496 KAKENH Grant Number JP21H04624. The synchrotron radiation experiments were performed at
- 497 SPring-8 with the approval of Japan Synchrotron Radiation Research Institute through proposal
- 498 number 2020A1796/1084.

499 Additional information

500 The authors declare no competing financial interests. Correspondence and requests for 501 materials should be addressed to Prof. Toda.

502 **Data availability**

- 503 The data that support the findings of this study are available from the corresponding author 504 upon request.
- 505 Figure Captions

506 **Figure 1 Excellent trapping energies of IMC particles to hydrogen**. The trapping energies of 507 various intermetallic compound particles to hydrogen were calculated by first-principles density 508 functional theory (DFT). The hydrogen trapping energies of various defect structures, such as 509 dislocations [28], grain boundaries [29], vacancies [30], precipitates [22,31], and micro pores [32], were derived from references. Al₁₁Mn₃Zn₂ shows advantages in hydrogen trapping compared to
other IMC particles and defect structures.

512 Figure 2 Superior HE resistance results from introducing IMC particles. a, 513 Stress-displacement curves for standard-state, Sc-added and Mn-added Al alloys after 514 low-hydrogen (LH) and high-hydrogen (HH) treatments, Mn-added Al alloys exhibited a superior HE resistance without sacrificing the ductility after HH treatment. b, Fracture morphology of 515 516 standard-state and Mn-added Al alloys after HH treatment, suppressing hydrogen-induced QCF 517 and IGF by introducing IMC particles was achieved. c, Fractional areas of QCF and IGF for 518 standard-state, Nb-added, Mo-added, V-added, Ca-added, Y-added, Sr-added, Ni-added, Zr-added, 519 Fe-added, Cr-added, Ti-added, Co-added, Sc-added and Mn-added Al alloys under HH treatments. 520 d, The relationships between the added elements, maximum trapping energies and fractional areas 521 of QCF fracture are summarized, the higher trapping energy with hydrogen, the lower fractional 522 area of QCF, which indicates that QCF was effectively suppressed by introducing IMC particles, 523 especially the IMC particles formed by adding Mn.

Figure 3 Fracture surfaces after preliminary tensile tests of HH and LH specimens. a,
Standard state, Sc-added and Mn-added Al alloys after HH treatments. b, Standard state, Sc-added
and Mn-added Al alloys after LH treatments.

527 Figure 4 Hydrogen redistribution driven by IMC particles. a, b, Distribution and crystal structure of MgZn₂ precipitates and Al₁₁Mn₃Zn₂ particles in Mn-added Al alloys observed by 528 529 HRTEM. c, 3D perspective views of IMC particles and micro pores obtained by high-resolution 530 X-ray tomography in Zr-added and Mn-added Al alloys. d, Hydrogen trapping occupancy before 531 and after adding Mn/Zr elements derived from thermodynamic equilibrium condition and 532 first-principles calculations, the formed Al₁₁Mn₃Zn₂ particles after Mn addition caused hydrogen 533 occupancy of the η_2 interfaces and grain boundaries to decrease by approximately 5 orders of 534 magnitude compared to that of the standard-state Al alloys.

- 535 Figure 5 APT results. a, Mn-added Al alloys. b, Standard-state Al alloys.
- 536 Figure 6 EBSD results. a, Standard-state Al alloys. b, Mn-added Al alloys.

537 Figure 7 Damage behavior of Al₁₁Mn₃Zn₂ particles and hydrogen partitioning behavior

- 538 during deformation after HH treatment. a-d, CT images of the Mn-added Al alloys at each
- 539 nominal strain on a x-z virtual cross section. e, f, Corresponding to the partial enlarged view in a

and **b**. **g**, **h**, 3D imaging around crack tip in Zernike imaging in Mn-added Al alloys. **i**, Hydrogen partitioning calculations for the regions ahead of hydrogen-induced quasi-cleavage crack tip and near the crack ridge in Mn-added Al alloys under the in-situ tensile strain of 3.6%. **j**, Hydrogen partitioning behavior among lattice, vacancies, dislocations, grain boundaries, IMC particles, η_2 precipitates and micro pores in Mn-added Al alloys before and after loading.

545 Figure 8 Micro-deformation mechanisms of Al₁₁Mn₃Zn₂ particles under HH and LH 546 conditions. a, Phase, KAM and TEM images of Mn-added Al alloys before tensile deformation, 547 TEM image was taken from 011 crystal zone axis of Al matrix. b, Phase, KAM and TEM images 548 taken from the crack tip region of HH-treated Mn-added Al alloys after tensile failure, strain 549 concentration around $Al_{11}Mn_3Zn_2$ particles mainly resulted from dislocations piled-up. c, Phase, 550 KAM and TEM images taken from the crack tip region of LH-treated Mn-added Al alloys after 551 tensile failure. d, KAM distributions in Al matrix and Al₁₁Mn₃Zn₂ particles of HH/LH-treated 552 Mn-added Al alloys before and after tensile failure. The strain distributions around $Al_{11}Mn_3Zn_2$ 553 and in Al matrix remained almost unaffected when changing treatment conditions from LH to HH. 554 Figure 9 Suppression of hydrogen-induced dislocations piled-up at grain boundaries by 555 Al₁₁Mn₃Zn₂. a, e, Microstructures near the fracture surfaces of Mn-added and standard-state Al 556 alloys treated by HH after tensile failure, respectively. The cracks mainly propagated along the 557 grain interiors in Mn-added Al alloys, while they mainly propagated along the grain boundaries in 558 standard-state Al alloys. b, TEM BF image cut from area b in a. c, d, TEM BF and DF images corresponding to the area c in b. f, TEM BF image cut from area f in e. g, h, TEM BF and DF 559 560 images corresponding to the area g in f. The piled-up dislocations at grain boundaries were 561 significantly reduced during tensile failure after adding Mn in standard-state Al alloys, indicating 562 that hydrogen-induced strain localization at grain boundaries was suppressed by Al₁₁Mn₃Zn₂ 563 particles.

Figure 10 Suppression of hydrogen-induced strain localization at η_2 phase interfaces by Al₁₁Mn₃Zn₂. a, h, Microstructures near the fracture surfaces of Mn-added and standard-state Al alloys treated by HH after tensile failure, respectively. b, TEM BF image cut from area b in a. c, d HRTEM images corresponding to the area c and d in b. e, f, Corresponding strain maps of geometric phase analysis (GPA) (in-plane rigid-body rotation, ω_{xy}) for HRTEM images c and d,

569 respectively. g, Inverse FFT and FFT patterns corresponding to HRTEM image c, inverse FFT pattern derives from $(\overline{111})$ reflection. i, TEM BF image cut from area i in h. j, k, HRTEM images 570 corresponding to the area **j** and **k** in **i**. **l**, **m**, Corresponding strain maps of geometric phase analysis 571 572 (GPA) (in-plane rigid-body rotation, ω_{xy}) for HRTEM images j and k, respectively. n, inverse 573 FFT and FFT patterns corresponding to HRTEM image j. Strain maps were plotted with respect to an internal reference lattice by $g_1 = (11\overline{1})_{matrix}$ and $g_2 = (200)_{matrix}$. Dislocations mainly 574 575 bypassed η_2 precipitates in Mn-added and standard-state Al alloys treated by HH. Density of edge 576 dislocations around the η_2 phase interfaces in Mn-added Al alloys is significantly lower than that 577 in standard-state Al alloys after tensile failure, moreover, the results of the strain distributions 578 originating from strain maps are also similar to that. The strain localization at η_2 phase interfaces 579 was effectively inhibited during tensile failure after adding Mn in standard-state Al alloys.

580 Extended Data Figure S1 Schematic diagrams of the thermomechanical treatment process. a,
581 High-hydrogen treatment. b, Low-hydrogen treatment.

582 Extended Data Figure S2 Desorption curves of the Mn-added Al alloys after HH and LH 583 treatments.

584 Extended Data Figure S3 Experimental setup for the imaging X-ray microtomography. a,
585 Schematic diagram. b, A pneumatic material test rig specially designed for X-ray
586 micro-tomography at the synchrotron radiation facility.

587 Extended Data Figure S4 Fracture surfaces after preliminary tensile tests of HH specimens.

Nb-added, Mo-added, V-added, Ca-added, Y-added, Sr-added, Ni-added, Zr-added, Fe-added,
Cr-added, Ti-added and Co-added Al alloys after HH treatments.

590 Extended Data Figure S5 TEM results. a, Standard-state Al alloys. b, Corresponding to the 591 partial area b in a. c, Mn-added Al alloys. d, Corresponding to the partial area d in c.

592 Extended Data Figure S6 Hydrogen partitioning behavior in HH-treated Mn-added Al alloys 593 during deformation. a, Hydrogen partitioning calculations for the region ahead of 594 hydrogen-induced quasi-cleavage crack tip under the in-situ tensile strain of 3.6%. b, Trapped 595 hydrogen contents among lattice, vacancies, dislocations, grain boundaries, IMC particles, η_2 596 precipitates and micro pores before and after loading. 597 Extended Data Figure S7 Suppression of hydrogen-induced cracks by Al₁₁Mn₃Zn₂. a, 598 Microstructures near the fracture surfaces of Mn-added Al alloys treated by HH after tensile 599 failure. The cracks mainly propagated along the grain interiors in Mn-added Al alloys. b, 600 Corresponding to the partial magnification images of the area 1 and 2 in a. c, d, Corresponding to 601 the partial magnification and mapping images of the area c and d in b, the IMC-rich region can 602 effectively blunt and suppress hydrogen-induced cracks.

603 Extended Data Figure S8 3D Mapping of equivalent plastic strain, vacancy concentrations

- and dislocation densities in the Mn-added Al alloys at load strain 0.0 to 3.6% on a x-z virtual
- 605 section. a, Equivalent plastic strain. b, Vacancy concentration. c, Geometrically necessary
- 606 dislocations (GND). **d**, Statistically stored dislocations (SSD).
- 607
- 608
- 609
- 610
- 611