New Interpretation to Disputed Matter of 300 to 400 Degrees Celsius Peak in Hydrogen-detecting Thermal Desorption Spectrum and Potentiality of Hydrogen Generation through this Phenomenon

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This paper provides an answer to the ever-lasting discussion concerning the interpretation of the hydrogen-related peak at 300 to 400°C in hydrogen-detecting Thermal Desorption Spectrum which often appears in a hydrogen-introduced steel sample, particularly when it has undergone an outdoor exposure or wet/dry cyclic accelerated-corrosion test and/or by acid immersion. Up to now, the focus of the discussion of the peak has been held with respect to whether the reaction is caused by diffusible hydrogen or not. However, the peak does not correspond either to a typical diffusible hydrogen peak at around 200°C or to a typical non-diffusible peak at over 400°C. Rather, the 300 to 400°C peak in TDS comes from the formation of hydrogen through a rust-related chemical reaction where iron (III) oxide-hydroxide (oxyhydroxide) of goethite “FeOOH” reacts into iron (III) oxide of hematite “Fe₂O₃”. It should be noted that the origin of FeOOH comes from rust which is too small to detect and which, even if you are sure that the surface is perfectly smooth and cleaned off by hand-polishing, is not removed due to deeply-rooted and small-diameter rust particles, and cannot be detected by the naked eye. In addition, following on from this new interpretation of the controversial spectrum, this phenomenon is re-interpreted as a new process to produce hydrogen gas, and the potentiality and the mechanism are also discussed.

KEY WORDS: hydrogen; thermal desorption spectrum; TDS; gas chromatograph; rust-based TDS signal; 300 degree celsius to 400 degree celsius peak; diffusible hydrogen; non-diffusible hydrogen; rust; iron oxyhydroxide; iron oxide-hydroxide; goethite; FeOOH; hematite; Fe₂O₃; cyclic corrosion test; hydrogen generation.

1. Introduction

This paper targets the interpretation of the 300 to 400°C peak in Thermal Desorption Spectroscopy (referred as “TDS” below) when measured the hydrogen content in steel.1,2)

Hydrogen is still one of important research targets in materials science. Hydrogen embrittlement remains as an unsolved problem. In hydrogen research, the techniques to measure hydrogen have been studied, and a glycerin-displacement-based volumetric method in Japan and a Mercury-displacement-based volumetric method have already been developed, and have been widely used since the early 1970s.3-6) As it is now, Thermal Desorption Spectroscopy3) has come to be commonly used to measure hydrogen content. This paper is concerned with an answer to the puzzle of the 300 to 400°C peak in the TDS spectrum.

Since TDS was introduced to hydrogen research, various steel samples have been evaluated, especially a hydrogen-introduced steel samples. Although it rarely appears in cathodically-charged samples, the peak in TDS is sometimes observed at 300 to 400°C,7-9) especially in corroded samples through outdoor exposure or wet/dry cyclic accelerated-corrosion test. In the following report, it should be noted that the TDS is based on the signals measured by using Gas Chromatograph (referred as “GC” below, and “GC-TDS” is also used below for GC-based TDS), unless otherwise stated.

Figure 1 is an example of the 300 to 400°C peak, of which there has been no widely-agreed interpretation. This sample is from a corroded sample with hand-polished “clear” surface after corroded in wet/dry cyclic accelerated-corrosion test of SAE-J2334,10) in which 80-cycle (80-day) is equivalent to a 9-year on-vehicle corrosion on a public roadway in melting-snow-agent spraying area for an automobile outer (coated) panel, e.g. for door, hood, trunk and so on.
Figure 2 is a typical TDS spectrum of “a normal steel sample”, i.e., as-received steel sheet sample which is annealed without any plating or coating. The area from room temperature (referred as “RT” below) to around 200°C has no peaks, and the area at over 400°C has hydrogen signals. In contrast, Fig. 3 is another typical example of cathodically-charged steel. Two kinds of hydrogen peaks are observed at both RT to 200°C and at around over 400°C. It is evident that the first peak is a result of hydrogen uptake by cathodic-charging by comparison with Fig. 2, and the peak has been regarded as diffusible hydrogen and is believed to be related to hydrogen embrittlement. In contrast, the spectrum at over 400 to 500°C used to be regarded as non-diffusible hydrogen and to be interpreted as independent of hydrogen embrittlement, but the signal is now often regarded only as background at high temperature range in GC-TDS and has been considered to be dependent on each GC apparatus itself and/or on each analyzed sample.

The issue in this paper is shown in Fig. 1. The signal of the spectrum at 300 to 400°C, strictly speaking, does not correspond either to the “diffusible hydrogen”-based peak at RT to 200°C, or to the “high-temperature background (or non-diffusible hydrogen)” peak at over 400°C. Although some people have proposed various ways of thinking about the unexplained spectrum, any comprehensive theories have not been widely acknowledged. Therefore, the objective of this paper is to present a new interpretation of the 300 to 400°C signal in TDS. Furthermore, by looking at this phenomenon from a slightly different angle, a new potential for hydrogen generation is advocated through the method of heat-treating corroded steel and/or rust at 300 to 400°C.

2. Experimental Procedure and Samples

First, the technical terms and wording are defined as below. In this paper, three peaks in TDS are topics at issue. Peaks and strong signal areas are categorized as three types. For the sake of convenience, they are called “RT to 200°C”, “300 to 400°C” and “over 400°C” by using temperature-based terms, because these three categories have strong signals mainly at each temperature range, although some signals sometimes overflow beyond the lower limit and/or the upper limit of each temperature-based wording. In addition, it should be noted that these temperatures do not have any physical meaning. The temperature to appear signals in the TDS spectrum depends on thickness of a sample and the rate of elevating temperature. Therefore, these classifications are expediential, but will be used throughout this paper. As shown in Fig. 1, the important point is that the peak at “300 to 400°C” in TDS has not been yet interpreted correctly. Another point is the wording to describe “TDS spectrum” and “raw chart”. The former is used throughout the paper, and the latter is used only in the Discussion section. As shown above in Figs. 1 to 3, the former is the spectrum to plot hydrogen desorption rate as a parameter of temperature. As shown far below in Fig. 14, the latter is the raw data to show thermal conductivity as a parameter of time at gas separation in TDS measurement. The hydrogen content corresponds to the peak at around 0.7 sec in our GC’s gas separation system, and the value of integral around the peak is calibrated to the hydrogen desorption rate and each data is plotted as a parameter of sampled temperature in “TDS spectrum”.

Experiments in this paper are based on a 2-step approach. The first step is TDS measurement of corroded samples roughly to learn the origin of the hydrogen signal at 300 to 400°C, and the second step is the investigation to understand the mechanism and to search for the potentiality to generate hydrogen at 300 to 400°C by re-interpreting this phenomenon as the reaction to emit hydrogen through heat-treatment at 300 to 400°C.

2.1. First Step Experiments

In the first step, steel samples were taken from 2 mm-thick “ordinary” sheet steel (cold-rolled and then annealed without any coatings), and were cut into samples with the dimensions of 2 mm thick, 50 to 70 mm long and 12 mm wide. The chemical composition is typical mild steel (0.02%carbon-based) without any intentional additions of alloy elements. The size of samples is smaller than regularly-tested ones in a wet/dry accelerated corrosion test, because the size is limited by the inner diameter of the sample holder in the GC-TDS. The samples were put in a cyclic wet/dry test chamber to control humidity, temperature and test duration. As shown in Fig. 4, the samples were corroded according to the SAE-J2334 test procedure, which is composed of weekday sessions and holiday sessions: the former is a three-stage wet/dry cycle of salt-dipping at 25°C for 15 min, a dry-period in 50%RH (Relative Humidity: referred as “RH” below) at 60°C for 17.75 h and a humid-period at 100%RH at 50°C for 6 h, and the latter session is to keep at 50%RH at 60°C.

The test samples and conditions are listed on Table 1. The test conditions are (1) as-received (no corroded sample for reference), (2) to (5) for each stage in the 10-day cycle and (6) a heavily-corroded sample to proceed rust-layer-delaminated stage (approximately 80 to 100-cycle). Only the (2) to (4) samples of 10-day at SAE-J2334 corrosion test were kept in liquid nitrogen as soon as it was picked out from the cyclic wet/dry test chamber. Hydrogen in these samples was measured after the rust and corrosion products were removed as thoroughly as possible. One condition of “(5) after humid” in 10-day corroded sample in SAE-J2334 test was used for hydrogen measurement of a rust-covered sample for comparison. They were also kept in liquid nitrogen, but the TDS measurement was carried out after the temperature returned to nearly RT. The “(6) heavily-corroded” sample was kept at room temperature inside the acceleration-corrosion test chamber (switched-off) until the hydrogen measurement was started, which is equivalent to keeping at room temperature. Figure 5 shows the appearance after having been corroded in the SAE-J2334 corrosion test: (a) a typical example of 10-day corroded sample and (b) samples corroded for over-80days. Since the latter is rust-layer-delaminated and is too fragile to take out as a sheet-formed sample, the rust was powderized and then was used for TDS measurement.

In regard to hand-polishing, a corroded sample was picked out from the liquid nitrogen and then the rust and corrosion products were carefully removed with abrasive papers by hand-polishing being held with a low-temperature-proof glove, and when the sample had been unavoidably heated up
Fig. 1. Typical example of the 300 to 400°C peak in TDS spectrum, which is based on a corroded steel sample with hand-polished surface after wet/dry cyclic accelerated corrosion test.

Fig. 2. TDS spectrum of an as-received steel sheet sample which is annealed without any plating or coating.

Fig. 3. TDS spectrum of cathodically-charged steel sample.

Fig. 4. SAE-J2334 test procedure of wet/dry-cyclic accelerated corrosion test.

Fig. 5. Typical appearance of as-corroded samples in SAE-J2334’s wet/dry cyclic accelerating corrosion test of (a) after 10-day (10-cycle) exposure and (b) over 80-day (80-cycle). (a) As-corroded after 10-day (10-cycle) exposure in SAE-J2334. (b) As-corroded after over 80-day (80-cycle) exposure in SAE-J2334.

Fig. 6. Typical appearance of (a) as-corroded sample and (b) as-rust-removed sample, whose surface looked metallic-sheeny and the spots could not be seen, due to frost, during polishing and/or just after polishing for removal of rust and corrosion product. (a) As-corroded after picked from wet/dry cyclic corrosion chamber. (b) As-rust-removed (as thoroughly as possible) after hand-polishing while cooled down below zero°C.

Fig. 7. GC-TDS spectra of “rust-removed” 10-day exposure samples in SAE-J2334’s wet/dry cyclic accelerated corrosion test: (a) after salt dip, (b) after dry stage and (c) after humid stage, which correspond to (2), (3) and (4) in Table 1, respectively. (a) After Salt dip (25°C). (b) After Dry stage (60°C, 50%RH). (c) After Humid stage (50°C, 100%RH).
to nearly zero°C by friction (rubbing), it was immediately returned to the liquid nitrogen to cool down. The polishing routine was repeated until the surface returned to a metallic sheen as thoroughly as possible. However, samples were so cold that the surface was often covered with frost, no matter how much care was paid to minimize it even by polishing in a dry condition. Figure 6 shows typical surfaces of (a) a corroded sample and (b) a rust-removed sample. In Fig. 6(b), small black spots are now confirmed by sight, but during polishing and/or just after polishing, the surface looked metallic-sheeny and the spots could not be seen, due to frost. The exceptions were rust-free as-received samples ((1) in Table 1), heavily-corroded samples ((6) in Table 1) and one of the 10-day samples (after humid stage: (5) in Table 1).

As for the test condition of TDS, GC-based TDS was used for all the tests in this paper. Each parameter in GC-TDS is listed on Table 2. Gas separation was based on molecular-sieve 5A (MS-5A) and hydrogen was detected by the thermal conductivity detector (TCD). The elevating temperature rate was 200°C/h and the sampling time was every 2.5 min. Hydrogen was evaluated as the mass fraction of hydrogen in the steel sample by “mass of desorbed hydrogen/sample weight measured just before TDS”. It should be noted that the carrier gas at the GC-TDS was high-purity Argon gas, and that any contaminated components, especially oxygen, did not enter the test chamber through the Argon gas.

### 2.2. Second Step Experiments

Advanced experiments were intended to discover the fundamental mechanism for signals to appear at 300 to 400°C in the TDS spectrum, and also to search for a new mechanism for hydrogen generation by re-interpreting the hydrogen signals at 300 to 400°C in TDS as a reaction which results in the emission of hydrogen through the heat-treatment at 300 to 400°C.

Although the results in the first-step experiment will be reported later, we would say at this stage that this reaction is related to the unavoidably-remaining rust. Therefore, commercially-available test chemicals of three kinds of rust were prepared for GC-TDS: (1) FeOOH (Goethite), (2) Fe₂O₃ (Hematite) and (3) Fe₃O₄ (Magnetite), whose catalog-based purities are over 95%, 99–101% and over 95%, respectively. These powdered materials were evaluated by GC-TDS, and one of them was evaluated by X-ray Diffraction (referred as “XRD” below) to know the phase change before/after TDS measurement. Also, the one of rust

### 3. Results

#### 3.1. Rust-removed 10-day Exposed Samples in SAE-J2334 Test at First-step Experiment

Figure 7 shows GC-TDS spectra of (a) after salt dipping, (b) after dry stage and (c) humid stage of 10-day exposed samples in SAE-J2334 test, which correspond to (2) to (4) in Table 1. For your reference, Fig. 1, which has already been mentioned above as a typical example of 300 to 400°C peaks, shows 30-day (after humid stage) corroded sample in SAE-J2334 test. Each sample has two peak-based 300 to 400°C signals in whose region hydrogen is (a) 0.39 ppm, (b) 0.19 ppm and (c) 0.31 ppm, respectively. In terms of the signal at RT to 200°C, which is thought to be diffusible hydrogen, the 10-day samples in Fig. 7 have only tiny signals. This means that the samples do not have any diffusible hydrogen. In terms of the signal at over 450°C, it is also at a tiny level. This experimental fact indicates two things: (1) The amount of hydrogen which is introduced by cathodic-charging in Fig. 3 is larger than that by the corrosion-based method, and (2) the 300 to 400°C signal is dependent on something which is related to the corrosion-based hydrogen reaction, but not to the cathodic-charging. The former fact of (1) is also supported in the literature.12) Regarding the latter fact of (2), as one possibility, it can be expected that Fe²⁺-related reaction may be linked, because cathodic-charging is based only on the reaction of “H⁺ + e− → H” to enter hydrogen into steel (electron is provided through potentiostat), but corrosion-based reaction requires both

| Table 1. List of test samples. |
|-----------------------------|
| rust removal | SAE-J2334 |
| (1) zero-day (reference) | – |
| (2) GC-TDS | (3) GC-TDS |
| (4) GC-TDS | --› kept in Liquid N₂ until TDS measurement |
| (5) GC-TDS | --› kept in RT until TDS measurement |
| (6) >80days | (No) |
| heavy-corrosion to rust-layer-delaminated | --› kept in Liquid N₂ until TDS measurement |

| Table 2. TDS’s test conditions. |
|-------------------------------|
| GC-TDS                         |
| Gas Chromatography Thermal Desorption Spectroscopy |
| Separation         | MS-5A column |
| Detector             | TCD (Thermal Conductivity Detector) |
| Flow gas             | high-purity Ar |
| Temperature          | RT to 600°C |
| Elevating Temp rate  | 200°C/h      |
| Sampling             | every 2.5 min |
| Sample size          | w12 mm × L50 mm to 70 mm × t2 mm |
“Fe → Fe²⁺ + 2e⁻” for anodic reaction and “H⁺ + e⁻ → H” for cathodic reaction, and Fe²⁺ ion is easily changed into rust and corrosion products.

As mentioned above, when hydrogen in the wet/dry cyclic corrosion test samples was measured by TDS, rust and corrosion products were cleaned off in advance as thoroughly as possible while the test sample was cold enough not to get out as hydrogen gas, in order to know hydrogen content correctly. Figure 8 shows the surface appearance at pre-and post-TDS measurement. After TDS measurement, blackish and grayish stains appear and seem to have seeped and spread out. This fact suggests that deeply-rooted rust is formed in corroded samples and cannot be removed completely even by a careful hand-polishing, and that these stains seeps from invisible deeply-rooted spot-like corrosion during TDS measurement. In the next section, the authors focus on rust-related phenomenon.

3.2. As-corroded Samples (10-day and Over 80-day) in SAE-J2334 Test at First-step Experiment

Figure 9 shows the GC-TDS spectrum of “rust-covered” as-corroded samples (10-day/after humid stage/(5) in Table 1) in SAE-J2334 test, and the appearance before and after TDS measurement. It should be noted that this sample is “as-corroded” without any hand-polishing. The appearance (surface color) changed from reddish brown at pre-TDS (as-corroded) to blackish brown at post-TDS. This suggests that rust is directly related to the signals at 300 to 400°C in TDS spectrum, and that the rust on the surface may be changed into a different phase. Figure 10 shows GC-TDS spectrum of over-80day sample in SAE-J2334 test ((6) in Table 1), and also the appearance of “before TDS”. As shown in Fig. 5(b), the over-80day sample was corroded heavily into a multi-layered delaminated form, and so the sample for GC-TDS was broken into a powder-like form when picked out from the comb-like holder. These data in Figs. 9 and 10 mean that rust is the origin of hydrogen at 300 to 400°C. The hydrogen at 300 to 400°C is 7.47 ppm in Fig. 9 and 10.45 ppm in Fig. 10.

However, a careful attention has to be paid to interpret these phenomena. Authors point out some interesting and notable points in Figs. 9 and 10 in comparison to Figs. 1 and 7: (1) the vertical axis in TDS spectra, (2) the definition of hydrogen content (concentration), (3) the one-peak and the two-peak at 300 to 400°C region in TDS spectrum.

The first topic is that the vertical axis in Figs. 9 and 10 are about 100 times higher than in Figs. 1, 3 and 7. Since the 300 to 400°C signal is based on unavoidably-remaining rust, it can be qualitatively explained that the amount of rust is much smaller inside rust-removed samples in Figs. 1 and 3 than rust-covered samples in Figs. 9 and 10. The second topic is that the definition of hydrogen content in Fig. 10 is different from the ones in any other examples, and it is related to the result at the above first topic in Fig. 10. The hydrogen content is evaluated as the following mathematical formulas of (1) to (3), because the denominator in the formula corresponds to the weight measured just before TDS analysis.

For a rust-removed or rust-free sample in Figs. 1–3 and 7;

*hydrogen / steel .......................... (1),

which is abbreviated to “H/S”, because the rust in rust-removed sample is a negligible small quantity.

For a rust-covered sample in Fig. 9;

*hydrogen / (steel + rust) ..................... (2),

which is abbreviated to “H/(S + R)”.

For a rust sample in Fig. 10 and in yet-undescribed Figs. 11 and 13;

*hydrogen / rust ........................... (3),

which is abbreviated to “H/R”.

The hydrogen contents in Figs. 9 and 10 are almost the same, but the physical meaning is completely different. Specifically, “HFig. 9/(SFig. 9 + RFig. 9)” in Fig. 9 looks the same as “HFig. 10/RFig. 10” in Fig. 10. As mentioned above, the 300 to 400 signal is not related to steel matrix but rather to rust itself, and therefore, it has to be described as “HFig. 9/RFig. 9” in Fig. 9 for a right interpretation to understand hydrogen generation at 300 to 400°C, that is, “HFig. 10/RFig. 10” = “HFig. 9/(SFig. 9 + RFig. 9)” < “HFig. 9/RFig. 9” in Fig. 9. It should be noted that SFig. 9 is a larger value. It suggests that the co-existence of steel and rust emits more hydrogen at 300 to 400°C than only rust. This topic will be mentioned later again in the following discussion section.

The last topic is that only Fig. 9 shows one-peak TDS spectrum, although other spectra in Figs. 1, 7 and 10 consist of two peaks at 300°C and 400°C. Also, each spectrum in Figs. 7(a) to 7(c) is two-peak-based signal but slightly different each other. Although authors can make mention of these phenomena, we do not have any supporting experimental data to explain it logically and it is an on-going issue for us.

From the results at this section and at the former section, the signal at 300 to 400°C in TDS is related to hydrogen generation by rust on the surface of the corroded sample. When we return to the original objective of “what is the signal at 300 to 400°C in TDS spectrum, especially of corroded steel (rust-removed) sample?”), deep-rooted rusts unavoidably remain as small-dotted stains in corroded samples and the remaining rust will be the origin of the hydrogen signal at 300 to 400°C; no matter how thoroughly the rust is removed by hand-polishing.

In the next section, the authors focus on what kind of rust (iron oxide) plays an important role in hydrogen emission, and discuss the potential to generate hydrogen by using rust, that is, by proposing the phenomenon of the hydrogen signal at 300 to 400°C in the TDS spectrum as a new method to make hydrogen.

3.3. TDS Spectra of Rust, and Change of Appearance before/after TDS at Second-step Experiment

In the following section, as in the above two results sections, it should be noted that the technical term of “TDS spectrum at 300 to 400°C” means that strong signals exist mainly at 300 to 400°C and some signals overflow at below 300°C and/or at over 400°C.

Three kinds of commercially-available rust were prepared for GC-TDS: (1) Goethite: FeOOH, (2) Hematite: Fe₂O₃, and (3) Magnetite: Fe₃O₄. The authors could not prepare wustite: FeO, because it is not stable enough to remain as wustite. Figure 11 shows these three GC-TDS spectra and...
Fig. 9. TDS spectra of “as-corroded (rust-covered)” 10-day exposure sample in SAE-J2334’s wet-dry cyclic accelerated corrosion test, and the appearance of before and after TDS measurements, which corresponds to (5) in Table 1.

Fig. 10. TDS spectra of over-80days exposure sample (mostly powdered rust) in SAE-J2334’s wet-dry cyclic accelerated corrosion test, and the appearance of before and after TDS measurements, which corresponds to (6) in Table 1.

Fig. 11. (a) TDS spectra of FeOOH, Fe₃O₄ and Fe₂O₃, and “before/after TDS measurement” appearance of (b) FeOOH (Goethite), (c) Fe₃O₄ (Magnetite) and (d) Fe₂O₃ (Hematite).

Fig. 12. XRD patterns of FeOOH as (a) before and (b) after TDS measurement, and these appearance of (c) FeOOH (before TDS), (d) after TDS of FeOOH and (e) hematite Fe₂O₃ for reference. (a) FeOOH (before TDS). (b) After TDS of FeOOH. (c) “Before TDS” of FeOOH (Fe₂O₃·H₂O) Goethite. (d) “After TDS” of FeOOH. (e) Fe₂O₃ Hematite (For reference).

Fig. 13. TDS spectrum of 2nd-measured FeOOH (precisely speaking, starting material is phase-transformed Fe₂O₃).

Fig. 14. TDS raw charts of (a) the initial stage (just after a sample is set, and air still remains) and of (b) typical example after over-1-hour TDS measurement (air is automatically ejected during measurement). (a) Initial stage (just after a sample is set). (b) Typical example after over 1 hour TDS measurement (air has been ejected during measurement).

the appearance before and after TDS measurement. Goethite “FeOOH” (over 95% purity), Magnetite “Fe₃O₄” (over 95% purity) and Hematite “Fe₂O₃” (99–101% purity) have the signals at 300 to 400°C, and the total of hydrogen is 3.1 ppm, 0.69 ppm and 0.06 ppm, respectively. The experimental fact proves that the origin of hydrogen at this temperature range is the rust of Goethite “FeOOH”. Magnetite and Hematite are not related to the hydrogen generation. The
data could suggest that Magnetite “Fe₃O₄” has a small effect on hydrogen generation, but commercially-available rust unavoidably has impurities of other rusts (over 95% purity is only guaranteed) and it is estimated that Goethite “FeOOH” as an impurity may be related to a small amount of hydrogen by Magnetite “Fe₃O₄” as a commercially-available reagent. The other point is the color of “before TDS (as-received)” and “after TDS” in Fig. 11. Goethite “FeOOH” changed from yellowish brown into reddish brown after TDS measurement. Hematite “Fe₂O₃” and Magnetite “Fe₃O₄” did not change in color at “before” and “after” testing. “After TDS” of Goethite “FeOOH” shows the same color of “before/after TDS” of Hematite “Fe₂O₃”. Therefore, Goethite “FeOOH” was evaluated by XRD to compare the phase of rust “before” and “after TDS” measurement. As shown in Fig. 12, “before TDS” is composed mainly of Goethite “FeOOH” with a small portion of Magnetite “Fe₃O₄”, and “after TDS” consists of Hematite “Fe₂O₃”. The experiment data in Figs. 11 and 12 imply that Goethite “FeOOH” is the origin of the emission of hydrogen at 300 to 400°C in TDS spectrum.

Figure 13 shows the second-time measurement by measuring “TDS-measured” Goethite “FeOOH”. Judging from Figs. 11 and 12, the first-time measurement is based on the change from Goethite “FeOOH” to Hematite “Fe₂O₃”, and the second time is based on the measurement of the phase-transformed Hematite “Fe₂O₃”. After the first measurement was finished, the rust sample stayed inside the sample holder about a half day under Argon gas flow until it was cooled down to RT and waited for the second measurement. As shown in Fig. 13, the spectrum is completely flat without any signals, and the color did not change at all, although the appearance is not shown here.

The topic in this paper focuses on the controversial signals at 300 to 400°C, and our experiment has given an answer to explain the mechanism that hydrogen comes from deeply-rooted rust-like rust (unavoidably remaining tiny spots of rust) of Goethite “FeOOH”, no matter how thoroughly the rust on the corroded steel sample is removed by hand-polishing. Precisely speaking, when the rust (Goethite FeOOH) is heat-treated at 300 to 400°C under Argon flow in TDS, hydrogen is generated. In the following discussion, rust-based hydrogen generation is dealt with as a potential procedure to make hydrogen.

4. Discussion

We have already understood in this paper that hydrogen is generated when Goethite “FeOOH” is heat-treated in Argon flowing (O₂-free) at 300 to 400°C inside TDS. Specifically, two kinds of hydrogen generation phenomena have been reported. One is hydrogen generation by Goethite “FeOOH” in Fe-existing environment as the main topic, and the other is hydrogen generation only by commercially-available rust reagent of Goethite “FeOOH” in Fe-free condition as a comparative experiment. In this section, the mechanism and the chemical reaction are discussed below to give a reasonable explanation to the phenomena in Figs. 1, 7, 9–11 and 13. Especially, the quantity and the efficiency of the emission of hydrogen are also discussed, and the potential to make hydrogen is also reported by re-interpreting this phenomenon.

Judging from the graph in Fig. 11, 1 kg of Goethite “FeOOH” makes just only 3.1 mg of hydrogen on weight basis (when theoretically evaluated on molar basis, 0.2%H₂ generation for 1 mol of FeOOH). In addition, this reaction is not at all a direct decomposition reaction of H₂O, which requires a large amount of energy. However, the generation efficiency in Fig. 11 is rather small. When reviewed from a different viewpoint, even if it is at rather low efficiency, there may be high potentiality to produce hydrogen easily without spending money by using redundant heat and scrap (rust) which lie around in a steelmaking plant. You can see a lot of reddish scrap and energy-filled conditions, for example, steam, fire and heat-exchangers, especially at the upper-stream process of ironmaking, steelmaking and coke furnaces. The authors imagine that such phenomenon may have been already used unconsciously in steel companies. Emitted gas in upper stream of steel plants is collected into a gas holder and used as an energy source to create electricity and heat. It is known that the collected gas is composed mainly of CO gas with a small portion of about 10% hydrogen. Although nobody may have paid attention, some of the hydrogen may also come from heated iron ore (rust on surface) and heated rusted scrap at the upper-stream process. As another viewpoint, we want to point out the potentiality to improve the efficiency of hydrogen generation when Goethite “FeOOH” and steel co-exists. As explained from the viewpoint of the definition of hydrogen calculation in Eq. (2) in Result 3.2., the rust-covered case in Fig. 9 is at a higher efficiency to make hydrogen than the rust-based powderized case in Fig. 10, which may slightly include iron component, and the latter amount in Fig. 10 shows a higher value than commercially-available test chemical of Goethite “FeOOH” in Fig. 11 (Fe-free condition). The above facts are discussed below in terms of the mechanism of hydrogen generation.

At first, we deny the possibility of well-known Schikorr reaction of (4) below that Fe(OH)₂ is heat-treated at 200 to 400°C under an O₂-free condition into magnetite, water and hydrogen. In our targeting reaction, the starting material is Goethite “FeOOH” and the final material is Hematite “Fe₂O₃”. Therefore, our Hydrogen-generating reaction is completely different from Schikorr reaction. Hydrogen is generated by the phase transformation at 300 to 400°C in an O₂-free condition (Argon flowing as the carrier gas). The proposed chemical reaction is tentatively given below as (5). The contents of Fe and hydrogen need to be balanced on the left-hand side and the right-hand side, and the oxygen is balanced on the right-side by using (omit) oxygen molecule, as described in brackets below in (5). From a classical point of view, the reaction from FeOOH to Fe₂O₃ can be also regarded as dehydration reaction and is described below as (9). Furthermore, when iron exists in the reaction system of (9), it has been known as (10) that dehydration-based H₂O is reduced to H₂ by iron and the iron is changed to hematite Fe₂O₃. Also, a simple H₂O decomposition of (11) is estimated not to occur at 300 to 400°C, because the reaction needs as high temperature as 1 500 to 2 000°C.

\[
\text{(Schikorr reaction)} \quad 3\text{Fe(OH)}_2 \rightarrow \text{Fe}_3\text{O}_4 + 2\text{H}_2\text{O} + \text{H}_2 \quad \ldots (4)
\]
FeOOH → l/2Fe₂O₃ + l/2H₂ + (+1/4O₂) ........................ (5)

(molar ratio) 1 mol (rust) 1/2 mol (hydrogen) .......................... (6)

(expected hydrogen) 100 g 100/121/2 × 2 = 0.83 g ... (7)

(in reality in Fig. 11) 1 g (100 g) 3.1 μg (310 μg) ... (8)

(classically defined) FeOOH → l/2Fe₂O₃ + l/4H₂O ... (9)

(Fe-oxidation-based reaction) 2Fe + 3H₂O → Fe₂O₃ + 3H₂ .......................... (10)

(Simple H₂O decomposition) H₂O → H₂ + l/2O₂ ... (11)

If the above chemical reaction of (5) is correct, oxygen must be observed as well as hydrogen. However, oxygen is observed only as a background level in the GC-TDS raw chart. Figure 14(a) is a typical TDS raw chart of a hydrogen-introduced sample when started TDS measurement. The signal at about 0.7 min corresponds to hydrogen, and the ones at 1.2 min and at 2.1 min correspond to oxygen and nitrogen, respectively. It should be noted that air inevitably remains inside the GC-TDS system, because the sample has been set in the sample holder tube and Air-originated N₂ and nitrogen, respectively. It is measured at RT to 600 °C by sampling every 2.5 min at an elevating temperature rate of 200°C/h. As shown in Fig. 14(b), when the sampling at 300 to 400°C is carried out, air-based signals have reached to a background level, and such a low level of air-based signals stays intact to the end of measurement. Therefore, the above tentative chemical formula of (5) may look wrong, but the authors think that the reaction of (5) still remains as another possibility to give a logical explanation to hydrogen generation by Goethite “FeOOH” under Fe-free condition, because the Fe-coexisting phenomena in Figs. 1, 7 and 9 can be explained by a classical concept of (9) and (10), but Fe-free phenomena (below a detection limit of Fe in XRD) in Figs. 10 and 11 cannot be interpreted only by this classical viewpoint. As another possibility, even if O₂ gas emits in addition to H₂ gas, it can be thought that most of hydrogen and oxygen are combined together into water and that the remaining hydrogen is detected and the remaining oxygen may be combined with neighboring Hematite “Fe₂O₃”. Surely, the O₂ signal may be rarely confirmed in TDS raw graph, because iron rust often comprises a non-stoichiometric composition and the rust may soak up the O₂ into oxide. However, the above explanation still includes a difficult point to explain why O₂ peak does not appear strongly, and our theory still remains a matter of discussion. We have not yet had any supporting data to explain the gap between the theory and the reality.

Judging from the discussion so far, the hydrogen generation in Figs. 1, 4, 7, 9, 10 and 11 may be related to two possible reactions of the tentative chemical formulae of (5) and/or the classical concept of “(9) and (10)”, although the Fe-coexisting hydrogen phenomena in Figs. 1, 4, 7 and 9 can be explained (omit) by the above classical concept of (9) and (10). If we persist in this classical concept to the Fe-free hydrogen phenomena in Figs. 10 and 11, we have to consider that even an impurity level of iron below the detection limit in XRD can play an important role to the reaction of (10). However, it would be unlikely to occur it.

The next topic is a quantitative discussion of hydrogen generation by the rust of Goethite “FeOOH” without including iron component. Regarding the above-mentioned chemical reaction and calculated quantity of materials of (6) to (8), we can see that if the chemical formula of (5) is right, hydrogen has to generate 1/2 mol for 1 mol of FeOOH in a molar basis. “100 g of FeOOH” corresponds to about 1 mol (0.83 mol) of FeOOH by calculating as 100/(56+64+1), and 1/2 mol of generated Hydrogen corresponds to 0.83 g by calculating as 100/(56+64+1)/2×2. The weight-based efficiency is theoretically 8.3×10⁻⁶, approximately 0.01. However, in the real reaction of (8), 3.1 ppm of hydrogen gas is generated, and the weight-based efficiency is 3.1×10⁻⁶. The difference between theory-based calculation and the real measurement is about 4000-fold. Therefore, it can be thought that most of the hydrogen disappears and/or the efficiency is much lower than expected in the estimated reaction of (5) or “(9) and (10)”. The authors think that most of hydrogen in (5) is combined with oxygen into H₂O, or that most of H₂O by dehydrating from FeOOH in (9) still stay as H₂O and the quantity of hydrogen is limited by the Fe-oxidation-based reaction of (10). As mentioned in the context to compare with Figs. 9 to 11, more hydrogen is confirmed at a rust-related sample with co-existing steel component. It is thought that the classically-recognized Fe-oxidation-based reaction of (10) occurs after the dehydration reaction of (9).

Although the discussion remains in terms of a key mechanism, we can conclude that the origin of the hydrogen peak at 300 to 400°C in GC-TDS is based on the iron rust transformation from Goethite “FeOOH” to Hematite “Fe₂O₃” at 300 to 400°C under Ar gas flow (O₂-free). Even the remaining stain-like rust, which is difficult to remove completely by hand-polishing, becomes an origin of hydrogen. There is a high possibility that H₂ is directly generated by the transformation from Goethite “FeOOH” to Hematite “Fe₂O₃” without any Fe component, in addition to the classically-accepted reaction that dehydrated FeOOH-based H₂O is changed to H₂ under co-existing Fe component. Some people may say that commercially-available FeOOH in Fig. 11 and heavily-corroded sample include a small amount of iron component, but we have already checked that the former sample does not show any Fe-based signal at XRD in Fig. 12(a). Therefore, it is reasonable that the reaction to emit hydrogen possibly exists under the condition without any Fe component. As mentioned above, (omit), we are only saying about two possible estimated reactions of (4) or “(9) and (10)”. The detailed mechanism in Fe-free environment is an issue in the future.

In addition, the authors want to point out that some similar but inherently-different ideas have been reported, and pick out one example below from the registered Japan patents. In similar reports of rust-related hydrogen generation, in many cases, iron-related materials tend to be
regarded as catalysts for the hydrogen generation. The patent has mentioned a hydrogen emission method by reacting a waste of foamed polystyrene and water under the existence of “iron oxyhydroxide (FeOOH)” or “iron oxide” as a catalyst. Although hydrogen is generated, they say that the origin of hydrogen is based on a waste of foamed polystyrene and water (including chemisorbed water, physisorbed water or crystallization water), but not iron-related material itself. In contrast, our report intrinsically includes a new concept in terms of “FeOOH” as an origin of hydrogen generation through the heat-treatment of 300 to 400°C peak in TDS spectrum. Our interpretation to this phenomenon gives an answer to the ever-lasting 300 to 400°C mysterious peak in TDS spectrum, and also suggests a new concept of hydrogen generation by the thermal reaction of iron rust “FeOOH”, although the reaction efficiency is rather low and the chemical reaction formula has not logically been described so far.

5. Summaries

The following things are summaries in this paper.

(1) It is proved that the TDS peak at 300 to 400°C is based on the hydrogen generation by the reaction at 300 to 400°C that Goethite “FeOOH” is changed into Hematite “Fe₂O₃” inside a TDS measurement apparatus.

(2) The signal at 300 to 400°C in the TDS spectrum often appears in as-corroded sample with corrosion products and corrosion layer, and even in hand-polished corroded sample. The origin of hydrogen is based on deeply-rooted rusts of Goethite “FeOOH” which cannot be removed and unavoidably remain as small dots, no matter how thoroughly it is hand-polished with abrasive papers.

(3) When this phenomenon is re-interpreted as a potential method to make hydrogen by using iron rust of FeOOH through the thermal reaction of 300 to 400°C, there may be high potentiality to produce hydrogen but the real efficiency is as low as “1 kg of FeOOH” can generate only “3.1 mg of H₂ gas”. From another viewpoint of steelmaking plant, especially at the upper-stream process, the heat source of around 300 to 400°C exists, and a lot of reddish scrap is relatively easily obtained in the whole area of a steelmaking factory. The authors think of FeOOH-based hydrogen generation as another potential way to get hydrogen easily through the thermal treatment of 300 to 400°C.

(4) As for the efficiency to emit hydrogen, the coexistence of iron rust of FeOOH and steel is more highly efficient than only FeOOH. In the coexistence case, in addition to an estimated rust-transformation-based direct reaction to emit hydrogen from Goethite “FeOOH” to Hematite “Fe₂O₃” (chemical reaction of (5)), it is reasonable that a widely-acknowledged H₂O-reduction-based reaction occurs preferentially or simultaneously with the above direct reaction, that is, dehydrated FeOOH-based H₂O is changed into H₂ when co-existing with Fe and Fe is oxidized to Fe₂O₃.

(5) The comprehensive reaction to emit hydrogen from the rust has not completely described as chemical formulae. It still remains an on-going subject.

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