Introduction

Wide distribution of high-purity hydrogen is anticipated to make it available as a clean secondary energy source: many primary energy resources can be hydrogen sources; and water is the only product of hydrogen combustion. Through the years, various hydrogen distribution systems such as compressed hydrogen, hydrogen liquefaction, and chemical storage using liquid compounds including ammonia and organic hydrides (cycloalkanes) have been examined. Nevertheless, compressed hydrogen requires large-scale systems and high-pressure storage tanks. Liquefied hydrogen is unsuitable for long-term storage because of its attendant evaporation loss (boil-off). Although ammonia has been anticipated as a promising hydrogen carrier with a high hydrogen capacity (17.6 wt%), it has high toxicity. Organic hydrides are more practical media for long-term storage and for mass transportation in terms of cost effectiveness and safety.

Using organic hydrides requires reversible hydrogenation and dehydrogenation reactions proceeding between aromatic and naphthene compounds (i.e. cycloalkanes). At hydrogen generation sites, aromatics are converted to naphthene compounds by catalytic hydrogenation. The reverse reaction: dehydrogenation of naphthene to aromatics, is conducted at hydrogen consumption sites. Among various liquid organic hydrides, the most well-known and promising system is the methylcyclohexane (MCH)–toluene cycle (hydrogen capacity: 6.1 wt%) by virtue of its nontoxicity and wider range of liquid phase at 178–374 K.

Dehydrogenation of MCH is an endothermic reaction (as shown in eqn (1)); an energy-consuming process. This reaction has been performed over a Pt-supported catalyst. Results show high activity for dehydrogenation above 623 K of reaction temperatures. However, Pt-based catalysts are easily deactivated with coke formation and/or toluene fouling during the reaction. Recently, Ir/USY, partially reduced Mo/SiO2, Pt–Mo/SiO2 (ref. 11 and 18) catalysts have been reported as effective catalysts for dehydrogenation of MCH.

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\begin{align*}
\text{C}_7\text{H}_{14} & \rightarrow \text{C}_7\text{H}_8 + 3\text{H}_2, \quad \Delta H_{298}^0 = 205 \text{ kJ mol}^{-1} \quad (1) \\
\text{C}_7\text{H}_8 + \text{H}_2 & \rightarrow \text{C}_6\text{H}_6 + \text{CH}_4, \quad \Delta H_{298}^0 = -42 \text{ kJ mol}^{-1} \quad (2)
\end{align*}
\]

Feasibility of the organic hydride system depends on the product selectivity and lifetime of MCH dehydrogenation catalysts. A side-reaction of MCH dehydrogenation, hydro-demethylation of toluene (as shown in eqn (2)), and coking on the catalyst are key difficulties to be overcome. They lead to production of benzene and methane as by-products, thereby.

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The methylcyclohexane (MCH)–toluene cycle is a promising liquid organic hydride system as a hydrogen carrier. Generally, MCH dehydrogenation has been conducted over Pt-supported catalysts, for which it requires temperatures higher than 623 K because of its endothermic nature. For this study, an electric field was applied to Pt/TiO2 catalyst to promote MCH dehydrogenation at low temperatures. Selective dehydrogenation was achieved with the electric field application exceeding thermodynamic equilibrium, even at 423 K. With the electric field, ‘inverse’ kinetic isotope effect (KIE) was observed by accelerated proton collision with MCH on the Pt/TiO2 catalyst. Moreover, Pt/TiO2 catalyst showed no methane by-production and less coke formation during MCH dehydrogenation. DRIFTS and XPS measurements revealed that electron donation from TiO2 to Pt weakened the interaction between catalyst surface and π-coordination of toluene. Results show that the electric field facilitated MCH dehydrogenation without methane and coke by-production over Pt/TiO2 catalyst.
increasing the costs of hydrogen purification.\textsuperscript{19} Furthermore, benzene is toxic for humans.\textsuperscript{21} For this study, an electric field was applied over Pt/TiO\textsubscript{2} catalyst to promote MCH dehydrogenation at lower temperatures without methane or coke by-production. This report is the first of low-temperature and selective catalytic dehydrogenation of MCH over Pt/TiO\textsubscript{2} catalyst in an electric field.

**Experimental**

**Catalyst preparation**

For catalyst preparation, platinum was supported on TiO\textsubscript{2} (JRC-TIO-16) and CeO\textsubscript{2} (JRC-CEO-1) using a wet-impregnation method. After catalyst supports were drenched in 25 mL of water for 120 min, Pt precursor, \textit{i.e.} Pt(NH\textsubscript{3})\textsubscript{4}(NO\textsubscript{3})\textsubscript{2} (Sigma-Aldrich Corp.) was added with 15 mL of diluted water and was dissolved for 120 min in an evaporator. An adequate amount of platinum precursor was introduced to make 3 wt\%Pt/TiO\textsubscript{2} and 3 wt\%Pt/CeO\textsubscript{2}. The obtained solution was dried and calcined at 773 K for 120 min. The resultant catalyst powder was pressed and sieved to 355–500 μm. Prepared catalyst was used for the reaction in the electric field without any pre-treatment.

**Catalytic activity tests**

A fixed-bed flow type reactor was packed with 200 mg of Pt-supported catalysts to conduct MCH dehydrogenation. An electric field was applied using two stainless steel electrodes set on upper side and bottom side of the catalyst bed. Catalytic dehydrogenation in the electric field was performed by application of 3.0 mA of constant current between two electrodes. The reaction temperature was set between 423 to 773 K with and without the electric field. A thermocouple was placed on the catalyst bed for monitoring the real temperature of the catalyst because Joule heat is expected to increase the catalyst temperature during dehydrogenation in the electric field. \(T_{ic}\) stands for the real catalyst temperature measured by the thermocouple. The reaction gas consisted of C\textsubscript{7}H\textsubscript{14} : Ar = 6.4 : 30 (total gas flow rate: 36.4 mL min\textsuperscript{-1}) at ambient pressure of 0.1 MPa. Reactions were conducted in a kinetic region, \textit{i.e.} diffusion is not a rate-determining factor. The gaseous products of the reaction were analyzed using GC-TCD (GC-8A; Shimadzu Corp.) and GC-FID (GC-8A; Shimadzu Corp.). The hydrogen yield was defined using the MCH feed rate (μmol min\textsuperscript{-1}) and H\textsubscript{2} formation rate (μmol min\textsuperscript{-1}).

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\text{Hydrogen yield (％) = } \frac{\text{H}_2}{(\text{C}_7\text{H}_{14} \text{ feed rate}) \times 3} \times 100
\]  

The apparent activation energy was estimated over Pt/TiO\textsubscript{2} from Arrhenius plots during MCH dehydrogenation with and without the electric field. Reaction rates were estimated in kinetic control. The reaction rates were calculated from the H\textsubscript{2} formation rate in the same manner as that used for activity tests.

Deposited coke amounts were evaluated using temperature programmed oxidation (TPO) measurements estimating CO or CO\textsubscript{2} formation caused by the oxidation of coke. The temperature was increased from 298 K to 1173 K at 10 K min\textsuperscript{-1} in the gas composition of O\textsubscript{2} : He = 10 : 90 (mL min\textsuperscript{-1}). The product components and coke formation amount indicated a credible mass balance (almost 100%).

**Partial pressure dependency and isotope effects**

Partial pressure dependencies of MCH, toluene, and hydrogen on reaction rates were evaluated over Pt/TiO\textsubscript{2} catalyst to elucidate the role of the electric field. Results were obtained with and without the electric field at the same MCH-conversion-level \textit{i.e.} (a) at 423 K with the electric field and (b) at 498 K without the electric field. For this experiment, the total feed gas flow was set as 56.4 mL min\textsuperscript{-1} at ambient pressure of 0.1 MPa. Then, the gas ratio was modified as C\textsubscript{7}H\textsubscript{14} : Ar = (3.2, 6.4, 9.6, 11.3) : (53.2, 50.0, 46.8, 45.1), C\textsubscript{7}H\textsubscript{14} : C\textsubscript{3}H\textsubscript{8} : Ar = 6.4 : (1.0, 2.0, 3.0, 4.0) : (49.0, 48.0, 47.0, 46.0) and C\textsubscript{2}H\textsubscript{14} : H\textsubscript{2} : Ar = 6.4 : (4.0, 7.0, 9.0, 12.0) : (46.0, 43.0, 41.0, 38.0). Reaction rates were determined by the H\textsubscript{2} formation rate measured using GC-TCD (GC-8A; Shimadzu Corp.).

MCH dehydrogenation was conducted by supplying isotopes of MCH and H\textsubscript{2} to investigate isotope effects with and without the electric field. Accordingly, the reaction gas consisted of C\textsubscript{7}H\textsubscript{14} (or C\textsubscript{2}D\textsubscript{14}) denoted as MCH\textsubscript{3}: H\textsubscript{2} (or D\textsubscript{2}): Ar = 6.4 : 4.0 : 46.0. The total gas flow rate was 56.4 mL min\textsuperscript{-1}. Reaction rates were assessed by the formation rates of H\textsubscript{2}, HD, and D\textsubscript{2}.

**DRIFTS measurements during toluene feed over Pt/TiO\textsubscript{2} and Pt/CeO\textsubscript{2}**

DRIFTS (diffuse reflectance for infrared Fourier transform spectroscopy) measurements were conducted to investigate toluene adsorption over Pt/TiO\textsubscript{2} and Pt/CeO\textsubscript{2} catalysts using a Fourier transform infrared spectrometer (FT/IR-6100; Jasco Corp.). Teflon-made DRIFTS cells were packed with sieved catalyst, on which two stainless steel electrodes were set to apply the electric field. Detailed information related to the apparatus of DRIFTS measurement is presented elsewhere.\textsuperscript{44} Although the electric field was stabilized over Pt/TiO\textsubscript{2} in a fixed-bed flow-type reactor, electrical discharge has occurred all the time over packed Pt/TiO\textsubscript{2} in the DRIFTS cell. Therefore, the experiment with the electric field was performed only over Pt/CeO\textsubscript{2} for safety reasons. Toluene gas was introduced into the measurement chamber for 30 min. Subsequently, in the case of Pt/CeO\textsubscript{2}, the feed gas was switched to argon. Then an electric field was applied for 30 min. Subsequently, purge treatment was conducted to extract toluene gas from the chamber atmosphere. The DRIFTS spectra were obtained after each process.

**Characterization of catalysts**

The metallic surface area of Pt metal was evaluated using CO pulse measurements (Autosorb iQ; Quantachrome Instruments). For vaporizing adsorbed water on the catalyst, pre-treatment was conducted at 673 K in helium. In addition, STEM images and EDX mapping results were obtained from scanning transmission electron microscopy (STEM; HF-2210; Hitachi Ltd.) to confirm the state of Pt over catalytic supports. The specific surface area of catalysts was examined using nitrogen adsorption with the BET method (Gemini VII; Micromeritics Instrument Corp.). For this experiment, pre-treatment
was conducted at 473 K in nitrogen atmosphere for 120 min. Additionally, the electric state of Pt was investigated using XPS measurements (VersaProbe2; ULVAC-PHI, Inc.) without an exposure to air. Orbital 4f$^7/2$ and 4f$^5/2$ of Pt were measured. For this experiment, X-ray source was Al Kα. The binding energies were referenced to the C 1s peak at 284.8 eV.

Results and discussion

Promotion effects of electric fields on catalytic activity during MCH dehydrogenation

After various screening tests, we performed MCH dehydrogenation over Pt/TiO$_2$ catalyst as an optimal catalyst at several temperatures from 423 K to 773 K with and without an electric field (3 mA of constant current). The resultant catalytic activities are presented in Table 1. Although the applied electric field produced little Joule heat, the increase of the catalyst bed temperature (thermocouple temperature, $T_{\text{tc}}$) was no more than 10 K during the reaction. Additionally, the electric field application drastically improved the catalytic activity. The electric field effect was more remarkable at lower temperatures. Especially at 423 K, the catalytic activity was 17.9%, even in a kinetic condition. That activity exceeded the equilibrium limitation (5%) at this temperature.

The values of apparent activation energy were evaluated with and without the electric field to elucidate the promoted catalytic performance of Pt/TiO$_2$ catalyst by the electric field. Fig. 1 shows Arrhenius plots over Pt/TiO$_2$ catalyst during MCH dehydrogenation with (filled plots) and without (blank plots) the electric field. According to these results, the respective values of apparent activation energy were 26.9 and 60.7 kJ mol$^{-1}$ with and without the electric field. Such experimental data indicate that the applied electric field facilitated the MCH dehydrogenation, and changed the reaction mechanism.

Kinetic analysis and isotope effect during MCH dehydrogenation in the electric field

To investigate the role of the applied electric field, partial pressure dependencies were evaluated. Fig. 2 presents results of partial pressure dependencies of MCH, toluene, and hydrogen. Generally speaking, the catalytic MCH dehydrogenation is a reversible endothermic reaction. The MCH dehydrogenation reaction rate is assumed as the following equation using MCH, toluene, and hydrogen partial pressures.

$$r_{\text{dehydrogenation}} = k[\text{MCH}]^a[\text{Toluene}]^b[\text{H}_2]^c$$ (4)

The dehydrogenation rate was correlated positively with the MCH partial pressure similarly in both cases: $a$ was 0.49 with the electric field and 0.47 without the electric field. As results over Pt/TiO$_2$ catalyst, the toluene partial pressure was unrelated to the dehydrogenation rate; $b$ was zero with and without the electric field. This tendency found for the toluene partial pressure, which differs from the well-known Pt catalyst supported on alumina, has been reported for Pt/TiO$_2$ catalyst. Accordingly, Pt catalyst supported on TiO$_2$ is more tolerant of inhibition by toluene adsorption during MCH dehydrogenation. Regarding results of hydrogen partial pressures found with and without the electric field, the MCH dehydrogenation rate was

![Fig. 1](image1.png)

![Fig. 2](image2.png)

Table 1 Catalytic activity during MCH dehydrogenation with/without the electric field$^a$

| Reaction | With EF | Without EF |
|----------|---------|------------|
| Temperature/K | $T_{\text{tc}}$/K | $T_{\text{tc}}$/K | $T_{\text{tc}}$/K | $T_{\text{tc}}$/K |
| 423       | 432     | 17.9       | 423         | 1.2         |
| 473       | 475     | 31.7       | 473         | 10.9        |
| 498       | —       | —          | 498         | 19.0        |
| 523       | 523     | 58.2       | 523         | 32.6        |
| 573       | 573     | 78.1       | 573         | 63.9        |
| 623       | 623     | 94.4       | 623         | 76.1        |
| 673       | 673     | 99.6       | 673         | 88.3        |
| 723       | 723     | 100.0      | 723         | 94.5        |
| 773       | 773     | 99.9       | 773         | 98.8        |

$^a$ $T_{\text{tc}}$: catalyst-bed temperature measured with a thermocouple.
negatively correlated by hydrogen without the electric field, $c$ was $-0.21$. One can infer that hydrogen promoted reverse toluene hydrogenation without the electric field. However, the hydrogen partial pressure was positively correlated with the MCH dehydrogenation rate in the electric field, $c$, was 0.15, which differs from common results. In earlier studies, such positive values of hydrogen partial pressure dependence are apparent if proton ($H^+$) hopping occurs on the catalyst surface.26–28 In the case of methane steam reforming in the electric field, protons hopped on Pd/CeO$_2$ catalyst surface and facilitated methane activation by proton collision with methane.24,26 Therefore, protons can function as a reaction promoter during MCH dehydrogenation in the electric field. In the low temperature and low conversion region, toluene and hydrogen, which were products of dehydrogenation had no negative influence on dehydrogenation rate. Therefore, dehydrogenation might proceed with an irreversible process.

To elucidate the role of proton species, isotope effects were confirmed by supplying isotopic MCHD and D$_2$ during MCH dehydrogenation with and without the electric field (Table 2). Here, isotope effects were expressed by the ratio of rate

### Table 2 Isotope effect during MCH dehydrogenation over Pt/TiO$_2$$^{a}$

| Condition | $T_{tc}$/K | $H_2$ production rate/μmol min$^{-1}$ | $H_2$ yield/% | $k_{D}/k_{H}$/— |
|-----------|-----------|-------------------------------|--------------|----------------|
| With EF (423 K) | | | | |
| MCH$_3$/H$_2$ | 437 | 109 | 13.8 | — |
| MCH$_3$/D$_2$ | 435 | 134 | 17.1 | 1.24 |
| MCH$_3$/H$_2$ | 434 | 135 | 17.1 | 1.24 |
| MCH$_3$/D$_2$ | 433 | 160 | 20.3 | 1.47 |
| Without EF (498 K) | | | | |
| MCH$_3$/H$_2$ | 496 | 169 | 21.5 | — |
| MCH$_3$/D$_2$ | 495 | 182 | 23.2 | 1.08 |
| MCH$_3$/H$_2$ | 497 | 111 | 14.2 | 0.66 |
| MCH$_3$/D$_2$ | 496 | 117 | 14.9 | 0.69 |

$^{a}$ $T_{tc}$: catalyst-bed temperature measured with a thermocouple.
constants ($k$) with/without isotope, i.e. $k_2/k_1$ ratio. Generally, $k_2/k_1$ is a value lower than 1 because chemical bonding of heavier isotopes is more stable (lower zero-point energy: ZPE). Without the electric field, the results follow these general trends when isotope MCHD was introduced, $k_2/k_1$ was 0.66 with MCHD/H2 and $k_2/k_1$ was 0.69 with MCHD/D2. Accordingly, the kinetic isotope effect (KIE) was confirmed without the electric field. Nevertheless, with the electric field, $k_2/k_1$ increased as a larger proportion of deuterium was introduced into the reactor, as shown in Table 2. This inverse KIE has been observed when C–H–H configuration was formed through proton collision, explained by the ZPE discrepancy on isotopes. During MCH dehydrogenation in the electric field, inverse KIE are observed by the collision of accelerated proton with H (or D)-atoms of MCH to form [H (or D)–H (or D)–C7H13]+ over Pt/TiO2 catalyst, to advance the reaction of MCH dehydrogenation further. After dehydrogenation, the resultant state of C7H13+ + H2 has much lower energy level than the three-atom transition and the initial physisorption state (Fig. 3). In this case, reverse hydrogenation has larger activation energy, making it difficult to proceed. Thereby, results suggest that the applied electric field promoted proton hopping and MCH dehydrogenation to show higher catalytic activity beyond the equilibrium limitation (Fig. 3).

### Methane by-production suppression and the electric state of platinum on Pt/TiO2 catalyst

Methane by-production rate and coke deposition over Pt/TiO2 catalyst were investigated during MCH dehydrogenation at 423 K in the electric field. Only slight to no coke deposition occurred in this catalyst system (ESI Table S1†). Also, the structure of catalyst was not changed even after the reaction in the electric field (ESI Table S2†). Fig. 4 presents results of the catalytic activity and methane by-production rate during 120 min of dehydrogenation. For better comparison, results over Pt/CeO2 catalyst were also presented in Fig. 4. Results show that Pt/TiO2 had no methane by-production after 10–25 min of dehydrogenation as the applied electric field being stabilized. According to the partial pressure dependency of toluene, Pt/TiO2 catalyst showed high tolerance against inhibition by toluene adsorption during MCH dehydrogenation both with and without the electric field. Methane by-production reportedly occurs with hydrodemethylation of produced toluene during MCH dehydrogenation. Consequently, the toluene behavior over Pt/TiO2 is regarded as related to the methane by-production rate during dehydrogenation.

DRIFTS measurements were conducted over Pt/TiO2 and Pt/CeO2 catalysts supplying toluene on the catalyst surface, as shown in Fig. 5. In the case of Pt/TiO2, the electric field cannot be stable and applicable in the DRIFTS system because of the cell structure. Therefore, in situ DRIFTS with the electric field was conducted over Pt/CeO2 instead of Pt/TiO2. Regarding the Pt/CeO2 catalyst, toluene adsorption was observed both with and without the electric field (C–H stretching around 3000 cm−1). However, the IR peaks for toluene are barely visible over Pt/TiO2 catalyst, even under a toluene atmosphere. Furthermore, the peaks disappeared completely after purge treatment. These results demonstrate that Pt/TiO2 catalyst can suppress toluene adsorption and methane by-production during MCH dehydrogenation.

### Conclusions

Dehydrogenation of methycyclohexane (MCH) was conducted over Pt/TiO2 catalyst by applying an electric field. The electric field promoted the reaction to achieve 17.9% hydrogen yield, even at 423 K and a kinetic condition, under which the catalytic activity exceeded the equilibrium limitation (5%). Despite the reversibility of dehydrogenation, the reaction rate was positively correlated with the partial pressure of hydrogen with the electric field being stabilized. These results demonstrate that Pt/TiO2 catalyst can suppress toluene adsorption and methane by-production during MCH dehydrogenation. The electric field promoted the reaction to achieve 17.9% hydrogen yield, even at 423 K and a kinetic condition, under which the catalytic activity exceeded the equilibrium limitation (5%).

| Catalyst | Condition | Binding energy/eV | Relative peak area/% |
|----------|-----------|-------------------|----------------------|
|          |           | Pt0   | Pt2+  | Pt4+  | Pt0   | Pt2+  | Pt4+  |
| 3 wt% Pt/TiO2 | As made | 70.8  | 72.4  | 74.2  | 22.9  | 52.2  | 24.9  |
| 3 wt% Pt/TiO2 | After reaction with EF (423 K) | 70.9  | 72.5  | 74.3  | 43.5  | 27.1  | 29.5  |
| 3 wt% Pt/CeO2 | As made | 70.9  | 72.4  | 74.2  | 16.5  | 74.8  | 8.8   |
| 3 wt% Pt/CeO2 | After reaction with EF (423 K) | 70.8  | 72.5  | 74.3  | 39.8  | 50.0  | 10.3  |
field application. Additionally, an inverse kinetic isotope effect (KIE) was observed with isotope MCHD (C7D14) and D2 in the electric field. These results indicate that hydrogen-derived species, i.e. protons, promoted the abstraction of H-atoms from MCH for dehydrogenation. Furthermore, Pt/TiO2 catalyst, different from other catalysts, showed neither methane nor coke by-production during MCH dehydrogenation in the electric field at 423 K. DRIFTS measurements revealed that toluene was only slightly adsorbed over Pt/TiO2 catalyst. XPS measurements demonstrated that Pt on TiO2 was more metallic after applying the electric field. Actually, TiO2 is known to be capable of donating electrons to the empty orbital of Pt. Therefore, we inferred that more metallic Pt on TiO2 support weakened interaction between Pt and π-coordination of toluene. In conclusion, results show that Pt/TiO2 catalyst promotes MCH dehydrogenation selectively at low temperatures by virtue of proton hopping in the electric field.

Conflicts of interest
There are no conflicts to declare.

Acknowledgements
This study was supported by JST-CREST JPMJCR1423 and JSPS Grants-in-Aid for Scientific Research – KAKENHI.

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