Observation of an oxonium ion intermediate in ethanol dehydration to ethene on zeolite

Xue Zhou1,2,4, Chao Wang1,4, Yueying Chu1,4, Jun Xu1,3, Qiang Wang1, Guodong Qi1, Xingling Zhao1,2, Ningdong Feng1 & Feng Deng1

Zeolite-catalyzed dehydration of ethanol offers promising perspectives for the sustainable production of ethene. Complex parallel-consecutive pathways are proposed to be involved in the reaction network of ethanol dehydration on zeolites, where the initial step of ethanol dehydration is still unclear particularly for the favorable production of ethene at lower temperature. Here we report the observation of a triethyloxonium ion (TEO) in the dehydration of ethanol on zeolite H-ZSM-5 by using ex situ and in situ solid-state NMR spectroscopy. TEO is identified as a stable surface species on the working catalyst, which shows high reactivity during reaction. Ethylation of the zeolite by TEO occurs at lower temperature, leading to the formation of surface ethoxy species and then ethene. The TEO-ethoxide pathway is found to be energetically preferable for the dehydration of ethanol to ethene in the initial stage, which is also supported by theoretical calculations.
Ethene is one of the most important commodity chemicals, which is currently produced by cracking processes from petroleum. Due to the large availability of bioethanol from renewable biomass sources and the dwindling of fossil resource, the conversion of ethanol to ethene and higher hydrocarbons now is receiving increasing attention from both academia and industry. Among the heterogeneous catalysts studied, zeolites especially ZSM-5 or modified ZSM-5 are the promising catalysts as they can be tuned to exhibit higher activity than the traditional alumina-based catalysts. The reaction temperature is lower and a low concentration of ethanol aqueous solution can be used. The formation of ethene is the first step in ethanol dehydration; subsequent polymerization, cracking and aromatization by the secondary reaction of ethene leads to the formation of longer-chain hydrocarbons, very similar to the formation of ethene and diethyl ether in the conversion of methanol over zeolites. The detailed knowledge of the reaction mechanism of ethene formation in ethanol dehydration is important not only for understanding ethene formation as a model reaction of alcohol conversion but also for the development of improved catalysts for light olefins production.

The formation of ethene in the initial dehydration process has been experimentally and theoretically investigated. A parallel-consecutive route was proposed: dehydration of ethanol to ethene and diethyl ether in parallel and a consecutive decomposition of diethyl ether to ethene. In the unimolecular dehydration of ethanol to ethene, it was proposed that the acidic proton transfers from the zeolite to the OH group on ethanol, which subsequently releases water to form an intermediate such as carbocation or an alkoxide bound to zeolite. Solid-state NMR and infrared spectroscopic studies confirmed the formation of surface-bound ethoxy species as a stable intermediate in ethanol dehydration on H-ZSM-5 and HY zeolites. Ethene and hydrocarbons are formed by the subsequent decomposition of the surface ethoxy species.

Diethyl ether (DEE) is concurrently produced with ethene, and the kinetic and spectroscopic studies revealed that intermolecular dehydration of ethylene leads to DEE via dimeric ethanoic species or reaction of an ethoxy group with undissociated ethanol on zeolites. The following cracking of DEE produces ethene, which occurs preferably at lower reaction temperatures as compared to the unimolecular dehydration of ethanol to ethene. Note that the ethoxy species was generally proposed to be the direct precursor to ethene. However, the generation of such species is still not well understood. Thus, although different reaction schemes have been proposed, the detailed mechanism of the initial step of ethanol dehydration remains elusive, particularly for the favorable production of ethene from DEE at lower temperature.

Here we report the observation and identification of reaction intermediates in ethanol dehydration to ethene over zeolite H-ZSM-5. The ex situ and in situ solid-state NMR spectroscopy allows for exploration of the dehydration of ethanol at the initial stage. A triethylxonium ion (TEO) is observed as a potential active intermediate on the working catalyst. We provide evidence for the involvement of TEO in the formation of surface ethoxy species and then ethene on the zeolite. The elementary reactions leading to the formation of ethene are identified by combining experiments and theoretical calculations, which suggests that a TEO-mediated reaction route is operative for the dehydration of ethanol to ethene.

**Results**

**Observation of TEO.** H-ZSM-5 (Si/Al = 11.5, Zeolyst) was used for the dehydration of ethanol. The XRD and 27Al and 29Si solid-state NMR spectra show structure information of the H-ZSM-5 catalyst (Supplementary Fig. 1 and 2). The acidic property was examined by FT-IR and NH3-TPD (Supplementary Fig. 3 and 4). The reactions were conducted in a pulse-quench reactor, in which ethanol was pulse-injected at temperatures ranging from 160 to 350 °C and allowed to react for 4 s under continuous He carrier gas flow before the reaction was thermally quenched by pulsing liquid nitrogen onto the catalyst bed. The reaction effluent was analyzed by GC (Supplementary Fig. 5). Ethene is the only product at 160–200 °C, indicating that the dehydration of ethanol to ethene is favorable at lower temperature. With increasing reaction temperature, the secondary reaction of ethene leads to the formation of C3+ products including olefins and aromatics.

Figure 1 shows the ex situ 13C CP/MAS NMR spectra of trapped products from the dehydration of 13C-1-enriched ethanol (CH313CH2OH, 99% 13C) on H-ZSM-5 at 160–350 °C for 4 s. Ethanol and DEE are observed as reflected by the strong signals at 62.5 and 69.0 ppm respectively, due to the 13C-enriched methylene carbons; the methyl carbons produce the weak signals at 12–17 ppm. The adsorbed species formed at 200 °C were further analyzed by two-dimensional (2D) 1H–13C HETCOR NMR experiment (Supplementary Fig. 6). A weak signal at 72.8 ppm that was seriously overlapped by the ethanol signal produced broad signals at 72.6 ppm.

Interestingly, a well-resolved signal appears at 85.0 ppm (Fig. 1) and grows up at intermediate temperature. The oligomeric alkoxo species formed by the oligomerization of ethene on acidic zeolites were previously proposed to produce broad signals at 78.0 to
90.0 ppm in the literature\textsuperscript{20,26}. Since there is no any C$_{3+}$ species formed at this reaction condition, the oligomerization of ethene did not likely occur in our case. Furthermore, no methylene signals at 20.0 to 40.0 ppm can be observed that are supposed to be oligomerized compounds\textsuperscript{27}. Therefore, the 85.0 ppm signal should not be ascribed to the oligomeric alkoxide species. Instead, we proposed on the basis of its chemical shift the formation of alkyl-substituted oxonium ions. Munson and Haw reported the formation of trimethyloxonium ion (TMO) with a characteristic $^{13}$C NMR signal at 80.0 ppm by the reaction of dimethyl ether on H-ZSM-5\textsuperscript{28}, and they also confirmed that TMO was not an intermediate in the methanol to hydrocarbons conversion. Since DEE is readily formed and in equilibrium with ethanol (Fig. 1b), the 62.5 ppm signal of DEE (69.0 ppm) and ethanol (62.5 ppm) was determined by $^{13}$C NMR experiment, which provides an unambiguous identification of bond connectivity of carbon species. The INADEQUATE (Incredible Natural Abundance Double Quantum Transfer Experiment)\textsuperscript{29} MAS NMR experiment was performed, which provides an unambiguous identification of bond connectivity of carbon species. The INADEQUATE spectrum (Fig. 2) of H-ZSM-5 reacted with $^{13}$CH$_3^{14}$CH$_2$OH at 200 °C for 4 s exhibits a clear correlation peak pair between the methylene carbon at 85.0 ppm and the methyl carbon at 12.0 ppm, confirming the formation of TEO that is composed of ethyl groups. Two other correlations are identified, showing the connectivity of the methylene carbons of DEE (69.0 ppm) and ethanol (62.5 ppm) to the corresponding methyl carbons at 12.8 and 17.0 ppm, respectively. The structure of TEO was also theoretically optimized in the H-ZSM-5 channel by periodic density functional theory (DFT) calculations (Supplementary Fig. 7). The $^{13}$C chemical shift of the methylene carbon of TEO is predicted to be 84.6 ppm, in good agreement with the experimental data. Therefore, our NMR experiments and theoretical calculations provide solid evidence for TEO formation in the ethanol dehydration process.

Fig. 2 Identification of triethyloxoium ion by correlation NMR spectroscopy. 2D $^{13}$C-$^{13}$C INADEQUATE MAS NMR spectrum of trapped species obtained from the reaction of $^{13}$CH$_3^{14}$CH$_2$OH on H-ZSM-5 at 220 °C for 4 s

To gain insights into the structure of TEO, a $^{13}$C-$^{13}$C J-based refocused INADEQUATE (Incredible Natural Abundance Double Quantum Transfer Experiment)\textsuperscript{29} MAS NMR experiment was performed, which provides an unambiguous identification of bond connectivity of carbon species. The INADEQUATE spectrum (Fig. 2) of H-ZSM-5 reacted with $^{13}$CH$_3^{14}$CH$_2$OH at 200 °C for 4 s exhibits a clear correlation peak pair between the methylene carbon at 85.0 ppm and the methyl carbon at 12.0 ppm, confirming the formation of TEO that is composed of ethyl groups. Two other correlations are identified, showing the connectivity of the methylene carbons of DEE (69.0 ppm) and ethanol (62.5 ppm) to the corresponding methyl carbons at 12.8 and 17.0 ppm, respectively. The structure of TEO was also theoretically optimized in the H-ZSM-5 channel by periodic density functional theory (DFT) calculations (Supplementary Fig. 7). The $^{13}$C chemical shift of the methylene carbon of TEO is predicted to be 84.6 ppm, in good agreement with the experimental data. Therefore, our NMR experiments and theoretical calculations provide solid evidence for TEO formation in the ethanol dehydration process.

For the adsorbed species formed on H-ZSM-5 after reaction at 200 °C for 4 s, the amount of TEO (85.0 ppm), ethoxy (72.8 ppm), DEE (69.0 ppm) and ethanol (62.5 ppm) was determined by $^{13}$C NMR experiment to be 0.34, 0.46, 28.76 and 6.99 mmol/g, respectively. TEO and the ethoxy species are solely formed and located on Brønsted acid sites for charge compensation, while DEE and ethanol are trapped over zeolite via either chemical or physical adsorption. Considering the total Brønsted acid sites (about 1.03 mmol/g) on the catalyst, the amount (less than 0.23 mmol/g) of DEE adsorbed on Brønsted acid sites should be lower than that (0.34 mmol/g) of TEO at this reaction condition. At temperature above 300 °C (Fig. 2d, e), the 85.0 ppm signal is almost unobservable, which can be ascribed to the decomposition of TEO. In the meantime, the secondary reaction of ethene produces the dominating C$_{3+}$ hydrocarbons, reflected by the strong high-field signals below 30.0 ppm. The signals at 147.3–156.3, 243.6–255.7 and 47.6 ppm are characteristics of the cyclic catons such as dimethycyclopentenyl and ethylcyclcopentenyl ions\textsuperscript{30–32}, which are usually involved in the formation of aromatics. This implies that the aromatics-based cycle prevails in a similar route to the conversion of methanol to hydrocarbons\textsuperscript{33,34}.

Reactivity and intermediate role of TEO. In order to explore the reactivity of TEO, continuous-flow $^{13}$CH$_3^{14}$CH$_2$OH conversion over H-ZSM-5 was investigated in situ $^{13}$C NMR spectroscopy (see methods section for details of the in situ NMR experiments). Figure 3 shows the real-time monitoring of formed surface species on H-ZSM-5 in the ethanol dehydration process. At a constant reaction temperature of 220 °C (Fig. 3a), the appearance of 85.0 ppm signal after 200 s points to the formation of TEO. The broad signal centered at 58.5 ppm should come from the adsorbed ethanol, whose methyl group produces a high-field strong signal at 15.0 ppm. Note that the methylene group of DEE (69.0 ppm) is invisible in the in situ experiment. This is probably due to the strong adsorption of DEE on zeolite, leading to large anisotropic interactions (chemical shifts and nuclear dipole-dipole couplings) which cannot be efficiently removed by the low magic spinning speed (2 kHz). However, the formation of DEE is evidenced by the 12.5 ppm signal due to its methyl group featured by high mobility. TEO is observable with increasing the reaction time up to 1200 s (20 min), indicative of its stability at the moderate reaction temperature. The GC analysis of the effluent products obtained from the reaction with time-on-stream shows the formation of ethene and a small amount of higher hydrocarbons (Supplementary Fig. 8). Figure 3b shows the in situ $^{13}$C NMR spectra obtained at elevating temperatures. The TEO is observable over a wide temperature range. The formation of hydrocarbons prevails at higher temperatures as reflected by a set of high-field signals (8.7–32.6 ppm) while the TEO is significantly reduced due to its decomposition, consistent with the ex situ NMR result (Fig. 1). In comparison, the structurally similar TMO species formed on H-ZSM-5 was readily decomposed before the onset of hydrocarbon synthesis\textsuperscript{35,36}.

To confirm the intermediate role of TEO in the formation of ethene, the $^{13}$C NMR spectra of trapped species obtained from the pulse-quench reactions of CH$_3^{13}$CH$_2$OH were recorded during the reaction course of 4~64 s at 250 °C. Figure 4a shows the expanded spectra and their deconvolutions (see Supplementary Fig. 9 for the whole spectra), in which the formation and evolution of TEO (85.0 ppm) is evident. We know from above experiments that the ethoxy species (72.8 ppm) is simultaneously produced, showing the high-field signals of TEO (85.0 ppm) and the ethoxy species at 12.5 ppm. The 85.0 ppm signal after 200 s points to the formation of TEO. The broad signal centered at 58.5 ppm should come from the adsorbed ethanol, whose methyl group produces a high-field strong signal at 15.0 ppm. Note that the methylene group of DEE (69.0 ppm) is invisible in the in situ experiment. This is probably due to the strong adsorption of DEE on zeolite, leading to large anisotropic interactions (chemical shifts and nuclear dipole-dipole couplings) which cannot be efficiently removed by the low magic spinning speed (2 kHz). However, the formation of DEE is evidenced by the 12.5 ppm signal due to its methyl group featured by high mobility. TEO is observable with increasing the reaction time up to 1200 s (20 min), indicative of its stability at the moderate reaction temperature. The GC analysis of the effluent products obtained from the reaction with time-on-stream shows the formation of ethene and a small amount of higher hydrocarbons (Supplementary Fig. 8). Figure 3b shows the in situ $^{13}$C NMR spectra obtained at elevating temperatures. The TEO is observable over a wide temperature range. The formation of hydrocarbons prevails at higher temperatures as reflected by a set of high-field signals (8.7–32.6 ppm) while the TEO is significantly reduced due to its decomposition, consistent with the ex situ NMR result (Fig. 1). In comparison, the structurally similar TMO species formed on H-ZSM-5 was readily decomposed before the onset of hydrocarbon synthesis\textsuperscript{35,36}.

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These results demonstrate a direct correlation of TEO to ethoxy species and ethene product.

**Mechanism of ethene formation via TEO.** Our experimental observations indicate a strong potential of TEO serving as an active intermediate in the formation of ethene. A whole reaction network for ethanol dehydration to ethene was proposed (Fig. 5), which includes the previously theoretically predicted unimolecular and bimolecular processes\(^\text{17,19}\) and our proposed TEO-mediated routes. It consists of three types of mechanisms: direct ethanol dehydration to ethene (route 5, 6, 7 and 8), diethyl ether decomposition to ethene (route 3 and 4) and TEO-mediated ethene formation (route 1 and 2). DFT calculations were performed to explore the most favorable ethene formation route and the optimized transition state structures on the zeolite cluster model (Supplementary Fig. 12) are displayed in Supplementary Fig. 13. Our calculations show that all the kinetic steps of ethanol dehydration are endothermic processes (Supplementary Table 2). The calculated Gibbs free energy barrier (Supplementary Fig. 14–18 for the energy diagrams), pre-exponential factor, and rate coefficient at 473 K of the forward and reverse reactions are tabulated in Table 1. The comparison of energy barriers and rate coefficients indicates that the direct dehydration processes (steps 9, 10, 15 and 16) are energetically less favorable for ethene formation, in agreement with the previous work\(^\text{17}\).

Compared with the direct ethanol dehydration routes (route 5, 6, 7 and 8), the two molecules mediated route 4 including dimer-mediated etherification (\(\Delta G_{\text{act}}(0) = 129.0 \text{ kJ/mol}\), \(k_f = 5.54 \times 10^{-2} \text{s}^{-1}\), step 4) and the following DEE decomposition to ethene (\(\Delta G_{\text{act}}(0) = 131.1 \text{ kJ/mol}\), \(k_f = 9.24 \times 10^{-3} \text{s}^{-1}\), step 11) is favorable for ethanol dehydration to ethene, in agreement with the previous report that the reaction path via DEE is preferentially involved in ethene formation at temperatures lower than 500 K\(^\text{17}\). It is reasonable to assume that TEO would be readily formed due to the facile formation of DEE. This is confirmed by a free energy barrier of ca. 94.1 kJ mol\(^{-1}\) for the formation of TEO from DEE (step 6), which is much lower than that of DEE decomposition routes (\(\Delta G_{\text{act}}(0) = 136.1 \text{ kJ/mol}\) for step 11 in route 4 and \(\Delta G_{\text{act}}(0) = 155.9 \text{ kJ/mol}\) for step 12 in route 3). We have experimentally observed that the ethoxy species and DEE can be produced from TEO on H-ZSM-5 (Fig. 4). This can be understood by the fact that TEO ethylates the conjugate base site of zeolite to form a framework-bound ethoxy species with the release of a DEE, which shows the characteristic property of trialkyl oxonium ions acting as an powerful alkylating agent\(^\text{37}\). This process is calculated to have an activation energy of 73.6 kJ mol\(^{-1}\) (step 7, Supplementary Table 2), in good agreement with our experimental value (77.3 kJ mol\(^{-1}\)), strongly supporting the formation of ethoxy via TEO intermediate. The low free energy barrier of step 8 (107.6 kJ mol\(^{-1}\)) suggests the transformation of ethoxy species to ethene can readily occur. Moreover, the calculated electronic energy barrier for the conversion of ethoxy species to ethene (127.6 kJ mol\(^{-1}\), Supplementary Table 2) is similar to the experimentally estimated value (ca. 122 kJ mol\(^{-1}\)) for the decomposition of ethoxy species to ethene on H-ZSM-5\(^\text{38}\). The direct decomposition of TEO into ethene (step 13 in route 2) is also considered, which is however characterized by a higher free energy barrier (127.7 kJ mol\(^{-1}\)). A full comparison of free energy barriers and rate coefficients of the rate-determination steps in the proposed reaction network indicates that route 1 (steps 4, 6, 7, 8) and route 2 (steps 4, 6, 13) involving TEO intermediate are the most favorable routes. Thus, concerning the overall kinetics during the ethanol dehydration to ethene,

**Fig. 3 Real-time NMR monitoring of ethanol dehydration on zeolite. In situ \(^{13}\text{C}\) MAS NMR spectra of \(^{13}\text{CH}_3^{13}\text{CH}_2\text{OH}\) conversion on H-ZSM-5 with time on stream (a) and at elevating temperatures (b). Spectra (a) were recorded at every 30 s from 0 to 1200 s at 220 °C and 30 scans were accumulated for each temperature slice. Spectra (b) were recorded at every 10 °C from 160–260 °C and 900 scans were accumulated for each temperature slice. The 1D spectra on the top are recorded at 840 s (a) and 250 °C (b), respectively.**
**Fig. 4** Evolution of TEO in the formation of ethene. $^{13}$C CP MAS NMR spectra of trapped species on H-ZSM-5 obtained from pulse-quench reactions of CH$_3^{13}$CH$_2$OH at 250 °C for different time (a) and $^{13}$C MAS NMR spectra of TEO-ZSM-5 at room temperature (RT), heated at 80 °C for 5 min, and at 200 °C for 5 min (b). The deconvoluted components are indicated in color.

**Fig. 5** Proposed catalytic cycle for ethanol dehydration to ethene. The structures of transition states (TS) involved in these reaction steps are depicted in Supplementary Fig. 13. * indicates the adsorbed state.
the presence of TEO provides an energetically preferable process at lower reaction temperature.

**Discussion**

The dehydration of ethanol to ethene over H-ZSM-5 has been investigated by combined experiments and DFT calculations. Our ex situ and in situ solid-state NMR data provide the evidence for the formation of TEO in the dehydration process. Generally,onium ions that are often isolable as salts are well recognized as reaction intermediates\(^\text{37}\). It was reported that trimethyloxonium can be generated on H-ZSM-5 zeolite. However, mechanistic significance of trimethyloxonium is uncertain since its intermediate role for the initial C–C bond formation in methanol conversion was excluded based on experimental observations\(^\text{27,28}\).

Herein, TEO is formed as a stable surface species on the working catalyst in both continuous-flow and pulse-quench reactions. We calculated the host-guest interaction free energy of TEO confined in the H-ZSM-5 channel, which is as low as \(-360.2\) kJ mol\(^{-1}\) at 473 K, indicating that the zeolite framework can stabilize TEO intermediate effectively.

The implications of the observation of TEO in ethanol dehydration to ethene in present work are manifold. The ethoxy species is often experimentally observed during ethanol dehydration on zeolite, which is generally considered as the precursor to ethene in both processes of unimolecular dehydration and bimolecular dehydration of ethanol followed by DEE decomposition. DEE is largely formed in the initial stage of dehydration of ethanol and its coverage on acid sites is estimated to be comparable to TEO at lower temperature (e.g., 473 K). The presence of TEO provides an energetically favorable mechanism to link DEE and ethoxy species, which is viable for the formation of ethene at low temperature. In the onium-ylide mechanism proposed for the C–C bond formation from C1 compound such as methanol and DME on zeolite, Olah et al. predicted the intermediate formation of ethylidemethyloxonium, which could undergo \(\beta\)-elimination to yield ethene and DME\(^\text{39}\). Analogously, the direct deprotonation and decomposition of TEO could lead to ethane and DEE on zeolites (route 2), which, however, occurs with a relatively higher energy barrier. We have experimentally shown that TEO tends to alkylate basic site (Si–O–Al) of zeolite to form surface ethoxy species, which in nature resembles the alkylloxonium salt featured by strong alkylating ability in organic synthesis. This result indicates the reactivity of alkylloxonium ion could be altered by the zeolite catalyst, which leads to different reaction routes. Although the alkylloxonium ion (i.e., a protonated alcohol) has long been proposed as a key intermediate in the dehydration of alcohol to produce alkenes and ethers particularly in strong liquid acid systems, the detailed characterizations of TEO in the ethanol dehydration process in this work provide an example of uncovering the important role of this kind of intermediate in the formation of ethene on zeolite catalysts, which may open avenues for further experimental and theoretical exploration of the oxonium ions chemistry in alcohol conversion.

**Methods**

**X-ray powder diffraction (XRD).** The ammonium form ZSM-5 (Si/Al = 11.5, obtained from Zeolyst) was calcined in air at 823 K for 5 h to obtain the proton form H-ZSM-5. The structure and crystalline nature of H-ZSM-5 zeolites were examined by X-ray diffractometer (X’pert Powder) using a Cu-Kα radiation with a step of 0.02° at a respective voltage of 40 kV and a current of 40 mA.

**FT-IR of pyridine adsorption**. The FT-IR spectra of pyridine-adsorbed samples were measured at 313 K after evacuation at 373, 473, 573, 673 and 773 K, residual pyridine was removed by vacuum. The FT-IR spectra of pyridine-adsorbed samples were measured at 313 K after evacuation at 373, 473, 573, 673 and 773 K, respectively.

**FT-IR of pyridine adsorption**. The FT-IR of pyridine adsorption measurements were performed on a Bruker Tensor 27 spectrometer. The catalysts were first activated at 773 K under high vacuum (<10\(^{-5}\) Pa) for 2 h. After cooling down to 313 K, a background spectrum was collected. Excessive amount of pyridine was then introduced to the infrared cell and held for 2 h to allow equilibrium. The residual pyridine was removed by vacuum. The FT-IR spectra of pyridine-adsorbed samples were measured at 313 K after evacuation at 373, 473, 573, 673 and 773 K, respectively.

**Temperature-programmed desorption of ammonia (NH\(_3\)-TPD)**. The NH\(_3\)-TPD measurement was performed using a FINSORB-3010 chemisorption instrument. Typically, 100 mg of sample was pre-treated at 423 K for 1 h under 30 sccm of helium gas. After the sample cooled down to 323 K, NH\(_3\) was introduced and held for 1 h. The temperature was then elevated from 323 to 1023 K at a ramping rate of 10 K/min and the desorbed NH\(_3\) was detected by a thermal conductivity detector (TCD).

**Catalytic testing**. The H-ZSM-5 powder was pressed into pellets between 60–80 mesh. The pellets (0.2 g) were activated at 400 °C in flowing helium for 1 h prior to the ethanol dehydration reactions. A pulse-quench reactor was used to quench the reaction by reducing the reaction temperature with liquid nitrogen within a very short period (<1 s)\(^\text{35}\). Typically, when the reaction proceeded in a

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**Table 1** Reaction Gibbs free energy barrier (\(\Delta G_{\text{act}}\), kJ mol\(^{-1}\)), pre-exponential factor (\(A_f\), s\(^{-1}\)), and rate coefficient (\(k_f\), s\(^{-1}\)) at 473 K of the forward (\(f\)) and reverse reactions (\(r\)) for the elementary step associated with the eight ethene production routes

| Elementary step | \(\Delta G_{\text{act}}\) | \(A_f\) | \(k_f\) | \(A_r\) | \(k_r\) |
|-----------------|-----------------|--------|--------|--------|--------|
| Route 1 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 6.43 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
| Route 2 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 4.63 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
| Route 3 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 4.63 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
| Route 4 Route 1 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 4.63 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
| Route 5 Route 2 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 4.63 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
| Route 6 Route 3 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 4.63 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
| Route 7 Route 4 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 4.63 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
| Route 8 Route 5 C ↔ DEE.H\(_2\)O (4) | 129.0 | 79.4 | 4.63 × 10\(^{11}\) | 5.54 × 10\(^{-2}\) | 4.20 × 10\(^{12}\) | 1.70 × 10\(^{4}\) |
pulse-quinch reactor for a pre-set period, the reaction was thermally quenched by pulsing liquid nitrogen onto the catalyst bed, which was achieved by using high-speed liquid nitrogen generator. In each case, 1H resonance was pulsed into the reactor (heated at 160–350 °C) containing 0.2 g H-ZSM-5 and allowed to react for different time, before quenching by liquid nitrogen. The trapped surface species were analyzed by 1H solid-state NMR spectroscopy and the effluent products were on line determined by GC-MS analysis. For the continuous reaction, ethanol was used as a weight hourly space velocity (WHSV) of 2 h⁻¹ reacted over the H-ZSM-5 (0.2 g) pellets in a fixed bed reactor at the temperature range of 140–260 °C.

Gas chromatography. The effluent products drawn from the flowing gas were analyzed quantitatively by online GC-FID chromatograph (Shimadzu GC-2010 plus) which equipped with a flame-ionization detector and a Supelco Supel-Q™ PLOT capillary column (30 m × 0.32 mm × 15 μm). The initial temperature was programmed at 40 °C (maintained for 2 min), followed by a rate of 5 °C min⁻¹ to 132.5 °C and a rate of 10 °C min⁻¹ to the final temperature of 220 °C. The retained products in the catalyst was directly analyzed by solid-state NMR spectroscopy (see the following).

Preparation of triethyloxonium ion (TEO) on H-ZSM-5. A concentration of 0.2 g of H-ZSM-5 zeolite (Si/Al = 11.5) was deactivated on a vacuum oven (≤10⁻² Pa) at 405 °C for 12 h. After dehydroxylation, the sample was allowed to cool down to room temperature for subsequent use. A concentration of 0.3 g triethyloxonium tetrafluoroborate (Aldrich) was added to 3 g dry dichloromethane solvent, and then the deactivated zeolite sample was added into the solution. All the operations were carried out in a glovebox equipped with pure N₂. After ultrasonic treatment of the mixture in the ice water bath at 0 °C for 20 min, the mixture was filtrated and evacuted completely and the sample was dried at room temperature for 5 h by vacuum. The obtained triethyloxonium ion (TEO) exchanged ZSM-5 zeolite was denoted as TEO-ZSM-5.

Solid-State NMR experiments. After the reaction was quenched, the pulse-quinch reactor containing the catalyst was sealed. The sealed reactor was then transferred to a glove box for the glove box transfer. The rotor was synchronized and set to 3.24 ms. 64 transients were co-added using a RF-1H (1H) 90-degree pulse of 4.8 μs and 9.6 μs, respectively. The cross-polarization (CP) pulses were 4.8 μs and 9.6 μs, respectively. The CP/MAS NMR spectra in Fig. 3 were generated by the overlay of 1D NMR spectra as a function of reaction time or temperature. The correlation between 1H resonances and the reaction parameters (time and temperature) was followed in the 2D maps, which facilitates the monitoring of the fate of the organic species during the time or temperature evolutions.

The activation energy of the formation of ethoxy species from TEO on H-ZSM-5 was measured as follows. The sealed NMR rotor containing the TEO-ZSM-5 sample was heated at a specific temperature (from 328 to 348 K) for a period of time, and then the reaction was quenched by liquid N₂. A temperature measurement was performed at room temperature. At such low reaction temperature, only ethoxy species and diethyl ether were produced on the TEO-ZSM-5. The concentration of TEO during the reaction was measured by normalized integrated NMR signal at 85 ppm. Finally, the temperature dependent rate constant was obtained and the activation energy was derived from the Arrhenius equation.

Theoretical calculations. The host-guest interactions between the cations and zeolite framework would result in the 13C chemical shift moving due to the redistribution of electronic environment invoked by the H-ZSM-5 confined pore. Thus, to unambiguously assign the experimental NMR results, the 13C NMR chemical shifts of the cations are further calculated using the periodic boundary condition.

All the in situ 13C solid-state NMR experiments were performed at 11.7 T on a Bruker Avance III 500 spectrometer, equipped with a 7 mm H/E MASCAY probe (Fig. S19), with resonance frequencies of 500.38 and 125.37 MHz for 1H and 13C, respectively. The magic angle spinning rate was set to 2 kHz. Prior to the in situ NMR experiments, 0.2 g of pre-dehydrated H-ZSM-5 was pressed into a 7 mm NMR rotor. An axial hole of 2.5 mm in diameter was made into the sample with a special tool, by which an uniform annular catalyst bed was formed in the rotor. After inserting the injection tube into the MAS rotor via the injection tube and pass through the catalyst from the bottom to the top and leave the rotor via an exhaust tube in the rotor cap. The 1H/CP/MAS NMR spectra were recorded with a recycle time of 5 ms and a recycle delay of 1 s. 300 and 900 scans were accumulated for each spectrum at different time and temperature respectively. The 13C chemical shifts were referenced to HMB (a second reference to TMS). The in situ 13C MAS NMR spectra in Fig. 3 were generated by the overlay of 1D NMR spectra as a function of reaction time or temperature. The correlation between 1H resonances and the reaction parameters (time and temperature) was followed in the 2D maps, which facilitates the monitoring of the fate of the organic species during the time or temperature evolutions.

The geometry optimizations were performed by using the CASTEP program41 with the generalized gradient approximation (GGA) proposed by Perdew–Burke–Ernzerhof (PBE) functional. And the ultrasoft pseudo potential, fine plane wave cut-off energy (340 eV) and a default fine level Monkhorst-Pack k-point (1 × 1 × 1) were adopted to sample the Brillouin zone. DFT-D method was used in the structure optimization and the NMR calculation to accurately describe the weak interaction in H-ZSM-5. During optimization, the 22T active sites of the adsorbed cation were relaxed to their equilibrium positions. All of the NMR shielding calculations were performed by the GIPAW method in the MS CASTEP-NMR code at the GGA/PBE level based on the optimized zeolite structures. All of the 13C chemical shifts (δ 13C) cal were derived using the 13C NMR spectra in Fig. 3. The 13C calculated chemical shifts were further converted to δ 13C values, which were plotted on the absolute shielding of TMO, namely 80 ppm for the experimental values.

For the ethene formation pathway calculation, H-ZSM-5 zeolites are represented by a 72T model (Si72T1Al10179, 252 atoms), which were extracted from their crystallographic structural data (http://www.iza-structure.org/databases/). The 72T model is the complete description of the channel structures. All of the TS geometries were optimized based on the optimized structures of cation accommodation in the H-ZSM-5 zeolite. The final k-point (1 × 1 × 1) and cut-off energy of 550 eV were employed, combined with core–valence interactions described by ultrasoft pseudo potential generated on-the-fly. The 13C calculated chemical shifts were further converted to δ 13C values, which were plotted on the absolute shielding of TMO, namely 80 ppm for the experimental values.
reactant and product, respectively, and finally optimized the resulting structures to the minimum structures. These methods have been widely employed in the previous theoretical work.

A combined theoretical approach, namely ONIOM (mB97xd/6–31G(d,p):am1) was used for the geometry optimization of adsorption states and transition states (TS). The mB97XD hybrid density function was developed to consider long-range-corrected hybrid functions, which implicitly accounted for empirical dispersion and could describe long-range dispersion interactions well with respect to the traditional density functional theory methods. This functional was also recently found to perform very well for the description of adsorption and reactions on zeolites. All energies reported herein were predicted at the mB97XD/6–31G(d,p) level based on the optimized structures.

The harmonic frequency calculations employing a partial Hessian vibrational analysis (PHVA) including the 22T high layer active acid sites and organic species were performed to check whether the stationary points found exhibit the proper number of imaginary frequencies. In frequency calculations, besides the atoms in high level and organic fragments, the constraints of the zeolite framework were also kept as the same in geometry optimizations, so one negative frequency would be observed for transition state point and none for the corresponding reactant and product. The Gibbs free energies (G) at 423 K (T) were calculated from harmonic frequencies.

$$G = H - TS = (E + ZPVE + H_{\text{rot}} + H_{\text{vib}} + H_{\text{trans}} + H_{\text{trans}}) - TS$$ (1)

It can be seen that Gibbs free energies (G) include contributions from electronic energies (E), zero-point vibrational energies (ZPVE), rotational enthalpies (H_rot), translational (H_trans) and rotational enthalpies (H_vib) for reactant molecules and the effect of entropy (T*S) which can more accurately describe the mechanism of ethanol dehydration reaction.

And then, the reaction rate constants (k) at 473 K were further obtained by transition state theory (TST):  

$$k = A \exp\left(\frac{-\Delta H_{\text{act}}}{RT}\right) = \frac{b_k T^2}{h} \exp\left(\frac{-\Delta S_{\text{act}}}{R}\right) \exp\left(\frac{-\Delta G_{\text{act}}}{RT}\right) = \frac{b_k T^2}{h} \exp\left(\frac{-\Delta G_{\text{act}}}{RT}\right)$$ (2)

where A is a pre-exponential factor, b_k is Boltzmann's constant, h is Planck's constant and T is the reaction temperature. All the geometry optimizations and frequency calculations were performed using the Gaussian 09 package.

**Data availability**

All the data supporting the findings of this study are available within the article and its supplementary information file or from the corresponding author upon reasonable request.

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**Author contributions**

X.Z, C.W. and X.L. Z prepared the samples and carried XRD and catalytic testing. Q.W, G.D.Q, and N.D.F performed and analyzed the ex-situ solid state NMR spectra. Y.Y.C performed the DFT calculations. X. Z, J.X, and F.D collected and analyzed the in situ NMR spectra; X.Z, C.W., Y.Y.C, J.X and F.D. wrote the manuscript, and all authors discussed the experiments and final manuscript.

**Additional information**

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**Competing interests:** The authors declare no competing interests.

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