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Introduction

Heterogeneous catalysts with acid–base properties, such as zeolites have been widely used in the chemical and petrochemical industry for catalytic cracking,¹ isomerization,² alkylation,³ and disproportionation.⁴ In aluminosilicates-type zeolites, the Brønsted acid site (BAS) and Lewis acid site (LAS) are responsible for the active sites in numerous catalytic reactions. Therefore, understanding the structures and properties of the acidic sites in zeolites is essential to exploring the reaction mechanism and to the optimization of their catalytic performance.^{5–7} It is well-known that the BAS is associated with the tetrahedral framework aluminum (FAL) in the form of a bridging hydroxyl (Si–OH–Al) in zeolites. Furthermore, the

The acidic nature of "NMR-invisible" tricoordinated framework aluminum species in zeolites[†]

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The unambiguous characterization of different acid sites in zeolites is of great importance for understanding their catalytic performance and the rational design of highly efficient zeolite catalysts. In addition to various well-characterized extra-framework Al species, a tri-coordinated framework aluminum species can also serve as a Lewis acid site in zeolites, which is "NMR-invisible" owing to its extremely distorted local environment. Here we provide a feasible and reliable approach to elucidate the acidic nature of the tri-coordinated framework Al in dehydrated H-ZSM-5 zeolites *via* sensitivity-enhanced two-dimensional (2D) multiple nuclear correlation NMR experiments coupled with trimethylphosphine oxide (TMPO) probe molecules. Two types of tri-coordinated framework Al sites have been unambiguously identified, which amount to 11.6% of the total Brønsted and Lewis acid sites. Furthermore, it was found that synergistic effects arising from the close spatial proximity between the tri-coordinated framework Al site and the Brønsted acid site lead to the generation of superacidity (with an acid strength stronger than 100% H_2SO_4) in the zeolite.

LAS is generally recognized as an extra-framework or framework aluminum species, formed upon calcination or steaming of zeolites.8,9 The properties of BAS associated with the tetrahedral framework Al and LAS associated with various extra-framework Al (EFAL) species have been extensively studied, from their structural characteristics¹⁰⁻¹⁴ to their functions in catalytic reactions.15-18 In particular, the LAS in zeolites has been proved to play an important role in the catalytic transformations of hydrocarbons such as the hydrogen transfer process,7,19,20 and also generates the Brønsted/Lewis acid synergy owing to the vicinity of the corresponding Al species in the local architecture of dealuminated zeolites.^{11,21-23} Although the tri-coordinated FAL is commonly considered as a LAS in zeolites, only a few characterizations²⁴⁻²⁶ have been attempted to confirm the existence of the "moisture sensitive" tri-coordinated Al species in the zeolite framework. Up to now, no experimental approach has been reported to probe the intrinsic acidic properties of this important framework Al species in the zeolite.

Solid-state NMR has emerged as an important technique for the characterization of heterogeneous catalysts at the atomic scale.^{22,27-31} The coordination state of both FAL and EFAL species in zeolites can be determined on the basis of onedimensional (1D) ²⁷Al magic-angle spinning (MAS) and 2D ²⁷Al multiple-quantum magic-angle spinning (MQMAS) NMR spectroscopy. 2D ¹H-¹H and ²⁷Al-²⁷Al double-quantum singlequantum (DQ-SQ) homonuclear correlation MAS NMR methods can provide in-depth structural information on the

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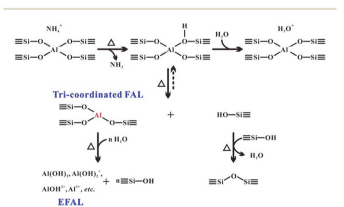
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spatial proximity of different acid sites in dealuminated zeolites, especially under high magnetic fields (>18.8 T).^{11,22,32} A recent report has shown that the perturbed aluminum species was attributed to terminal Al-OH in the hydrated zeolites, which was considered to be derived from hydrous tricoordination of the Al species.33 However, most of these measurements were performed under hydration conditions. In the case of non-hydrated zeolites, it is difficult to distinguish FAL species owing to the resolution/sensitivity issues from the quadrupolar nature of the ²⁷Al isotope. The stretch of the acidic proton in the BAS induces a larger distortion of the corresponding tetrahedral FAL in the H-form zeolites, leading to quadrupolar coupling constants ($C_{\rm O}$) up to 14-18 MHz.³⁴⁻³⁶ Generally, mild calcination and dehydration treatments of zeolites would lead to partial breaking (or hydrolysis) of the framework \equiv Al-O- bond and generation of a tri-coordinated FAL and framework silanol group as illustrated in Scheme 1.37-39 Although the neighboring framework Si-OH would interact with the tri-coordinated FAL to restore the tetrahedral FAL after cooling down of the sample to room temperature, partial Si-OH derived from the breaking of the aluminum oxygen bond that bears the proton may be further dehydroxylated by a neighboring defect site (Si-OH group), thus causing the irreversible formation of a tri-coordinated coordinated aluminum.²⁶ However, reducing the number of neighboring oxygen groups bound to FAL yields a dramatic distortion of its local environment, and the extreme asymmetry of its surrounding electric field results in a considerable line broadening of its ²⁷Al NMR resonance (usually with $C_{\rm Q} > 30$ MHz).^{33,40} Therefore, the direct observation of tri-coordinated FAL species in a dehydrated sample is generally impossible using conventional ²⁷Al MAS NMR owing to its huge quadrupolar broadening and relatively low concentration, and tri-coordinated FAL was suggested to be the "NMR-invisible" species in zeolites.26,33 After rehydration, the tri-coordinated FAL could be easily transformed into a symmetric tetra-coordinated FAL,33,38 which is usually indistinguishable from the FAL of BAS. Further hydrolysis of the framework Al-O bond leads to removal of the Al from the zeolite framework, forming EFAL species such as Al(OH)₃, $Al(OH)_2^+$, $AlOH^{2+}$, Al^{3+} , AlO^+ and so forth, which have been well characterized using solid-state NMR spectroscopy.11,22,32,36,41,42



Scheme 1 Formation route for the tri-coordinated FAL and EFAL species in zeolites.

The adsorption of basic probe molecules such as 2-¹³Cacetone and trimethylphosphine oxide (TMPO) is a feasible approach to investigate the acidity of zeolites.^{22,43-45} The great majority of studies utilized the chemical shift of certain elements (*e.g.* ¹³C or ³¹P) in probe molecules to determine the acidic features of zeolites, which were usually combined with density functional theory (DFT) calculations to reveal the relationship between chemical shifts and the corresponding acidic properties (*e.g.* type, distribution, and strength) on zeolites.^{22,45} However, it has previously been a great challenge to provide precise and localized information on the structure of the interface between the "guest" probe molecules and the specific acid sites of the "host" zeolite *via* analytical or spectroscopic techniques.

In this contribution, we provide a unique insight into the acidic nature of tri-coordinated FAL species on a H-ZSM-5 zeolite using solid-state NMR spectroscopy, and present a clear adsorption picture of the TMPO probe molecules on different framework Al species in a dehydrated H-ZSM-5 zeolite. In particular, the sensitivity-enhanced 2D ³¹P-²⁷Al heteronuclear correlation (HETCOR) MAS NMR technique⁴⁶ is employed to successfully discriminate the interactions between distinct adsorbed TMPO molecules and Al species in the zeolite framework. Thus, it is straightforward to ascertain the presence of LAS originating from the tri-coordinated FAL. Two tricoordinated FAL species are unambiguously identified using the advanced solid-state NMR technique. Further quantitative NMR analysis reveals that a considerable amount (ca. 11.6%) of this type of LAS exists in the framework of the dehydrated H-ZSM-5 zeolite even with moderate thermal treatments. Moreover, we demonstrate that the presence of the tri-coordinated FAL is responsible for the formation of superacidity (characterized by a down-field ³¹P chemical shift of the adsorbed TMPO up to 85-88 ppm) owing to the Brønsted/Lewis acid synergy in the zeolite framework which was further confirmed using 2D ²⁷Al-²⁷Al DQ-SQ homonuclear correlation spectroscopy and DFT calculations.

Results and discussion

Structural characterization of the dehydrated zeolites

The ²⁷Al MAS NMR spectra of the hydrated parent NH₄-ZSM-5 acquired at two different magnetic fields (18.8 and 11.4 T) shown in Fig. 1a and d both displayed a typical narrow line around 54 ppm, respectively, corresponding to relatively symmetrical tetrahedral Al sites in the zeolite framework. Usually a significantly reduced quadrupolar broadening of the ²⁷Al NMR line shape can be obtained at a high field ($B_0 = 18.8 \text{ T}$) and fast MAS speed ($v_R = 40 \text{ kHz}$) compared to that at a medium field ($B_0 = 11.7$ T) and moderate MAS speed ($\nu_{\rm R} = 10$ kHz). After calcination and dehydration, the ²⁷Al spectra of the dehydrated H-ZSM-5 zeolite exhibited a very broad resonance (Fig. 1b and e) with a full width of ca. 20 kHz even at 18.8 T, indicative of a dominant contribution from the large second-order quadrupolar broadening. Previous studies have demonstrated that this broad signal was mainly attributed to the more-distorted tetrahedral FAL species with a $C_{\rm Q}$ \approx 16 MHz from the

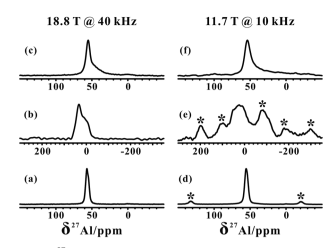


Fig. 1 1D ²⁷Al MAS NMR spectra of parent NH₄-ZSM-5 (a) and (d); calcined and dehydrated H-ZSM-5 (b) and (e); and TMPO-adsorbed H-ZSM-5 with P/Al = 0.42 (c) and (f); obtained at $B_0 = 18.8$ T (a)–(c) and $B_0 = 11.7$ T (d)–(f). The asterisks denote spinning sidebands.

Brønsted acidic site on the dehydrated zeolites.^{35,36} In fact, dealumination is inevitable in most sample preparation procedures, while the mild vacuum or flow temperature-programmed methods used here could make that framework dealumination rarely occurs, which would largely reduce the interference of EFAL species in the study of tri-coordinated FAL.

The XRD patterns (Fig. S1[†]) of the ZSM-5 zeolites indicated no apparent change in the crystallinity after these treatments. To estimate the amount of EFAL in our calcined sample, the parent and calcined forms of these zeolites were treated using 1 M aqueous NaNO₃ at 353 K three times. As the Na⁺ cations can balance the charge of the framework AlO_4^- tetrahedra, the Na/ Al ratio is a reliable parameter to assess the variation of the fraction of the tetrahedral framework Al.47 The Na/Al ratio measured using inductively coupled plasma (ICP) was 1.01 and 0.98 respectively for the parent NH₄-ZSM-5 and dehydrated H-ZSM-5 zeolites treated with Na⁺ exchange, suggesting that most of the Al species were still located in the zeolite framework. This is also supported by a slight loss (1%) of framework \equiv Si-O-Al moieties in the dehydrated zeolite via 1D ²⁹Si MAS and 2D ²⁹Si{¹H} HETCOR NMR analysis (Fig. S2 and S3[†]). The quantitative ²⁹Si MAS spectra shown in Fig. S3[†] indicates an increase of the Si-OH groups in the dehydrated H-ZSM-5 zeolite compared with those in the parent NH₄-form spectra. In addition, further ¹H MAS NMR analysis (Fig. S4[†]) showed that the number of Si-OH groups decreased when raising the dehydration temperature from 573 to 673 K, suggesting that the dehydroxylation of Si-OH groups may occur as depicted in Scheme 1.

Framework aluminum studied using 2D ²⁷Al 3QMAS

In our measurements, TMPO molecules were adsorbed onto the H-ZSM-5 zeolite at a concentration much lower than that of the framework acid sites, meaning the interference between the TMPO molecules themselves can be largely avoided. Interestingly, in addition to the narrow 27 Al signal at *ca.* 54 ppm, a new broad shoulder feature (at *ca.* 45 ppm) was visible in the 1D 27 Al

MAS spectra of the dehydrated zeolite loaded with TMPO (P/Al = 0.42, Fig. 1c and f). As the tri-coordinated FAL with Lewis acidity (if present) can directly interact with the oxygen atom of TMPO, which would significantly decrease the asymmetry of the local environment of the Lewis acidic FAL, yielding its "NMRdetectable" ²⁷Al resonance on non-hydrated zeolites. Then, a 2D ²⁷Al 3QMAS experiment at 18.8 T was employed to discriminate between specific aluminum sites in the dehydrated and TMPOloaded sample (Fig. 2a). Three signals, denoted as Al_a, Al_b and Al_c, were well-resolved with isotropic chemical shifts of 54.4, 51.6 and 58.8 ppm, respectively. For the dehydrated zeolite with a lower TMPO concentration (P/Al = 0.18), an obvious decline of the relative proportion of Al_c and Al_b signals (Fig. 2b, S10[†]) was evident. The NMR parameters deduced from the 2D spectrum are listed in Table 1. The Al_a signal with $P_{\rm O} = 2.3$ MHz could be attributed to the regular tetrahedral FAL species from Brønsted acidic sites analogous to that of the NH₄-form or hydrated zeolites. The local structure of the tetrahedral FAL in dehydrated H-ZSM-5 became more symmetrical stemming from

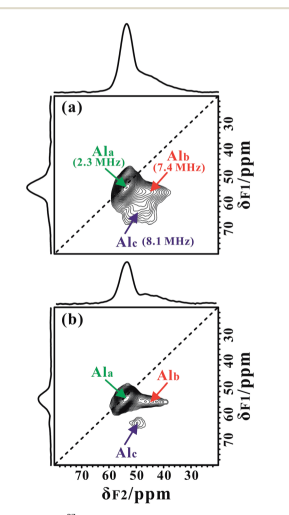


Fig. 2 Sheared 2D ²⁷Al 3QMAS NMR spectra of dehydrated H-ZSM-5 zeolites with different TMPO loadings, (a) P/Al = 0.42 and (b) P/Al = 0.18, obtained at $B_0 = 18.8$ T. Skyline projections are drawn along the F1 and F2 dimensions. The determined quadrupolar parameter $P_{\rm Q}$ are indicated in the brackets for different Al species.

Table 1 NMR parameters obtained from the ²⁷Al MQMAS spectra

Site	$\delta_{\mathrm{F2}}{}^{a}\!/\mathrm{ppm}$	$\delta_{\mathrm{F1}}{}^{a}/\mathrm{ppm}$	$\delta_{\rm CS}{}^b/{\rm ppm}$	$P_{\rm Q}^{\ b}/{\rm MHz}$	$C_{\rm Q}^{\ \ c}/{\rm MHz}$
Al _a	53.6	54.8	54.4	2.3	2.2
Al_b	44.1	56.0	51.6	7.4	7.1
Al_c	49.7	64.1	58.8	8.1	7.8

^{*a*} δ_{F2} and δ_{F1} represent the center of gravity of each particular species measured along the direct dimension F2 and isotropic dimension F1 on the sheared spectra, respectively. ^{*b*} Isotropic chemical shift δ_{CS} and quadrupole interaction product P_Q of each aluminum species were deduced from δ_{F2} and δ_{F1}. ^{*c*} The quadrupolar coupling constant (C_Q) was extracted from the slice of the 3QMAS spectra by fitting the corresponding second-order quadrupolar line shape ($\eta = 0.5$) with the DMFIT program.

a shrinkage of the Al–O(H) bond length in BAS after the TMPO adsorptions. According to their isotropic chemical shifts, Al_b and Al_c should both be associated to tetrahedral FAL species as well, while exhibiting a larger inhomogeneous second-order quadrupolar broadening ($P_Q > 7$ MHz). However, their assignments were ambiguous, as they could be from either the FAL in another type of BAS, or the tri-coordinated FAL of LAS directly bound to a TMPO molecule, yielding a new distorted tetrahedral FAL state. In particular, the Al_b and Al_c resonances contributed to a considerable fraction in the quantitative ²⁷Al NMR spectra of the dehydrated zeolites with different TMPO loadings (Fig. 1c). Therefore, further experiments are required to identify the host–guest correlations/interactions between the adsorbed TMPO probe molecules and the framework of zeolites.

Acid properties analyzed using 2D ³¹P{¹H} HETCOR

For the acidity characterization using TMPO probe molecules, dichloromethane (CH₂Cl₂) is usually used as the solvent in order to disperse the TMPO molecules into zeolite well.45,48 Although the evacuation at 323 K seems to be insufficient to dispose of all of the solvent, in the 2D ${}^{13}C{}^{1}H$ and ${}^{31}P{}^{1}H$ HETCOR spectra of the TMPO/H-ZSM-5 zeolite (Fig. S5[†]) acquired with a relatively short cross polarization (CP) contact time (4 ms or 1 ms) there is no correlation between the CH_2Cl_2 and TMPO molecules, indicating that the residual solvent and adsorbed TMPO molecules are well separated. Furthermore, the ²⁷Al MAS NMR spectrum of the H-ZSM-5 zeolite with only CH₂Cl₂ adsorbed (Fig. S6[†]) still exhibits a very broad resonance $(\Delta \delta \approx 90 \text{ ppm})$, indicating that the residual CH₂Cl₂ hardly interacts with the acidic sites on the zeolite, and thus has almost no influence on the acidity characterization using TMPO probe molecules. Note that, after removing the residual solvent on the TMPO/H-ZSM-5 zeolite at a higher temperature (363 K, Fig. S7[†]), the ²⁷Al MAS NMR spectrum of TMPO/H-ZSM-5 was the same as that treated at 323 K (Fig. S8[†]), which clearly indicated that the influence of the solvent (CH₂Cl₂) residues on the framework Al species can be negligible.

The 1D ³¹P MAS NMR spectra of the H-ZSM-5 zeolites with different TMPO loadings (P/Al = 0.18 and 0.42) are shown in Fig. 3a and c, respectively. Up to six characteristic peaks at 88, 85, 76, 69, 65 and 51 ppm were identified from the ³¹P MAS NMR spectra by using Gaussian deconvolutions (also see Table

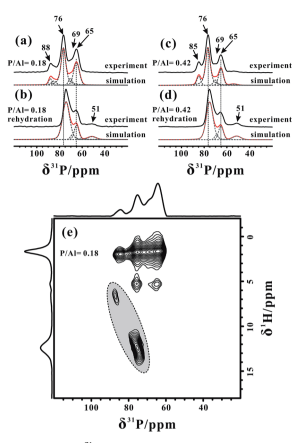


Fig. 3 1D one-pulse ³¹P MAS spectra of dehydrated H-ZSM-5 zeolites with different TMPO loadings (a) and (c), and subsequent rehydration (b) and (d). 2D ³¹P{¹H} CP HETCOR MAS NMR spectrum (e) of the dehydrated H-ZSM-5 zeolite with a low TMPO loading (P/Al = 0.18).

S1[†]). Indeed, quantitative analyses of ²⁷Al MAS NMR are difficult owing to either different quadrupolar broadenings of distinct FAL species or only partial acid sites interacting with the TMPO probe molecules, however, here the ${}^{31}P(I = 1/2)$ MAS NMR spectra can provide quantitative information on the framework BAS and LAS that react with the TMPO molecules. The 2D ³¹P{¹H} CP-HETCOR spectrum of the TMPO/H-ZSM-5 zeolite (P/Al = 0.18) acquired with a CP contact time of 8 ms in Fig. 3e exhibited two broad distributions of the correlation peaks in "sloped" ovals (dotted line) centered at (76, 12.2) ppm and (88, 6.4) ppm, respectively, which could be assigned to distinct protonated TMPOH⁺ ionic pair complexes. These correlations were also observable when a short CP contact time (1 ms) was used (see Fig. S5[†]). The interaction of the adsorbed TMPO molecules with BAS in the H-ZSM-5 zeolite will result in a downfield shift of the ¹H resonance of the acidic proton, and the electron cloud density surrounding the ³¹P nucleus in the TMPO molecule decreases with the increasing acidic strength of BAS, leading to movement of the ³¹P resonance towards a higher chemical shift (downfield).^{22,44,45} Furthermore, it was also found that the ³¹P chemical shift of the adsorbed TMPO molecules linearly increases with the increase of the Brønsted acid strength, and a ³¹P chemical shift of *ca.* 86 ppm was demonstrated for the threshold of superacidity (with acid strength

stronger than 100% H₂SO₄).⁴⁵ Therefore, the ³¹P peak at $\delta_{^{31}P} =$ 88 ppm reflected the strongest acid site with a strength over the threshold of superacidity in the zeolite. In addition, the correlation peaks between all ³¹P signals and methyl groups ($\delta_{^{1}H} =$ 1.8 ppm) of the adsorbed TMPO molecules can be observed at all times. As the ¹H–¹H spin diffusion will promote the detection of long range interactions in the case of a long CP contact time (8 ms), weak correlations between the protons of the residual CH₂Cl₂ solvent ($\delta_{^{1}H} \approx 5.4$ ppm) and the ³¹P of TMPO molecules were also detectable in the 2D spectrum.

However, as illustrated in the 2D ³¹P{¹H} HETCOR spectrum of Fig. 3e, even though the ¹H-¹H spin diffusion also contributed to the ¹H-³¹P correlations when the CP contact time was set to 8 ms, no correlation can be observed for the ³¹P signals around 65 ppm with the ¹H signals of BAS, which suggested that the ³¹P signals do not originate from the protonated TMPOH⁺ complexes. It is generally recognized that H₂O could competitively react with the Lewis acid sites, such as the tri-coordinated Al species, to form weak BAS.48 After exposing our samples to humidity for 2 h, the corresponding ³¹P MAS NMR spectra in Fig. 3b and d exhibited an apparent decline in the relative intensity of the signals at 65 and 69 ppm (also see Table S1⁺), and the signal at 88 ppm (or 85 ppm for the sample with P/Al =0.42) disappeared completely. In addition, the peak at 76 ppm was slightly shifted upfield with a broader distribution, accompanying a notable increase of the peak at 51 ppm from the TMPO adsorbed on the weak acid sites. Therefore, we can reasonably consider that the ³¹P signals at 65 and 88 ppm (or 85 ppm) are both associated with Lewis acid sites directly or indirectly. Although the peaks at ca. 88 ppm have proved to be from TMPO molecules adsorbed on Brønsted acid sites, it seems that BAS together with the neighboring LAS should contribute to the formation of the superacidity (e.g. $\delta_{^{31}P} = 88$ ppm), indicative of the presence of a synergy effect. Accordingly, after humidity exposure of the dehydrated zeolite, the LAS was preferentially hydrated, thus the influence on its neighboring BAS (the superacid feature) vanished, which in turn caused the BAS to revert back to its original acidity. The two ³¹P signals at 65 and 69 ppm were probably from the TMPO molecule adsorbed on the Lewis acidic FAL species. Concerning the abundance of Al_b and Al_c in Fig. 2 (also in Fig. S10[†]), we proposed that the aforementioned ³¹P and ²⁷Al species belong to the interacting phosphorus-aluminum pairs in the Lewis acid complexes, $(CH_3)_3PO\cdots Al \equiv$. In addition, we also acquired the 2D ³¹P{¹H} HETCOR spectrum (Fig. S9[†]) of the TMPO/H-ZSM-5 zeolite with the complete removal of solvent at 363 K, which exhibited similar ¹H-³¹P correlations between the TMPO molecules and the framework acid sites in the zeolite compared with that treated at 323 K, and further excluded the influence of the residual solvent.

Tri-coordinated framework Al characterized using ${}^{31}P{}^{27}Al$ HMQC and ${}^{31}P{}^{27}Al$ S-RESPDOR

To verify this assumption, the Lewis complex $(CH_3)_3PO\cdots Al \equiv$ was identified directly using our recently developed ${}^{31}P-{}^{27}Al$ population-transfer dipolar-mediated HMQC (PT-D-HMQC) experiment which can establish the direct correlation/ connectivity between TPMO and various Al species. As shown in Fig. S5,† the 1D ³¹P{²⁷Al} PT-D-HMQC MAS NMR spectra *via* ³¹P \rightarrow ²⁷Al \rightarrow ³¹P filtering on TMPO-loaded H-ZSM-5 zeolite (P/ Al = 0.42) revealed that the two main ³¹P resonances around 76 and 65 ppm were correlated with the ²⁷Al species in the zeolite. When using a short recoupling time ($\tau_{mix} = 1.33$ ms), the peaks at *ca*. 65 ppm became more intense compared to that at *ca*. 76 ppm (Fig. S11†). This indicated the stronger ³¹P-²⁷Al dipolar interaction of the corresponding spin pairs, further suggesting the presence of the Lewis complex [(CH₃)₃PO···Al \equiv].

Notably, the 2D ³¹P{²⁷Al} PT-D-HMQC spectrum acquired at 11.7 T for the TMPO-adsorbed H-ZSM-5 zeolite in Fig. 4b clearly illustrated two correlation groups. The correlation peaks at around (76, 54) ppm were associated with TMPO adsorbed on the bridging hydroxyl (Si-OH-Al) proton (i.e. the Brønsted acid sites). Importantly, other correlations indicated that the ³¹P resonance peaks at 65 and 69 ppm are both correlated with broad ²⁷Al resonances from distorted four-coordinated aluminum, reflecting the presence of two types of Lewis acid sites. We also conducted 2D ²⁷Al{³¹P} D-HMQC experiments at 18.8 T. Owing to the short transverse relaxation times (T_2) of the ²⁷Al resonances, we could only observe ²⁷Al-³¹P correlations from the Lewis complex $[(CH_3)_3PO\cdots Al \equiv]$ in Fig. 4c, in which the ³¹P signals at 65 and 69 ppm were correlated with Al_b and Al_c, respectively (Fig. 4d and e). Meanwhile, a reduction of the TMPO concentration led to a decrease in the proportion of the ³¹P signal at 69 ppm relative to that at 65 ppm (Table S1[†]), this was consistent with the relative proportion variation of Al_c and Al_{b} (Fig. 2). Therefore, we can confirm that Al_{c} and Al_{b} are both from tri-coordinated FAL species bound to the O atom of the TMPO molecules, which were discriminated well from their different Lewis acid strengths.

Owing to its low content and Brønsted acid feature, we did not observe the ³¹P-²⁷Al correlations from the ³¹P signal at 88 ppm (or 85 ppm) in Fig. 4. Fortunately, the symmetry-based saturation-pulse resonance-echo double-resonance (S-RESPDOR) method⁴⁹ can be employed here to estimate the ³¹P-²⁷Al heteronuclear dipolar interactions. The results are usually analyzed by plotting the signal fraction $\Delta S/S_0 = (S_0 - S')/(S_0 - S')$ S_0 (S' and S_0 represent the signal intensity with and without dipolar dephasing, respectively) as a function of the recoupling time τ . The ³¹P difference spectrum ($\Delta S = S_0 - S'$, shown in Fig. S12[†]) obtained using 1D ³¹P{²⁷Al} S-RESPDOR experiments still suggested spatial interactions/proximities between all of the ³¹P species of the adsorbed TMPO molecules (including those giving rise to the resonance at 88 ppm) and ²⁷Al species in the framework of H-ZSM-5. Furthermore, the reduced concentration of TMPO molecules (P/Al = 0.18) allowed us to estimate the distances between ³¹P atoms and ²⁷Al atoms, as we could approximately consider the Lewis complex $[(CH_3)_3PO\cdots Al \equiv]$ with ³¹P signals at 65 ppm or the Brønsted acid complex $[(CH_3)_3PO\cdots H-OA] \equiv]$ with ³¹P signals at 76 ppm as "isolated" phosphor-aluminum spin pairs in this case. Therefore, the P-Al distance of the former was determined to be 2.9 \pm 0.2 Å by fitting the dephasing curve (Fig. 5a), which was rationalized in terms of the structural model of the TMPO molecule adsorbed

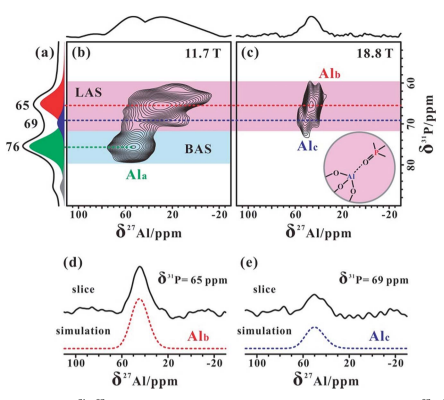


Fig. 4 1D one-pulse ³¹P MAS (a) and 2D ³¹P{²⁷Al} PT-D-HMQC MAS (b) NMR spectra recorded at 11.7 T, and 2D ²⁷Al{³¹P} D-HMQC MAS NMR spectrum (c) recorded at 18.8 T of the dehydrated H-ZSM-5 zeolite (Si/Al = 25) with adsorbed TMPO (P/Al = 0.42). Selected slices with simulations are shown in (d) and (e)

on the framework tri-coordinated aluminum species (Fig. 5c). The later complex had a reasonable P–Al distance of 4.1 ± 0.4 Å (Fig. 5b), in which the ³¹P atom was about four bonds away from the Al site (Fig. 5d). We also considered the dipolar coupling networks of multiple-spin systems (shown in Fig. S13[†]). The representative three spin systems are depicted in Fig. S13b-d.[†] In the normalized dipolar dephasing curve with respect to the first extremum (or maximum) in each spin system, it was found that the dephasing curves of the observed ³¹P signals in multispin systems have a similar upward trend to that of the isolated ³¹P-²⁷Al spin pair. This result suggests that the upward trend of the dipolar dephasing curve is dominated by the shortrange (strongest) ³¹P-²⁷Al dipolar interaction. The long-range couplings have little impact on the measurement of the short distance, which mainly affects the oscillation of the dephasing curves after reaching its first extremum. Moreover, the ³¹P-²⁷Al distances extracted using the dipolar dephasing curves were in good agreement with the DFT calculations, in which the theoretically predicted ³¹P-²⁷Al distances of the Lewis acid complex $[(CH_3)_3PO\cdots Al \equiv]$ and Brønsted acid complex $[(CH_3)_3PO\cdots H OAl \equiv$] were 3.1 Å and 4.1 Å (Fig. 5c and d), respectively.

Although it is difficult to calculate the accurate amount of the tri-coordinated FAL in our sample, an estimation of the lower limit of the concentration via the quantitative analysis of the ³¹P MAS NMR spectrum (Fig. 3) is possible using the following equation:

$$\Omega(\text{FAL}^{\text{III}}) = \chi(\text{P/Al}) \times \Omega(^{31}\text{P}_{\text{LAS}})$$

In which $\Omega(FAL^{III})$ is the fraction of the tri-coordinated FAL species which interacted with the TMPO in the total Al content, $\Omega(^{31}P_{LAS}) = 27.0\%$ is the proportion of the ^{31}P signals at 65 and 69 ppm in the quantitative ³¹P MAS NMR spectrum (see Table S1[†]), and $\gamma(P/Al) = 0.42$ is the atomic ratio of the P to Al of the sample obtained using ICP. Therefore, the calculated $\Omega(FAL^{III})$ = 11.6% suggested that a considerable amount of LAS formed and was located in the framework of the dehydrated H-ZSM-5 zeolite, this was in agreement with the relative content of the Lewis acid sites (13.3%) determined using our Fourier transform infrared (FT-IR) analysis of the pyridine adsorption measurements (Table 2 and Fig. S14[†]).

Brønsted/Lewis synergy effect studied using 2D ²⁷Al DQ-SQ NMR and DFT calculations

We have successfully obtained a high-resolution ²⁷Al-²⁷Al DQ-SQ homonuclear correlation spectra of the hydrated zeolites (HY, MOR, ZSM-5, etc.) using our developed BR2¹₂ recoupling scheme at a high magnetic field,^{32,41} in which only Al species from the framework BAS and extra-framework LAS were considered. In the case of non-hydrated zeolite catalysts, the broadening and overlap of the ²⁷Al quadrupolar patterns usually results in considerable barriers to the implementation of the 2D ²⁷Al DO–SO experiment. Fortunately, the improved resolution of the ²⁷Al MAS NMR spectra of the dehydrated H-ZSM-5 zeolite after TMPO adsorptions offers an opportunity to investigate the

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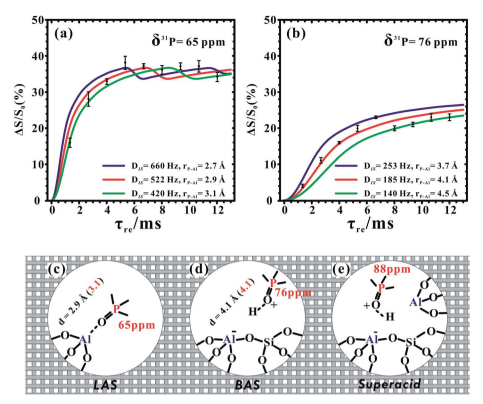


Fig. 5 ${}^{31}P{}^{27}Al$ S-RESPDOR built-up curves of TMPO adsorbed on (a) LAS (tri-coordinated FAL) and (b) BAS of dehydrated H-ZSM-5 zeolite, fitted using an analytical formula. D_{IS} and r_{P-Al} represent the dipolar interaction constant and internuclear distance of the ${}^{31}P{}^{-27}Al$ spin pair, respectively. Three adsorption structure models of: (c) TMPO adsorbed on tri-coordinated FAL, forming a distorted tetrahedral FAL; (d) TMPO adsorbed on a BAS; and (e) TMPO adsorbed on a BAS with a tri-coordinated FAL in close proximity. The red numbers in parentheses are the distances obtained using DFT calculations.

intrinsic structure of different framework Al species under the dehydrated state, the acidic natures of which have been clearly characterized. Fig. 6a displays the ²⁷Al-²⁷Al DQ-SQ MAS NMR spectrum of the TMPO-loaded H-ZSM-5 with P/Al = 0.42 at 18.8 T. Two auto-correlation peaks can be observed, in which the intense blue part indicates that the tetra-coordinated framework Al species (Al_a) is in close proximity to another (Al_a) , and the weak orange part is ascribed to the spatial proximity of a low content of Al_b-Al_b or Al_c-Al_c pairs. Note that, it is still hard to discriminate the Al_b and Al_c in the ²⁷Al DQ-SQ MAS NMR spectrum. In addition, the appearance of an intense cross-peak pair between Al_a and Al_b (red part in Fig. 6a) provided direct evidence of the spatial proximity between the framework Brønsted and Lewis acid sites, in which Al_a and Al_b had been confirmed from the Brønsted acid complex [(CH₃)₃PO···H- $OAl \equiv]$ and the Lewis acid complex $[(CH_3)_3PO\cdots Al \equiv]$ in the TMPO-loaded H-ZSM-5, respectively. Therefore, it implied that the superacidity characterized by the ³¹P NMR signals at 8588 ppm (Fig. 3) on the specific BAS could be derived from the influence of the neighboring tri-coordinated FAL (LAS), generating a Brønsted/Lewis synergy effect. The assignment was also confirmed by the absence of the ³¹P signals at 85–88 ppm on a quantitatively TMPO-adsorbed H-ZSM-5 zeolite with a low Si/ Al of 140 (see Fig. S15†), which has a significantly reduced probability of a Al–Al pair being in close proximity in the zeolite framework owing to the much lower Al content.

In order to gain further insights into the Brønsted/Lewis synergy effect, we also performed DFT calculations (see Fig. S16†). As shown in the local theoretically optimized structures (Fig. 6b and c), for TMPO adsorbed on the BAS with a neighboring tri-coordinated FAL site, a ³¹P chemical shift of 83 ppm was theoretically predicted, while for TMPO adsorbed on the isolated BAS, a ³¹P chemical shift of 76 ppm was achieved, this was consistent with our experimental observations. The calculated ³¹P and ²⁷Al isotropic chemical shifts for different adsorption models are listed in Table S3.† Except for

Table 2Concentration of BAS and LAS in the H-ZSM-5 (Si/Al = 25) sample							
Sample	$n (BAS + LAS)^a \mu mol g^{-1}$	$n (BAS)^b \mu mol g^{-1}$	$n (LAS)^c \mu mol g^{-1}$				
H-ZSM-5/25	647.2	511.4	86.1				
^a Calculated using ICP-	OES. ^b Determined using ¹ H MAS NMR. ^c Calc	lated from pyridine-FTIR.					

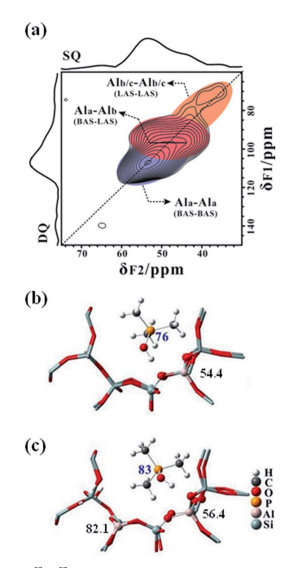


Fig. 6 2D ²⁷Al–²⁷Al DQ–SQ MAS NMR spectrum (a) of dehydrated and TMPO-loaded H-ZSM-5 zeolite (P/Al = 0.42), recorded at 18.8 T with a spinning speed of 20 kHz. Local theoretically optimized structures of TMPO adsorbed at the isolated Brønsted site (b) and the Brønsted/Lewis acid synergetic site (c) with a tri-coordinated FAL in close proximity. The predicted ³¹P chemical shifts of the TMPO (blue numbers) and ²⁷Al chemical shifts of the corresponding acid sites (black numbers) are labeled in ppm.

the unreacted "NMR-invisible" framework LAS (tri-coordinated FAL), the chemical shift parameters extracted from the DFT calculations were consistent with our experimental results. These results indicate that the synergistic effect between the framework BAS and the framework LAS (tri-coordinated FAL) owing to their spatial proximity (Fig. 5e) leads to an enhancement of the Brønsted acidity and thus generates the super-acidity in the H-ZSM-5 zeolite.

Conclusions

Tri-coordinated framework aluminum species can also serve as Lewis acid sites in principle, however, one of the fundamental

challenges is to observe and distinguish these tri-coordinated framework aluminum species in zeolites. Their relatively low concentration and largely distorted local environment make them "NMR-invisible" in zeolites, thus experimental studies on their acidic properties are rarely reported. Benefiting from current state-of-the-art multi-nuclear and multi-dimensional solid-state NMR spectroscopy, unique insights into the structure and properties of tri-coordinated FAL species acting as Lewis acid sites in H-ZSM-5 have been achieved in conjunction with TMPO probe molecule adsorptions. As the unsaturated tricoordinated FAL species can act with the basic TMPO probe molecules to form distorted tetrahedral FAL sites, two tricoordinated FAL species with different Lewis acidities were discriminated using sensitivity-enhanced 2D ³¹P-²⁷Al D-HMQC MAS NMR experiments. In particular, the adsorption models of the TMPO probe molecules on distinct acidic sites in H-ZSM-5 were clearly illustrated, and the strong ³¹P-²⁷Al dipolar interaction from the Lewis complex $[(CH_3)_3PO\cdots Al \equiv]$ was confirmed using measurements of the distance between the corresponding ³¹P and ²⁷Al atoms. Furthermore, the remarkably improved resolution and sensitivity of the ²⁷Al MAS NMR spectrum of the TMPO-adsorbed H-ZSM-5 zeolite allowed us to obtain a wellresolved 2D ²⁷Al DQ-SQ homonuclear correlation spectrum, and thus to understand the structure of the different framework acid sites in the non-hydrated zeolite. Consequently, the superacidity of BAS (with acid strength stronger than 100% H₂SO₄) induced by the synergistic effect between the framework BAS and framework LAS (tri-coordinated FAL) was evidenced by their close spatial proximity in dehydrated H-ZSM-5 zeolites and the observation of a ³¹P chemical shift of up to 85-88 ppm for the adsorbed TMPO. Notably, apart from the extra-framework Al as a LAS in the zeolite, the existence of a considerable amount of tri-coordinated framework Al serving as LAS in the zeolite framework suggests that further studies are needed to understand its catalytic function, which may be essential to rationally design highly efficient zeolite catalysts. The detailed characterization of the tri-coordinated framework Al in H-ZSM-5 in this work provides a useful strategy to explore the acidic nature of this type of "moisture-sensitive" species in zeolite catalysts, and may shed light on the experimental exploration of its structure-function relationship in zeolite catalysis.

Experimental section

Sample preparation

The parent NH₄-ZSM-5 with Si/Al = 25 and Si/Al = 140 was purchased from Zeolyst International. The sample was calcined at 773 K under a dry air atmosphere for 10 h to remove the template and the H-form of ZSM-5 was obtained. Prior to the adsorption of TMPO, the sample was dehydrated on a vacuum line. The temperature was gradually increased at a rate of 1 °C min⁻¹ and the sample was kept at a final temperature of 673 K at a pressure below 10⁻³ Pa overnight. The detailed procedures involved in introducing the TMPO probe molecule onto the sample can be found elsewhere.⁴⁵ In brief, a known amount of TMPO dissolved in anhydrous CH₂Cl₂ was first added into a vessel containing the dehydrated H-ZSM-5 sample

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in a N₂ glovebox, followed by removal of the CH₂Cl₂ solvent by evacuation on a vacuum line at 323 or 363 K. To ensure a uniform adsorption of the TMPO molecules in the pores/ channels of the zeolites, the sealed vessel was further subjected to a thermal treatment at 453 K for 2 h. Finally, the sample was transferred into a ZrO_2 MAS rotor in the N₂ glovebox prior to the solid-state NMR experiments.

ICP experiment

Atomic emission spectroscopy (AES-ICP) was used to determine the relative amounts of silicon, aluminum and phosphorus in the TMPO-loaded H-ZSM-5 samples. The Na and Al contents were measured using AES-ICP for a 10 mg sample dissolved in 50 mL HF solution.

FT-IR of pyridine adsorption

The FT-IR analysis of the pyridine adsorption measurements were conducted on a Bruker Tensor 27 spectrometer. About 20 mg of the sample was pressed into a self-supported wafer with a diameter of 13 mm. Prior to the measurement, the catalysts were evacuated to 10^{-2} Pa at 673 K for 2 h. After cooling down to room temperature, pyridine vapor was introduced into the sample cell at room temperature for 30 min to allow it to reach equilibrium, the residual pyridine was removed by vacuum. The FT-IR spectra of the pyridine-adsorbed samples were measured at 423, 523, and 623 after evacuation for 30 min, respectively.

Solid-state NMR experiments

For all 2D HETCOR experiments, the isotope within the braces indicates that it is indirectly detected in the F1 dimension. 1D ¹H MAS NMR and ³¹P MAS NMR experiments were performed on an 9.4 T Bruker AVANCE-III spectrometer using 4 mm rotors at a spinning frequency of 10 kHz (operating at a Larmor frequency of 399.3 MHz and 161.7 MHz, respectively). The ¹H single-pulse acquisition was employed with a pulse width of 3.9 μ s (*ca.* $\pi/2$ pulse), a recycle delay of 3 s, and 32 scans. The ³¹P single-pulse acquisition was employed with a pulse width of 3 μ s (*ca.* $\pi/2$ pulse), a recycle delay of 30 s, and 1024 scans, in which the RF amplitude of ¹H TPPM decoupling was set to *ca.* 50 kHz.

The ²⁹Si MAS NMR experiments were performed on an 11.7 T Bruker AVANCE-III spectrometer using 7 mm rotors at a spinning frequency of 6 kHz (operating at a Larmor frequency of 99.4 MHz). The ²⁹Si single-pulse acquisitions were employed with a pulse width of 8.4 μ s (*ca.* $\pi/2$ pulse), a recycle delay of 30 s, and 320 scans, in which the RF amplitude of ¹H TPPM decoupling was set to *ca.* 38 kHz.

The single-pulse ²⁷Al MAS NMR data were collected on an 11.7 T Bruker AVANCE-III spectrometer with a 4 mm doubleresonance probe at a spinning rate of 10 kHz, a pulse width of 0.4 μ s (*ca.* π /6 pulse), a recycle delay of 1 s, and 2048 scans. To improve the resolution, the single-pulse ²⁷Al MAS NMR data were also collected on an 18.8 T Bruker AVANCE-III spectrometer (corresponding to a ²⁷Al Larmor frequency of 208.6 MHz) with an 1.9 mm double-resonance probe at a spinning rate of 40 kHz, a pulse width of 0.3 μs (ca. $\pi/9$ pulse), a recycle delay of 1 s, and 4096 scans.

The 2D ²⁷Al triple-quantum (3Q) MAS z-filtering NMR experiments with fast amplitude modulation (FAM) enhancement⁵⁰ were also carried out at 18.8 T. The pulse duration was set to 3.6 µs for the first strong pulse with $\nu_{\rm RF} \approx 125$ kHz. A fast amplitude-modulated (FAM) pulse train consisting of four loops of two short pulses ($\nu_{\rm RF} \approx 125$ kHz) and two delays with the same duration of 0.7 µs was used to convert $p = \pm 3$ to ± 1 coherences. The "soft" $\pi/2$ pulses were set to 10.4 µs.

The 1D ³¹P{²⁷Al} S-RESPDOR and 2D ³¹P{²⁷Al} PT-D-HMQC MAS NMR experiments were performed on an 11.7 T Bruker AVANCE-III spectrometer using commercial 7 mm rotors with an o-ring cap at a spinning frequency of 6 kHz, tuned and matched to the ¹H, ³¹P and ²⁷Al Larmor frequencies (500.6 MHz, 202.6 MHz and 130.5 MHz, respectively). ¹H-³¹P CP with a contact time of 8 ms was employed to prepare the initial ³¹P signal.

For the ³¹P{²⁷Al} S-RESPDOR experiments, a saturation pulse on the ²⁷Al channel with an amplitude of *ca.* 25 kHz and a length of 166.67 μ s ($T_{\rm R}$) was irradiated at *ca.* 30 ppm to transfer the ³¹P-²⁷Al interactions. For the 2D ³¹P{²⁷Al} PT-D-HMQC experiments, repetitive sideband-selective (SS) WURST-80 adiabatic pulses with a length of 166.67 µs were employed on the Al channel during SR4 recoupling to accelerate the coherence transfers between ³¹P and ²⁷Al. The values for the peak RF amplitude and offset of WURST-80 were optimized with $(v_1^{\text{max}}, v_{\text{offset}}) = (7.5, 175) \text{ kHz on the }^{27}\text{Al channel. For the 2D }^{27}\text{Al}$ {³¹P} D-HMQC and ²⁷Al DQ-SQ MAS NMR experiments, the central transition (CT) enhancement of the initial magnetizations (27Al) was obtained by using a SS-WURST-80 irradiation with a length of 1 ms. More details on the NMR parameters of the 2D or double-resonance experiments can be found in Table S2.†

All simulations of the 1D MAS NMR spectra or slices were performed with the DMFIT software.⁵¹ The chemical shifts for ¹H, ³¹P, ²⁹Si and ²⁷Al were referenced to adamantane, 85% H_3PO_4 , kaolinite and 1 M aqueous Al(NO₃)₃, respectively.

DFT calculations

All of the structures of the ZSM-5 zeolite were represented by a 64 T model (containing the complete double 10-membered ring (MR) intersection pores formed by 10-MR straight and 10-MR sinusoidal pore channels), which were extracted from their crystallographic structural data (http://www.iza-structure.org/ databases/). The terminal Si-H was fixed at a bond length of 1.47 Å, oriented along the direction of the corresponding Si–O bond. The geometries of TMPO adsorbed at the isolated Brønsted acid site and the Brønsted/Lewis synergetic site of ZSM-5 were optimized over the 64 T cluster model by the combined theoretical ONIOM method. In the calculations, the TMPO and the high layer atoms of the zeolite framework were allowed to fully relax with the dispersion-corrected ω B97XD functional⁵² with a 6-31G (d, p) basis set, whereas the remaining atoms were fixed at their crystallographic positions using the AM1 method. Based on the optimized structures, the ³¹P and

 ^{27}Al NMR chemical shifts were then calculated at the $\omega\text{B97XD}/$ TZVP level using the GIAO method.

Conflicts of interest

The authors declare no conflicts of interest.

Acknowledgements

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References

- 1 J. Biswas and I. E. Maxwell, Appl. Catal., 1990, 63, 197-258.
- 2 M. Moliner, Y. Roman-Leshkov and M. E. Davis, *Proc. Natl. Acad. Sci. U. S. A.*, 2010, **107**, 6164–6168.
- 3 A. Corma, V. Martínez-Soria and E. Schnoeveld, *J. Catal.*, 2000, **192**, 163–173.
- 4 J. Wahlen, S. Dehertogh, D. Devos, V. Nardello, S. Bogaert, J. Aubry, P. Alsters and P. Jacobs, *J. Catal.*, 2005, 233, 422–433.
- 5 A. Vjunov, J. L. Fulton, T. Huthwelker, S. Pin, D. H. Mei, G. K. Schenter, N. Govind, D. M. Camaioni, J. Z. Hu and J. A. Lercher, *J. Am. Chem. Soc.*, 2014, **136**, 8296–8306.
- 6 T. Y. Liang, J. L. Chen, Z. F. Qin, J. F. Li, P. F. Wang, S. Wang,
 G. F. Wang, M. Dong, W. B. Fan and J. G. Wang, *ACS Catal.*, 2016, 6, 7311–7325.
- 7 K. Z. Chen, M. Abdolrhamani, E. Sheets, J. Freeman, G. Ward and J. L. White, *J. Am. Chem. Soc.*, 2017, **139**, 18698–18704.
- 8 H. M. Kao, C. P. Grey, K. Pitchumani, P. H. Lakshminarasimhan and V. Ramamurthy, *J. Phys. Chem. A*, 1998, **102**, 5627–5638.
- 9 S. M. Maier, A. Jentys and J. A. Lercher, *J. Phys. Chem. C*, 2011, 115, 8005–8013.
- 10 W. O. Haag, R. M. Lago and P. B. Weisz, *Nature*, 1984, **309**, 589–591.
- S. H. Li, A. M. Zheng, Y. C. Su, H. L. Zhang, L. Chen, J. Yang,
 C. H. Ye and F. Deng, *J. Am. Chem. Soc.*, 2007, 129, 11161–11171.
- 12 D. E. Perea, I. Arslan, J. Liu, Z. Ristanovic, L. Kovarik, B. W. Arey, J. A. Lercher, S. R. Bare and B. M. Weckhuysen, *Nat. Commun.*, 2015, 6, 7589.
- 13 H. Koller, T. Uesbeck, M. R. Hansen and M. Hunger, *J. Phys. Chem. C*, 2017, **121**, 25930–25940.
- 14 C. Schroeder, M. R. Hansen and H. Koller, *Angew. Chem., Int. Ed.*, 2018, 57, 14281–14285.

- 15 M. Milina, S. Mitchell, P. Crivelli, D. Cooke and J. Perez-Ramirez, *Nat. Commun.*, 2014, 5, 3922.
- 16 Z. Wang, L. Wang, Y. Jiang, M. Hunger and J. Huang, ACS Catal., 2014, 4, 1144–1147.
- 17 S. Prodinger, M. A. Derewinski, A. Vjunov, S. D. Burton, I. Arslan and J. A. Lercher, *J. Am. Chem. Soc.*, 2016, 138, 4408–4415.
- 18 I. Yarulina, K. De Wispelaere, S. Bailleul, J. Goetze, M. Radersma, E. Abou-Hamad, I. Vollmer, M. Goesten, B. Mezari, E. J. M. Hensen, J. S. Martinez-Espin, M. Morten, S. Mitchell, J. Perez-Ramirez, U. Olsbye, B. M. Weckhuysen, V. Van Speybroeck, F. Kapteijn and J. Gascon, *Nat. Chem.*, 2018, **10**, 804–812.
- 19 R. Weingarten, G. A. Tompsett, W. C. Conner and G. W. Huber, *J. Catal.*, 2011, 279, 174–182.
- 20 M. Orazov and M. E. Davis, Chem. Sci., 2016, 7, 2264-2274.
- 21 P. Batamack, C. Doremieuxmorin, R. Vincent and J. Fraissard, *Microporous Mater.*, 1994, 2, 515–524.
- 22 A. Zheng, S. Li, S. B. Liu and F. Deng, *Acc. Chem. Res.*, 2016, **49**, 655–663.
- 23 C. Wang, Y. Y. Chu, J. Xu, Q. Wang, G. D. Qi, P. Gao, X. Zhou and F. Deng, *Angew. Chem., Int. Ed.*, 2018, 57, 10197–10201.
- 24 A. Zecchina, S. Bordiga, G. Spoto, D. Scarano, G. Petrini,
 G. Leofanti, M. Padovan and C. O. Arean, *J. Chem. Soc., Faraday Trans.*, 1992, 88, 2959–2969.
- 25 F. Collignon, P. A. Jacobs, P. Grobet and G. Poncelet, *J. Phys. Chem. B*, 2001, **105**, 6812–6816.
- 26 J. A. van Bokhoven, A. M. J. van der Eerden and D. C. Koningsberger, J. Am. Chem. Soc., 2003, 125, 7435– 7442.
- 27 W. P. Zhang, S. T. Xu, X. W. Han and X. H. Bao, *Chem. Soc. Rev.*, 2012, **41**, 192–210.
- 28 S. E. Ashbrook and S. Sneddon, J. Am. Chem. Soc., 2014, 136, 15440–15456.
- 29 Z. C. Wang, Y. J. Jiang, O. Lafon, J. Trebosc, K. D. Kim, C. Stampfl, A. Baiker, J. P. Amoureux and J. Huang, *Nat. Commun.*, 2016, 7, 13820.
- 30 C. Coperet, W. C. Liao, C. P. Gordon and T. C. Ong, J. Am. Chem. Soc., 2017, 139, 10588–10596.
- 31 D. Xiao, S. T. Xu, X. W. Han, X. E. Bao, Z. M. Liu and F. Blanc, *Chem. Sci.*, 2017, 8, 8309–8314.
- 32 Z. W. Yu, A. M. Zheng, Q. Wang, L. Chen, J. Xu, J. P. Amoureux and F. Deng, *Angew. Chem., Int. Ed.*, 2010, 49, 8657–8661.
- 33 J. Brus, L. Kobera, W. Schoefberger, M. Urbanova, P. Klein, P. Sazama, E. Tabor, S. Sklenak, A. V. Fishchuk and J. Dedecek, *Angew. Chem.*, *Int. Ed.*, 2015, 54, 541–545.
- 34 C. P. Grey and A. J. Vega, *J. Am. Chem. Soc.*, 1995, **117**, 8232-8242.
- 35 A. P. M. Kentgens, D. Iuga, M. Kalwei and H. Koller, J. Am. Chem. Soc., 2001, **123**, 2925–2926.
- 36 J. Jiao, J. Kanellopoulos, W. Wang, S. S. Ray, H. Foerster, D. Freude and M. Hunger, *Phys. Chem. Chem. Phys.*, 2005, 7, 3221–3226.
- 37 P. A. Jacobs and H. K. Beyer, J. Phys. Chem., 1979, 83, 1174– 1177.

- 38 Q. L. Wang, G. Giannetto, M. Torrealba, G. Perot, C. Kappenstein and M. Guisnet, J. Catal., 1991, 130, 459–470.
- 39 G. L. Woolery, G. H. Kuehl, H. C. Timken, A. W. Chester and J. C. Vartuli, *Zeolites*, 1997, **19**, 288–296.
- 40 E. Lam, A. Comas-Vives and C. Copéret, *J. Phys. Chem. C*, 2017, **121**, 19946–19957.
- 41 Z. W. Yu, S. H. Li, Q. Wang, A. M. Zheng, X. Jun, L. Chen and F. Deng, *J. Phys. Chem. C*, 2011, **115**, 22320–22327.
- 42 X. F. Yi, K. Y. Liu, W. Chen, J. J. Li, S. T. Xu, C. B. Li, Y. Xiao, H. C. Liu, X. W. Guo, S. B. Liu and A. M. Zheng, *J. Am. Chem. Soc.*, 2018, **140**, 10764–10774.
- 43 J. F. Haw, J. B. Nicholas, T. Xu, L. W. Beck and D. B. Ferguson, *Acc. Chem. Res.*, 1996, **29**, 259–267.
- 44 E. F. Rakiewicz, A. W. Peters, F. Wormsbecher, K. J. Sutovich and K. T. Mueller, *J. Phys. Chem. B*, 1998, **102**, 2890–2896.
- 45 A. Zheng, S. B. Liu and F. Deng, *Chem. Rev.*, 2017, **117**, 12475–12531.

- 46 Q. Wang, Y. X. Li, J. Trebosc, O. Lafon, J. Xu, B. W. Hu,
 N. D. Feng, Q. Chen, J. P. Amoureux and F. Deng, *J. Chem. Phys.*, 2015, 142.
- 47 A. Janda and A. T. Bell, *J. Am. Chem. Soc.*, 2013, **135**, 19193–19207.
- 48 Q. Zhao, W. H. Chen, S. J. Huang, Y. C. Wu, H. K. Lee and S. B. Liu, *J. Phys. Chem. B*, 2002, **106**, 4462–4469.
- 49 L. Chen, X. Y. Lu, Q. A. Wang, O. Lafon, J. Trebosc, F. Deng and J. P. Amoureux, *J. Magn. Reson.*, 2010, **206**, 269–273.
- 50 P. K. Madhu, A. Goldbourt, L. Frydman and S. Vega, *Chem. Phys. Lett.*, 1999, **307**, 41–47.
- 51 D. Massiot, F. Fayon, M. Capron, I. King, S. Le Calve, B. Alonso, J. O. Durand, B. Bujoli, Z. H. Gan and G. Hoatson, *Magn. Reson. Chem.*, 2002, **40**, 70–76.
- 52 J. D. Chai and M. Head-Gordon, J. Chem. Phys., 2008, 128.