Monomolecular Cracking of Propane: Effect of Zeolite Confinement and Acidity

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ABSTRACT: The effect of zeolite pore geometry and intrinsic acidity on the activation energy of propane monomolecular cracking was investigated for six topologically distinct zeolites with different pore sizes. Periodic density functional theory calculations were used to calculate the activation energy, while cluster models were used to calculate deprotonation energies. The computed intrinsic activation energies showed a smaller variation with topology than the adsorption energies. No correlation was found between the computed deprotonation and ammonia adsorption energies at the acid site and the intrinsic activation energy. Detailed analysis of the computed structures and properties suggests that acid sites with different pore topologies impose geometrical constraints on the ion-pair formed by the ammonium molecule, which differs significantly from those that affect the propane reaction.

INTRODUCTION

Zeolites are microporous materials composed of corner-sharing AlO$_4^-$ and SiO$_4$ tetrahedra linked in a superlattice of cages and channels. When the negative charge on the AlO$_4^-$ site is compensated by a proton, a (≡Al−OH−Si≡) Brønsted acid site is formed. For any particular zeolite cage, the specific geometric structure of the Brønsted acid sites affects their ability to accommodate guest species. Guest-site congruity imparts differences in the stability of reactive species, transition states, and intermediates, giving rise to shape selectivity. In addition to acidity and microporosity, the high thermal stability and low production cost facilitate the use of zeolites in many industrial processes. Fluid catalytic cracking is a key process in the conversion of crude oil to transportation fuels and petrochemical feedstock. The mechanism of cracking depends on the reaction conditions. The bimolecular mechanism dominates at low temperatures, high partial alkanes pressure, and high conversion. This mechanism involves a chain of reactions propagated by the carbenium ions. The alternative monomolecular Haag-Dessau mechanism is preferred at high temperature (T > 600 K) and low partial alkanes pressure. This mechanism represents the direct protonation of the alkane at the Brønsted acid site, creating a carbocation intermediate, which then dissociates into a shorter alkene and alkane:

$$C_6H_{14} \rightarrow C_5H_6 + C_4H_8$$ (1)

Monomolecular cracking has been subject to numerous experimental and theoretical studies based on density functional theory (DFT) using periodic and cluster models. Most studies agree that the rate-limiting step of monomolecular cracking is the protonation of the alkane by attacking the C–C bond.

The experimental measurement of the apparent activation energy, $E_{\text{app}}$, inevitably also includes the contributions of both site adsorption energy and the intrinsic activation energy:

$$E_{\text{app}} = E_{\text{ads}} + E_{\text{int}}$$ (2)

where $E_{\text{ads}}$ is the alkane adsorption energy at the reaction site and $E_{\text{int}}$ is the intrinsic activation energy. The intrinsic activation energy is the energy required to cross the barrier between the adsorbed reactant state and the product state. Experimental activation enthalpies of monomolecular cracking decrease as the pore diameter of the zeolite framework decreases. The analysis of experimental data suggests that the decrease in the apparent activation energy with confinement is caused by the decrease in the adsorption energy and that the intrinsic activation energy is rather insensitive to framework type. However, the analysis that distinguishes the intrinsic activation energy from the apparent activation energies has some shortcomings. The adsorption measurements are usually conducted at temperatures well below the actual cracking temperatures and include the contribution of alkanes adsorption at non-Brønsted acid sites. Recent progress has been made both in experimental and theoretical studies on the effect of confinement on the intrinsic activation energy.
computationally and experimentally in addressing these issues. Theoretically, adsorption energies were calculated by Janda et al. using Monte Carlo simulations of the Bronsted acid sites and at realistic reaction temperatures. This facilitated the calculation of the intrinsic activation energies from the experimentally measured activation energies for butane cracking and dehydrogenation. They demonstrated that the intrinsic activation energy might be affected by confinement in 10-ring zeolites, depending on the location of the cracked bond. The intrinsic activation energy of cracking at the terminal carbon atom decreased as confinement increased. On the other hand, the intrinsic activation energy of cracking at the central bond of butane was found to be invariant with respect to the level of confinement. Experimentally, Kadam et al. obtained the apparent rate constants, \( k_{app} \) and adsorption constants, \( K_{ad} \) at the same temperature using IR operando spectroscopy. They concluded that confinement mainly influences the adsorption energies and that the intrinsic activation energies in different zeolites are similar. More recently, Berger et al. calculated the intrinsic activation energy of propane cracking in MFI, FER, MVE, CHA, and FAU zeolites using a hybrid QM-QM method and confirmed that adsorption energies dominate the predicted apparent activation energies.

Since monomolecular cracking involves proton transfer from the acid site to the alkane, zeolite acidity has been the subject of extensive theoretical studies. There are two proxies commonly used to quantify the acidity of a site: the deprotonation energy or the adsorption energy of basic molecules. Deprotonation energy is defined as the energy required to release the proton from the Bronsted site. It is now accepted that the intrinsic acidity of the Bronsted sites in different zeolite frameworks is similar. Deprotonation energies were calculated using the QM-Pot method for CHA, TON, FER, MFI, MOR, and FAU and were found to lie within a range <30 kJ mol\(^{-1}\). This conclusion was confirmed by Jones and Iglesia who calculated the ensemble average deprotonation energy, \( \langle DPE \rangle \), for the 12 distinct T sites in MFI and found them to be similar to a value of 1201 ± 11 kJ mol\(^{-1}\).

Ammonia, NH\(_3\), adsorption is a convenient base also commonly used to probe the acidity of zeolites. While ammonia adsorption is well suited for Bronsted acid site quantification, it has been suggested that the correlation between the intrinsic acid site strength and the adsorption energy is not unambiguous. For instance, Jones and Iglesia have shown that NH\(_3\) adsorption energies do not correlate monotonically with the DFT calculated ensemble-averaged deprotonation energies (DPE), suggesting that the measured adsorption energies are indicative of the confinement effect. Boronat and Corma also concluded that the adsorption energies of strong bases include the contribution of multiple interactions between the protonated base and the surrounding anionic framework O atoms. Even though ammonia adsorption energies do not correlate with deprotonation energies, Liu et al. reported a correlation between NH\(_3\) adsorption energy and the intrinsic activation energy of propane cracking in FAU type zeolites.

In the present study, the monomolecular cracking activation energy and the acidity will be investigated with DFT methods in 6 frameworks (FAU, MFI, MEL, CHA, AEI). The influence of confinement and acidity on the intrinsic activation energy, apparent activation energy, and adsorption energies will be investigated.

### METHODS

**Periodic DFT Calculations.** Periodic density functional theory (DFT) calculations were performed with the CRYSTAL software with the effects of electronic exchange and correlation described by the ob97X-D functional, which has been reported to take noncovalent interactions into account and was used for similar systems. London dispersion interactions were estimated using Grimme’s dispersion correction.

Triple valence 6-311\(G(d,p)\) Gaussian basis sets were used throughout the calculations. The Brillouin zone sampling was restricted to the \( \gamma \) point. The reaction paths and the transition states were determined with the climbing image nudged elastic band method as implemented in the atomic simulation environment package (ASE) package. Geometry optimization of the initial, final, and path images was performed with fixed cell parameters. Vibrational frequencies were computed to confirm the nature of the transition state. Unconstrained geometry optimization of all coordinates was performed to find local minima with convergence assumed to be reached when the forces on each atom are less than 0.02 eV Å\(^{-1}\).

**Activation and Adsorption Enthalpies.** The enthalpy of a system \( X \), \( H_X \), was calculated as

\[
H_X = E_0 + E_T + PV
\]

where the zero-point vibrational energy, \( E_0 \), and the vibrational contribution to the thermal energy, \( E_T \), were calculated within the quasi-harmonic approximation. The intrinsic activation energy was then calculated as

\[
\Delta H_{\text{act}} = H_{T_1} - H_{Z-P}
\]

where \( H_{T_1} \) denotes the enthalpy of the transition state, and \( H_{Z-P} \) denotes the enthalpy of the adsorbed propane. The adsorption enthalpy was calculated as

\[
\Delta H_{\text{ad}} = H_{Z-M} - (H_Z + H_M)
\]

where \( H_{Z-M} \) denotes the enthalpy of the adsorbed system, \( H_Z \) denotes the enthalpy of the zeolite crystal, and \( H_M \) denotes the enthalpy of the molecule in the gas phase. Adsorption enthalpies were then corrected for the basis set superposition error (BSSE) using the counterpoise correction method.

**Cluster DFT Calculations.** Using periodic models allows for an accurate and systematic comparison of the intrinsic activation energies and the ammonia adsorption energies in different zeolites. However, unambiguous assignment of the energy reference for the potential energy in charged cells means that it is problematic to compare descriptors such as the deprotonation energy between zeolites frameworks. Here this is achieved by using cluster models of the reaction sites using the same basis set, exchange-correlation functional, and numerical tolerances as the periodic calculation. Suitably converged clusters must be defined. A widely used method is to include all atoms within N bonds from the Al atom. The clusters generated by this method are termed N-bond clusters. Jones et al. examined the effect of the cluster size on deprotonation energies using N-bond clusters. Their results showed that the deprotonation energy varies systematically with the cluster size. The variation in the calculated deprotonation energy converges for clusters with more than 20T atoms, where T stands for either of the tetrahedrally coordinated Si or Al atoms. Hence, Jones et al. deduced that the negative charge that remains upon deprotonation is localized within the second coordination sphere of the oxygen atoms from the aluminum atom. In this...
study selecting $N = 6$ generated clusters with more than 30T atoms for all of the examined frameworks and thus was used to generate the clusters to evaluate the deprotonation energy. Deprotonation energies were then calculated according to the following equation:

$$E_{DPE} = E_{H^+} + E_{H^-} - E_{ZH}$$  \(6\)

Zeolite Frameworks. Six frameworks with varying pore sizes were used to sample a wide range of confinement and acidity strengths. Those frameworks are FAU, AFI, MFI, MEL, AEI, and CHA. Details of these structures and the optimized cell parameters are listed in Table 1. AFI has a lattice constant of just 8.58 Å in the crystallographic $c$-direction, which may result in interactions between periodic images. AFI was therefore described with a doubled unit cell along this direction. The periodic cell models are illustrated in Figure 1. Zeolites frameworks can have several crystallographically distinct T sites, and each T atom is coordinated to four oxygen atoms.

Table 1. Optimized Cell Parameters for Studied Zeolites’ Periodic Models Used in DFT Calculations and LCD

| framework type | LCD\(^*\) (Å) | Al position | Si/Al | cell | cell parameters |
|----------------|----------------|-------------|-------|------|----------------|
| FAU 1          | 11.9           | T1          | 47    | rhombohedra | $a = b = c = 17.37$ |
| FAU 2          | 11.9           | T1          | 47    | rhombohedra | $a = b = c = 17.37$ |
| AFI            | 8.1            | T1          | 47    | hexagonal   | $a = 13.89, b = 13.89, c = 17.22$ |
| MFI            | 7              | T12         | 95    | orthorhombic| $a = 20.36, b = 19.78, c = 13.36$ |
| MEL            | 8.4            | T6          | 47    | rhombohedral| $a = b = c = 15.83$ |
| CHA            | 8              | T1          | 35    | trigonal    | $a = b = 13.84, c = 14.45$ |
| AEI            | 8              | T1          | 47    | orthorhombic| $a = 13.79, b = 12.63, c = 18.50$ |

\(^*\)Largest cavity diameter (LCD) calculated by First et al.\(^{50}\)

Figure 1. Unit cells of the periodic models used in the DFT calculations represent zeolites. FAU (48 T atom), AFI (48 T atom), MFI (96 T atom), MEL (48 T atom), CHA (36 T atom), AEI (48 T atoms).
acidic proton can hop between those oxygen atoms.32 In the present study, we are focused on the trends in activation energies with zeolite topology. We, therefore, adopt a model of the reaction site based on a single T site in each zeolite where the acidic hydrogen is bonded to a single oxygen position as indicated in Table 4. This choice facilitates the extraction of trends but prevents the direct comparison of computed data to experiment, which is best achieved with an ensemble average over all of the kinetically accessible T sites and oxygen positions. Absolute values for experimental results are quoted below for qualitative comparison with those computed.

Table 2. Selected Interatomic Distances (Å) of Adsorbed Propane Geometries in Zeolites: FAU, AFI, MFI, MEL, CHA, AEI

|           | H_b−C1 | H_b−C2 | H_b−C3 | [H_{propane−O_{zeolite}}]_{shortest} |
|-----------|--------|--------|--------|-------------------------------------|
| FAU (T1)(O4) | 2.69   | 2.49   | 3.07   | 2.73                                |
| FAU (T1)(O1) | 2.95   | 3.00   | 3.12   | 2.62                                |
| AFI (T1)(O2) | 3.33   | 2.41   | 2.43   | 2.69                                |
| MFI (T12)(O25) | 3.58   | 2.50   | 2.25   | 2.68                                |
| MEL (T4)(O1) | 3.19   | 2.48   | 2.57   | 2.67                                |
| CHA (T1)(O1) | 3.24   | 3.23   | 3.62   | 2.45                                |
| AEI (T1)(O4) | 4.5    | 3.05   | 3.38   | 2.66                                |

*H_b denotes the Bronsted hydrogen.

Results and Discussion

Propane Adsorption Energies. Adsorbed propane geometries are displayed in Figure 2. Propane interacts with the zeolite through multiple hydrogen bonds with the framework oxygen atoms. The main geometrical parameters have been supplied in Table 2. These parameters are the distances between the Bronsted hydrogen atom and the propane carbon atoms, and the shortest distance between the propane hydrogen atoms and the zeolite oxygen atoms, [H_{propane−O_{zeolite}}]_{shortest}. The reactant state geometries suggest that the oxygen atoms in the acidic site act as a Lewis base anchoring the propane molecule prior to the proton attack on the carbon−carbon bond.

The calculated adsorption energies and the BSSE corrected energies are displayed in Table 3. The results show that the BSSE correction is significant at this basis set level. The calculated adsorption enthalpies are then displayed in Table 4 along with the experimental values. The calculated adsorption enthalpies range from −33 to −47 kJ mol⁻¹, and there is a good agreement between the computed and experimental values.
Table 3. Calculated Propane Adsorption Energies, $\Delta E$(BSSE), and BSSE Corrected Energies, $\Delta E$, in kJ mol$^{-1}$ in Zeolites: FAU, AFI, MFI, MEL, CHA, AEI

| framework | $T$[K] | $\Delta H_{ab}$ | Si/Al | $\Delta H_{ab}$ |
|-----------|--------|----------------|-------|----------------|
| FAU (T1)(O4) | 323 | −30 | 2.7 | −31$^{15}$ |
| FAU (T1)(O1) | 323 | −33 | 2.7 | −31 |
| AFI (T1)(O2) | 323 | −39 | 2.7 | −31 |
| MFI (T12)(O2S) | 323 | −44 | 35 | −46$^{11}$ |
| MEL (T4)(O1) | 323 | −47 | 14 | −38$^{12}$ |
| CHA (T1)(O1) | 313 | −37 | 14 | −38$^{12}$ |
| AEI (T1)(O4) | 323 | −35 | 2.7 | −31 |

Table 4. Calculated Adsorption Enthalpies and Experimental Adsorption Enthalpies in kJ mol$^{-1}$ in Zeolites: FAU, AFI, MFI, MEL, CHA, AEI

| framework | $T$[K] | $\Delta H_{ab}$ | Si/Al | $\Delta H_{ab}$ |
|-----------|--------|----------------|-------|----------------|
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| CHA (T1)(O1) | 313 | −37 | 14 | −38$^{12}$ |
| AEI (T1)(O4) | 323 | −35 | 2.7 | −31 |

agreement in general between the experimental values and the calculated values. The calculated adsorption energies generally reduce (become more negative) with the decrease of the LCD of the framework channels or cages except for propane adsorption energy in MEL. This may be due to the incomplete sampling of the configuration space. Bucic et al. demonstrated that adsorption energies could vary by as much as 12 kJ mol$^{-1}$ within a single framework.

Activation Energies. The transition state geometries and the main geometrical parameters are displayed in Figure 3 and Table 5. The distances of the cracked bond at the transition state range from 2.12 to 2.60 Å, indicating a late transition state. This big distance range can be attributed to the high mobility of the framework channels or cages except for propane adsorption to the acid site. Swisher et al. have also reported late transition states with C–C bond distances of 2.56 and 2.66 Å for propane cracking in FAU and MFI, respectively, obtained from B3LYP calculations on T₄₅-clusters. Berger et al. on the other hand, reported the distances of the cracked C–C bond in three cracking mechanisms using PBE+D2 periodic calculations. They obtained 1.8 Å for the C–C bond when cracking is initiated by the protonation of the terminal carbon atom, 1.8 Å for the protonation of the C–C bond, and 2.5 Å for the concerted path, in which no alkoxide intermediate is formed.

Table 6 shows the calculated and experimental activation enthalpies. Experimental results are affected by extra-framework species, surface acidity, and internal silanol groups. Those factors may result in differences between the calculated activation enthalpies and the experimental activation enthalpies. The range of the calculated apparent activation energies for all frameworks, <20 kJ mol$^{-1}$, is smaller than the experimental range of the apparent activation energies (<52 kJ mol$^{-1}$) for the FAU, MFI, and CHA frameworks, which is probably due to the wide variability of the experimental samples in terms of Si/Al ratios, extra-framework species, surface acidity, and morphologies. The calculated apparent activation enthalpies for FAU lie within the range of the reported experimental values. However, for CHA, the exact agreement between the calculated and experimental values is probably accidental, considering the differences between real crystals and our idealized model. There is an 11 kJ mol$^{-1}$ difference between the calculated apparent activation enthalpy for MFI and the experimental value.

The calculated intrinsic activation energies fall between 193 and 207 kJ mol$^{-1}$ with a range of 14 kJ mol$^{-1}$ for the 6 frameworks. It is also interesting to note that the intrinsic activation energies calculated for FAU, at the same acidic site starting from a different propane orientation and Bronsted H position, differ by 10 kJ mol$^{-1}$. Bucic et al. have also reported that the orientation of the propane molecule relative to the framework can cause the intrinsic activation energy of propane cracking to vary by as much as 23 kJ mol$^{-1}$ in the CHA framework. The residual negative charge upon site deprotonation is localized within the second coordination sphere of O atoms from the Al atom. Therefore, the charge does not extend to the whole ring, and the electrostatic interactions are confined to the region that is close to the Al ion. Moreover, van der Waals forces are similar for the transition states and the reactant states. Hence, the observed differences in the intrinsic activation energies at a single site are due to the different geometrical constraints that the propane experiences at different orientations to the framework, which can either contribute to strengthening or weakening the ionic interactions between the cationic transition state and the anionic framework. This explanation also suggests that the pore diameter does not influence the intrinsic activation energies. This conclusion is supported by the results of Kadam et al. and Berger et al. who reported a narrow range of 4 kJ mol$^{-1}$ and 8 kJ mol$^{-1}$ for the intrinsic activation energies, respectively, in several zeolite frameworks.

Deprotonation Energies. The link between the intrinsic acidity and the intrinsic activation energy can be illustrated by the thermochemical cycle of the acid-catalyzed activation of alkanes, which is depicted in Scheme 1. The thermochemical cycle links the activation energy with the energies of the hypothetical elementary steps that lead to the transition state. Those steps are adsorption to the acid site $E_{ads}$, deprotonation of the acid site $E_{DPB}$, protonation of the reactant $E_{HPB}$, and finally stabilization of the transition state $E_{stab}$. The calculated deprotonation energies of the studied frameworks range from 1211 kJ mol$^{-1}$ to 1230 kJ mol$^{-1}$, see Table 7, which is in agreement with previous cluster DFT studies. It is evident in Figure 5b that there is no correlation between the calculated apparent activation energies calculated for FAU, at the same acidic site starting from a different propane orientation and Bronsted H position, differ by 10 kJ mol$^{-1}$. It is evident in Figure 5b that there is no correlation between the calculated apparent activation energy for MFI and the experimental value.

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Ammonia Adsorption Energies. The interaction between the ammonia molecule and the Bronsted acid site results in the formation of ammonium $\text{[NH}_4^+\text{]}$, which then interacts with the negatively charged framework through several hydrogen bonds as shown in Figure 4. Table 8 lists the shortest two hydrogen bonds between the ammonium molecule and the negatively charged zeolite. According to Jones and Iglesia, the N–H–O bonds may contribute up to 60 kJ mol$^{-1}$ to the adsorption energy depending on the bond length and angle, and it is sensitive to the local geometrical variations of the acid site.
Table 8 shows variable bond distances between the ammonium and the framework oxygen atoms.

The calculated adsorption energies and the BSSE corrected adsorption energies are displayed in Table 9. In the case of ammonia adsorption, the discrepancy between the experimental adsorption energies and the calculated adsorption energies is

Figure 3. Optimized geometries of the transition states inside the main cages and channels or of acidic zeolites; FAU (a,b), AFI (c), MFI (d), MEL (e), CHA (f), AEI (g).

Table 5. Selected Interatomic Distances (Å) of Transition States in Zeolites: FAU, AFI, MFI, MEL, CHA, AEI

| zeolite  | C−C | H_b−C | H_a−C | [H_T−O_{zeolite}]_{shortest} |
|----------|-----|-------|-------|-------------------------------|
| FAU (T1)(O4) | 2.3 | 1.16  | 1.42  | 2.14                          |
| FAU (T1)(O1) | 2.12| 1.20  | 1.34  | 2.15                          |
| AFI (T1)(O2) | 2.51| 1.14  | 1.55  | 2.20                          |
| MFI (T12)(O25) | 2.60| 1.12  | 1.70  | 2.20                          |
| MEL (T4)(O1) | 2.40| 1.16  | 1.46  | 1.98                          |
| CHA (T1)(O1) | 2.60| 1.13  | 1.7   | 1.90                          |
| AEI (T1)(O4) | 2.16| 1.19  | 1.36  | 2.18                          |

“H_b denotes the Bronsted hydrogen, H_T denotes the hydrogen atom in the transition state structure.

Table 6. Calculated Apparent and Intrinsic Activation Energies and Experimental Apparent and Intrinsic Activation Energies in kJ mol⁻¹ in Zeolites: FAU, AFI, MFI, MEL, CHA, AEI

| framework  | T[K] | ΔH_{app} | ΔH_{tot} | Si/Al | ΔH_{app} | ΔH_{tot} |
|------------|------|----------|----------|-------|----------|----------|
| FAU (T1) (O4) | 823  | 181      | 207      | 2.6, 3.3, 3.6 | 201, 179, 169 | 231, 209, 199 |
| FAU (T1) (O1) | 823  | 172      | 197      | 2.6, 3.3, 3.6 | 201, 179, 169 | 231, 209, 199 |
| AFI (T1) (O2) | 773  | 161      | 201      | 35    | 149 ± 10 | 192 ± 10 |
| MFI (T12) (O25) | 773 | 164      | 207      |       |          |          |
| MEL (T4) (O1) | 773  | 164      | 201      | 12    | 164      | 200      |
| CHA (T1) (O1) | 773  | 165      | 195      |       |          |          |
| AEI (T1) (O4) | 773  | 165      | 195      |       |          |          |

Table 8 shows variable bond distances between the ammonium and the framework oxygen atoms.
Scheme 1. Hypothetical Elementary Steps of Acid-Catalyzed Activation of Alkanes in Zeolites Bronsted Acid Sites

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ZEOLITE + H+ + ALKANE → [ALKANE-H]+ + EINT + Eias
```

**Table 7. Deprotonation Energies (DPE) in kJ mol⁻¹ of Zeolites: FAU, AFI, MFI, MEL, CHA, AEI**

| framework      | DPE  |
|----------------|------|
| FAU (T1)(O4)   | 1228 |
| FAU (T1)(O1)   | 1215 |
| AFI (T1)(O2)   | 1230 |
| MFI (T12)(O25) | 1211 |
| MEL (T4)(O1)   | 1219 |
| CHA (T1)(O1)   | 1223 |
| AEI (T1)(O4)   | 1225 |

**Table 8. Shortest O–H Bonds Distances (Å) in Adsorbed Ammonia Geometries in Zeolites: FAU, AFI, MFI, MEL, CHA, AEI**

|          | H–O  | H–O  |
|----------|------|------|
| FAU (T1)(O4) | 1.66 | 2.16 |
| FAU (T1)(O1) | 1.70 | 1.73 |
| AFI (T1)(O2) | 1.63 | 1.97 |
| MFI (T12)(O25) | 1.63 | 2.03 |
| MEL (T4)(O1) | 1.70 | 1.87 |
| CHA (T1)(O1) | 1.82 | 1.86 |
| AEI (T1)(O4) | 1.65 | 1.95 |

**Table 9. Calculated Ammonia Adsorption Energies, ΔE(BSSE), and BSSE Corrected Energies, ΔE, in kJ mol⁻¹ in Zeolites; FAU, AFI, MFI, MEL, CHA, AEI**

|          | ΔE(BSSE) | ΔE   |
|----------|----------|------|
| FAU (T1)(O4) | −147     | −123 |
| FAU (T1)(O1) | −166     | −135 |
| AFI (T1)(O2) | −153     | −115 |
| MFI (T12)(O25) | −144   | −113 |
| MEL (T4)(O1) | −145     | −116 |
| CHA (T1)(O1) | −169     | −135 |
| AEI (T1)(O4) | −176     | −145 |

Therefore, uncorrected adsorption energies will be used in the following analysis.

Ammonia adsorption enthalpies are listed in Table 10 along with the experimental values. As noted above, the experimental values are a statistical average of the ammonia adsorption energies at all the T sites and possible [NH₄⁺] orientations. For instance, ammonia adsorption enthalpy at FAU (O4) differs by 16 kJ mol⁻¹ compared to the adsorption enthalpy at FAU (O1). To compare directly with experiments, *ab initio* molecular dynamics simulations may be required to sample the full configuration space of adsorbed [NH₄⁺]. With this consideration, the calculated adsorption enthalpies lie within a reasonable range from the reported experimental values of ammonia adsorption. As discussed earlier, it has been suggested that quantification of acidity using the binding energy of a basic molecule includes contributions from both the intrinsic acidity and confinement.⁵⁷ If one accepts deprotonation energies as a measure for the intrinsic acidity, it is evident that ammonia adsorption energies do not correlate with deprotonation energies due to the dominant effect of the electrostatic interactions and dispersion forces.

Probe molecules that integrate acidity and ion pair interaction energy were potentially suggested for predicting the intrinsic activation energy. However, the dispersion contribution to the ammonia adsorption energy is not desirable for predicting the intrinsic activation energy as the long-range dispersion forces may be expected to be similar in the reactants and transition states and thus provide poor differentiation. Recalculating the ammonia adsorption energies explicitly excluding the long-range dispersion contribution, ΔE(D)ads([NH₃]), produces the data displayed in Table 10. Figure 5a shows no correlation between the calculated ammonia adsorption energies and the intrinsic activation enthalpies for the zeolites topologies studied here. This indicates that the ion-pair interaction energies are specific to the molecule-site pair and that the intrinsic activation energy cannot, therefore, be approximated reliably by the adsorption of the ammonia molecule, which experiences different geometrical constraints to propane. A similar conclusion was made by Liu et al.,⁵⁷ who reported that MFI and CHA did not follow the scaling relation obtained for FAU and the acid strength determined by the NH₃ adsorption energies for the conversion of π-adsorbed isobutene into alkoxy species.

**CONCLUSION**

The influence of the acid site geometry on the activation energy of propane cracking was investigated using periodic DFT calculations for a number of topologically distinct zeolite frameworks. Acid sites were characterized with the LCD,
ammonia adsorption energies, and deprotonation energies. The calculations were performed with the ωB97X-D functional, 6-311G(d,p) Gaussian basis set, and Grimme’s dispersion correction. Deprotonation energies were calculated using cluster models to achieve unambiguous referencing of energies in charged systems. A single reaction site in each framework was adopted to facilitate the examination of trends due to the varying confinement. The computed data suggest that the intrinsic activation energies are not significantly affected by the zeolite pore diameter. The computed intrinsic activation energies did

Figure 4. Optimized geometries of ammonia inside the main cages and channels or acidic zeolites; FAU (a,b), AFI (c), MFI (d), MEL (e), CHA (f), AEI (g).

Figure 5. Plots of the intrinsic activation energy $\Delta E_{\text{int}}$ obtained by the periodic DFT calculation and the NH$_3$ adsorption energy $\Delta E_{\text{ads}}$[NH$_3$] (a), deprotonation energy DPE (b).
not correlate with deprotonation energies due to the ionic interaction at the transition state, which attenuates the deprotonation energies. Intrinsic activation energies also did not correlate with the ammonia adsorption energies because the ammonia ion-pair interaction is subject to different geometrical constraints to that of the protonated propane. We, therefore, conclude that neither deprotonation energies nor ammonia adsorption energies provides a reliable proxy for intrinsic activation energies in aluminosilicate frameworks with different topologies.

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**Notes**

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