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DFT Study on the Effect of Aluminum Position in Zn-Exchanged MFI on Methane Activation

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ABSTRACT

The position of aluminum atoms in ion-exchanged zeolites is known to affect the reactivity of active sites. In this work, we used DFT calculations to systematically quantify the effect of Al-atom position within the α-ring of Zn-exchanged MFI (Zn-MFI) on the activation of methane. Our DFT results indicate that the most stable configuration for the Zn-exchanged cluster of the α-ring is obtained when the Al atoms are located at the T11-T2 crystallographic sites. For each Al-atom configuration, we analyzed the reaction pathways for methane activation. Our results suggest that the activation of methane yields the formation of a Brønsted acid site, which can be formed at an oxygen atom within the α-ring or at an oxygen atom that lies outside the α-ring, and that the lowest reaction energy for methane activation is obtained when the Brønsted acid site is formed at the oxygen atom in which the HOMO of the isolated cluster is located. Furthermore, our results indicate that the partial atomic charge of the Zn atom within the α-ring of MFI can be correlated to the transition state energy of methane activation, which ranges from 87 to 131 kJ/mol depending on the location of Al atoms. The fundamental studies conducted in this work contribute to the elucidation of essential parameters and correlations, based on electrostatic and electron density, for the activation of methane on Zn-MFI zeolites.
1. INTRODUCTION

Increased methane (CH₄) availability has sparked interest in its use as a raw material to produce chemicals, such as methanol and acetic acid.¹ ² According to the U.S. Energy Information Administration (EIA), coalbed methane reserve has reached 11.8 trillion cubic feet in the United States.³ Currently, methane is mainly used in industry for the production of syngas at temperatures higher than 973 K, which is then converted into chemicals, such as hydrocarbons or alcohols.² The current challenge, however, is the direct conversion of methane to value-added products due to its highly stable C-H bond, which has a dissociation energy of 438.8 kJ/mol.² ⁴

The MFI zeolite has been shown to activate methane at low temperatures when it is exchanged with zinc cations (Zn²⁺).⁵ Therefore, experimental⁶–⁹ and theoretical¹⁰–¹² studies have focused on understanding the mechanism of hydrogen atom abstraction from methane within the pore of this zeolite. By using ¹H and ¹³C NMR techniques, Kolyagin et al.⁶ demonstrated that methane dissociates over Zn-exchanged MFI (Zn-MFI), and this dissociation yields the formation of a hydroxyl group in the zeolite, which is in agreement with other previous studies.⁶,¹³–¹⁵ Furthermore, they found that these hydroxyl groups are stable and cause a reduction of the catalytic activity. Similarly, through in situ NMR, Stepanov et al.¹⁶ also determined that Zn-MFI facilitates C-H bond activation in methane and ethane. They also found that the hydrogen atom is strongly attracted to an oxygen atom, thus leading to the formation of Brønsted acid sites.¹³,¹⁶,¹⁷

Different studies have been also carried out on methane activation with MFI zeolites exchanged with either oxo-clusters or metal cations, such as Cu, Fe, and Zn.⁹,¹⁸–²⁰ For the Cu-based MFI catalysts, it has been demonstrated that the oxo-clusters [M–O–M]²⁺ yield lower energy barriers than metal cations. For instance, the energy barrier for methane dissociation on [Cu-O-Cu]-
exchanged MFI is 82 kJ/mol,\textsuperscript{18} while on Cu-MFI is 129 kJ/mol.\textsuperscript{21} However, for methane dissociation on Zn catalysts, it has been demonstrated that Zn-exchanged MFI is more active than the zinc oxides.\textsuperscript{17}

Previous studies have aimed to analyze the Al distribution in zeolites and its possible effects in catalysis.\textsuperscript{22,23} For instance, in an early work by Grunder and Iglesia, they concluded that the distribution of Al atoms in the T-sites of MFI can affect the energy barrier for alkane activation.\textsuperscript{24} Subsequent studies have then confirmed this conclusion, and various studies have also reported that the Brønsted acidity of the MFI zeolite varies depending on the distribution of the aluminum atoms within the framework.\textsuperscript{24–26} Thus, the catalyst activity can be compromised depending on this distribution, and this can yield low conversion rates. Previous studies have also aimed to pinpoint the MFI crystallographic sites in which the Al atoms are located.\textsuperscript{15} For instance, Sklenak et al.\textsuperscript{27} combined high resolution 27aluminum (27Al) NMR spectroscopy with density functional theory (DFT) / molecular mechanics calculations, and they established which of the 12 distinguishable T-sites of MFI zeolite are occupied by Al-atoms. Furthermore, Yokoi et al.\textsuperscript{28} were able to control the location of Al-atoms on MFI zeolites by using different organic molecules during the synthesis of the zeolite. For instance, when tetrapropylammonium cations were used, Al-atoms are primarily located at the intersection of MFI zeolite channels.

While efforts have been put forth in studying and controlling the Al-atom distribution in various zeolites, the specific effects of this distribution on methane activation has not been thoroughly studied. Recently, Zhao et al.\textsuperscript{29} conducted DFT studies to analyze the effect of Al positions in Cu-MOR for the oxidation of methane to methanol. They reported that methane adsorption energies ranged from 43 to 65 kJ/mol and activation energies ranged from 63 to 124 kJ/mol depending on
the position of the Al atoms within the ring. To our knowledge, a similar study on MFI zeolite, which has been more widely used for methane activation, still lacks in the literature.

In this work, we have conducted a systematic study of the effect of Al-atom location within the α-ring of Zn exchanged MFI on the energy barrier for methane activation, as described in Figure 1. Our work unravels that: (i) Al atoms distribution produces a variation in the energy barrier of methane activation, (ii) the activation of methane yields the formation of a Brønsted acid site at the oxygen in which the HOMO of the isolated cluster is located, and (iii) there is a correlation between the transition state energy of methane activation and the partial atomic charge of the Zn atom within the α-ring of MFI.

2. METHODOLOGY

Methane activation was analyzed on MFI zeolite by creating a cluster based on the crystallography data available in the Database of Zeolite Structures. Here we analyzed the α-ring in MFI, which has been reported as the most active site in Zn-exchanged MFI. Figure 1(a) shows the location of the ring within the zeolite pore. This ring is composed of seven silicon atoms, and each silicon atom corresponds to a different crystallographic T-site. The crystallographic sites are identified in Figure 1(b). In this work, the location of each Al atom within the α-ring is identified with the T-sites shown in Figure 1(b).
Figure 1. (a) MFI zeolite framework. The α-ring of the zeolite framework is highlighted with yellow for silicon atoms and red for oxygen atoms. (b) The 74-atom cluster of the α-ring of MFI used in our calculations with the corresponding crystallographic T-sites. Hydrogen atoms are shown in white.

The ZnAl2Si17O22H32 cluster analyzed in this work is composed of 74 atoms. The cluster was constructed by cutting the periodic structure, and the boundary silicon atoms were saturated with hydrogen atoms. These hydrogen atoms were aligned along the same direction of the oxygen atoms that were removed, and the Si−H bond lengths were set to 1.50 Å. Two Al atoms were substituted within the α-ring to stabilize the Zn2+ charge on the cluster. As illustrated in Figure 1(b), the α-ring of MFI has seven silicon atoms. We systematically analyzed the clusters that can be formed by replacing two Si atoms within the α-ring with two Al atoms. We further performed geometry optimizations on the clusters, and no symmetry constraints was imposed in any of the systems. Thus, we considered all the different configurations in which the Al atoms could be located within the α-ring. The only constraint that we followed was the Loewenstein rule, which specifies that there cannot be two consecutive AlO4 tetrahedra. Therefore, clusters containing Al-O-Al sequences were not considered in this work. As shown in Figure 2(b), this results in 13 different
configurations. All geometrical optimizations were performed on neutral charge clusters with singlet spin state.

All calculations conducted in this work used ωB97XD as the exchange-correlation functional, which is a long-range corrected functional. The basis set LANL2DZ was used for the Zn atom, while 6-31+g** was used for Al, O, Si, H, and, C atoms. In all geometry optimization calculations, all atoms within the cluster were relaxed, except for the terminal H-atoms that were kept fixed. The methodology that we employed in this work has been previously applied and validated by different authors for the computational study of zeolites, such as MFI, BEA, and FAU. The partial atomic charges were obtained with the natural bond orbital (NBO) population analysis. Vibrational frequencies were calculated for each optimized structure, within the rigid-rotor, harmonic-oscillator approximation. All the transition states were corroborated by displaying only one imaginary frequency along the reaction coordinate. All calculations were performed with Gaussian 09.

3. RESULTS AND DISCUSSION

The relative energies of the MFI α-ring isolated clusters with respect to the most stable Al atom configuration are compared in Figure 2(a). The T11-T2 structure is the most stable configuration, followed by T11-T1. The most unstable cluster is T5-T4, and it is 113 kJ/mol higher in energy than T11-T2. By comparing some of the geometrical parameters among the optimized α-ring clusters, we note that the Zn atom is always located at the center of the α-ring, and it is coordinated to four oxygen atoms in all configurations. On the isolated clusters, the Zn atom has a square planar arrangement with respect to the oxygen atoms on the α-ring. Our calculated Zn-O distances in the isolated clusters are comparable to the Fe-O distances reported on Fe-exchanged MFI. The
differences on the bond lengths range from 0.7 % when Al-atoms are located on the T7-T1 sites up to 2.2 % on the T11-T2 Al-configuration. Similarly, the Zn-O distances are also comparable to those reported by Montejo-Valencia et al. on Cu-exchanged MFI. The reported Cu-O bond lengths are just 3.4% lower than those that we obtained for Zn-O. The Al-Zn distances in each cluster, which range from 2.75 to 3.36 Å, are reported in Table S1 in the Supporting Information. We note that in T11-T2, which is the lowest energy configuration, the two Zn-Al distances are equal, with a value of 2.91 Å. We also note that the O-Al-O angles formed on that configuration are very similar, with values of 89.2° in T11, and 91.0° in T2 site. Thus, the inclusion of the Al atoms at T11 and T2 crystallographic sites yields a quasi-symmetric structure with similar distances and angles, in contrast to the other configurations, in which the angles and distance parameters are not symmetric.
Figure 2. (a) Relative energy of the MFI α-ring cluster with respect to the lowest energy configuration, T11-T2. (b) Schematic representation of the 13 Al configurations of the α-ring. Atoms beyond the α-ring were removed from the figures for clarity.
Methane activation involves two elementary steps: (1) methane adsorption and (2) detachment of a hydrogen atom to form methyl (CH3) and an –OH Brønsted site. The –OH Brønsted site can be formed in any of the oxygen atoms within the α-ring or in an oxygen atom adjacent to the α-ring. It is known that the location of the –OH Brønsted site yields different reaction energies in methane activation. Hence, we have systematically analyzed two plausible reaction paths for methane activation on all the 13 Al-atom configurations of the α-ring of Zn-MFI, depending on whether the Brønsted site is formed on an external oxygen atom, Oe, or an internal oxygen atom, Oi. For example, Figure 3 shows the location of the oxygen atoms in which the –OH Brønsted sites can be formed on the T11-T2 Al configuration. In that figure, we labeled the oxygen atoms within the α-ring as Oi-n, where n is a number from one to four, and the external oxygen atoms that are located outside the α-ring as Oe-n.

**Figure 3.** Oxygen atoms in which the –OH Brønsted acid can be formed in the T11-T2 configuration. Oe, highlighted in light blue, denotes the oxygen atoms located outside the α-ring. Oi, highlighted in light green, denotes the oxygen atoms within the α-ring.
Once the internal and external oxygen atoms were identified, we analyzed the minimum energy path. For each of the 13 Al configurations shown in Figure 2(b), we calculated the reaction energies for methane dissociation, by considering all possible oxygen atoms in which the -OH Brønsted site can be formed on each cluster. In this work, we considered the formation of hydroxyl groups on the oxygen atoms that are directly bonded to the Al atoms. Thus, the total number of Brønsted sites that can be formed varies depending on the Al configuration. For instance, Figure 4 shows the reaction energies when the Al atoms are located on the $\text{T8-T5}$ crystallographic sites. For this configuration, the reaction energies for methane dissociation range from 5 kJ/mol up to 93 kJ/mol. All the reaction energies obtained for H abstraction are endothermic. The reaction energies for the 13 Al atom configurations shown in Figure 2(b) are tabulated in Figure S2 in the Supporting Information.

**Figure 4.** Reaction intermediates that can be formed upon methane activation with the various oxygen atoms on the $\alpha$-ring of Zn-MFI when Al atoms are in $\text{T8-T5}$ crystallographic sites, and the HOMO of the isolated Zn-MFI cluster. Reaction energies for the formation of the intermediates were calculated with the energy of the isolated species as reference, i.e. $\Delta E_{\text{CH4-DISS}} = E_{\text{complex}} - (E_{\text{methane}} + E_{\text{Zn-MFI-T8-T5}})$.

In Figure 4, we also illustrate the highest occupied molecular orbital (HOMO) for the isolated cluster when the Al atoms are located on the crystallographic sites $\text{T8-T5}$. Our results indicate that for a specific Al configuration, the lowest reaction energy for methane dissociation is obtained when the Brønsted acid site is formed on the oxygen atom in which the HOMO of the isolated
cluster is located. For the configuration shown in Figure 4, the lowest reaction energy was obtained when the Brønsted site is formed on **Oe-1**. Figure S3 in the Supporting Information illustrates the HOMO’s for each isolated cluster along with all the reaction energies. Based on these results, the location of the HOMO orbital can be used to identify the oxygen atom in which the formation of the Brønsted site would yield the lowest reaction energy for methane dissociation.

We analyzed two plausible reaction paths for methane dissociation on each of the clusters on the \( \alpha \)-ring configurations of Zn-MFI, as shown in Scheme 1. Complex **I** corresponds to the isolated species, i.e. methane and the cluster of the Zn-MFI zeolite. The first step in the reaction mechanism is methane adsorption to form Complex **II**, in which methane is physisorbed on the zeolite. The carbon atom of methane is attracted to the positive charge of the Zn atom, but no chemical bond is formed. For all the Al configurations that were analyzed, methane physisorption is exothermic with minimal variations among adsorption energies. The values of these energies range between -36 to -48 kJ/mol corresponding to **T7-T5** and **T8-T5**, respectively. Our calculations demonstrate that upon methane physisorption, the geometrical parameters of the \( \alpha \)-ring in Complex **II** are nearly identical to the isolated counterparts. More pronounced differences, however, are obtained when comparing the distances between the zinc and carbon atom in Complex **II** on the different Al configurations. These distances vary from 2.49 to 2.70 Å, as shown in Table S2 in the Supporting Information.
Scheme 1. Reaction mechanisms studied for methane activation on Zn-MFI α-ring for the clusters in which the Al atoms are located on T8-T5 crystallographic sites. The most energetically favorable path involves the formation of the Brønsted acid in an external oxygen atom of the α-ring. Atoms beyond the α-ring were removed from the figures for clarity.

In complex III-Oe, the Brønsted site is formed with an external oxygen atom (Oe), whereas in complex III-Oi the Brønsted site is formed inside the α-ring, with an internal oxygen atom (Oi) as depicted in Scheme 1. The energies of these two complexes are different. The energies, relative to the isolated species in complex I, are 5 kJ/mol for complex III-Oe and 39 kJ/mol for complex III-Oi. The transition states to form complexes III-Oe and III-Oi (i.e., TS-Oe and TS-Oi, respectively) are also different. The energy of TS-Oe is 87 kJ/mol, while the energy of TS-Oi is 121 kJ/mol. Thus, when the Al atoms are located on the T8-T5 crystallographic sites, the minimum energy path for methane activation is through the formation of a -OH Brønsted acid site on an external oxygen atom. Figure 5 shows the minimum energy reaction pathways for the 13 Al configurations of the α-ring of Zn-MFI. Figure 5(a) illustrates the results for the systems in which the minimum energy path for methane activation occurs through an external oxygen atom of the α-ring. Figure 5(b) illustrates the results of the systems in which methane activation is more
favorable to occur through an internal oxygen atom within the $\alpha$-ring. We included in Table S3 in the Supporting Information the energy barriers, which take into account the energy of methane adsorption.

Our calculations demonstrate that the geometrical parameters of the $\alpha$-ring of the clusters in complex III are modified upon methane activation. For instance, the distance between C and Zn atoms is decreased in all Al-arrays, resulting in the bond formation between these two atoms. For all the Al configurations, the Zn-C bond length was on average 2.01 Å, suggesting that the Zn-C distance is independent of the location of the Brønsted site. This bond length is in good agreement with the experimental value reported by Haaland et al., with an error of 2.5%. The formation of the Zn-C bond causes the Zn atom to lose a coordination with one of the oxygen atoms within the $\alpha$-ring, and this is reflected through the increase of the Zn-O distance. For instance, in the T8-T5 Al-configuration the Zn-O distance increased from 2.11 to 3.03 Å. The Zn-O distances for the isolated clusters, for the transition states, and for complex III are tabulated in Tables S4, S5 and S6 in the Supporting Information.
Figure 5. Minimum energy path for methane activation on the α-ring of Zn-MFI for all the Al configurations illustrated in Figure 2(b). For each Al configuration, the zero energy corresponds to the isolated cluster and methane. Complexes I, II, III, TS-Oe and TS-Oi are identified in Scheme 1. (a) Energy diagrams for the systems in which the hydrogen abstraction occurs through an external oxygen atom, Oe, and (b) through an internal oxygen atom, Oi, of the α-ring.

As shown in Figure 5(a) and (b), the energies barriers to form the transition states TS-Oe and TS-Oi vary significantly depending on the Al atom configuration on the α-ring. TS energies to form TS-Oe range from 87 to 122 kJ/mol, and TS energies to form TS-Oi range from 89 to 131 kJ/mol. We note that the most stable Al configuration of the isolated cluster, T11-T2, yields an activation barrier of 122 kJ/mol, which is the highest barrier obtained. In contrast, the lowest barrier to form TS-Oi was 89 kJ/mol on T7-T5, whereas the lowest barrier to form TS-Oe was 87 kJ/mol on T8-T5. The energy barriers that we have obtained are in good agreement with the energy barrier reported in the literature of 96 kJ/mol.21 Nevertheless, we note that the energy barriers that
we calculated differ to those reported by Arzumanov et al.\textsuperscript{7} The discrepancy in the energies is probably due to the differences in the material. Arzumanov et al. reported that the Brønsted acid sites in the zeolite could act as cocatalysts by weakening the C-H bond in methane. In this work, however, we calculated the energy barriers on a catalyst that has no Brønsted acid sites prior to the reaction. Future work should address the impact of the Brønsted acid sites on the kinetics of methane activation.

We also note that the energy barriers that we obtained are comparable to the energy barrier that has been reported for methane dissociation on Cu-exchanged MFI with the same Al-atoms configuration, with a difference of only 10 kJ/mol.\textsuperscript{21} On the other hand, the energy barriers on Zn-MFI are significantly lower than the energy barrier reported for Fe-MFI of 169 kJ/mol when oxo-clusters [Fe–O–Fe] are the active site.\textsuperscript{20}

In all transition states geometries, the Zn-C distance decreases when compared to those obtained in complex II, and it ranges from 2.07 to 2.21 Å. Furthermore, we note that transition state structures involving an external oxygen atom yield higher deformations in the zeolite than TS-Oi. For instance, when an external oxygen atom is involved in the transition state there is an increase of up to 16 % in the O-Al-O angle on TS-Oe when compared to the angle on complex II. On the other hand, on TS-Oi the maximum increase in that angle is 7 %. As shown in Figure 5(a) and (b), the overall reaction energies for the formation of complexes III-Oe and III-Oi are different for each Al configuration. Reaction energies range from 5 kJ/mol to 70 kJ/mol when an external oxygen atom is involved, and from 35 kJ/mol to 104 kJ/mol when an internal oxygen atom participates in the reaction. The T8-T5 structure yields the lowest reaction energy for methane activation, with an overall reaction energy of 5 kJ/mol.
The acidity of the zeolites has been widely suggested to affect its catalytic activity.\textsuperscript{47–49} While a
debate still exists on how to accurately measure zeolite acidity, previous works have related the
energy of the lowest occupied molecular orbital (E_{LUMO}) to the Lewis acidity of zeolites.\textsuperscript{50} In order
to verify if E_{LUMO} can be correlated to the catalytic activity of the zeolite, we plotted the TS
energies with the corresponding LUMO energy of the isolated clusters as shown in Figure S4 in
the Supporting Information. Our calculations demonstrate that the LUMO energies are not
appropriate descriptors of the TS energies for this particular system, since no clear trend is obtained
in the two reaction pathways analyzed in this work. However, we note that the lowest LUMO
energy corresponds to the T8-T5 Al configuration, which is the configuration with the lowest TS
energy.

The Al-atom distribution within the active site of Zn-MFI can significantly affect the energy
required for the abstraction of the H atom of methane. Thus, we further analyzed the effect of the
Al position on the active site of Zn-MFI, and our calculations demonstrate that the partial atomic
charge of the Zn atom (q_{Zn}) varies depending on the position of the Al atoms within the α-ring of
MFI. The partial atomic charge q_{Zn} ranged from 1.57 for T5-T4 to 1.63 for T8-T5. All the values
are shown in Table S1 in the Supporting Information. Our calculations further revealed that these
partial charges, q_{Zn}, could be correlated to the energy barrier of methane dissociation. Figure 6(a)
shows the trends for the Al configurations in which the minimum energy path leads to the
formation of complex TS-Oi. For the clusters in which it is energetically more favorable to form
the Brønsted site within the α-ring of Zn-MFI, the energy of the transition state decreases as the
charge of the Zn atom decreases. However, for the clusters in which it is favorable to form the
Brønsted site outside the α-ring, the transition state energy decreases with increasing charge of the
Zn atom, as shown in Figure 6(b). Trends are also maintained with enthalpies and Gibbs free
energies, as shown in Figure S5 in the Supporting Information. The location of the Al-atoms has an impact on the electronic density along the α-ring, which is manifested through the partial atomic charges and the electrostatic potential of the isolated clusters. The NBO charges and the electrostatic potentials of the 13 isolated clusters are illustrated in Table S7 in the Supporting Information. We note that the different electrostatic environments directly affect the strength through which the Zn and the C atoms interact, and also this impacts the geometry of the transition state. For instance, Zn-C distance in the methane physisorption in complex II varies depending on the partial atomic charge of the Zn atom. Furthermore, the distance between the H-atom and the oxygen that is involved in the transition state also varies depending partial atomic charge of the Zn atom. Thus, the Al atom position within the active site of Zn-MFI affects the partial atomic charge of the Zn atom, which also impacts the energy barrier in these catalytic systems. Therefore, these results suggest that the Al-atom position could be a design parameter that could be optimized to improve the catalysis of methane activation.

Figure 6. Transition state energies relative to the isolated species as function of the partial atomic charge of the Zn atom in the isolated cluster (I). (a) Transition state energies for H-atom abstraction
through an Oi atom in the α-ring (▲) and (b) transition state energies for H-atom abstraction through an Oe atom in the α-ring (♦).

4. CONCLUSIONS

We have conducted a systematic DFT study to analyze the effect of Al atom distribution within the α-ring of Zn-MFI on the activation of methane. We analyzed 13 different clusters of the α-ring of Zn-MFI. Our results demonstrate that the most stable configuration for the Zn-exchange cluster of the α-ring is obtained when the Al atoms are at the T11-T2 crystallographic sites. Our results indicate that physisorption energies of methane on the active site of the Zn-exchanged MFI cluster are independent of the configuration of Al atoms in the α-ring, with physisorption energies that range from -36 to -48 kJ/mol. The activation of methane takes place on Zn, and upon activation a hydrogen atom binds to an oxygen atom of the zeolite framework. This reaction yields the formation of a Brønsted –OH acid site, which can be formed at an oxygen atom within the α-ring or at an oxygen atom that lies outside the α-ring. For each of the 13 clusters, we analyzed the reaction pathways with all the possible oxygen atoms. Our calculations revealed that for all the Al-configurations, the lowest reaction energies were obtained when the Brønsted-OH site is formed at the oxygen atom in which the HOMO of the isolated cluster is located. Furthermore, our results revealed that the Al-atom configurations affects the partial atomic charge of the Zn-atom, and this partial charge can be correlated to the transition state energy of methane activation. The fundamental studies conducted in this work aims to provide a correlation based on the electron density for the activation of methane on Zn-MFI zeolites. This work contributes to the rational design of catalytic materials for methane activation, which can provide experimentalist with insights on the synthesis of highly active zeolite catalyst for methane activation.

Supporting Information
Geometric parameters of the configurations on the reaction mechanism, reaction energies for methane activation, and HOMOs and LUMOs of each isolated Zn-MFI cluster (PDF) xyz coordinates of stable intermediates and transition states (ZIP)

Notes
The authors declare no competing financial interests.

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