1. Introduction

Enhancing the efficiency for the selective activation of carbon–hydrogen bonds is linked to the success in generating new or improving existing catalysts.\textsuperscript{3–5} To this end, great efforts have been undertaken to reveal the mechanisms of bond activation processes at a molecular level.\textsuperscript{3–6} Among the various catalysts so far applied in industry, quite a few employ transition metals. Zinc-based catalysts are also in use, for example in the oxidative conversion of CH\textsubscript{4}, C\textsubscript{2}H\textsubscript{6} and C\textsubscript{3}H\textsubscript{8}.\textsuperscript{9} Further, zinc-doped zeolites are known to be effective catalysts for promoting dehydrogenation and aromatization of light alkanes, and Zn species including [Zn(OH)]\textsuperscript{+} are believed to play a key role in these mechanistically rather complex transformations.\textsuperscript{10–12} Also, pure or Li doped zinc oxides act as catalyst for the C–H bond activation of light alkanes, e.g. in the oxidative coupling of methane or the oxidative dehydrogenation of ethane and propane.\textsuperscript{13} While in all these reactions, Zn species are considered as the active ingredients, the reaction mechanisms as well as the precise structure and exact composition of the active sites of the catalysts are still under debate.\textsuperscript{12,14,15} In this respect, gas-phase experiments have proven useful because they provide in an unperturbed way rather detailed insight into the elementary steps of numerous transformations mediated by zinc-containing catalysts.\textsuperscript{16–18}

There have been several experimental and theoretical studies on the gas-phase reactions with various zinc species.\textsuperscript{19–23} As shown by Georgiadis and Armentrout, C–C bond cleavage of alkanes can be achieved by atomic [Zn].\textsuperscript{24} Further, the interaction of neutral [ZnO] with CH\textsubscript{4} has been investigated by theoretical methods, and possible pathways yielding syngas, CH\textsubscript{2}O, and H\textsubscript{2}O were postulated.\textsuperscript{25} Kretschmer et al. reported N–H bond activation of NH\textsubscript{3} by [Zn(OH)]\textsuperscript{+}, and CO\textsubscript{2} activation has been brought about in the reaction of [L\textsubscript{n}Zn(OH)]\textsuperscript{+} (L = imidazole and pyridine; n = 1, 2) in analogy with the Lipscoumb mechanism for carbonic anhydrase.\textsuperscript{18} However, the activation of C–H bonds of light alkanes with zinc hydroxide is much less investigated. This is rather surprising given the fact that well-designed gas-phase processes of transition-metal fragments using advanced mass-spectrometric techniques in conjunction with theoretical studies have greatly helped in uncovering mechanistic aspects underlying C–H bond activation.\textsuperscript{5,7,8,26–33}

Herein we present a combined experimental/theoretical investigation of the gas-phase reactions of cationic zinc hydroxide...
with alkanes. While $[\text{Zn(OH)}]^+$ does not activate methane and ethane, mechanistically rather remarkable processes with propane are observed. As will be shown, studying the mechanistic aspects of C-H bond activation by zinc species proves helpful to understand the role of zinc in catalysis in a broader context.

2. Methods

Experiments were performed with a VG BIO-Q mass spectrometer of QHQ configuration (Q: quadrupole, H: hexapole) equipped with an ESI source, as described previously in detail.34 To this end, Q(1) is used for mass-selection of the ion of interest and then, in the r-f only hexapole, the ion/molecule reactions are conducted. Ionic products are analyzed by scanning Q(2). Further, for ESI, millimolar solutions of $\text{Zn(NO}_3\text{)}_2$ in pure methanol were introduced through a fused-silica capillary to the ESI source by a syringe pump (ca. 4 $\mu$L min$^{-1}$) to produce the $[\text{Zn(OH)}]^+$ cations. Nitrogen was used as a nebulizing and drying gas at a source temperature of 80 °C. Maximal yields of the desired complexes were achieved by adjusting the cone voltage ($U_c$) to 80 V; $U_c$ determines the degree of collisional activation of the incident ions in the transfer from the ESI source to the mass spectrometer. The identity of the ions was confirmed by comparison with the calculated isotope patterns, which also assisted in the choice of the adequate precursor ion to avoid coincidental mass overlaps of isobaric species in the mass-selected ion beam.35 Here, we selected $[^{64}\text{Zn(OH)}]^+$ as the reactant ions by means of Q(1). In the hexapole, the ion/molecule reactions with $\text{CH}_4$, $\text{C}_2\text{H}_6$, $\text{C}_2\text{D}_6$, $\text{C}_3\text{H}_8$, $\text{C}_3\text{D}_8$, $\text{CD}_3\text{CH}_2\text{CD}_3$, and $\text{CH}_3\text{CD}_2\text{CH}_3$ were probed at a collision energy ($E_{\text{coll}}$) set to nominally 0 eV; this, in conjunction with the kinetic energy width of about 0.4 eV of the parent ion at peak half-height, allowed the investigation of quasi-thermal reactions, as demonstrated previously.36

Since absolute rate constants cannot readily be determined by using the experimental setup of the VG BIO-Q mass spectrometer, the rate constant and the branching ratios of the reaction of $[\text{Zn(OH)}]^+$ with $\text{C}_3\text{H}_8$ have been determined by using a Spectropac CMS 47X Fourier Transform Ion Cyclotron Resonance (FT-ICR) mass spectrometer; details of the instrument have been described previously.37,38 Atomic $[\text{Zn}]^+$ ions were generated by laser ablation of pure Zn metal disks using a Nd:YAG laser operating at 1064 nm in the presence of helium carrier gas. The $[^{64}\text{Zn}]^+$ isotope was isolated and allowed to react with a mixture of $\text{N}_2\text{O}$ and $\text{H}_2\text{O}$ (ca. 5:1) to give $[^{64}\text{Zn(OH)}]^+$. The so-formed product ions are subsequently quenched by collisional thermalization with the buffer gas (argon, ca. 2 x 10$^{-6}$ mbar). After collisional thermalization, the $[^{64}\text{Zn(OH)}]^+$ species were mass-selected and exposed to react with $\text{C}_3\text{H}_8$ by introducing the substrate through a leak-valve. For the thermalized ions a temperature of 298 K was assumed.36,37 The branching ratios have been determined by extrapolating the ratios at different reaction times to $t = 0$ s. Note, that somehow different branching ratios are obtained by using the two types of mass spectrometers applied in this study, cf. Fig. 1a (branching ratios of reactions (a), (b) and (c) are 78%, 12% and 10%) and eqn (a)-(c); these differences may reflect the lack of proper collisional thermalization in the experiments using the VG BIO-Q mass spectrometer.

Calculations were carried out by using the Gaussian 09 program suite.39 Potential energy surfaces (PESs) are calculated by using the Möller–Plesset second-order perturbation MP2 method35,40 employing a triple-zeta level basis set with diffuse and polarization functions 6-311++G(2d,2p) for all atoms.41 To obtain even more accurate energies of the relevant structures, the coupled-cluster CCSD(T) method42,43 with single, double, and perturbative treatment of triple excitations in conjunction with the correlation-consistent polarized valence triple-zeta basis sets cc-pVTZ was used.44,45 The MP2/6-311++G(2d,2p) optimized geometries were employed for the single-point coupled cluster calculations without reoptimization at the CCSD(T)/cc-pVTZ levels, and the results are in line with the MP2 calculations (Table 1). All geometries were fully optimized without symmetry constraints. Vibrational frequency calculations were performed to identify the nature of reaction intermediates, transition states (TSs) and products. To corroborate which minima are linked by the considered transition states, normal coordinate analyses were performed on these TS structures by intrinsic reaction coordinate (IRC) routes in both reactant and product directions.46–48 Additional geometry optimizations starting from the last IRC structures were carried out when the IRC calculations did not converge. Unscaled vibrational frequencies were used to calculate zero-point energy (ZPE) corrections. To demonstrate the applicability of the MP2 method selected for this study, test calculations were performed at the MP2/6-311++G(2d,2p) level of theory (Table S1, ESi); the results are in agreement, within 0.36 eV, of experimental values.21,22,49

The relative energies of P1 and TS3/4 in Fig. 2 have also been calculated by DFT using different functionals50–59 and the 6-311++G(2d,2p) basis set, as well as by single-point energy calculations using the CCSD(T) method; the results are listed in Table S2 (ESi). MP2 calculated relative energies of high-spin and low-spin products of the reaction of $[\text{Zn(OH)}]^+$ with $\text{C}_3\text{H}_8$, $\text{C}_2\text{H}_6$, and $\text{CD}_3\text{CH}_2\text{CD}_3$.
and 

Table 1 The relative energies (in eV) of intermediates, transition states and products obtained by MP2/6-311+G(2d,2p) and CCSD(T)/cc-pVQZ (relative to separated C\(_3\)H\(_8\) and [Zn(OH)]\(^+\))

<table>
<thead>
<tr>
<th>Path 1</th>
<th>I1</th>
<th>TS1/2</th>
<th>P1</th>
<th>I2</th>
<th>TS3/4</th>
<th>I4</th>
<th>P2</th>
</tr>
</thead>
<tbody>
<tr>
<td>MP2</td>
<td>-1.70</td>
<td>-0.38</td>
<td>-0.39</td>
<td>-0.20</td>
<td>-1.70</td>
<td>-0.40</td>
<td>-1.78</td>
</tr>
<tr>
<td>CCSD(T)</td>
<td>-1.65</td>
<td>-0.31</td>
<td>-0.32</td>
<td>-0.13</td>
<td>-1.65</td>
<td>-0.38</td>
<td>-1.62</td>
</tr>
<tr>
<td>TS5/4</td>
<td>I5</td>
<td>P3</td>
<td>Path 2</td>
<td>I6</td>
<td>P4</td>
<td>I7</td>
<td>TS7/8</td>
</tr>
<tr>
<td>-1.70</td>
<td>-2.22</td>
<td>-1.23</td>
<td>MP2</td>
<td>-1.64</td>
<td>0.06</td>
<td>-1.82</td>
<td>-0.12</td>
</tr>
<tr>
<td>-1.75</td>
<td>-2.23</td>
<td>-1.12</td>
<td>CCSD(T)</td>
<td>-1.58</td>
<td>0.19</td>
<td>-1.80</td>
<td>-0.17</td>
</tr>
<tr>
<td>P5</td>
<td>Path 3</td>
<td>I9</td>
<td>TS9/10</td>
<td>I10</td>
<td>P6</td>
<td>TS1</td>
<td>TS2</td>
</tr>
<tr>
<td>-1.07</td>
<td>MP2</td>
<td>-0.17</td>
<td>0.16</td>
<td>-0.16</td>
<td>0.06</td>
<td>0.16</td>
<td>0.30</td>
</tr>
<tr>
<td>-0.89</td>
<td>CCSD(T)</td>
<td>-0.17</td>
<td>0.06</td>
<td>-0.07</td>
<td>0.08</td>
<td>0.18</td>
<td>0.20</td>
</tr>
</tbody>
</table>

To obtain additional insight in the mechanisms of the reactions of [Zn(OH)]\(^+\) with C\(_3\)H\(_8\), MP2/6-311+G(2d,2p) calculations have been performed, and the corresponding PESs are shown in Fig. 3 and Fig. S1 (ESI\(^\ddagger\)). Overall, these reactions are controlled by the Lewis-acid character of the metal center interacting with the electron donating C–H bonds of propane. Some pertinent details of the most favorable PESs are shown in Fig. 3a and b; possible pathways which involve the OH moiety interacting with C–H bonds of propane have also been tested (Fig. 3c) but turned out to be higher in energy. Four different encounter complexes have been located on the PES for the initial interaction of the metal center with C–H bonds of C\(_3\)H\(_8\). In the iso-energetic I1 and I3, the metal interacts with two hydrogen atoms of the secondary position, and H–Zn bond lengths amount to 187/185 pm for I1 and 193 pm for I2, respectively. In I6 two hydrogens from one methyl group participate while in I7, one C–H bond of each of the two methyl groups is involved. All four intermediates form remarkably stable ion/molecule complexes [Zn(OH)(C\(_3\)H\(_8\))]\(^+\), and these complexes profit from the electron donation from the C–H bonds into the empty 4s–4p hybrid orbital of zinc (see also below). These interactions are indicated by the fact that the coordinating C–H bonds are slightly elongated (from 109 pm in free propane to 112 pm in I1 and I3, and to 110–111 pm in I6 and in I7, respectively). Regarding the hydride transfer from a methyl group in intermediate I6 to the Zn atom, no barrier has been located in this step I6 → P4 ([i-C\(_3\)H\(_7\)]\(^+\)/[HZn(OH)]\(^+\)); in contrast, the hydride transfer from I1 to P1 ([i-C\(_3\)H\(_7\)]\(^+\)/[HZn(OH)]) proceeds via transition structure TS1/2 and intermediate I2. However, TS1/2 and I2 are almost iso-energetic, i.e. this pathway proceeds in a quasi barrier-free process. Notably, while the formation of product P1 is exothermic (−0.20 eV) and thus accessible under thermal conditions, product P4 is much higher in energy (0.06 eV); this is in line with the labeling experiments clearly favoring the former reaction (see above). As shown in Fig. S1a (ESI\(^\ddagger\)), formation of [i-C\(_3\)H\(_7\)]\(^+\) is also accessible via the more complex pathway R → I1′ → TS1′/2′ → I2′ → P1. In the formation of [i-C\(_3\)H\(_7\)]\(^+\) (Fig. 3a and Fig. S1a, ESI\(^\ddagger\)), neutral HZn(OH) is co-generated; the alternative to produce a neutral water complex Zn(H\(_2\)O) is less favorable both kinetically and thermodynamically (product P6 of path 3, Fig. 3c). Likewise, the combined formation of Zn(H\(_2\)O) and [i-C\(_3\)H\(_7\)]\(^+\) is more endothermic than generating HZn(OH) and [i-C\(_3\)H\(_7\)]; the former product pair is 0.30 eV above the entrance channel (P8, Fig. S1c, ESI\(^\ddagger\)). Thus, the interaction of the OH group of [Zn(OH)]\(^+\) to C–H bonds of C\(_3\)H\(_8\) (Fig. 3c and Fig. S1c, ESI\(^\ddagger\)) respectively to form neutral Zn(H\(_2\)O) cannot compete with the initial coordination of the metal center to secondary or primary C–H bonds of propane, respectively (Fig. 3a and b). These findings are also confirmed in single point energy calculations using the CCSD(T) method.

For the regiospecific ODH process, the sequence I3 → TS3/4 → I4 → P2 (Fig. 4a) constitutes the energetically most favorable pathway. Formation of propene takes place via TS3/4, in which neutral HZn(OH) interacts with [i-C\(_3\)H\(_7\)]\(^+\) resulting in intermediate I4; the NBO atomic charge of the HZn(OH) moiety in TS3/4 corresponds to 0.06 [\(\epsilon\)] and the associated IRC paths are shown.

[Diagram of Reaction Pathways]
The weakly bound propene ligand in I4 can easily be liberated to yield the experimentally observed ODH product [HZn(H₂O)]⁺ (P2). In competition, I5 is generated in which the Zn atom is coordinated to the C=C double bond of C₃H₆. The weakly-bound H₂O group can be eliminated yielding the cationic product [HZn(CH₃CH₂)]⁺ (P3), which is experimentally observed in reaction (c). A kinetically less favorable pathway for the elimination of water and formation of [Zn(n-C₃H₇)]⁺ of reaction (c) is shown in Fig. 2b (see below). The energetic requirement to produce P2 and P3 are comparable (−1.12 versus −1.23 eV); this is consistent with similar branching ratio of reactions (b) and (c) as observed experimentally. As to the energetics of the competitive productions of P1 versus P2 and P3, the relative energy of P1 (−0.20 eV) is higher than that of TS3/4 (−0.40 eV). The same order of relative energies of P1 and TS3/4 are also obtained by DFT calculations using different functionals as well as single point energy calculations using the CCSD(T) method (see Table S2, ESI†). Taking into account the errors as well as previous work,⁶⁷–⁶⁹ our calculations are in agreement with the branching ratio of the production of [i-C₃H₇]⁺/[HZn(OH)] (reaction (a)) versus reactions (b) and (c) (ca. 54% : 46% by using the FT-ICR mass spectrometer). In addition, the direct dissociation I₁ → P₁ is kinetically favored over a more complex rearrangement/dissociation path proceeding via the tight transition state TS3/4.

For the alternative water formation pathway of reaction (c), as shown in Fig. 2b, the generation of [Zn(n-C₃H₇)]⁺ occurs along the route I₇ → TS7/8 → I₈ → P₅ with the intermediate formation of a propyl–water complex I₈, [Zn(C₃H₇)(H₂O)]⁺;
mechanistically, this is a σ bond metathesis reaction. The relative energy of TS7/8 amounts to ~0.12 eV, which is higher in energy as compared to TS3/4 (~0.40 eV), and thus unlikely to compete efficiently under thermal conditions. A brief comparison of structural features of the transition states for the σ-metathesis reaction, i.e. TS7/8 versus TS1 (Fig. S1b, ESI) is indicated. In TS7/8, the metal center is not only a constituent of the four-membered ring which is essential for a σ-bond metathesis but is also coordinated by a C–H bond of the distal methyl group of propane; this results in a slight but clearly discernible elongation of the C–H bond from 109 pm to 113 pm. Quite likely, this agostic interaction is beneficial for the stabilization of TS7/8 versus TS1; the latter one lacks this stabilization.

Finally, with regard to the C–H bond activation in [Zn(OH)]⁺, the 4s orbital hybridizes with a 4p orbital of Zn leading to the formation of a σ bond metathesis reaction. The highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO), respectively (Fig. 3a and b). While the former is used to bind to the OH ligand, the latter can accept two electrons from a C–H bond of propane resulting in hydride transfer and carbocation formation; this view is supported by the similarity of the HOMO of neutral [HZn(OH)] with the LUMO of [Zn(OH)]⁺ (Fig. 3b and c).

On the triplet PES, all reaction channels described above for the [Zn(OH)]⁺–C₃H₈ system have been calculated to be endothermic (Table S3, ESI) with the conversion of alkane by zinc catalysts.⁷²–⁷⁵ Thus, the identification of the active sites of ZnOH species helps to unravel part of the enigma associated with the conversion of alkane by zinc catalysts.⁷²–⁷⁵

4. Conclusion

Here, we report and analyze the thermal gas-phase reactions of [Zn(OH)]⁺ with C₃H₈ by using experimental and theoretical methods. The reactivity of cationic zinc hydroxide [Zn(OH)]⁺ toward C₃H₈ is characterized by C–H bond activation; the main reaction channel corresponds to a hydride transfer from the hydrocarbon to the Lewis acidic metal center resulting in the generation of [HZn(OH)]/ [i-C₃H₇]⁺. Homolytic C–H bond activation give rise to an ODH channel (generation of propene) as well as the competitive formation of [HZn(CH₃CH=CH₂)]⁺/H₂O. Our study may prove helpful to further understand the industrially relevant, catalytic conversion of small alkanes by Zn species.

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