How Lewis Acids Catalyze Ring-Openings of Cyclohexene Oxide

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ABSTRACT: We have quantum chemically studied the Lewis acid-catalyzed epoxide ring-opening reaction of cyclohexene epoxide by MeZH (Z = O, S, and NH) using relativistic dispersion-corrected density functional theory. We found that the reaction barrier of the Lewis acid-catalyzed epoxide ring-opening reactions decreases upon ascending in group 1 along the series Cs⁺ > Rb⁺ > K⁺ > Na⁺ > Li⁺ > H⁺. Our activation strain and Kohn–Sham molecular orbital analyses reveal that the enhanced reactivity of the Lewis acid-catalyzed ring-opening reaction is caused by the reduced steric (Pauli) repulsion between the filled orbitals of the epoxide and the nucleophile, as the Lewis acid polarizes the filled orbitals of the epoxide more efficiently away from the incoming nucleophile. Furthermore, we established that the regioselectivity of these ring-opening reactions is, aside from the “classical” strain control, also dictated by a hitherto unknown mechanism, namely, the steric (Pauli) repulsion between the nucleophile and the substrate, which could be traced back to the asymmetric orbital density on the epoxide. In all, this work again demonstrates that the concept of Pauli-lowering catalysis is a general phenomenon.

INTRODUCTION

Cyclohexene oxides are valuable building blocks in synthetic chemistry, finding applications in many organic reactions. They can undergo chemical transformations with a broad scope of nucleophiles, making cyclohexene oxides useful building blocks for the synthesis of complex molecules. It is well known that nucleophilic ring-opening reactions of cyclohexene oxides can proceed with excellent regioselectivity. The Förster–Plattner rule attributes this regiochemical control to a large preference for the reaction pathway that follows the more stable chair-like transition state (i.e., attack at the α-position; Scheme 1) compared to the one proceeding through the unfavored twist boat-like transition state (i.e., attack at the β-position; Scheme 1). However, despite recent efforts, limited quantitative data is available regarding the origin of the regioselectivity in epoxide ring-opening reactions.

In general, this class of ring-opening reactions is extremely slow and requires a form of activation in order to proceed with efficient conversion. Lewis acids based on group 1 cations (e.g., LiClO₄) are popular catalysts to accelerate the reactivity of cyclohexene oxides, and their enhanced reactivity is generally ascribed to the stabilization (i.e., lowering) of the LUMO of the epoxide, which in turn leads to a smaller HOMO_nucleophile−LUMO_epoxide gap and hence a lower ring-opening reaction barrier. In contrast to the current rationale, we have shown recently that Lewis acids do not catalyze organic transformations by enhancing the orbital interactions (i.e., reducing the HOMO−LUMO gap) but instead by diminishing the steric (Pauli) repulsion between the reactants. The complexation between a Lewis acid and the reactant induces a polarization of the occupied orbital densities of the reactant away from the reactive region, resulting in less repulsive occupied−occupied orbital overlap and hence a lower reaction barrier. We envision that this concept of Pauli-repulsion lowering catalysis might also be the driving force in Lewis acid-catalyzed epoxide ring-opening reactions.

To ascertain the universality of this novel Pauli-repulsion lowering catalysis mechanism, we have here performed a comprehensive computational study to unravel the driving force behind Lewis acid-catalyzed cyclohexene oxide ring-opening reactions. In addition, we also studied the physical factors controlling the regioselectivity of this reaction. To this end, we have analyzed the potential energy surfaces of the uncatalyzed and Lewis acid-catalyzed ring-opening reaction of epoxide 1-Y⁺ (Y⁺ = none, Cs⁺, Rb⁺, K⁺, Na⁺, Li⁺, and H⁺) by MeZH (Z = O, S, and NH) using relativistic dispersion-corrected density functional theory (Scheme 1). The activation strain model (ASM) of reactivity in combination with Kohn–Sham molecular orbital (KS-MO) theory and the matching...
Scheme 1. α-Attack (i.e., Chair-Like TS) and β-Attack (i.e., Twist Boat-Like TS) of the Lewis Acid-Catalyzed Cyclohexene Oxide Ring-Opening of MeZH with Epoxide-Y⁺ (1-Y⁺), Where Z = O, S, and NH and Y⁺ = none, Cs⁺, Rb⁺, K⁺, Na⁺, Li⁺, and H⁺.

Table 1. Activation Strain Analysis, Energy Decomposition Analysis (in kcal mol⁻¹), and Key Geometrical Details (in Å) for the Interaction between the Lewis Acid Y⁺ and Epoxide 1 in the 1-Y⁺ Complex

| Y⁺ | ΔE     | ΔEstrain | ΔEint | ΔVskew | ΔEpush | ΔEstat | r (O···Y⁺) | r (Cα-O) | r (Cβ-O) |
|----|--------|----------|-------|--------|--------|--------|-----------|---------|---------|
| Cs⁺ | −17.7  | 0.2      | −17.9 | −14.9  | 4.7    | −7.7   | 2.830     | 1.467   | 1.461   |
| Rb⁺ | −19.5  | 0.2      | −19.7 | −15.8  | 4.6    | −8.5   | 2.680     | 1.468   | 1.462   |
| K⁺  | −23.4  | 0.5      | −23.9 | −22.0  | 8.2    | −10.1  | 2.521     | 1.470   | 1.464   |
| Na⁺ | −31.5  | 0.7      | −32.2 | −28.3  | 10.1   | −14.0  | 2.126     | 1.475   | 1.469   |
| Li⁺ | −44.9  | 1.1      | −46.0 | −34.8  | 13.3   | −24.5  | 1.760     | 1.485   | 1.479   |
| H⁺  | −201.3 | 10.8     | −212.1| −40.6  | 0.0    | −171.5 | 0.983     | 1.575   | 1.566   |

Electronic energies computed at ZORA-M06-2X/TZ2P//B3LYP-D3(BJ)/6-31G+(d), whereas Cs⁺ and Rb⁺ atoms were treated with the def2-TZVP basis set. In the isolated epoxide 1, Cα−O = 1.445 Å and Cβ−O = 1.438 Å.

RESULTS AND DISCUSSION

First, we have investigated the strength and nature of the interaction between epoxide (1) and the Lewis acid (Y⁺), adopting the most stable half-chair conformation of the complex 1-Y⁺ (see Figure S1 for conformational energy landscape (CEL) maps) using a combined activation strain and energy decomposition analysis approach (Table 1).

Interestingly, all computed epoxides are asymmetric, that is, the Cα−O bond is longer than the Cβ−O bond, which will, as we show later, play a prominent role in the regioselectivity of these epoxide ring-opening reactions. The complexation energies become more stabilizing on ascending in group 1, from −17.7 kcal mol⁻¹ for Cs⁺ to −201.3 kcal mol⁻¹ for H⁺, which is exclusively determined by the interaction energy. The strain energy, on the other hand, is marginal and only plays a role when Y⁺ = H⁺, where a significant Cα/β−O elongation in 1-H⁺ occurs. Notably, the trend in complexation energies (i.e., ΔE) is in line with the trend in alkali metal cation affinities (AMCA) studied by Boughlala et al. The O···Y⁺ bond length becomes systematically shorter from Cs⁺ to H⁺, which is in line with the decreasing effective size of the Lewis acid upon ascending in group 1. Both the electrostatic and orbital interactions play an important role in the 1-Y⁺ interaction and become increasingly more stabilizing along Cs⁺ to H⁺. The stronger electrostatic interaction is a result of a less diffuse electron density and a shorter O···Y⁺ bond length. The trend in orbital interactions, which is important for the strength of the catalysis (vide infra), can be rationalized by the use of a Kohn–Sham molecular orbital analysis (see Figure S2 in the Supporting Information). The formation of the O···Y⁺ bond involves a key donor–acceptor interaction between the oxygen lone pair orbital of 1 (HOMO−1 for Y⁺ = Cs⁺, Rb⁺, K⁺, Na⁺, and Li⁺; HOMO for Y⁺ = H⁺) and the empty ns atomic orbital (AO) of Y⁺. The HOMO of the epoxide is a lone pair orbital on the oxygen that is oriented perpendicular to the epoxide ring. The HOMO−1 of the epoxide, on the other hand, is a lone pair orbital on the oxygen oriented inplane with the epoxide ring (see schematic MOs in Figure S2). H⁺ can, due to its lack of electrons and hence no steric (Pauli) repulsion with the Cα and Cβ of the epoxide, interact with the higher energy HOMO of the epoxide. The larger group 1 cations, however, do have, if they interact with the HOMO of the epoxide, a significant steric (Pauli) repulsion with the Cα and Cβ of the epoxide. To minimize this steric repulsion, the group 1 cation will move away from the Cα and Cβ of the epoxide toward the HOMO−1 of the epoxide. Besides reducing the steric repulsion, this displacement of the cation...
Table 2: Energies Relative to Reactants (in kcal mol$^{-1}$) of the Stationary Points of the Uncatalyzed and Lewis Acid-Catalyzed Epoxide Ring-Opening Reactions between MeZH ($Z = O, S,$ and NH) and 1-$Y^+$ ($Y^+ =$ none, Cs$^+$, Rb$^+$, K$^+$, Na$^+$, Li$^+$, and H$^+$)$^a$

| MeZH   | species | none | Cs$^+$ | Rb$^+$ | K$^+$ | Na$^+$ | Li$^+$ | H$^+$ |
|--------|---------|------|--------|--------|-------|--------|--------|-------|
| MeOH   | RC      | −3.4 | −7.0   | −7.1   | −7.4  | −8.0   | −8.9   | −13.2 |
|        | TS-$\alpha$ | 53.0 | 21.5   | 21.1   | 19.1  | 16.1   | 10.0   | −11.7 |
|        | TS-$\beta$ | 57.1 | 28.3   | 28.1   | 26.0  | 19.6   | 13.1   | −8.1  |
|        | INT-$\alpha$ | b    | 19.2   | 18.9   | 16.4  | 11.6   | 1.2    | −30.3 |
|        | INT-$\beta$ | 57.0 | 23.8   | 23.5   | 21.7  | 18.6   | 7.7    | −23.8 |
| MeSH   | RC      | −3.6 | −5.6   | −5.9   | −6.1  | −6.5   | −7.1   | −38.1 |
|        | TS-$\alpha$ | 46.5 | 20.2   | 20.1   | 18.4  | 15.3   | 8.9    | −9.2  |
|        | TS-$\beta$ | 50.8 | 24.9   | 24.7   | 22.8  | 19.7   | 13.3   | −9.2  |
|        | INT-$\alpha$ | b    | 12.5   | 12.3   | 9.7   | 5.0    | −5.6   | −31.0 |
|        | INT-$\beta$ | 43.6 | 19.2   | 19.0   | 16.3  | 11.5   | 0.8    | −6.8  |
| MeNH$_2$ | RC     | −3.4 | −6.8   | −6.9   | −7.2  | −7.8   | −8.7   | −8.7  |
|        | TS-$\alpha$ | 34.4 | 12.0   | 11.7   | 10.0  | 6.6    | 0.5    | 0.5   |
|        | TS-$\beta$ | 38.5 | 16.0   | 15.6   | 13.9  | 10.6   | 4.4    | 4.4   |
|        | INT-$\alpha$ | 24.6 | −10.7  | −10.9  | −13.7 | −18.5  | −29.4  | −62.2 |
|        | INT-$\beta$ | 31.1 | −4.2   | −4.4   | −7.1  | −12.0  | −22.9  | −55.0 |

$^a$Electronic energies computed at M06-2X/6-311++G(d,p)//B3LYP-D3(BJ)/6-31G+(d), whereas Cs$^+$ and Rb$^+$ atoms were treated with the def2-TZVP basis set (see Figure 1 for designation of species). $^b$Nonexistent: formation of the zwitterionic INT-$\alpha$ is unstable, which decomposes barrierless in a cyclohexanone-like species and H$_2$. $^c$Nonexistent: barrierless process toward an INT.

Figure 1. Structures and key distances (in Å) of stationary points of the Lewis acid-catalyzed epoxide ring-opening reactions between MeOH and 1-Li$^+$ computed at B3LYP-D3(BJ)/6-31G+(d).

also results in a loss of a favorable orbital overlap between the empty ns atomic orbital (AO) of the group 1 cation and the HOMO of the epoxide, but it will gain a stabilizing orbital overlap between the ns AO of the group 1 cation and the HOMO−1 of the epoxide (see Figure S2). The key orbital overlap increases from 0.06 for Cs$^+$ to 0.33 for H$^+$ due to the more compact nature of the empty ns AO of Y$^+$ when ascending in group 1. In addition, the orbital energy gap also becomes smaller, going from Cs$^+$ to H$^+$, because of the consistently more stable empty ns AO of Y$^+$, which further contributes to a more stabilizing orbital interaction in the case of lighter Lewis acids.

Table 2 and Figure 1 summarize the computed reaction profiles and structural data of the Lewis acid-catalyzed epoxide ring-opening reactions between MeZH ($Z = O, S,$ and NH) and 1-$Y^+$, forming the $\alpha$- and $\beta$-intermediate. Three distinct trends can be observed. In the first place, the nucleophile attack can occur at both the $\alpha$- and $\beta$-carbon of the epoxide ring, of which, for all computed systems, the $\alpha$-attack occurs with a 3−7 kcal mol$^{-1}$ lower reaction barrier than the $\beta$-attack. Second, the reaction barrier systematically decreases when $Y^+$ ascends group 1 (Cs$^+$ > Rb$^+$ > K$^+$ > Na$^+$ > Li$^+$ > H$^+$). This is in good agreement with the experimentally determined reactivity trends for Lewis acid-catalyzed epoxide ring-opening reactions, which showed that the H$^+$-catalyzed epoxide ring-opening reaction can already be performed at room temperature within 30 min, while the Li$^+$- and Na$^+$-catalyzed analogs require an elevated temperature and a significantly longer reaction time. The reactions catalyzed by K$^+$, Rb$^+$, and Cs$^+$, on the other hand, are experimentally not used due to their poor catalytic ability. For the more nucleophilic MeSH and MeNH$_2$, the reaction barrier decreases along the series of $Y^+$ to such an extent that the epoxide ring-opening reaction catalyzed by H$^+$ is barrierless. Third, when changing the nucleophile from MeOH to MeSH and to MeNH$_2$, the reaction barriers of all epoxide ring-opening reactions become lower due to the enhanced nucleophilicity whereby the nucleophile can engage in a stronger acid−base-like interaction with the substrate 1-$Y^+$.
Next, we turn to the activation strain model (ASM)\(^8\) of reactivity to gain quantitative insights into the physical factors controlling the regiochemical preference for the epoxide ring-opening at the \(\alpha\)-position. In Figure 2a, we focus on the Li\(^+\)-catalyzed epoxide ring-opening reaction. Note that the nucleophile and 1-Li\(^+\) experience less steric repulsion than for the attack at the \(\beta\)-position. The attractive electrostatic and orbital interactions are, on the other hand, slightly more stabilizing for the \(\beta\)-attack, showing a trend opposite of the trend in interaction energy, and hence not responsible for the observed regioselectivity.

The origin of the less destabilizing Pauli repulsion for the attack at the \(\alpha\)-position compared to the \(\beta\)-position was further investigated by performing a Kohn–Sham molecular orbital analysis.\(^9\) The occupied orbitals of MeOH and 1-Li\(^+\), for both the attack at the \(\alpha\)- and \(\beta\)-position, were quantified at transition state-like, consistent geometries with a \(C^{\alpha}_{O}⁻\) bond stretch of 0.54 \(\text{Å}\) (Figure 3). The most important occupied molecular orbitals (MOs) that dictate the trend in Pauli repulsion, that is, the occupied orbitals responsible for the differences in steric repulsion between the reactant along the different regiochemical reaction pathways, are the HOMO\(_\text{MeOH}\) and HOMO\(_{\text{Li}^+}\). The HOMO\(_{\text{Li}^+}\) is an oxygen lone pair orbital and the HOMO\(_{\text{MeOH}}\) is an all in-phase \(\sigma\)-orbital, whereas the HOMO\(_{\text{Li}^+}\) is a filled \(\sigma\)-orbital delocalized over the cyclohexane ring of 1-Li\(^+\) (Figure 3b). As shown in Figure 3a, the occupied–occupied orbital overlap between these orbitals is nonexistent for the \(\alpha\)-attack (\(S = 0.00\) and \(S = 0.00\)), while it is present, and hence destabilizing, for the \(\beta\)-attack (\(S = 0.03\) and \(S = 0.02\)). The asymmetry in the \(C^{\alpha}_{O}⁻\) bond lengths of 1-Li\(^+\) induces an asymmetry in the HOMO\(_{\text{MeOH}}\) (Figure 3b), namely, there is more orbital density located on the carbon atom participating in the shorter epoxide \(C^{\beta}_{O}⁻\) bond (\(\beta\)-carbon) than on the carbon atom involved in the longer epoxide \(C^{\alpha}_{O}⁻\) bond (\(\alpha\)-carbon). Thus, when MeOH attacks the \(\beta\)-position, it encounters and hence overlaps with the large occupied orbital amplitude of the HOMO\(_{\text{MeOH}}\) located on the \(\beta\)-carbon, manifesting in more destabilizing Pauli repulsion and a higher reaction barrier compared to the \(\alpha\)-attack.

After having established that the nucleophile preferentially opens the epoxide at the \(\alpha\)-position, we analyze the reactivity trends of this reaction pathway for all six Lewis acids (\(Y^+ = \text{Cs}^+, \text{Rb}^+, \text{K}^+, \text{Na}^+, \text{Li}^+, \text{and H}^+\)) and for the uncatalyzed reaction (\(Y^+ = \text{none}\)). Figure 4a shows the activation strain model (ASM) results from the reactant complexes to the transition states for the uncatalyzed and \(\text{Cs}^+\)- and \(\text{H}^+\)-catalyzed epoxide ring-opening reactions (see Table S3 and Figure S4 in the Supporting Information for all ASM results). The uncatalyzed reaction (black) goes with the highest reaction barrier, and coordinating a \(\text{Cs}^+\) cation (orange) lowers the barrier, which then consistently decreases upon ascending in...
MeOH of 0.59 Å. To assess the potential in flH⁺ conditions, we showed that the results of the EDA are also epoxide ring-opening reactions under basic and acidic trend in interaction energy. Note, however, that these EDA behind this catalysis, only have a minor contribution to the interactions, which are commonly seen as the driving force destabilizing) than the black curve (Figure 4b). The orbital reaction coordinate, signi fi cantly lower in energy (Figure 5a). In line with our previous work,7 we found that by coordinating a Lewis acid to 1-Y⁺ distances (Figure 5; vide supra). The FMO₃⁻ and HOMO or HOMO−1⁻ that are decisive for the trend in Pauli repulsion, namely, with the HOMOMeOH and HOMO−6MeOH. The respective orbital overlap and hence repulsion are the largest and most destabilizing for the uncatalyzed reaction (S = 0.07 and 0.09) and the smallest and least destabilizing for the H⁺-catalyzed reaction (S = 0.01 and 0.02) (Figure 5a). In line with our previous work, we found that by coordinating a Lewis acid to 1, the σ-orbital of 1 becomes polarized away from the incoming MeOH due to both the positive potential of the cationic Lewis acid and the

**Figure 3.** (a) Molecular orbital diagram of the most important occupied—occupied orbital overlap for the Lewis acid-catalyzed ring-opening reactions of the α- and β-attack for MeOH + 1-Li⁺. (b) Key occupied orbital (isovalue = 0.03 Bohr−1/2) computed at consistent TS-like geometries with a C¹−O bond stretch of 0.54 Å, in which 1-Li⁺ is depicted from the bottom side. Computed at ZORA-M06-2X/TZ2P//B3LYP-D3(BJ)/6-31G+(d).
catalyzed epoxide ring-opening reaction decreases upon ascending in group 1 along the series Cs⁺ > Rb⁺ > K⁺ > Na⁺ > Li⁺ > H⁺. Furthermore, we found that the reaction pathway following a chair-like transition state goes, for all studied systems, with a significantly lower reaction barrier than the twist boat-like analog.

Our activation strain and Kohn–Sham molecular orbital analyses showed that, in contrast to the generally accepted belief, the catalytic ability of the Lewis acids is caused by the reduction of steric (Pauli) repulsion between the Lewis acid-activated epoxide 1-Y⁻ and the nucleophile and not by the enhanced stabilizing orbital interactions as is currently the prevailing mechanism in the literature. The reduction in steric (Pauli) repulsion can be traced back to the Lewis acid-induced polarization of the occupied orbitals on 1-Y⁻ away from the incoming nucleophile. This effect gets more pronounced ascending in group 1 because the positive potential of the lighter cations polarizes due to the shorter O···Y⁻ bond, the occupied orbitals on 1-Y⁻ to a greater extent and their lower-energy valence ns AO sets engage in a stronger donor–acceptor interaction with the occupied orbitals of epoxide 1-Y⁻. This again demonstrates that Pauli-lowering catalysis is a general phenomenon, which is not only limited to Michael addition reactions and cycloadditions.

In addition, we discovered a novel physical mechanism that is responsible for the regioselectivity of the cyclohexene oxides ring-opening that acts aside from the Fürst-Plattner rule, which proposes that the regioselectivity is solely determined by the strain of the associated reaction path. Instead, we found that regioselectivity for nucleophilic attack at the Cβ of the epoxide 1-Y⁻ is, to a substantial degree, controlled by the steric interaction with the incoming nucleophile attacking the β-Carbon and hence a significant steric interaction with the incoming nucleophile attacking the β-Carbon. These findings will equip experimentalists with the mechanistic insight to understand and rationalize the trends in reactivity as well as regioselectivity of both uncatalyzed and Lewis acid-catalyzed epoxide ring-opening reactions.

**METHODS**

**Computational Details.** Computations were performed using Gaussian 09 Rev. D.01.14 using the hybrid functional B3LYP-D3(BJ)15 with 6-31+G(d)16 as a basis set for geometry optimization. More accurate electronic energies were obtained by a single-point calculation with M06-2X17 with 6-311+G(dp)18 for all atoms, except for Cs⁺ and Rb⁺, which were treated with def2-TZVP.16 This approach proved to give an excellent trade-off between accuracy and computational time.19 The geometry convergence criteria were set to tight (max. force = 1.5×10⁻⁵, max. displacement = 6.0×10⁻⁵), and an internally defined superine grid size was used (Int = veryfinegrid), which is a pruned 175,974 grid for first-row atoms and a 250,974 grid for all other atoms. These parameters were chosen as a recent paper indicated a significant dependence of the computed frequencies on the molecule orientation when a smaller grid size is used.20 Geometries were optimized without symmetry constraints. The quasi-harmonic correction21 was applied to all frequencies by modifying all vibration below 100 cm⁻¹ to 100 cm⁻¹. All computed stationary points have been confirmed by performing a vibrational analysis calculation. The energy minima had no imaginary frequencies, while the transition states had only one imaginary frequency has been inspected to ensure that it corresponds to the reaction of interest. The potential energy

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**Figure 4.** (a) Activation strain model, where ΔE = solid lines, ΔEstrain = dashed lines, and ΔEΔσ = dotted lines; (b) Energy decomposition analysis, where ΔVrel = dotted lines, ΔEΔσ = solid lines, and ΔErel = dashed lines, for the uncatalyzed and Lewis acid-catalyzed ring-opening reactions of MeOH +1-Y⁻ via the σ-attack (Y⁻ = none, black; Cs⁺, orange; and H⁺, red), where the energy values are plotted from the reactant complex to the transition state and projected onto the Cα···O bond stretch. The transition states are indicated by a dot. Computed at ZORA-M06-2X/TZ2P//B3LYP-D5(Bj)/6-31G+(d), whereas the Cs⁺ atom is treated with the def2-TZVP basis set.

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**CONCLUSIONS**

Our computational study reveals that Lewis acids (Y⁻) efficiently catalyze the nucleophilic ring-opening reactions of cyclohexene oxides. The reaction barrier of the Lewis acid-catalyzed epoxide ring-opening reaction decreases upon ascending in group 1 along the series Cs⁺ > Rb⁺ > K⁺ > Na⁺ > Li⁺ > H⁺. Furthermore, we found that the reaction pathway following a chair-like transition state goes, for all studied systems, with a significantly lower reaction barrier than the twist boat-like analog.

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Figure 5. (a) Molecular orbital diagram of the most important occupied–occupied orbital overlap for the uncatalyzed and Lewis acid-catalyzed ring-opening reactions of MeOH + I-Y (Y' = none, Cs', Rb', K', Na', Li', and H'). (b) Key occupied orbital (isovalue = 0.03 Bohr⁻¹) computed at consistent TS-like geometries with a C°−O stretch of 0.59 Å and MeOH−C° bond length of 1.98 Å. The key filled orbital lobe of 1-Y', which has an overlap with the filled orbitals of the incoming MeOH, is highlighted with a blue dashed outline. Computed at ZORA-M06-2X/TZ2P//B3LYP-D3(BJ)/6-31G+(d), whereas Cs', Rb', and H' atoms were treated with the def2-TZVP basis set.

**Activation Strain and Energy Decomposition Analysis.** The activation strain model (ASM)³ of reactivity, also known as the distortion/interaction model, is a fragment-based approach based on the idea that the energy of an interaction system, \( \Delta E \), can be described and understood by looking at the unperturbed charge distributions of the (deformed) reactants and is usually attractive. The Pauli repulsion, \( \Delta\text{Pauli}(\zeta) \), describes the destabilizing interaction between fully occupied closed-shell orbitals of both fragments due to Pauli’s principle. The orbital interaction energy, \( \Delta\text{Int}(\zeta) \), accounts for polarization and charge transfer between the fragments, such as HOMO−LUMO interactions.

In the activation strain and accompanied energy decomposition diagrams presented in this work, the various energy terms are projected onto the carbon−oxygen (C°−O) distance. This critical reaction coordinate undergoes a well-defined change during the reaction from the reactant complex via the transition state to the product and is shown to be a valid reaction coordinate for studying nucleophilic substitution reactions.¹⁰

**Thermochemistry.** Bond enthalpies, i.e., bond dissociation energies (BDE), are calculated at 298.15 K and 1 atm (\( \Delta H_{\text{BDE}} \)) from electronic bond energies (\( \Delta E \)) and vibrational frequencies using standard thermochemistry relations for an ideal gas [eq 3]²⁹:

\[
\Delta H_{\text{BDE}} = \Delta E + \Delta E_{\text{trans,298}} + \Delta E_{\text{rot,298}} + \Delta E_{\text{vib,0}} + \Delta(\Delta E_{\text{vib,0}})_{298}
\]

(3)

Here, \( \Delta E_{\text{trans,298}} \), \( \Delta E_{\text{rot,298}} \), and \( \Delta E_{\text{vib,0}} \) are the differences between the epoxide and the ring-opened diradical that results from breaking either the C°−O or C°−O bond in translational, rotational, and zero-point vibrational energy, respectively. The last term, \( \Delta(\Delta E_{\text{vib,0}})_{298} \) is the change in the vibrational energy difference when going from 0 K to 298.15 K.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.joc.0c02955.

Additional computational results and Cartesian coordinates, energies, and number of imaginary frequencies of all stationary points (PDF)
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Notes
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