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Systematic investigation of the role of the epoxides as substrates for CO₂ capture in the cycloaddition reaction catalysed by ascorbic acid†

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This work establishes a comprehensive theoretical framework for synthesizing cyclic organic carbonates, crucial for the polymer industry, through the organocatalytic cycloaddition of carbon dioxide (CO₂) to epoxides under mild pressure and temperature conditions. Using advanced computational techniques, the study examines the thermodynamic and kinetic aspects of the reaction, with a particular focus on epoxide substrates featuring diverse substituents. Detailed analysis reveals activation energy barriers and identifies the rate-determining step (rds), offering crucial insights into the molecular processes governing the reaction. An automated data-driven workflow revealed that the buried volume of the epoxide O atoms was among the most influential molecular features affecting reaction barriers. Overall, the findings align with experimental data, offering insights into substrate design for optimized CO₂ utilization. This work calls for a systematic exploration of ascorbic acid-based catalyst modifications to optimize energy barriers and improve overall reaction performance, paving the way for rational catalyst design and predictive catalysis in CO₂ valorization. The computational study is not limited to basic research or ascorbic acid but is applicable to most catalysts capable of carrying out this reaction in the polymer industry.

Keywords: Epoxide; CO₂ activation; Sustainable catalysis; Data-driven workflows; DFT calculations; Predictive catalysis; Cycloaddition.

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1 Introduction

The valorisation of CO₂ as a renewable source of carbon is fundamental according to the raised atmospheric CO₂ levels. While carbon capture and storage offer mitigation potential, their deployment is hindered by high energy demands,¹ economic constraints, and reliance on energy-intensive reagents.^{2,3}

Nevertheless, CO₂ capture and utilization remains a privileged approach for further development.^{4–8} In CO₂ capture,

various techniques, including chemical and physical absorption^{9,10} by metal organic frameworks^{11,12} and membrane separation¹³ are being explored. Ionic liquids (ILs) show promise as green solvents for CO₂ absorption,^{14–17} yet current materials have limited efficiency and pose environmental concerns like high regeneration costs and not enough stability.^{9,10,18} Different approaches have been utilized to enhance the characteristics of ionic liquids for carbon capture,¹⁹ utilization, and storage (CCUS).²⁰ These include the development of ionic liquids with favourable properties like amino acid ILs,^{21–23} amine-functionalized ILs,^{24–26} phosphonium-based ILs,^{22,27} and protic ILs.^{28–30} Certain types of ionic liquids are reusable, offering a potential solution to mitigate environmental concerns linked to their usage.^{31–35} On the other hand, CO₂ capture alone is limited by the need for suitable storage.³⁶ Therefore, researchers are actively seeking methodologies and materials with improved properties to address the challenge of CO₂ capture and conversion.^{37–41} In this context, the main nonreductive approach to CO₂ conversion is represented by the cycloaddition of CO₂ to highly available epoxide substrates that can be produced synthetically or even have biobased origin.^{42–44} The functionalization of CO₂ by

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epoxides to afford cyclic carbonates leads to a wide selection of applications, as aprotic polar solvents⁴⁵ that can be found in electrolytes in lithium ion batteries,⁴⁶ polymer precursors^{47,48} and fuel additives. Furthermore, they are vital organic intermediates or precursors in the production of pharmaceutical, agricultural, and fine chemical products.⁴⁹ Moreover, the cycloaddition of CO₂ to epoxides can be carried out under mild conditions by using extremely simple biobased compounds⁵⁰ mainly as hydrogen bond donors such as amino acids,⁵¹ polysaccharides,⁵² lignin,^{53,54} and even ascorbic acid³⁸ when used in the presence of a nucleophilic compound (see Scheme 1 for mechanistic pathway using ascorbic acid and tetrabutylammonium iodide, TBAI, as the catalytic system).⁵⁵ Ascorbic acid, in particular, serves for the cycloaddition of CO₂ to epoxides under ambient conditions and it has been recently developed into a single component system for application as a recoverable ionic catalyst in biphasic media.⁵⁶ Recent works have recognized ascorbic acid as a rare example of viable biobased catalysts for CO₂-epoxide cycloaddition⁵⁷ and it is often used as the hydrogen bond donor of choice for the preparation of challenging multicarbonate monomers as precursors of sustainable polyhydroxyurethanes.^{58,59}

In parallel to the development of new catalytic systems and concepts, researchers have been looking into gaining a deeper understanding in the mechanistic features of the cycloaddition reaction through mechanistic^{60,61} and/or computational studies.^{62,63} Generally, in such computational studies, just a few epoxide substrates are used, most typically, ethylene and

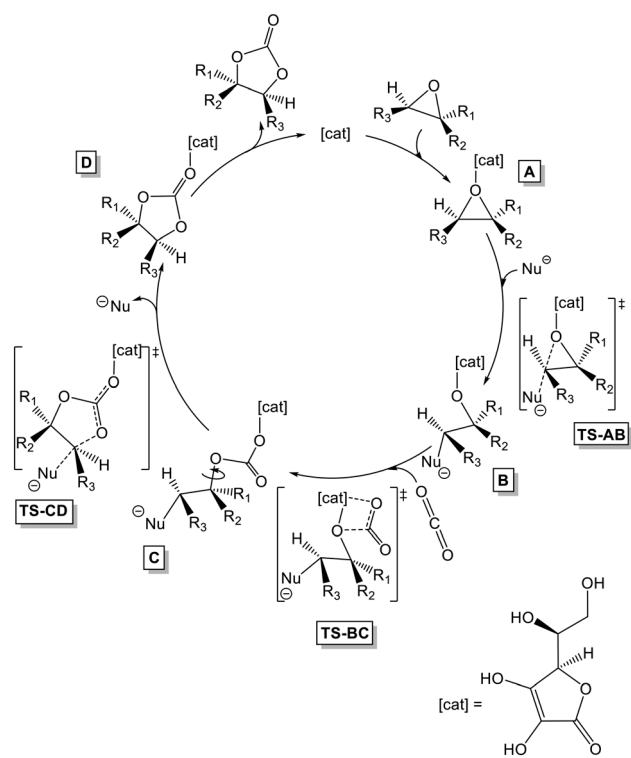
propylene oxide for the sake of simplicity, *i.e.*, some of the most reactive epoxides, together with epichlorohydrin and glycidyl ethers, *a priori*, due to limited steric hindrance. However, such an approach does not render the whole picture of the extremely different reactivity of the diverse range of available epoxides that ultimately depends on their substitution pattern and which, in turn, leads to very different conversion values from substrate to substrate.⁶⁴ Studies involving a systematic investigation of the reactivity of multiple (six) substrates are extremely rare,⁶⁵ and yet limited to the case of metal-based catalytic systems, without including other crucially important epoxides for various applications such as those bearing halogen substituents⁶⁶ and glycidyl ethers.⁶⁷ Here, by density functional theory (DFT) calculations, we carry out a systematic study involving a large library of epoxides to investigate the effect of the substrate structure and electronic features on the efficiency of the catalytic process by using an organocatalytic systems (ascorbic acid/TBAI). This work explores as well the role of the counter-cation of the nucleophilic halide, which is generally overlooked in literature-reported calculations. Additionally, we have used a data-driven workflow including automated featurization and machine learning (ML) modelling to analyse the most important molecular parameters influencing reaction barriers. Through this comprehensive approach, the project aims to provide an efficient, sustainable methodology with practical applications in cyclic carbonate synthesis, and the results are extrapolable to any of the broad range of studies that lead to those compounds from CO₂, from ionic liquids^{68,69} to the complexity and encapsulation of metallic organic frameworks (MOFs).⁷⁰

2 Results and discussion

As commonly reported in the literature,⁷¹ our investigation uses ascorbic acid as the hydrogen bond donor and primarily an iodide ion as the nucleophile, initially neglecting the role of the counter-cation. Specifically, a quaternary ammonium salt was omitted from the reaction mechanism due to its negligible coordinating ability. This study focuses on various epoxide substrates included in Table 1 interacting with CO₂.

Table 2 provides a comprehensive breakdown of the relative energies associated with the pathway of the cycloaddition reaction, as illustrated in Scheme 1. Table 2 provides a comprehensive overview of the relative energies associated with the different steps of the reaction of all the epoxide substrates (see Table 1), offering insights into the thermodynamics and kinetics of the molecular processes involved, referenced from the catalyst, *i.e.*, ascorbic acid (all the xyz coordinates can be found in the ESI†).

For the **EPO-H** (unsubstituted epoxide; ethylene oxide) the binding of ascorbic acid to the epoxide (**A**), despite the presence of two hydrogen bonds between the acid and the epoxide oxygen, has a cost of 1.7 kcal mol⁻¹, attributed to the entropic component. Subsequently, the nucleophilic agent, represented exclusively by the iodide ion in this case, initiates the reaction by attacking the less substituted position of the epoxide, resulting in its ring-opening. The corresponding



Scheme 1 Cycloaddition of epoxides (R = substituent; cat = ascorbic acid; Nu = nucleophile, being F, Cl, Br, I, or TBAI) and CO₂.



Table 1 Substituents of each epoxide system, as presented in Scheme 1

System	R ₁	R ₂	R ₃
EPO-H	-H	-H	-H
EPO-CH ₃	-H	-CH ₃	-H
EPO-(CH ₃) ₂	-CH ₃	-CH ₃	-H
EPO- <i>cis</i> (CH ₃) ₂	-H	-CH ₃	-CH ₃
EPO- <i>trans</i> (CH ₃) ₂	-CH ₃	-H	-CH ₃
EPO- <i>t</i> Bu	-H	-C(CH ₃) ₃	-H
EPO-CH ₂ - <i>n</i> Bu	-H	-(CH ₂) ₄ -CH ₃	-H
EPO-OH	-H	-OH	-H
EPO-CH ₂ -OH	-H	-CH ₂ -OH	-H
EPO-Glyc	-H	-CH ₂ -O-C(=O)-(=CH ₂)CH ₃	-H
EPO-Ph	-H	-Ph	-H
EPO-(Ph) ₂	-Ph	-Ph	-H
EPO- <i>cis</i> (Ph) ₂	-H	-Ph	-Ph
EPO- <i>trans</i> (Ph) ₂	-Ph	-H	-Ph
EPO-CH ₂ Ph	-H	-CH ₂ -Ph	-H
EPO-CF ₃	-H	-CF ₃	-H
EPO-C(CF ₃) ₃	-H	-C(CF ₃) ₃	-H
EPO-CH ₂ F	-H	-CH ₂ F	-H
EPO-CH ₂ Cl	-H	-CH ₂ Cl	-H
EPO-CH ₂ Br	-H	-CH ₂ Br	-H
EPO-Cyc	-H	-(CH ₂) ₄ -	-H

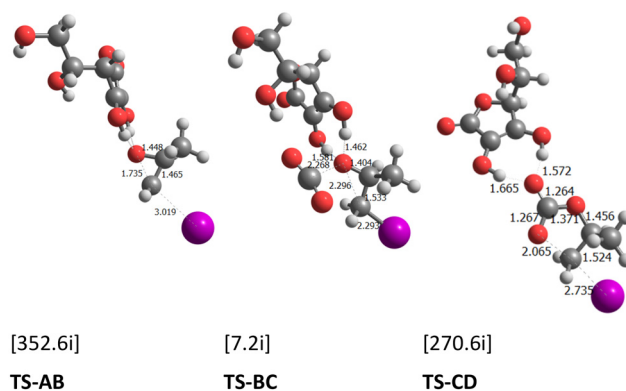
Table 2 Relative Gibbs energies (in kcal mol⁻¹) obtained at various stages of the reaction of different epoxide substrates with CO₂, catalysed by ascorbic acid and an iodide ion

Substrate	A	TS-AB	B	TS-BC	C	TS-CD	D	E
EPO-H	1.7	17.9	8.0	—	5.7	21.2	-1.5	-0.1
EPO-CH ₃	1.3	18.3	10.7	19.8	8.6	23.1	-0.8	0.0
EPO-(CH ₃) ₂	1.2	18.9	13.2	23.3	14.2	27.5	-0.2	-0.2
EPO- <i>cis</i> (CH ₃) ₂	0.9	20.8	14.5	23.4	12.6	27.8	1.0	-1.3
EPO- <i>trans</i> (CH ₃) ₂	1.3	21.9	16.3	21.5	13.5	27.3	0.6	-0.9
EPO- <i>t</i> Bu	0.2	19.8	14.0	—	11.9	25.7	-0.1	-2.0
EPO-CH ₂ - <i>n</i> Bu	1.7	19.5	8.6	22.4	9.4	25.5	0.9	-6.1
EPO-OH	1.5	17.9	7.8	—	7.5	23.0	-0.1	5.9
EPO-CH ₂ -OH	0.9	18.0	11.6	—	10.0	24.3	5.5	-0.7
EPO-Glyc	-0.5	—	9.9	—	11.5	21.4	0.0	-0.9
EPO-Ph	2.3	20.8	12.7	—	9.4	24.6	2.3	1.3
EPO-(Ph) ₂	3.8	20.6	16.6	—	22.6	31.9	4.9	3.5
EPO- <i>cis</i> (Ph) ₂	5.0	26.0	17.0	19.2	18.2	29.3	5.6	5.4
EPO- <i>trans</i> (Ph) ₂	3.3	21.2	16.8	—	17.2	24.2	6.7	3.7
EPO-CH ₂ Ph	1.8	18.7	14.2	14.6	11.3	26.6	3.5	1.4
EPO-CF ₃	3.1	17.2	8.0	—	8.6	25.4	2.3	3.2
EPO-C(CF ₃) ₃	2.2	21.8	12.2	—	15.6	32.7	4.2	3.2
EPO-CH ₂ F	2.1	18.2	10.5	—	9.5	22.7	-0.5	1.0
EPO-CH ₂ Cl	2.1	18.1	11.7	—	8.3	23.4	0.2	1.4
EPO-CH ₂ Br	2.0	18.2	12.1	—	10.7	24.0	0.4	1.9
EPO-Cyc	0.9	23.0	20.0	—	10.6	29.1	1.2	-2.4

transition state corresponds to an energy barrier of 16.2 kcal mol⁻¹, but speaking of the kinetic cost, it should be referenced to the lowest energy intermediate, which in this case would correspond to the substrate and catalyst separately, *i.e.*, the point of 0.0 kcal mol⁻¹. Thus, following the scheme described by Kozuch and Shaik,^{72,73} the kinetic cost of **TS-AB** is 17.9 kcal mol⁻¹. This transition state leads to an intermediate **B**, relatively unstable, specifically 6.3 kcal mol⁻¹ relative to intermediate **A**. The following step in the reaction mechanism involves the insertion of CO₂ from intermediate **B**. It is noteworthy that for the **EPO-H**, the

corresponding transition state could not be identified, although the subsequent intermediate **C** was successfully located. This insertion step is exergonic, with an energy release of 2.3 kcal mol⁻¹. The inability to locate the CO₂ insertion transition state is not uncommon, as it typically features very low negative frequencies that are often challenging to identify and are therefore frequently omitted in reaction mechanism descriptions. From there, the **TS-CD** for the closing of the cyclic carbonate, with the expansion of the initial epoxide ring from 3 to 5 members, guides us in an exergonic step of 7.2 kcal mol⁻¹, that is still also exergonic relative to the starting materials in agreement with the thermodynamically favourable nature of the cycloaddition process.⁷⁴ It involves a kinetic cost of 21.2 kcal mol⁻¹, and therefore, it is the rate-determining state (rds) of the reaction, although relative to intermediate **C**, it would only have an energy barrier of 15.5 kcal mol⁻¹, therefore, lower than the corresponding barrier for **TS-AB**. Finally, additionally, intermediate **E** is simply obtained from **D** eliminating the iodide nucleophilic ion that is no longer bonded to any carbon.

As for the methylated epoxide (**EPO-CH₃**), the binding of the co-catalyst (**TS-AB**) presents a similar high barrier, but slightly higher, with a value of 18.3 kcal mol⁻¹. This small increase of 0.4 kcal mol⁻¹ may already suggest that steric hindrance could be the reason. Then for this substrate, **TS-BC** was located, thus ruling out the insertion of CO₂ as a rate-limiting step since it is 3.3 kcal mol⁻¹ lower than **TS-CD**, which is 1.9 kcal mol⁻¹ higher than for **EPO-H**. Since also **TS-BC** was located for **EPO-CH₃**, the three corresponding transition states for this substrate are displayed in Fig. 1 to investigate their nature (and in Fig. S1† including the intermediates, as well). It should be highlighted that ascorbic acid primarily stabilizes the epoxide through two hydrogen bonds *via* its enediol hydroxyls to the epoxide oxygen. However, one aspect to note is that once the iodide attack cleaves the C-O bond of the epoxide in **TS-AB**, it entails the transfer of a hydrogen from ascorbic acid, thus protonating the oxygen of the

**Fig. 1** Transition states of the cycloaddition of carbon dioxide (CO₂) to epoxide **EPO-CH₃** (selected distances in Å and imaginary frequencies included within brackets in cm⁻¹).

epoxide momentarily in intermediate **B**. It is interesting to note that the possible protonation of the alkoxide intermediate by hydrogen bond donors, which is further validated by the difference in pK_a of the species, was considered in previous studies. The protonation of the alkoxide was suggested as a possible factor affecting the catalytic activity of strongly acidic hydrogen bond donors such as carboxylic acids.⁷⁵ It is interesting to note that the possible protonation of the alkoxide intermediate by the hydrogen bond donor, despite being seldom discussed in the literature, is expected due to the difference in pK_a of the species.⁶³ On the other hand, it should be noted that already in the next transition state, **TS-BC**, the proton automatically returns to ascorbic acid. In fact, Fig. S1† also includes an adduct of intermediate **B** with a CO_2 molecule, called **B** + CO_2 , where ascorbic acid regains the proton. This species also serves to show that the CO_2 entry barrier is mostly minimal, and in fact for **EPO-CH₃**, it is only 0.2 kcal mol⁻¹ in gas-phase optimization, *i.e.*, only by geometry, but once the solvent effect is incorporated, it becomes negative by 1.1 kcal mol⁻¹. This also explains why the transition state **TS-BC** was only observed for a few epoxide substrates. As previously commented, the imaginary frequency for this transition state is very small just 7.2i cm⁻¹. In intermediate **C**, it is notable that ascorbic acid continues to form two hydrogen bonds through two of its hydroxyl groups. However, these interactions now involve both oxygen atoms of the bonded CO_2 molecule, rather than being limited to a single oxygen atom. In the transition state corresponding to the cyclization of the cyclic carbonate (**TS-CD**), ascorbic acid once again establishes two hydrogen bonds, but this time exclusively with a single oxygen atom of the forming carbonate (see Fig. 1 and S2†).

The explored substituents on the epoxide, which increase the steric hindrance, include $-(\text{CH}_3)_2$, $-t\text{Bu}$ and $-\text{Ph}_2$. Specifically, for **EPO-CH₃**, the reaction presents larger energy barriers. The nucleophilic attack of the iodide anion (**TS-AB**) presents a kinetic cost of 18.9 kcal mol⁻¹, the insertion of CO_2 (**TS-BC**) of 23.3 kcal mol⁻¹, but the most significant increase is found in **TS-CD**, with a value of 27.5 kcal mol⁻¹. The rigidity of the epoxy backbone is considered to be somewhat significant, as the rds barrier with **EPO-Cyc** increases to 29.1 kcal mol⁻¹, at least 1.3 kcal mol⁻¹ higher than the analogous system, **EPO-*cis*(CH₃)₂**, for comparison.

Regarding **EPO-*t*Bu**, this **TS-CD** is slightly more disfavoured compared to **EPO-CH₃**, up to 2 kcal mol⁻¹. Therefore, the effect of disubstitution at the same carbon atom seems less detrimental than increasing the bulkiness of a single substituent. However, this finding cannot be generalized, as the **EPO-Ph** and **EPO-Ph₂** systems exhibit energy barriers of 24.6 and 31.9 kcal mol⁻¹, respectively. Therefore, disubstitution emerges as a critical factor in some cases.

EPO-CH₂Cl shows an energy barrier comparable to **EPO-CH₃**, while **EPO-CH₂Br** has a slightly larger barrier than **EPO-CH₂Cl** — less than 1 kcal mol⁻¹. **EPO-CH₂F** with a relatively smaller and highly electronegative fluorine atom has a even

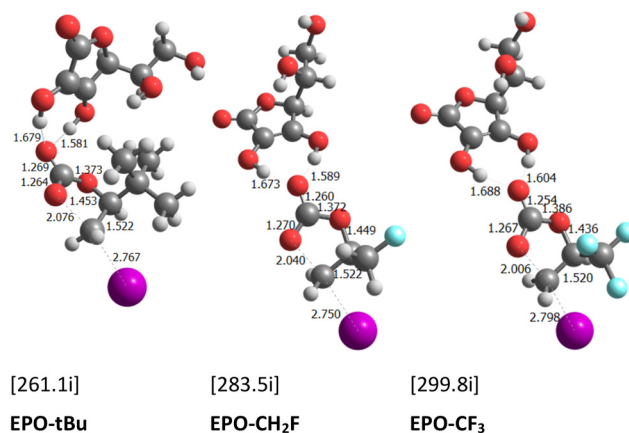


Fig. 2 Transition state **TS-CD** of the cycloaddition of CO_2 for **EPO-*t*Bu**, **EPO-CH₂F** and **EPO-CF₃** (selected distances in Å and imaginary frequencies included within brackets in cm⁻¹).

lower upper reaction barrier than **EPO-CH₃** confirming the high reactivity of this substrate.⁵⁶ This effect of the halide substituent becomes even clearer when comparing the values of **TS-CD** of 22.7, 23.4, and 24.0 kcal mol⁻¹ for **EPO-CH₂F**, **EPO-CH₂Cl**, and **EPO-CH₂Br**, respectively. In fact, comparing the experimental reactivity of the last two substrates, both clearly emerge at a disadvantage compared to other more structurally simple substrates, such as **EPO-H** or **EPO-CH₃**. Therefore, the DFT computed energy barriers confirm the past experimental results.^{76,77}

In Fig. 2, the **TS-CD** structure for some of the other studied epoxides is displayed (all included in Fig. S2†). For the formation distance of the $\text{C}\cdots\text{O}$ bond and the breaking $\text{C}\cdots\text{I}$ bond, it can be observed how bulky groups tend to modify the corresponding energy, but no substantial difference is observed. Therefore, in the following sections, an additional effort must be made in the structural study, as well as to provide insights into the electronic factors influencing the rds.

Since the transition state for the closure of the cyclic carbonate ring (**TS-CD**) is found as the most energy-demanding step of the reaction we plotted in Fig. 3 how the kinetic cost ranges between the different epoxide candidates.

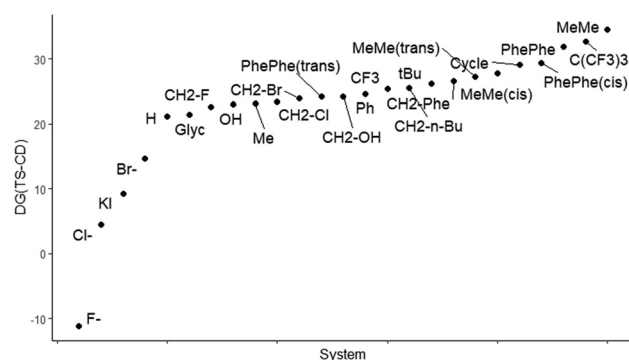


Fig. 3 Plot of all the studied epoxides with respect to the relative energy of **TS-CD** with respect to reactant complex **A** (Gibbs energies in kcal mol⁻¹).



Table 3 Selected distances for the free epoxide and for **TS-CD** (in Å)

Substrate	Distances				% V_{Bur}	
	$d(\text{C-O})$ epoxide	$d(\text{C-C})$ epoxide	$d(\text{C}\cdots\text{O})$ TS-CD	$d(\text{C}\cdots\text{X})$ TS-CD	Epoxide	TS-CD
EPO-H	1.430	1.478	2.060	2.721	32.1	40.6
EPO-CH₃	1.434	1.480	2.065	2.735	36.9	46.0
EPO-(CH₃)₂	1.446	1.483	2.090	2.749	41.9	50.9
EPO-cis(CH₃)₂	1.440	1.485	2.098	3.002	46.5	55.1
EPO-trans(CH₃)₂	1.440	1.481	2.121	2.886	46.8	55.3
EPO-tBu	1.437	1.478	2.076	2.767	42.5	51.1
EPO-CH₂-nBu	1.434	1.479	1.996	2.832	38.7	49.1
EPO-OH	1.445	1.469	2.033	2.765	35.8	44.8
EPO-CH₂-OH	1.437	1.476	2.089	2.719	38.3	47.5
EPO-Glyc	1.441	1.477	1.951	2.839	41.7	46.5
EPO-Ph	1.430	1.488	2.039	2.750	39.9	49.2
EPO-(Ph)₂	1.447	1.490	2.042	2.807	48.1	57.2
EPO-cis(Ph)₂	1.433	1.502	2.133	2.956	58.1	66.3
EPO-trans(Ph)₂	1.435	1.502	2.046	3.042	59.0	67.7
EPO-CH₂Ph	1.436	1.477	2.010	2.828	40.7	50.8
EPO-CF₃	1.436	1.476	2.006	2.798	40.5	49.4
EPO-C(CF₃)₃	1.439	1.477	2.028	2.815	49.2	56.6
EPO-CH₂F	1.436	1.477	2.040	2.750	38.1	46.1
EPO-CH₂Cl	1.438	1.476	2.038	2.752	39.2	46.2
EPO-CH₂Br	1.437	1.476	2.035	2.753	39.6	46.3
EPO-Cyc	1.436	1.486	2.133	2.934	50.9	59.4

One key factor in explaining the high energy barriers observed in Fig. 3. for some of the substrates at this transition state is the bond distance between the carbon of the epoxide bearing the iodine and the oxygen of the inserted CO_2 . Based on the data in Table 3 and the Hammond postulate,⁷⁸ it is expected that the smaller the C-I distance is, the more reactant-like the **TS-CD** and the more favoured the ring closure would be, and consequently, the energy barrier at this step of the reaction would be smaller. At the same time, the C-C distance of the former epoxide should be considered. For the **TS-CD**, the distances of the forming C-O bond and the breaking C-I bond are inconclusive and do not generate a clear trend, although it is perceived that the shorter both distances are, the more feasible the closure of the cyclic carbonate is. Using the SambVca2.1 program,^{79,80} steric maps have been generated for the different substrates. Table 3 also includes the values of the overall % V_{Bur} (by quadrants in Tables S1 and S2†) of the free epoxide and **TS-CD**. It should be noted that quantitatively the hindrance increases in the transition state, but qualitatively by the steric maps collected in Fig. 4 the analysis leads to similar conclusions. Thus, globally, the % V_{Bur} increases noticeably with the presence of bulky groups on the carbon that facilitates the C-O bond formation. This effect is even more pronounced on the alpha carbon, especially when this position is doubly substituted.

Additionally, the asymmetry of the steric maps should be highlighted, as this can help determine whether the steric hindrance is confined to a specific quadrant. Fig. 4, with the steric maps, clarifies that although there may be a heavily hindered quadrant, the rest is accessible,^{81,82} especially in cases of monosubstitution rather than disubstitution of the carbon atom, as seen in **EPO-(Me)₂** or **EPO-(Ph)₂**.

Moreover, the nature of the nucleophilic ion has also been studied. As shown in Table 4, the rds remains the same for Br, Cl, and F, and the kinetics also detect **TS-CD** as the rds, with barriers changing for **EPO-CH₃** from 23.1 kcal mol⁻¹ for I^- to 16.2, 20.4, and 29.5 kcal mol⁻¹ with Br^- , Cl^- , and F^- , respectively (see Tables S3-S5† for further structural and electronic details). Thus, it is revealed that Cl^- and even more so Br^- would perform as better nucleophiles than I^- . However such an observation is at variance with what is generally observed experimentally in the cycloaddition reaction catalysed by hydrogen bond donors and halide salts where the iodide anion is the most efficient nucleophile,^{63,75,83} and calls for further clarification, probably in terms of solvation in the reaction mixture. Thus, there must be this or some other factor/s. Actually, smaller halides are supposed to be better nucleophiles in polar aprotic solvent, but epoxides do not seem to follow such a trend experimentally, and the inclusion of the counteraction TBA^+ might be crucial.

On the other hand, it should be mentioned that what changes the most in our calculations when varying the halide is the exergonicity of the process, and thus, the adduct of ascorbic acid and the epoxide, when the halide is at a non-coordination distance, opens a stability difference window of over 40 kcal mol⁻¹. It is important to clarify at this point that the origin of the kinetic barrier is not the same for the computational set. Due to the endothermicity revealed by the CO_2 cycloaddition process using I^- as nucleophile for all the substrates included in our setup, the kinetic cost of the process was calculated as the difference between the energy of the transition state **TS-CD** with respect to the reference (**A**). The barrier from the intermediate **C** to the **TS-CD** is slightly lower but rigorously speaking the larger energy difference must be taken as the kinetic cost of the process. Nevertheless, the use of F^- , Cl^- , and Br^- as nucleophiles



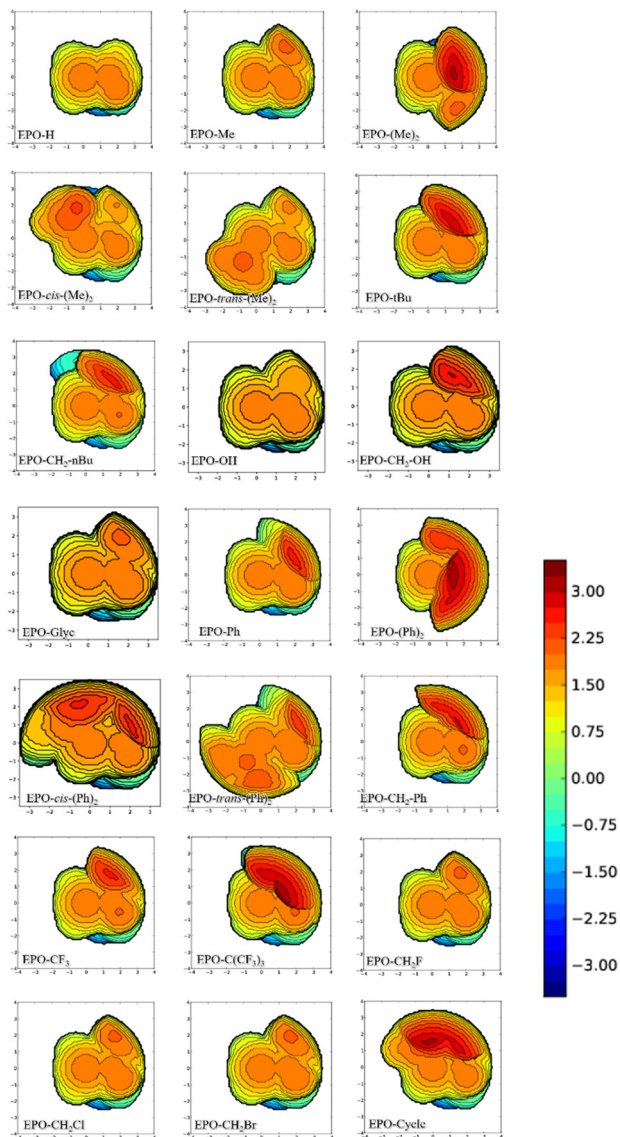


Fig. 4 % V_{Bur} and topographic steric maps in Å (XY plane) for TS-CD, using the carbon forming the C–O bond as the center, with a radius of 3.5 Å and with the Z axis defined by the C–O bond, the other carbon of the original epoxide in the XZ plane.

Table 4 Relative Gibbs energies (in kcal mol⁻¹) obtained at various stages of the reaction of EPO-CH₃ substrate with CO₂, catalysed by ascorbic acid and different ions or salts as nucleophiles

Nucleophile	A	A + X ⁻	TS-AB	B	TS-BC	C	TS-CD	D
I ⁻	0.0	1.3	18.3	10.7	19.8	8.6	23.1	-0.8
Br ⁻	0.0	8.6	14.0	-0.1	—	-1.5	14.7	-2.0
Cl ⁻	0.0	-15.9	-22.7	-12.8	—	-14.9	4.5	-8.7
F ⁻	0.0	-40.6	-8.3	-41.3	—	-39.7	-11.1	-12.7
KI	0.0	-11.7	9.0	-6.5	—	-5.7	9.2	-20.8
TBAI	0.0	8.2	—	5.9	—	9.7	—	-0.2

implies changes in the thermodynamics. Anions of more electronegative halides stabilize intermediates and transition states along the reaction mechanism and as reported in Table 4

the energy difference between A and TS-CD becomes much lower, even negative. In this scenario, the kinetic cost of the reaction must be accounted from intermediate C, since the rds of the process is to overcome the TS-CD from C intermediate.

The large difference in the reaction thermodynamics as function of the selected nucleophile prompted us to try to reduce the anionic character weight of the halide by studying what happens when instead of using the I⁻ ion, we use a salt such as KI. To note, in the discussion above, the counteraction of the nucleophilic halide, that is TBA⁺, was omitted. This omission has two reasons, firstly, the high computational cost of including the tetra-*n*-butylammonium cation, but also to avoid distorting the results since this cation lacks coordinating nature and hence could be located at almost any position. It is worth noting that such unpredictability of the cation position greatly complicates its inclusion in DFT static calculations, and dynamic calculations would likely be necessary. Nevertheless, the disparity in its position throughout the reaction mechanism could also hinder the proper comparison between different epoxide substituents. However, we have simplistically attempted to include the whole TBAI molecule in the calculations in Table 4, as well. In detail, in the presence of TBAI, the intermediates were located, but attempts to locate the connecting transition states failed. Therefore, the much simpler KI was used, for which it is indeed demonstrated that the TS-CD is maintained as the rds with almost the same kinetic cost (see Fig. 5), decreasing only from 23.1 to 20.9 kcal mol⁻¹, while the opening of the epoxide requires 20.9 kcal mol⁻¹ compared with 18.3 kcal mol⁻¹ when using the I⁻ anion alone (see Tables S6–S9[†] for further energetic, structural, and electronic details).

The role of the potassium counteraction is not insignificant, as Fig. 5 demonstrates how the structure of ascorbic acid folds towards the epoxide, while being strongly bound to the iodide anion. It should be noted that TBAI would diminish this counteraction strength, given its less binding nature compared to a K cation. In addition, Fig. 5 also shows the evidence of the role of the side chain of hydroxyl groups.

Next, we analysed reactivity trends using electronic and steric descriptors derived from the starting epoxide substrates. These parameters were generated from SMILES strings using an automated featurization workflow implemented within the

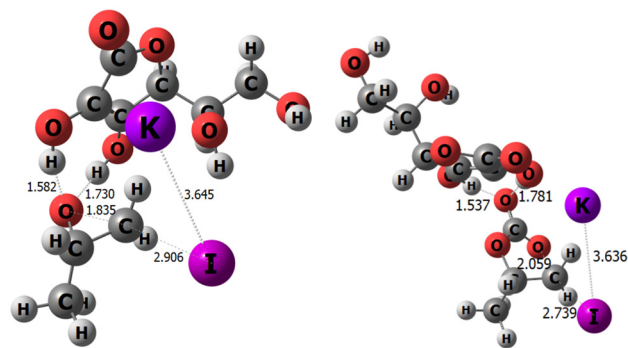


Fig. 5 Transition states TS-AB (left) and TS-CD (right) of the cycloaddition of CO₂ for EPO-CH₃ (selected distances in Å).



AQME program.⁸⁴ In this protocol, a wide array of 34 electronic properties was generated using the PTB and GFN methods of χ TB,⁸⁵ combined with 4 steric descriptors from MORFEUS,⁸⁶ and the octanol/water partition coefficient (MolLogP) obtained with RDKit (Fig. 6a).⁸⁷

The collection of descriptors in Fig. 6a integrates a diverse range of molecular properties along with specific features of the oxygen atom in the epoxide, which is central to the rds and thus expected to influence reaction kinetics. The descriptor database was employed to build machine learning models using the ROBERT software,⁸⁸ including random forest, neural network, gradient boosting and linear regression. After excluding two preliminary outliers (**EPO-(CH₃)₂** and **EPO-trans(Ph)₂**), we obtained a linear model with 19 data points, utilizing % V_{Bur} as

the sole parameter to estimate activation barriers. As shown in Fig. 6b, the model shows high coefficients of determination between predicted and measured values (R^2 of 0.90 and 0.86 in the training and validation sets) and low root mean squared errors (RMSE of 1.0 and 1.3 in the training and validation sets).

No significant overfitting was observed in the leave-one-out cross-validation test (LOOCV; Fig. 6c, left). Additionally, to ensure the model captures meaningful trends in the data, three “flawed” models were evaluated: (i) y -mean, which assesses accuracy when predicted y values are fixed to the mean y value, (ii) y -shuffle, which evaluates accuracy using a model trained on randomly shuffled y values, and (iii) one-hot, which tests accuracy when all descriptors are replaced with binary values (0 s and 1 s).⁸⁹ In all cases, the resulting RMSE was significantly higher than those of the original model and the LOOCV (Fig. 6c, right).

The predictive and interpretative performance of the model was further assessed using the ROBERT score (ROBERT_report PDF file of the ESI[†]).⁸⁸ This score, rated out of ten, evaluates models based on multiple criteria, including correlation between predictions and measurements, human interpretability, error distribution, sensitivity to features, and resilience against overfitting and underfitting.⁹⁰ The score of 7 achieved in this case is a satisfactory result for this low-data scenario, indicating that % V_{Bur} is likely among the most influential descriptors affecting reactivity. This automated SMILES-based workflow also underscores the potential of combining cost-effective descriptors with automation to analyse reactivity trends even in datasets with limited datapoints.⁹¹

3 Conclusions

This study reports a detailed and enhanced investigation of the reaction pathway of CO₂ cycloaddition to epoxides, focusing on the dual catalytic role of ascorbic acid and the iodide ion as the primary nucleophile. The investigation focuses on the interaction of various epoxides substrates with CO₂ with a particular emphasis on the thermodynamics and kinetics of the molecular processes. Through comprehensive computational analysis, the findings highlight the interplay of thermodynamic and kinetic factors governing the reaction pathway, with particular emphasis on the transition states and intermediate species. For unsubstituted ethylene oxide (**EPO-H**), the cyclization of the carbonate is confirmed as the rds, with a substantial energy barrier that aligns with experimental observations. Comparatively, methyl-substituted (**EPO-CH₃**) and bulkier epoxides show an increase in steric hindrance, leading to a higher kinetic cost, particularly at the rds. In detail, steric maps reveal that the steric hindrance increases from the initial free epoxides to transition states (**TS-CD**). The % V_{Bur} values rise with bulkier groups on the carbon facilitating the C–O bond formation, especially when the alpha carbon is doubly substituted. Notably, steric maps exhibit asymmetry, indicating that steric hindrance may be confined to specific quadrants. In monosubstituted cases,

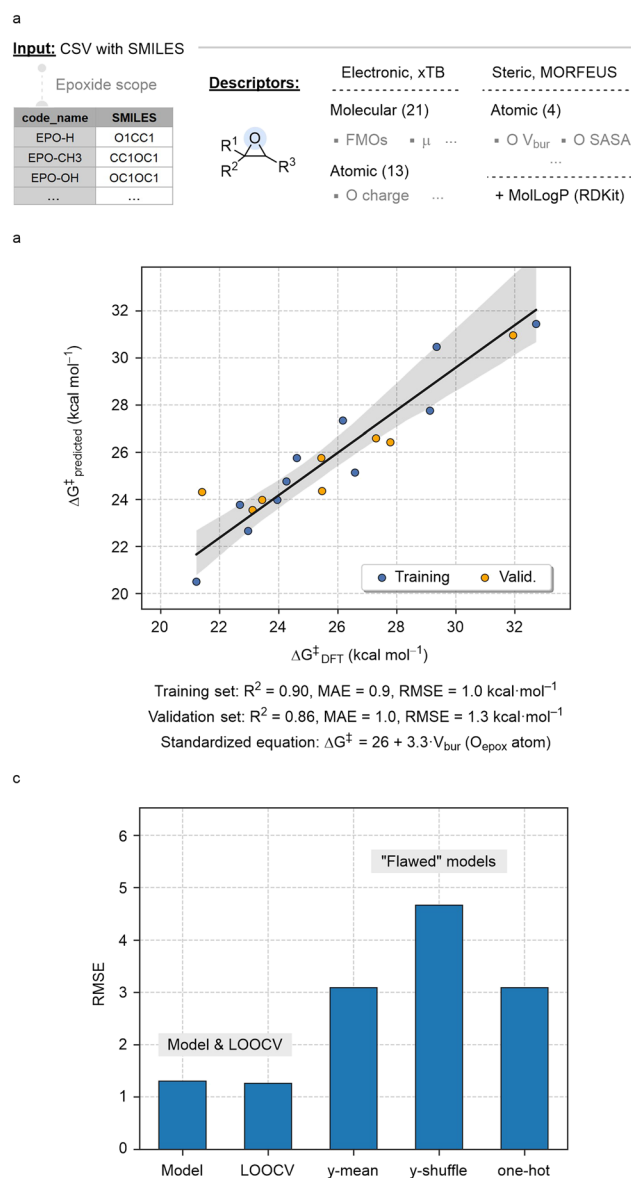


Fig. 6 (a) Summary of the molecular featurization workflow; (b) calculated vs. predicted ΔG^\ddagger (kcal mol⁻¹) obtained with the optimal linear model; (c) RMSE of model, LOOCV and “flawed” models.



certain quadrants remain accessible, unlike in disubstituted scenarios such as **EPO-(Me)₂** or **EPO-(Ph)₂**.

The study demonstrates the successful integration of electronic and steric descriptors into a machine learning framework to predict reactivity trends in epoxide substrates. Utilizing the AQME program for molecular featurization, a broad set of descriptors was compiled, emphasizing features of the O atom critical to the rds. This dataset facilitated the construction of a linear model *via* the ROBERT software,⁸⁸ achieving a strong predictive performance and no significant overfitting. The use of % V_{Bur} as the sole parameter underscores its significance in governing activation barriers. These results illustrate the potential of combining automated, cost-effective molecular descriptors with machine learning to analyze reactivity trends, even with limited datasets.

4 Computational details

4.1 Density functional theory (DFT) calculations

Geometry optimizations using DFT were conducted employing the BP86 level of the generalized gradient approximation,^{92,93} implemented in the Gaussian16 software.⁹⁴ To account for dispersion corrections, the D3 version of Grimme was incorporated.⁹⁵ The electronic structure of the systems was modelled using the valence double- ζ with polarization (SVP) basis set for main group atoms,⁹⁶ while for iodine the quasi-relativistic SDD effective core potential of Stuttgart/Dresden was utilized.⁹⁷ Gibbs energies were determined from single-point energy calculations on the BP86-D3/SVP~sdd geometries using the B3LYP functional,⁹⁸ Grimme's dispersion, and the cc-pVTZ basis set.⁹⁹ Solvent effects, with propylene oxide (PO) as the solvent, were included using the polarization continuum model (PCM).¹⁰⁰ Electronic energies from the latter single point energy calculations in solvent were then converted to Gibbs energy, incorporating zero-point energy and thermal corrections from gas-phase frequency calculations at the BP86/SVP~sdd level.

4.2 Molecular featurization

The AQME program (v1.7.1) was used to generate Boltzmann-averaged descriptors from a CSV file containing SMILES strings of the epoxide substrates.⁸⁴ This automated workflow initially performs conformational searches with RDKit, geometry optimization with xTB,⁸⁵ and descriptor generation with xTB (GFN2 and PTB), MORFEUS,⁸⁶ and RDKit.⁸⁷ Both molecular and atomic descriptors were considered, and all the data was stored in a CSV database. The command line used was:

```
“python -m aqme --qdescp --input “epox.csv” --qdescp_atoms “[O1CC1]””
```

In this command, epox.csv refers to the initial CSV file of SMILES strings and substrate names (SMILES and code_name columns, respectively). The option “--qdescp_atoms “[O1CC1]”” specifies the creation of atomic descriptors for the three elements forming the epoxide rings. The program generated three CSV files with varying number of descriptors (“denovo”, “interpret”, and “full”), from which the “interpret” level was selected as our descriptor database. The atomic descriptors for

the O atom were kept, while the descriptors for the two C atoms were removed (see discussion below).

4.3 ML workflow

The ROBERT program (v1.3.0)⁸⁸ used the “interpret” descriptor database created with AQME as an input, including also the target values (activation barriers, g_barr). ROBERT performed data curation, ML model screening, testing model reliability, and feature importance analysis, among other protocols. The command line used to execute ROBERT was:

```
“python -m robert --csv_name “AQME-ROBERT_interpret.csv” --ignore “[SMILES]” --names “code_name” --y “g_barr””.
```

ROBERT generated a PDF report containing comprehensive information about the models, enhancing transparency and providing instructions for reproducibility. Both this PDF file and the input descriptor database are available as part of the ESI.†

Data availability

The data supporting this article have been included in the main manuscript and as part of the ESI.†

Conflicts of interest

The authors declare no conflict of interest.

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