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Aromatic sulfonation with sulfur trioxide: mechanism and kinetic model†

Samuel L. C. Moors,^{*a} Xavier Deraet,^a Guy Van Assche,^b Paul Geerlings^a and Frank De Proft^a

Electrophilic aromatic sulfonation of benzene with sulfur trioxide is studied with *ab initio* molecular dynamics simulations in gas phase, and in explicit noncomplexing (CCl_3F) and complexing (CH_3NO_2) solvent models. We investigate different possible reaction pathways, the number of SO_3 molecules participating in the reaction, and the influence of the solvent. Our simulations confirm the existence of a low-energy concerted pathway with formation of a cyclic transition state with two SO_3 molecules. Based on the simulation results, we propose a sequence of elementary reaction steps and a kinetic model compatible with experimental data. Furthermore, a new alternative reaction pathway is proposed in complexing solvent, involving two SO_3 and one CH_3NO_2 .

Introduction

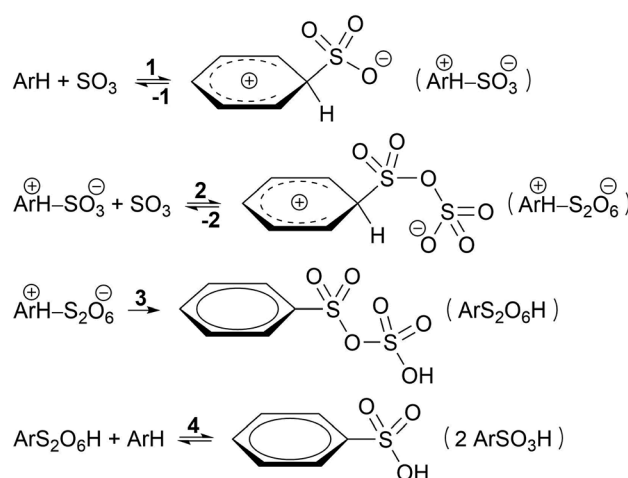
Aromatic sulfonation is a very important chemical transformation of organic compounds.^{1,2} It belongs to the well-known class of electrophilic aromatic substitution ($\text{S}_{\text{E}}\text{Ar}$) reactions alongside nitration, halogenation, acylation and alkylolation. Sulfonation is a key reaction step in large industrial applications including pharmaceuticals, detergents, surfactants, dyes, and pesticides.^{3–5} The most commonly used sulfonating agents are sulfur trioxide (SO_3), oleum, sulfuric acid and chlorosulfuric acid.⁶

Traditionally, $\text{S}_{\text{E}}\text{Ar}$ reactions are explained by a two-step $\text{S}_{\text{E}2}$ or arenium ion mechanism.⁷ In the first and rate-determining step, an electrophile attacks the electron-rich aromatic ring to form a σ -complex or wheland or arenium ion intermediate, which is stabilized by mesomery. Aromaticity is restored in the second step by elimination of H^+ . Recently, alternative pathways to the arenium mechanism have been proposed by Schleyer and coworkers in various electrophilic aromatic substitution reactions⁸ including halogenation,^{9,10} nitration,¹¹ and sulfonation.¹² These studies highlight the diversity in reaction mechanisms for $\text{S}_{\text{E}}\text{Ar}$ and their dependence on substrate and reaction conditions.

Based on kinetic experiments on chlorobenzene and 1,4-dichlorobenzene, Cerfontain and coworkers proposed a three-step kinetic scheme for the sulfonation of arenes with SO_3 in aprotic solvents (Scheme 1).^{13–16} The first step is a reversible

reaction wherein a σ -complex is formed between the arene and a SO_3 molecule. In the second step a second SO_3 reversibly binds to the first SO_3 . Finally, proton transfer from the arene to the second SO_3 in the third step restores aromaticity and drives the reaction toward arenepyrosulfonic acid ($\text{ArS}_2\text{O}_6\text{H}$), which can be readily hydrolyzed in aqueous media. In apolar noncomplexing CCl_3F solvent sulfonation kinetics are first order in SO_3 , and thus step 1 was thought to be rate-limiting.¹⁷ In contrast, in polar SO_3 -complexing¹⁸ CH_3NO_2 the rate is second order in SO_3 , and step 2 was taken as the rate-limiting step.¹⁷

If less than two equivalents of SO_3 are used per mole arene, two sulfonation stages can be distinguished. A fast primary stage purportedly proceeds *via* the reaction steps 1–3 (Scheme 1). A much slower secondary stage was proposed to



Scheme 1 Kinetic model proposed by Cerfontain and coworkers for sulfonation with SO_3 .

^aEenheid Algemene Chemie (ALGC), Vrije Universiteit Brussel (VUB), Pleinlaan 2, 1050 Elsene, Brussels, Belgium. E-mail: samuel.moors@vub.ac.be

^bPhysical Chemistry and Polymer Science (FYSC), Vrije Universiteit Brussel (VUB), Pleinlaan 2, 1050 Elsene, Brussels, Belgium

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proceed through a reaction of arenepyrosulfonic acid with arene, forming two molecules of arenesulfonic acid (ArSO_3H , step 4).^{13,19} This study, however, is focused on the primary sulfonation stage.

The mechanism proposed by Cefontain and coworkers involving a σ -complex intermediate with one SO_3 has been challenged by recent theoretical studies. Morley *et al.*^{20,21} performed quantum chemical calculations at the Hartree-Fock level on the sulfonation of toluene with SO_3 in gas phase, and concluded that formation of a toluene- SO_3 σ -complex was unlikely due to the high energy change required. Instead, they proposed the initial formation of a toluene- $\cdots\text{SO}_3\cdots\text{SO}_3$ π -complex followed by a toluene- S_2O_6 σ -complex with almost the same energy, and a cyclic proton rearrangement yielding toluenepyrosulfonic acid.

Schleyer and coworkers studied the sulfonation of several arenes with static density functional and SCS-MP2 calculations in implicit solvent.¹² The sulfonation of benzene, 1,4-dichlorobenzene, toluene, and naphthalene, in gas phase and in apolar solvent were found to proceed *via* a concerted pathway involving two SO_3 molecules forming a cyclic σ -complex transition state (TS), without intermediate (Scheme 2). In CH_3NO_2 , the cyclic σ -complex became an intermediate state, but with low stability. With only one SO_3 , no intermediate σ -complex was formed in both solvents, and very high energy barriers were needed for sulfonation.

The absence of an intramolecular primary hydrogen kinetic isotope effect (KIE)¹⁷ led Cefontain and coworkers to conclude that the reaction step involving proton transfer (step 3 in Scheme 1) is not rate-limiting. However, the KIE may also be small if the C-H bond is only partially broken at the TS.²² In the cyclic transition state structures involving two SO_3 molecules, as calculated by Schleyer and coworkers,¹² the C-H bond is only slightly elongated at the TS in both CCl_3F (0.08 Å) and CH_3NO_2 (0.10 Å at the second TS). The small bond elongations are consistent with ratios $k_{\text{H}}/k_{\text{D}} = 1.2\text{--}1.3$, suggesting that step 3 may indeed be rate-limiting.

Importantly however, it is not clear how the participation of two SO_3 molecules, as suggested previously,^{12,21} fits in with the first order rate dependence on the SO_3 concentration in apolar solvent. Morkovnik and Akopova²³ have proposed an alternative low-barrier mechanism with one SO_3 in apolar solvent. In their

proposal, a sulfuric acid molecule acts as a catalyst by transferring the proton from benzene to SO_3 *via* a relay-race mechanism (Scheme 2).

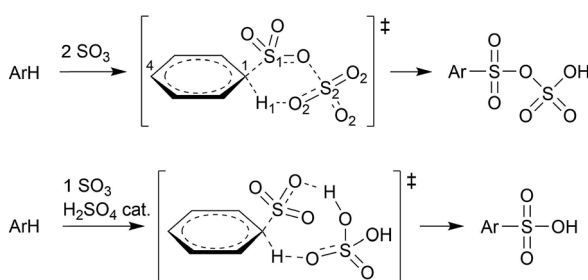
In this study, we aim to further elucidate the reaction mechanism and explain the experimental data. To fully account for solvation and dynamic effects, *ab initio* molecular dynamics (AIMD) is the preferred method of choice. The article is organized as follows. We start with the proposal of a new kinetic model, which is validated by a variety of AIMD simulations in gas phase as well as fully solvated in CCl_3F and CH_3NO_2 solvent. Metadynamics (MTD) simulations are performed to study the stability and reactivity of benzene- SO_3 and benzene- S_2O_6 σ -complexes, and to estimate the free energy surface (FES) of the sulfonation reaction. From unbiased MD simulations, we analyze intermolecular interactions between benzene, SO_3 , and solvent, and we investigate the stability of benzene- $\cdots\text{SO}_3$ and benzene- $\cdots\text{SO}_3\cdots\text{SO}_3$ π -complexes in each environment. Finally, restrained MD (rMD) simulations are performed to evaluate the reactivity of the benzene- S_2O_6 σ -complex, and of benzene- $\text{SO}_3 + 1$ catalytic H_2SO_4 .

Results and discussion

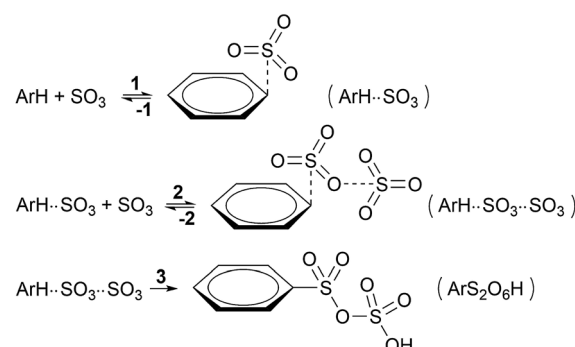
Mechanism and kinetic model

To enhance readability of the study, we start with the proposal of a new sequence of reaction steps given in Scheme 3 based on our calculation results, which are presented in the following sections, and in agreement with experimental data.

Here, step 1 represents formation of $\text{ArH}\cdots\text{SO}_3$, a π -complex between the arene and a first SO_3 molecule (hereafter named the primary SO_3), step 2 is the formation of the $\text{ArH}\cdots\text{SO}_3\cdots\text{SO}_3$ π -complex between $\text{ArH}\cdots\text{SO}_3$ and a second SO_3 (the assisting SO_3), and step 3 is the actual sulfonation reaction with formation of arenepyrosulfonic acid $\text{ArS}_2\text{O}_6\text{H}$, the main reaction product. Step 3 requires a significant reorientation of the two SO_3 molecules toward the plane of the benzene ring to make a cyclic proton rearrangement possible. This mechanism differs from the proposed mechanism of Cefontain and coworkers¹³⁻¹⁶ in that it does not account for an intermediate σ -complex state. Although its presence is not ruled out, our simulations confirm previous experimental and theoretical data^{12,24} that the stability



Scheme 2 Concerted sulfonation mechanisms in gas phase or apolar solvent with 2 SO_3 molecules according to Schleyer and coworkers,¹² and with 1 $\text{SO}_3 + \text{H}_2\text{SO}_4$ catalyst according to Morkovnik and Akopova.²³



Scheme 3 New proposed kinetic model for the primary sulfonation stage with SO_3 .



of σ -complexes is generally low and may be discarded from the kinetic model. Using steady-state conditions, the reaction rate is given by the rate equation (see Appendix):

$$\frac{d(ArS_2O_6H)}{dt} = \frac{k_1 k_2 k_3 [ArH][SO_3]^2}{(k_{-2} + k_3)(k_{-1} + k_2[SO_3])} \quad (1)$$

Depending on the solvent, the reaction may be first order or second order in SO_3 ,^{13,14} which dictates limiting condition 1 (Table 1). In apolar CCl_3F solvent, the reaction is first order in SO_3 , and thus $k_{-1} \ll k_2[SO_3]$, which implies that the $ArH \cdots SO_3$ complex must remain stable long enough that it can associate with an assisting SO_3 before the primary SO_3 dissociates from the arene. In polar CH_3NO_2 , the reaction is second order in SO_3 and $k_{-1} \gg k_2[SO_3]$, thus the arene exists as mostly uncomplexed with SO_3 . Whereas the rate-limiting step could in principle be step 1 or 3 in CCl_3F , and step 2 or 3 in CH_3NO_2 (Table 1, condition 2), our results suggest that (i) the calculated free energy barriers of sulfonation are relatively high, and (ii) the assisting SO_3 detaches rapidly from $ArH \cdots SO_3 \cdots SO_3$, which strongly suggests that $k_{-2} \gg k_3$, thus step 3 is rate-limiting in both solvents.

In the following sections we validate the proposed mechanism and kinetic model by analyzing a carefully chosen set of advanced AIMD simulations in gas phase and fully solvated in CCl_3F and CH_3NO_2 .

Reaction of benzene + 1 SO_3

We first examine if a stable σ -complex can be formed between benzene and a single SO_3 molecule, as proposed by Cerfontain and coworkers,^{13–16} and if aromatic sulfonation is possible with only 1 SO_3 . MTD simulations are performed with one collective variable (CV), the coordination number CN_{CS} that measures bond formation between a benzene carbon and the SO_3 sulfur. The resulting potential of mean force (PMF) in gas phase, in CCl_3F , and in CH_3NO_2 is shown in Fig. 1. The benzene $\cdots SO_3$ complex is located at $CN_{CS} \approx 0.085$ ($r_{CS} = 2.9 \text{ \AA}$). Large free energy differences are observed between the three environments, the σ -complex ($CN_{CS} \approx 0.5$ –0.6) being most stabilized in the polar CH_3NO_2 . However, no intermediate σ -complex between benzene and SO_3 can be discerned from the PMF, in agreement with previous static calculations.¹²

The MTD simulations are continued by further adding Gaussian hills to the reactant state until reaction. No reaction is observed until a barrier height $\Delta F^\ddagger \approx 244 \text{ kJ mol}^{-1}$ is reached,

Table 1 Experimental order in SO_3 (x) in solvents CCl_3F and CH_3NO_2 , corresponding limiting conditions and rate-limiting step (RLS) derived from eqn (1)

Solvent	$[SO_3]^x$	Condition 1	Condition 2	RLS
CCl_3F	1	$k_{-1} \ll k_2[SO_3]$	$k_{-2} \ll k_3$ $k_{-2} \gg k_3$	1 3
CH_3NO_2	2	$k_{-1} \gg k_2[SO_3]$	$k_{-2} \ll k_3$ $k_{-2} \gg k_3$	2 3

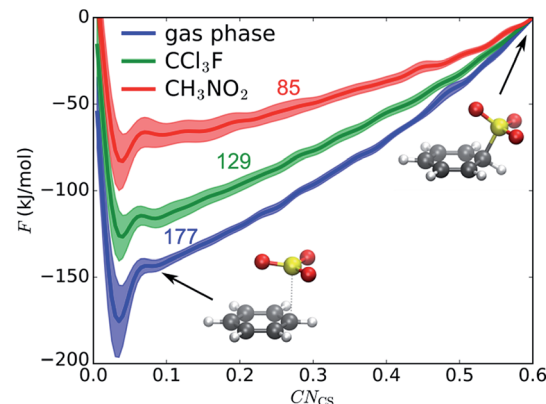


Fig. 1 Reaction between benzene and 1 SO_3 in gas phase, CCl_3F , and CH_3NO_2 . PMF of 1D MTD with error bars representing the standard deviation. Numbers above the curves indicate free energy ΔF (kJ mol^{-1}) of the σ -complex at $CN_{CS} = 0.6$ relative to the reactant state. Structures refer to the gas phase curve.

which is incompatible with the fast experimental reaction kinetics.¹⁵ In conclusion, the MTD simulations suggest that sulfonation of benzene with a single SO_3 is not a viable pathway, and can be discarded in the kinetic analysis.

Stability of benzene $\cdots SO_3 \cdots SO_3$ and benzene $\cdots SO_3$ π -complexes

To assess the validity of steps 1 and 2 of our kinetic model (Scheme 3), the stability of the π -complexes is studied with a series of independent (different starting structures and velocities) and unbiased AIMD simulations starting from the benzene $\cdots SO_3 \cdots SO_3$ complex. The results are listed in Table 2 and summarized in Fig. 2.

In gas phase, the benzene $\cdots SO_3 \cdots SO_3$ complex remains stable throughout the simulation. A clear distinction can be made between the two interacting SO_3 molecules (Fig. 3a). The first (primary) SO_3 is tightly bound to benzene by strong π interaction between its S atom and the π -conjugated ring system located above the benzene C atoms. The second (assisting) SO_3 is loosely bound to the primary SO_3 and occasionally strays away from the complex, while remaining at the same side of the ring plane most of the time. In both CCl_3F and CH_3NO_2 solvents, however, one SO_3 dissociates from the complex and drifts into the bulk solvent. The resulting benzene $\cdots SO_3$ complex remains stable in CCl_3F , whereas in CH_3NO_2 this complex dissociates into solvated benzene and SO_3 .

Overall, the unbiased MD simulations described here show that the benzene $\cdots SO_3 \cdots SO_3$ π -complex is stable in gas phase but unstable in both solvents. The benzene $\cdots SO_3$ complex appears relatively stable in CCl_3F , consistent with the limiting condition $k_{-1} \ll k_2[SO_3]$ (Table 1), and in agreement with the experimental first order rate in SO_3 . In contrast, strong interactions with CH_3NO_2 solvent shift the equilibrium toward separate benzene and SO_3 reactants, in accord with $k_{-1} \gg k_2[SO_3]$ (Table 1), in correspondence with the second order rate



Table 2 Overview of MD and rMD simulations

Initial conditions	Environment	t^a (ps)	Reaction ^b
Unbiased MD			
Benzene \cdots SO ₃ \cdots SO ₃	Gas phase	120	Remains stable
	CCl ₃ F	200	\rightarrow Benzene \cdots SO ₃ + SO ₃ (30 ps)
		200	\rightarrow Benzene \cdots SO ₃ + SO ₃ (140 ps)
	CH ₃ NO ₂	70	\rightarrow Benzene \cdots SO ₃ + SO ₃ (30 ps) \rightarrow Benzene + 2 SO ₃ (70 ps)
		180	\rightarrow Benzene \cdots SO ₃ + SO ₃ (60 ps) \rightarrow Benzene + 2 SO ₃ (180 ps)
Restrained MD			
Benzene-S ₂ O ₆	Gas phase	40	\rightarrow Benzene-S ₂ O ₆ H
		60	\rightarrow Benzene-S ₂ O ₆ H
	CCl ₃ F	5	\rightarrow Benzene-S ₂ O ₆ H
		10	\rightarrow Benzene-S ₂ O ₆ H
Benzene-SO ₃ \cdots H ₂ SO ₄	CH ₃ NO ₂	200	Remains stable
	Gas phase	6	\rightarrow Benzene-SO ₃ H \cdots H ₂ SO ₄
	CCl ₃ F	170	\rightarrow Benzene-SO ₃ H \cdots H ₂ SO ₄
	CH ₃ NO ₂	90	\rightarrow Benzene-SO ₃ H \cdots H ₂ SO ₄

^a Total simulation time. ^b Time at which the event takes place is given between parentheses.

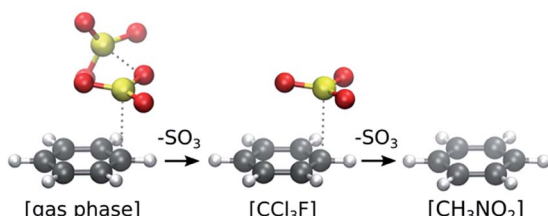


Fig. 2 Stability of π -complexes in different solvation models, showing the decrease in number of SO₃ molecules complexed with benzene.

in SO₃. In summary, our simulation results suggest that the reactant species is benzene \cdots SO₃ in CCl₃F and benzene CH₃NO₂, in agreement with steps 1 and 2 of our kinetic model (Scheme 3).

Specific solvation of reactants

The low stability of the benzene \cdots SO₃ π -complex in CH₃NO₂ is attributed to specific solvent interactions. Electrostatic interaction between the positively charged S atom of SO₃ and the negatively charged O atoms of CH₃NO₂ strongly increases upon dissociation of the complex. This solvent interaction is demonstrated in Fig. 3b by the red probability isosurface of CH₃NO₂ O atoms around SO₃, where larger volumes indicate a higher probability of finding an O atom inside the isosurface. Likewise, electrostatic interactions between the benzene π electron cloud and the positively charged methyl group of CH₃NO₂ (grey), and between the positively charged H atoms of benzene and the O atoms of CH₃NO₂ (red) markedly increase upon dissociation (Fig. 3c).

Reaction of benzene + 2 SO₃

The sulfonation reaction starting from the benzene \cdots SO₃ \cdots SO₃ π -complex, corresponding to step 3 in our kinetic model (Scheme 3), is investigated with MTD and two collective

variables, CN_{CS} and CN_{CO}; the latter is the coordination between a benzene C and the O atom of the assisting SO₃. In Fig. 4a–c, the resulting 2D FES is shown as the average of three

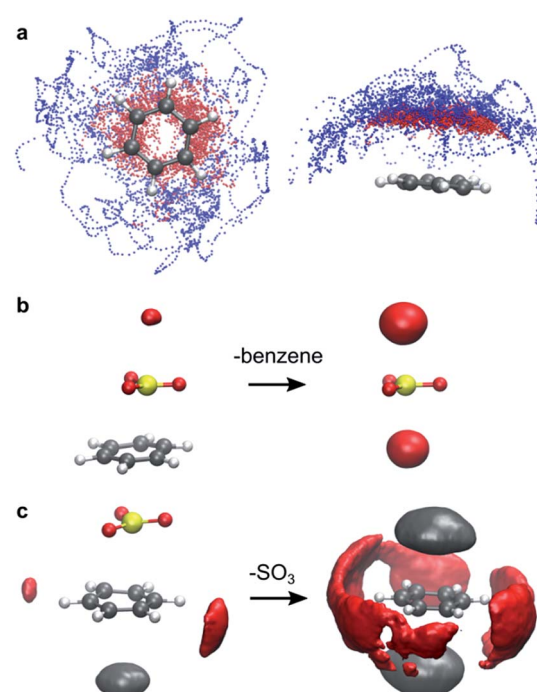


Fig. 3 Intermolecular interactions observed during AIMD simulations. (a) Benzene-SO₃ interactions in gas phase. Superposition of snapshots of the benzene \cdots SO₃ \cdots SO₃ complex in top view (left) and side view (right). Dots represent the S atom positions of the primary SO₃ (red) and assisting SO₃ (blue). (b) Specific SO₃-CH₃NO₂ interactions. Probability isosurface (red) of CH₃NO₂ O atoms around the SO₃ S atom in a benzene \cdots SO₃ complex (left) and in separated SO₃ (right). (c) Specific benzene-CH₃NO₂ interactions. Probability isosurface of solvent O (red) and C (grey) atoms around benzene C atoms, in a benzene \cdots SO₃ complex (left) and in separated benzene (right). Deviations from symmetry in the isosurfaces are due to finite sampling.



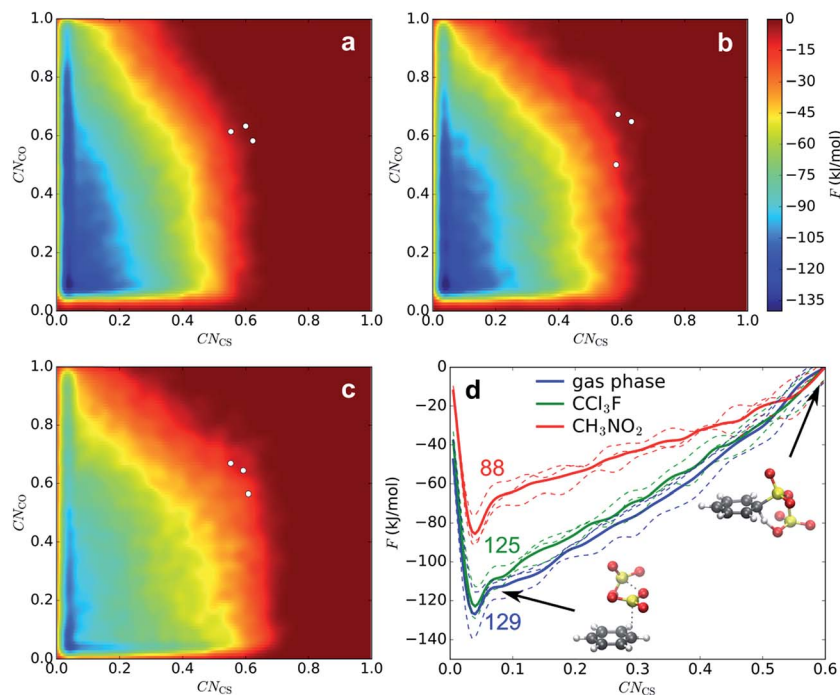


Fig. 4 FES of 2D MTD in gas phase (a), CCl_3F (b), and CH_3NO_2 (c), and corresponding projected PMFs along CN_{CS} (d). The given numbers indicate average ΔF^\ddagger values (kJ mol^{-1}).

MTD simulations. Again, no intermediate σ -complex is observed in the three environments. The surface is very similar in gas phase and CCl_3F , whereas in CH_3NO_2 the reactant valley is much shallower due to stabilization of TS by the electrostatic field. The position of TS, as indicated with white dots, is similar in the three environments: $\text{CV1} \cong \text{CV2} \cong 0.6$, which corresponds to a cyclic transition state with two SO_3 molecules at $r_{\text{CS}} \cong 1.84 \text{ \AA}$ and $r_{\text{CO}} \cong 2.7\text{--}3.2 \text{ \AA}$. In all cases benzene pyrosulfonic acid is formed in one step. In contrast to CV1, the free energy change along the CV2 axis is small. After projecting CV2 onto CV1, a 1D PMF is obtained (Fig. 4d). Mean geometric parameters at TS are shown in Table S2 (ESI†).

Low-temperature MD of the σ -complex in CH_3NO_2 shows that a metastable intermediate state may exist nonetheless (see ESI†). The stability of the σ -complex is however very low, which may in part be due to the BLYP density functional used in this study. Low stabilities of the intermediate state ($2\text{--}4 \text{ kJ mol}^{-1}$) have also been calculated previously at the M06-2X/6-311+G(2d,2p) level for benzene and 1,4-dichlorobenzene in implicit CH_3NO_2 .¹²

Reactivity of benzene- S_2O_6 σ -complex

Although we have now established that σ -complexes of benzene and SO_3 are fairly unstable, it is insightful to analyze the benzene- SO_3 interactions in the σ -complex state and to monitor the influence of the environment on their reactivity toward benzene- $\text{S}_2\text{O}_6\text{H}$ formation. Additionally, the obtained reactivity information will allow us to decompose the free energy barrier into several discrete contributions.

Restrained MD (rMD) simulations are performed of the benzene- SO_3 σ -complex + an assisting SO_3 , which forms a benzene- S_2O_6 σ -complex. To stabilize the σ -complex, the $\text{C}_1\text{--}\text{S}_1$ bond is restrained with a harmonic potential, which keeps CN_{CS} between 0.5 and 0.6 until sulfonation takes place. The results are summarized in Table 2. In gas phase (40–60 ps) and in CCl_3F solvent (5–10 ps), a concerted sulfonation reaction takes place by proton transfer to the assisting SO_3 with formation of benzene pyrosulfonic acid in one step (Fig. 5). In CH_3NO_2 solvent, however, no product formation is observed during 200 ps. Two factors contribute to the increased stability of the σ -complex in CH_3NO_2 : (i) stabilization of the zwitterionic TS by the strong solvent electrostatic field, and (ii) competition with the solvent for hydrogen bonding with H_1 . The latter effect is demonstrated in Fig. 6a, showing a stable intramolecular $\text{O}_2\text{--}\text{H}_1$

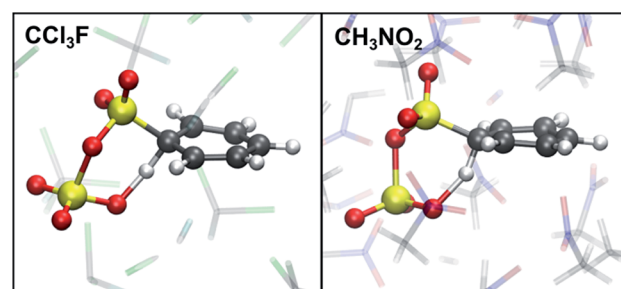


Fig. 5 TS of sulfonation of benzene with 2 SO_3 . In CCl_3F , the reaction takes place spontaneously during rMD. In CH_3NO_2 an additional barrier needs to be crossed with rMTD. Atom colors are: C (grey), Cl (green), F (cyan), H (white), N (blue), O (red), S (yellow).



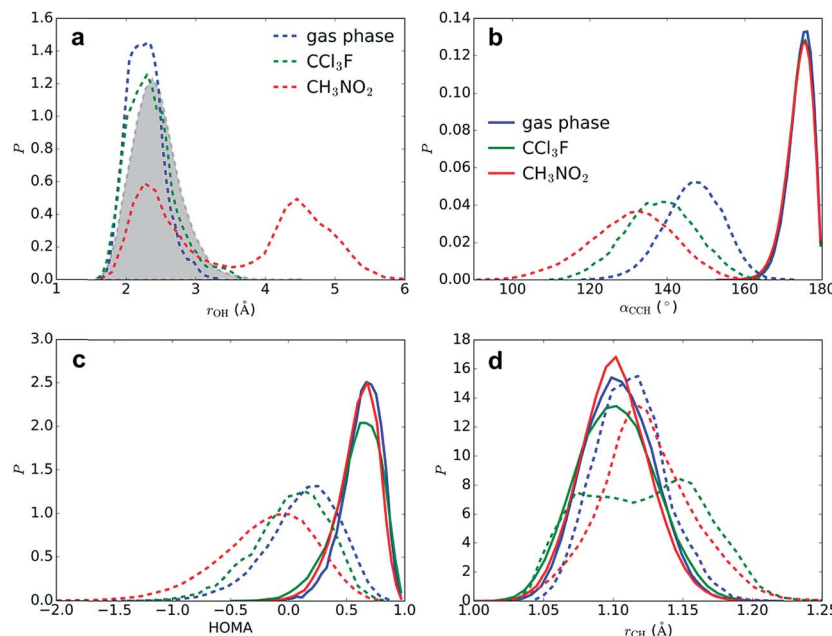


Fig. 6 Geometric probability densities (P) in reactant state (benzene- $\cdots\text{SO}_3$ π -complex, full lines) and close to TS (benzene- S_2O_6 σ -complex, dashed lines), in gas phase (blue), CCl_3F (green), and CH_3NO_2 (red). (a) minimal distance $r(\text{H}_1-\text{O}_2)$ and minimal distance $r(\text{H}_1-\text{O})$ with CH_3NO_2 (filled grey), (b) angle $\alpha(\text{C}_4-\text{C}_1-\text{H}_1)$, (c) HOMA aromaticity index, (d) distance $r(\text{C}_1-\text{H}_1)$.

hydrogen bond in gas phase and CCl_3F , whereas in CH_3NO_2 this hydrogen bond is present only $\sim 50\%$ of the time.

Solvation effects at the transition state

The segments of rMD trajectories before reaction are analyzed to determine how the transition state geometry is affected by solvation effects. The effects of the solvent on benzene geometry near TS are striking (Fig. 6b-d). The angle $\alpha(\text{C}_4-\text{C}_1-\text{H}_1)$ is calculated as a measure for the degree of sp^3 hybridization at C_1 and thus the stability of the σ -complex. In the reactant state, the mean angle $\langle\alpha\rangle = 174^\circ$ in all three environments, indicating that H_1 is located mostly in the plane of the benzene ring and hence C_1 is almost purely sp^2 hybridized, whereas near TS C_1 displays strong sp^3 character. The smallest $\langle\alpha\rangle$ is measured in CH_3NO_2 (131°), followed by CCl_3F (138°) and gas phase (147°). The strong stabilizing effect of CCl_3F is remarkable given its low polarity. The harmonic oscillator model of aromaticity index (HOMA)²⁵ fluctuates around a mean value $\langle\text{HOMA}\rangle = 0.59-0.63$ in the reactant state, but degrades near TS, indicating nearly complete loss of aromaticity. This effect is especially strong in CH_3NO_2 ($\langle\text{HOMA}\rangle = -0.19$), in comparison with CCl_3F (0.02) and gas phase (0.13). The C_1-H_1 bond is stretched from $\langle r_{\text{CH}} \rangle = 1.102-1.104$ Å to 1.124, 1.121, and 1.113 Å in CH_3NO_2 , CCl_3F , and gas phase, respectively. In CCl_3F , a bimodal distribution is observed due to frequent near-proton transfer events taking place.

An alternative pathway in CH_3NO_2

An estimate of the additional free energy barrier required for sulfonation of the benzene- S_2O_6 σ -complex in CH_3NO_2 is

obtained with restrained MTD (rMTD) (for computational details and PMFs, see ESI†), yielding an average additional $\Delta F = 12$ kJ mol⁻¹. In one of the rMTD runs, an alternative pathway is observed (Fig. 7). First, the C-H bond is weakened by hydrogen bonding interaction with the assisting SO_3 , which strongly lowers the activation free energy of proton transfer. Next, the proton is transferred to a nitromethane molecule (224 fs, TS1). Shortly thereafter, the proton is transferred from nitromethane to the primary SO_3 to form benzenesulfonic acid

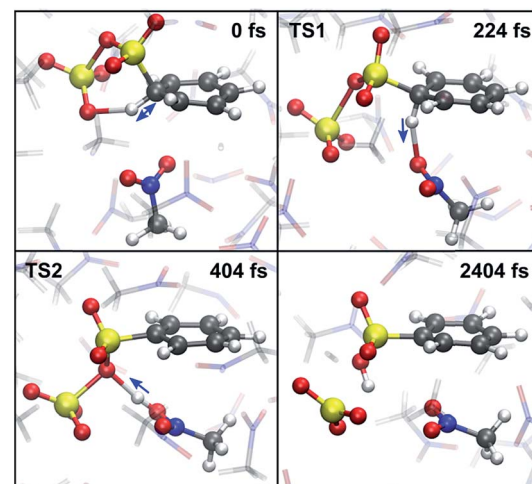


Fig. 7 Snapshots of the new CH_3NO_2 -mediated sulfonation pathway. (0 fs) H_1-O_2 hydrogen bond at maximal strength. (224 fs) TS1: proton transfer from benzene to CH_3NO_2 . (404 fs) TS2: proton transfer from CH_3NO_2 to the primary SO_3 . (2404 fs) stable benzenesulfonic acid product. Blue arrows indicate the direction of motion of the proton.

(404 fs, TS2). Thus, although in this mechanism the assisting SO_3 is not the proton acceptor, it still plays an essential role in the reaction by activating the $\text{C}_1\text{-H}_1$ bond. The calculated additional free energy barrier for this pathway $\Delta F^\ddagger = 14 \text{ kJ mol}^{-1}$, suggesting that this new mechanism is feasible and may compete with the cyclic proton transfer pathway.

Free energy decomposition

Combining the results of 1D and 2D MTD and the rMD simulations, the total ΔF^\ddagger can be decomposed into three contributions: (1) benzene- $\cdots\text{SO}_3$ π -complex to benzene- SO_3 σ -complex, (2) benzene- SO_3 σ -complex to benzene- S_2O_6 σ -complex, and (3) benzene- S_2O_6 to TS. In gas phase, the presence of an assisting SO_3 strongly reduces the free energy required to form a σ -complex from the π -complex (177 to 129 kJ mol^{-1}). Once the benzene- S_2O_6 σ -complex is formed, sulfonation proceeds spontaneously. In CCl_3F , the assisting SO_3 has a slight stabilizing effect on the σ -complex (129 to 125 kJ mol^{-1}). In CH_3NO_2 , the solvent already has a strong stabilizing effect, which causes the additional stabilizing effect of the assisting SO_3 to be rather small (85 to 76 kJ mol^{-1}). The combined stabilizing effects lead to an additional free energy barrier required for sulfonation (76 + 12 = 88 kJ mol^{-1}).

H_2SO_4 as a catalyst

As an alternative explanation for the first-order reaction in SO_3 in noncomplexing media, Morkovnik and Akopova suggested a relay-race mechanism involving a Brønsted acid as a catalyst (Scheme 2).²³ The plausibility of this mechanism is verified with rMD simulations of the benzene- SO_3 σ -complex, restrained at the $\text{C}_1\text{-S}_1$ bond, in the presence of a H_2SO_4 molecule. Spontaneous sulfonation takes place in all three environments within 170 ps (Table 2), although the mechanism differs from the mechanism proposed previously²³ and depends on the polarity

of the environment (Fig. 8). In gas phase and CCl_3F , H_1 is first transferred from benzene to H_2SO_4 at TS1, stabilized by one (gas phase) or two (CCl_3F) $\text{H}_2\text{SO}_4\cdots\text{SO}_3$ hydrogen bonds. Proton transfer from H_2SO_4 to SO_3 follows quickly thereafter (TS2, 16 fs). In CH_3NO_2 , first proton transfer occurs from H_2SO_4 to SO_3 (TS1), followed by benzene-to- H_2SO_4 proton transfer 106 fs later (TS2). We conclude that catalytic amounts of H_2SO_4 in CCl_3F may provide an alternative pathway consistent with first-order kinetics in SO_3 , however the participation of H_2SO_4 is not required in our kinetic model. In CH_3NO_2 , the stability of the benzene- $\cdots\text{SO}_3\cdots\text{H}_2\text{SO}_4$ π -complex is probably too low to meaningfully contribute to the overall reaction.

Conclusions

The mechanism and kinetics of electrophilic aromatic sulfonation have been investigated in great detail with various DFT-based first-principles MD and MTD simulations. Three different environments were compared: gas phase, and fully solvated in explicit apolar (CCl_3F) and polar (CH_3NO_2) solvents. Several alternative reaction mechanisms were evaluated, including with 1 or 2 SO_3 molecules, with a catalytic H_2SO_4 molecule, and with a participating solvent molecule. The kinetic model proposed by Cerfontain and coworkers, involving stable ArH-SO_3 and $\text{ArH-S}_2\text{O}_6$ σ -complexes in step 1 and 2, was examined. Our results suggest that both benzene- SO_3 and benzene- S_2O_6 σ -complexes are unstable at room temperature, thus ruling out the Cerfontain model. Our simulation data confirm the static gas phase and implicit solvent calculations performed by Schleyer and coworkers, which suggest that the free energy barrier for sulfonation with a single SO_3 is too high to be feasible. In all three environments, a low-energy concerted pathway starting from the benzene- $\cdots\text{SO}_3\cdots\text{SO}_3$ π -complex was found, involving a cyclic transition state with proton transfer from benzene to the assisting SO_3 , and with formation of benzene pyrosulfonic acid. This mechanism is in agreement with the calculations of Schleyer and coworkers, although in CH_3NO_2 an intermediate cyclic σ -complex state with low stability was found by them.

In order to bring the concerted mechanism into agreement with experimental kinetic data, a new kinetic model was proposed. Steps 1 and 2 represent the formation of ArH-SO_3 and $\text{ArH-SO}_3\cdots\text{SO}_3$ π -complexes, respectively, followed by cyclic proton transfer with formation of benzene pyrosulfonic acid in step 3. A rate equation was calculated using the steady-state approximation, and the limiting conditions were determined on the basis of the experimental rate order in SO_3 . In CCl_3F we found that $k_{-1} \ll k_2[\text{SO}_3]$, whereas in CH_3NO_2 the opposite condition $k_{-1} \ll k_2[\text{SO}_3]$ applies. The validity of steps 1 and 2 was confirmed with long timescale MD simulations, suggesting that the benzene- $\cdots\text{SO}_3$ π -complex is relatively stable in CCl_3F but quickly dissociates in CH_3NO_2 . The mechanism suggested by Morkovnik and Akopova involving one SO_3 and a catalytic H_2SO_4 molecule was shown to provide a viable alternative route to sulfonation in CCl_3F . Furthermore, a new and unanticipated mechanism was discovered in CH_3NO_2 , in which two SO_3 and one CH_3NO_2 cooperate in a stepwise proton transfer from benzene to CH_3NO_2 , followed by proton transfer

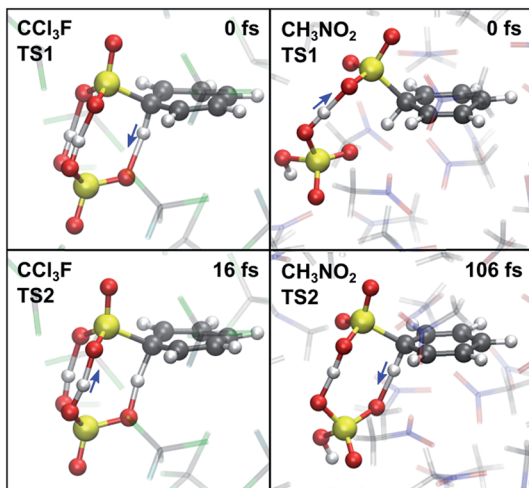


Fig. 8 TS1 (top) and TS2 (bottom) in the presence of catalytic H_2SO_4 , corresponding to the first and second proton transfer in CCl_3F (left) and CH_3NO_2 (right).



to the primary SO_3 , leading directly to benzenesulfonic acid. Future research will focus on sulfonation of substituted benzenes and benzene derivatives with SO_3 to investigate whether our kinetic model is applicable to a wide range of arenes.

Computational methods

Molecular dynamics

Born–Oppenheimer molecular dynamics simulations are performed at the DFT level with the gradient-corrected BLYP functional,^{26,27} the DZVP-GTH basis set,²⁸ and Grimme D3 dispersion corrections.²⁹ The BLYP functional has been shown to produce barriers for SN_2 and E_2 reactions in good agreement with high-level (MP2) calculations.³⁰ The integration time step is set at 1 fs, with snapshots taken every 2 fs. Simulations in solvent are performed in the NVT ensemble, using the canonical sampling through velocity rescaling thermostat³¹ with a time constant of 50 fs. The reactants are placed inside a periodic cubic box filled with 25 CCl_3F or 40 CH_3NO_2 molecules. The densities are set to the experimental densities of CCl_3F (1.48 g ml⁻¹) and CH_3NO_2 (1.13 g ml⁻¹) at 298 K, with corresponding box lengths 15.847/15.911 Å (CCl_3F) and 15.482/15.549 Å (CH_3NO_2), depending on the number of SO_3 molecules present. All AIMD simulations are performed with the CP2K simulation package (version 2.6).³²

Metadynamics

Most chemical reactions are not accessible in the timescale (typically < 1 ns) that can be reached with AIMD. In order to sample the relevant TS regions, enhanced sampling techniques need to be used. Metadynamics is a nonequilibrium MD method introduced by Laio and Parrinello.^{33,34} In recent studies, we have successfully used the MTD technique to estimate free energy surfaces in a variety of systems, including heterogeneous catalysis and reactions in solution.^{35–37} Sampling is advanced by adding Gaussian potential hills along a limited number of carefully chosen collected variables during the simulation, effectively flattening the FES by filling low-energy regions. The FES is then calculated as the opposite of the summation of the Gaussian hills.

The hills are added every 25 steps along one or two collective variables (CVs), defined by coordination numbers CN:

$$\text{CN} = \sum_{i,j} \frac{1 - (r_{ij}/r_0)^6}{1 - (r_{ij}/r_0)^{12}} \quad (2)$$

where the sum runs over two nonoverlapping sets of atoms i and j , r_{ij} is the distance between atoms i and j , and r_0 is a reference distance. CV1 is described by CN_{CS} between a benzene C_1 and S_1 (see Scheme 2 for atom numbering), with $r_0 = 1.964$ Å. In the 2D MTD simulations, CV2 corresponds to CN_{CO} , the sum of coordination numbers between C_1 and the three O_2 atoms, with $r_0 = 2.719$ Å. The width of the hills is set to 0.02. In the 1D MTD simulations, the hill height is initially set to $H = 1$ kJ mol⁻¹, and reduced to 0.5 kJ mol⁻¹ after 40 ps. In the 2D MTD, $H = 2$ kJ mol⁻¹. To limit sampling to a region close to the bound

state, half harmonic bias potentials are added to CV1 and CV2 at position 0.03 with a force constant $K_f = 100$ a.u. In the 2D MTD, an additional half harmonic potential is added to the $\text{S}_1\text{--S}_2$ distance at 4.5 Å with $K_f = 0.19$ a.u. The MTD simulations are ended once the product state is reached. To dampen excessive proton fluctuations, the mass of H_1 is increased to that of tritium.

In the 2D MTD, Gaussian hills are placed along two CVs simultaneously: CN_{CS} and CN_{CO} , which allows for a sulfonation with either one or two SO_3 molecules. Due to the involvement of proton transfer, however, it is impossible to simulate the reaction in the conventional way, *i.e.* sampling many forward and reverse reactions by filling both the reactant and product wells with Gaussian hills. Instead, three independent MTD simulations are initiated from different starting geometries and terminated as soon as the proton transfer has taken place. The TS is then taken as the last stationary point along the r_{CH} trajectory before proton transfer takes place.

Analysis

Probability isosurfaces are calculated with the volmap tool of VMD.³⁸ The probability isosurface of CH_3NO_2 O atoms within 4 Å of the SO_3 S atom is calculated with isovalue = 65%. Probability isosurfaces of CH_3NO_2 O and C atoms within 4 Å of any of the six benzene C atoms are calculated with isovalue = 30%. Conversion of the 2D FES into a 1D PMF is achieved by projecting CV2 onto CV1, and free energy barriers are calculated as the free energy difference between the TS and the reactant state.³⁶

Appendix

Here we derive the rate equation (eqn (1)). Let $\text{A} = \text{ArH}$, $\text{B} = \text{SO}_3$, $\text{C} = \text{ArH}\cdots\text{SO}_3$, $\text{D} = \text{ArH}\cdots\text{SO}_3\cdots\text{SO}_3$, $\text{E} = \text{ArS}_2\text{O}_6\text{H}$. Using the steady-state approximation, we have

$$\frac{d[\text{C}]}{dt} = k_1[\text{A}][\text{B}] - k_{-1}[\text{C}] - k_2[\text{B}][\text{C}] = 0 \quad (3)$$

$$\frac{d[\text{D}]}{dt} = k_2[\text{B}][\text{C}] - k_{-2}[\text{D}] - k_3[\text{D}] = 0 \quad (4)$$

From eqn (3)–(4):

$$[\text{D}] = \frac{k_1 k_2 [\text{A}][\text{B}]^2}{(k_{-2} + k_3)(k_{-1} + k_2[\text{B}])} \quad (5)$$

$$\frac{d[\text{E}]}{dt} = k_3[\text{D}] = \frac{k_1 k_2 k_3 [\text{A}][\text{B}]^2}{(k_{-2} + k_3)(k_{-1} + k_2[\text{B}])} \quad (6)$$

The form of this equation differs from the rate equation presented by Lammertsma and Cerfontain,¹⁷ who in the denominator neglect k_{-2} in comparison to k_3 in the term $(k_{-2} + k_3)k_2[\text{B}]$, but not in $(k_{-2} + k_3)k_{-1}$. No rationale was given for the partial neglect of k_{-2} , and it does not seem to simplify the kinetic analysis.



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References

- 1 E. A. Knaggs and M. J. Nepras, in *Kirk-Othmer Encyclopedia of Chemical Technology*, John Wiley & Sons, Inc., 2000, DOI: 10.1002/0471238961.1921120611140107.a01.
- 2 M. B. Smith and J. March, in *March's Advanced Organic Chemistry*, John Wiley & Sons, Inc., 2006, DOI: 10.1002/9780470084960.ch11, pp. 657–751.
- 3 J. S. Carey, D. Laffan, C. Thomson and M. T. Williams, *Org. Biomol. Chem.*, 2006, **4**, 2337–2347.
- 4 G. P. Dado, E. A. Knaggs and M. J. Nepras, in *Kirk-Othmer Encyclopedia of Chemical Technology*, John Wiley & Sons, Inc., 2000, DOI: 10.1002/0471238961.1921120611140107.a01.pub2.
- 5 D. W. Roberts, *Org. Process Res. Dev.*, 2003, **7**, 172–184.
- 6 Y. Chen, Y. Su, F. Jiao and G. Chen, *RSC Adv.*, 2012, **2**, 5637–5644.
- 7 C. M. Suter and A. W. Weston, in *Organic Reactions*, John Wiley & Sons, Inc., 2004, vol. 3, pp. 141–197.
- 8 B. Galabov, D. Nalbantova, P. v. R. Schleyer and H. F. Schaefer, *Acc. Chem. Res.*, 2016, **49**, 1191–1199.
- 9 J. Kong, B. Galabov, G. Koleva, J.-J. Zou, H. F. Schaefer and P. v. R. Schleyer, *Angew. Chem., Int. Ed.*, 2011, **50**, 6809–6813.
- 10 B. Galabov, G. Koleva, S. Simova, B. Hadjieva, H. F. Schaefer and P. v. R. Schleyer, *Proc. Natl. Acad. Sci. U. S. A.*, 2014, **111**, 10067–10072.
- 11 G. Koleva, B. Galabov, B. Hadjieva, H. F. Schaefer and P. v. R. Schleyer, *Angew. Chem., Int. Ed.*, 2015, **54**, 14123–14127.
- 12 G. Koleva, B. Galabov, J. Kong, H. F. Schaefer and P. v. R. Schleyer, *J. Am. Chem. Soc.*, 2011, **133**, 19094–19101.
- 13 J. K. Bosscher and H. Cerfontain, *Recl. Trav. Chim. Pays-Bas*, 2010, **87**, 873–887.
- 14 J. K. Bosscher and H. Cerfontain, *Tetrahedron*, 1968, **24**, 6543–6555.
- 15 J. K. Bosscher and H. Cerfontain, *J. Chem. Soc. B*, 1968, 1524–1526, DOI: 10.1039/j29680001524.
- 16 H. Cerfontain, *Recl. Trav. Chim. Pays-Bas*, 2010, **104**, 153–165.
- 17 K. Lammertsma and H. Cerfontain, *J. Chem. Soc., Perkin Trans. 2*, 1980, 28–32, DOI: 10.1039/p29800000028.
- 18 H. Cerfontain and A. Koeberg-Telder, *Recl. Trav. Chim. Pays-Bas*, 2010, **89**, 569–574.
- 19 A. Koeberg-Telder and H. Cerfontain, *Recl. Trav. Chim. Pays-Bas*, 2010, **90**, 193–206.
- 20 J. O. Morley and D. W. Roberts, *J. Org. Chem.*, 1997, **62**, 7358–7363.
- 21 J. O. Morley, D. W. Roberts and S. P. Watson, *J. Chem. Soc., Perkin Trans. 2*, 2002, 538–544, DOI: 10.1039/b109338j.
- 22 F. H. Westheimer, *Chem. Rev.*, 1961, **61**, 265–273.
- 23 A. S. Morkovnik and A. R. Akopova, *Dokl. Chem.*, 2013, **450**, 122–126.
- 24 F. Kučera and J. Jančář, *Polym. Eng. Sci.*, 1998, **38**, 783–792.
- 25 T. M. Krygowski and M. K. Cyranski, *Phys. Chem. Chem. Phys.*, 2004, **6**, 249–255.
- 26 A. D. Becke, *Phys. Rev. A: At., Mol., Opt. Phys.*, 1988, **38**, 3098–3100.
- 27 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B*, 1988, **37**, 785–789.
- 28 S. Goedecker, M. Teter and J. Hutter, *Phys. Rev. B: Condens. Matter*, 1996, **54**, 1703–1710.
- 29 S. Grimme, J. Antony, S. Ehrlich and H. Krieg, *J. Chem. Phys.*, 2010, **132**, 154104.
- 30 B. Ensing and M. L. Klein, *Proc. Natl. Acad. Sci. U. S. A.*, 2005, **102**, 6755–6759.
- 31 G. Bussi, D. Donadio and M. Parrinello, *J. Chem. Phys.*, 2007, **126**, 014101.
- 32 J. Hutter, M. Iannuzzi, F. Schiffmann and J. VandeVondele, *Wiley Interdiscip. Rev.: Comput. Mol. Sci.*, 2014, **4**, 15–25.
- 33 A. Laio and F. L. Gervasio, *Rep. Prog. Phys.*, 2008, **71**, 126601.
- 34 A. Laio and M. Parrinello, *Proc. Natl. Acad. Sci. U. S. A.*, 2002, **99**, 12562–12566.
- 35 S. L. C. Moors, B. Brigou, D. Hertsen, B. Pinter, P. Geerlings, V. Van Speybroeck, S. Catak and F. De Proft, *J. Org. Chem.*, 2016, **81**, 1635–1644.
- 36 S. L. C. Moors, K. De Wispelaere, J. Van der Mynsbrugge, M. Waroquier and V. Van Speybroeck, *ACS Catal.*, 2013, **3**, 2556–2567.
- 37 J. Van der Mynsbrugge, S. L. C. Moors, K. De Wispelaere and V. Van Speybroeck, *ChemCatChem*, 2014, **6**, 1906–1918.
- 38 W. Humphrey, A. Dalke and K. Schulten, *J. Mol. Graphics*, 1996, **14**, 33–38, 27–38.

