Chlorination of oxybenzone and prediction of transformation products using non-equilibrium “forced” molecular dynamics

Harry Ridgwaya,*, John Orbellb, Stephen Grayc

*Corresponding author.

AquaMem Scientific Consultants, PO Box 251, Rodeo, NM 88056, USA, Tel. +1 575-557-7777/+1 575-654-2813; email: ridgway@vtc.net
bInstitute for Sustainable Industries & Liveable Cities, Victoria University (Werribee Campus), PO Box 14428, Melbourne 8001, Australia, Tel. +61 3 9919 8066; email: john.orbell@vu.edu.au
cInstitute for Sustainable Industries & Liveable Cities, Victoria University, PO Box 14428, Melbourne 8001, Australia, Tel. +61 3 9919 8097/+61 434 605 847; email: stephen.gray@vu.edu.au

Received 21 February 2018; Accepted 5 April 2018

ABSTRACT

Oxybenzone (OxBZ) is a member of the benzophenone family of sunscreen filters. Despite its broad biotoxicity, OxBZ and its transformation products (TPs) have been detected in coastal seawater, wastewater, and drinking water supplies around the globe. Unfortunately, little is known about the fate of OxBZ during water treatment, especially when exposed to chlorine or ozone disinfection. In this report, a hybrid quantum/molecular mechanics (QM/MM) algorithm is described that simulates reactions between OxBZ and hypochlorous acid (HOCl) in an aqueous background. The algorithm uses non-equilibrium forced molecular dynamics (N_eqFMD) to force bi-molecular collisions between HOCl and OxBZ over a population of random reactant conformations and orientations. A semiempirical QM method (RM1) is used to control and manage reactant collisions and compute molecular orbital rearrangements leading to bond dissociation and formation, while explicit water is treated classically as a perturbation on the electronic wave function. The N_eqFMD method qualitatively emulated key aspects of aromatic ring activation and substituent directing effects (including an –OCH₃ π–σ* hyperconjugation) associated with OxBZ pendant groups. When applied to the solvated HOCl–OxBZ system, a diversity of potential TPs was observed, several of which have been documented in prior experimental studies, including hydroxy radical (•OH), chlorine radical (•Cl), methyl radical (•CH₃), formaldehyde (H₂C=O; or its di-radical precursor •CH₂O•), chloromethane (CH₂Cl), mono-chloro-oxybenzone or its meta-Wheland intermediate, as well as open-ring and ring-scission products such as mono-chlorobenzene, para-chloromethoxyphenol (chloro-guaiacol), and 4-methoxy-catechol. Halogen addition to the methoxy ring was preceded by homolytic cleavage of HOCl to form an •OH radical + a transitory “semi-bound” •Cl radical. The later underwent nascent covalent bonding to an ortho-carbon via π-resonance delocalization to form a meta-stable carbocation intermediate expected to devolve by hydrogen abstraction to the monochlorinated ortho-substituted OxBZ product observed in experimental studies. Chlorine electron density (which increased over the collision trajectory) delocalized inversely with the mean partial charge of the methoxy ring, as well as the total charge of the OxBZ molecule, consistent with an SEAr (arenium ion) mechanism. Kinetically induced conformational strain in OxBZ was accompanied by resonance fluctuations in conjugated π-bonds comprising the methoxy ring, as revealed by coupled interatomic oscillations in sp² aromatic carbon partial charges. Finally, an unusual ipso hydroxylation/di-radical mechanism not previously reported is described for the formation of formaldehyde in HOCl–OxBZ collisions. Results indicate that N_eqFMD simulations can complement and extend experimental approaches to predict reaction outcomes and assist in the elucidation of reaction mechanisms involved in disinfection by-product formation.

Keywords: Benzophenone; Chlorine; Disinfection; Disinfection by-product; Molecular dynamics; Molecular modeling; Oxybenzone; QM/MM; Reaction mechanism; Wastewater; Water reuse; Water treatment

* Corresponding author.
1. Background

Oxybenzone (OxBZ; 2-hydroxy-4-methoxyphenyl-phenylmethanone) is a member of the chemically related benzophenone family of UV-blocking sunscreen filters [1,2]. Due to its widespread use globally, OxBZ and its transformation products (TPs) have become widely dispersed into the environment, and they have been detected in coastal seawater, municipal wastewater, groundwater, and drinking water supplies [3,4]. OxBZ that has entered coastal seawater ecosystems by a variety of pathways including municipal wastewater discharges has been implicated in coral reef toxicity and bleaching [5].

OxBZ is an estrogen-disrupting compound [6-8] and phototoxic agent with a number of TPs of health significance formed after exposure to sunlight [2,3,9]. Structurally, OxBZ is similar to the plastic additive bis-phenol-A (BPA), which is a known human carcinogen [10]. Like BPA, OxBZ and related compounds can form mutagenic and genotoxic metabolic by-products through photo-oxidative stress responses [9] or DNA-adduct formation involving bioactivation by cytochrome P450 enzymes [11,12]. OxBZ and other benzophenones also increase tumor metastasis potential by inducing proliferation in cancer cell lines via estrogen-mediated [13,14] and non-estrogenic mechanisms [15].

Little is known about the fate of OxBZ during water treatment, particularly with respect to TP formation following exposure to chlorine disinfection or advanced oxidation processes. Despite several recent studies reporting a range of toxic halolipides [3,4]. OxBZ that has entered coastal seawater ecosystems by a variety of pathways including municipal wastewater discharges has been implicated in coral reef toxicity and bleaching [5].

In an effort to better understand the potential fate of OxBZ and its TPs formed during chlorination disinfection in wastewater reuse and potable water treatment processes, we have developed and applied a hybrid quantum/molecular mechanics (QM/MM) non-equilibrium “forced” molecular dynamics (N_FMD) approach that can be used to rapidly screen for and identify TPs of interest in simulated chlorination reactions. The in silico method was employed to detect potentially stable first-generation TPs and transient intermediates formed during the reaction of hypochlorous acid (HOCl) with OxBZ in an aqueous (Newtonian) background. The algorithm robotically executes bi-/multi-molecular collisions between reactive species, for example, an oxidant molecule + a small organic compound, controlled by a semiempirical QM method able to estimate molecular orbital (MO) energies and rearrangements, which, in turn, emulate bond dissociation and formation events leading to TP formation. Semiempirical calculations are employed in the N_FMD method because they are computationally less expensive than ab initio or density functional theory (DFT) methods while at the same time providing reasonable accuracy in forecasting all-valence electron MO rearrangements. Suitably parameterized for the molecular species under consideration, semiempirical QM/MM calculations have been successfully applied to solvated inorganic systems [21], as well as enzyme-catalyzed biochemical reactions [22–25], and they have been shown to yield results that are frequently comparable with those obtained by ab initio and DFT approaches [26–29].

Nevertheless, given the theoretical limitations of semiempirical calculations, which do not explicitly address electron correlation, the N_FMD algorithm employed in this study is primarily intended as a first-principles computational screening procedure for predicting potential TP types and their statistical distributions. An interesting benefit of this approach is that it can be utilized to survey large numbers of potential TPs and identify those of special interest (e.g., biototoxic compounds), as well as the precise reactive collision conditions that led to their formation. The N_FMD simulations may be replayed to elucidate mechanistic details leading to TP formation using the same or additional levels of QM theory. Although the N_FMD method yields only initial TPs resulting from first-generation collisions, it was recently, successfully applied to identify several key reaction products and intermediates in gas-phase transformations of sabinene and benzene by ozone [30].

2. Methods

2.1. Modeling reactant compounds

HOCl and OxBZ models were built and optimized using a combination of standard Newtonian mechanics (MM) force fields and self-consistent field (SCF) post-Hartree–Fock QM calculation methods provided in Hyperchem Version 8.0.10 (Hypercube, Inc., Gainesville, FL, USA). Following the initial 2D sketching, the compounds were converted to approximate 3D structures using Hyperchem’s heuristic-based auto-conformer method, and geometry optimization via the Polak–Ribiere conjugate-gradient method by Hyperchem’s MM + classical force field using bond dipoles to initially describe system electrostatics. Each compound was then re-optimized in vacuo using a single ground-state SCF unrestricted Hartree–Fock (UHF) semiempirical calculation method, that is, RM1/UHF [31] and/or an ab initio calculation using the 6-31G* polarizable basis set (spin multiplicity = 3). The QM calculations were also used to assign partial atomic charges to all atoms in the systems. Optimization convergence criteria were typically set at 0.1–0.01 kcal/mol. Depending on the length of the calculation and available computational resources. Following the QM optimization calculations, the structures were further refined by performing several additional alternating cycles of semiempirical (or ab initio) calculations and optimization by the AMBER99 force field using standard Coulomb electrostatics, a switched cutoff and distance-dependent dielectric for the non-solvated conditions. Electrostatic and Van der Waals 1–4 scale factors were each set at their default values of 0.5 for the AMBER force field. In a later step described below, TIP3P water models [32] were used to solvate the reactant molecules in preparation for carrying out the N_FMD simulations.

2.2. Description of N_FMD algorithm and simulations protocols

Reactive molecular collisions were managed according to a novel N_FMD algorithm [30] that was scripted using the Tool Command Language and Toolkit (https://www.tcl.tk/software/tcltk/). Reactants were introduced into a non-periodic “nano-solvation cage” (NSC) comprising ~100–200 explicit TIP3P water molecules adjusted to a density of ~1.0 g/cc at standard temperature and pressure.
(STP; 25°C, 1 atmosphere) (Figs. 1 and 2). Because solvent can influence reaction outcomes by dipole interactions, by steric hindrance of prescribed reactant trajectories, or by directly entering the reaction pathway itself, the \( N_{\text{FMD}} \) algorithm incorporated provisions to allow water proximal to one or both reactants (e.g., within a 5-Å shell radius) to enter into the RM1-controlled reaction coordinate. Reactants were initially separated from one another by \(~8–12\) Å and independently, randomly rotated on their XYZ axes to scramble their pre-collision orientations. Because the OxBZ molecule was non-spheroïdal and asymmetric (axial ratio \( >2.0 \)), the maximum allowable rotation along any given axis was restricted to \( \pm 72° \). This amount of rotational freedom was sufficient to permit collisions with the HOCl molecule that were unbiased by the OxBZ reactant orientation but not so great as to leave any portion of the rotated OxBZ extending outside the NSC. Orientation bias was further suppressed by randomly targeting four atom centers distributed across the major axis of the OxBZ molecule. These atoms included 1, 3, 10, and 15 (see OxBZ atom numbering scheme, Fig. 5), and at the beginning of each collision, immediately following axial rotations, only one of these atoms was randomly selected as the collision target for HOCl. Typical adjustable variables and settings used in the \( N_{\text{FMD}} \) simulations are summarized in Table 1.

Following the introduction of the reactant species, the complete system (i.e., water + reactants) was energy minimized to \(<0.1\) kcal/mol.Å using the AMBER99 force field with a constant non-scaled dielectric (\( \varepsilon = 80 \)). Water molecules >5 Å distant from the HOCl center-of-mass were designated as “mechanical atoms,” which rendered them immobilized (i.e., frozen or tethered) and only accessible in the SCF (MO) calculations as an electrostatic background. The HOCl molecule was subsequently accelerated toward the target organic compound at the user-specified velocity (75–90 Å/ps in this study).

To help avoid excessive (and unstable) repulsive energy gradients over the reaction coordinate and reduce the likelihood of collapsing the SCF calculations, the trajectory was split into 4–6 discrete “quantum runs” (QRs) controlled by the RM1 method using a spin multiplicity of 3 and “Pulay mixing”, that is, direct inversion of the iterative subspace [33,34] to improve energy convergence. Each QR corresponded to one segment of the total collision trajectory. To further relieve excessive interaction potentials, reactants were briefly re-optimized immediately prior to each QR cycle using the RM1 method with a maximum of 10 iterations or a convergence value of 10 kcal/mol.Å. At the end of each QR cycle, the reacting species were “frozen” (tethered), and the water phase was re-optimized by a suitable Newtonian force field (typically, AMBER99) to permit adaptation of the TIP3P water to the updated nuclear positions of the reactants. Following water re-optimization, the aqueous phase was again tethered, and the reactants were allowed to continue (i.e., restart using stored velocities from the previous QR cycle) along their pre-established collision trajectories and adapt to the immobilized water background, which served as a perturbation on the RM1 electronic (i.e., clamped-nucleus) Hamiltonian (Helec). The collision velocity \( (V_r...V_q) \), where \( r = \) the QR cycle number, \( \) duration \( (D_r...D_q) \) of each...
QR cycle, and the MD time step (\(T_S, \ldots, T_n\)) were user-adjustable parameters and could be autonomously decremented by ~10%–15% per QR cycle until a specified minimum "floor" was reached as the reactants approached one another, ostensibly to allow more time for reactant reorientation and relaxation of repulsive non-bonded (Lennard–Jones + Coulomb) interactions. Standard initial velocity and QR duration in this study were usually set at ~75–90 Å/ps and ~0.05–0.075 ps, respectively, using a 0.5-fs MD time step, resulting in a total reaction time of ~0.25–0.5 ps. Reaction durations could vary depending on reactant chemistries (see below). Typically, several \(N_{\text{FMD}}\) test runs were performed to calibrate the time of reactant “impact” (RIT), i.e., the point where the system reached a maximum “activation” (strain) potential due to MO distortions. The RIT was typically adjusted to approximately coincide with the end of QRI or very early in the QR2 cycle. This QM/MM cycling process was continued for each collision until the last QR cycle was completed, at which time the molecular system and trajectory files were saved for subsequent system re-bonding and analysis (see below).

Choice of a semiempirical quantum-chemistry method (RM1) to dynamically emulate small-organic reactive collisions in an aqueous background was based primarily on the large size of the molecular systems and applicability to the atom types involved. The nano-solvation collision cages typically comprised one organic molecule (e.g., OxBZ) + 100–200 explicit TIP3P water molecules + a single oxidant species, for example, HOCl, H\(_2\)NCl, or O\(_2\). Such systems were too large to perform iterative MD simulations over pragmatic time scales using higher levels of QM theory, including small-basis-set ab initio or DFT protocols. The RM1 method was also well parameterized for the main group organic/inorganic atom types encountered in this study (C, H, O, N, P, S, and Cl). Although some systematic inaccuracies and unpredictability is to be expected in the use of RM1 (or other semiempirical methods) to emulate chemical bonding in small organic systems (namely due to the lack of explicit electron correlation), the goal here was to rapidly generate a distribution of potential collision outcomes that could be statistically searched for reasonable reaction patterns that could be subsequently compared with and verified against published experimental data. Finally, it is noteworthy that many modern semiempirical methods, including RM1 and others that have undergone empirical corrections (i.e., reparameterization) and formal extensions, provide improved conformational energies, heats of formation, and hydrogen-bond enthalpies that approach those of ab initio or DFT calculations [35–38].

### 2.3. System re-bonding algorithm and collision product identification

Following completion of all collisions in a series, a second custom algorithm (also encoded in Tcl/Tk) was invoked to re-bond the molecular system based on the magnitude of departure of calculated bond distances from their thermodynamically ideal (normal mode) values (Fig. 3). Bonding and non-bonding distance criteria were independently user-adjustable variables, and their values were established based on

<table>
<thead>
<tr>
<th>RXN parameter</th>
<th>Value</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Semiempirical QM method</td>
<td>RM1</td>
<td>Parameterized for all elements in this study</td>
</tr>
<tr>
<td>SCF convergence</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>SCF iterations</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>UHF or RHF</td>
<td>UHF</td>
<td>UHF = unrestricted Hartree–Fock</td>
</tr>
<tr>
<td>Spin multiplicity</td>
<td>3</td>
<td>1 or 3 for closed shell systems; 2 for open shell</td>
</tr>
<tr>
<td>Net system charge</td>
<td>0(e)</td>
<td>Total charge of molecular system</td>
</tr>
<tr>
<td>DIIS invoked</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>Mechanics force field</td>
<td>AMBER99</td>
<td></td>
</tr>
<tr>
<td>Assigned velocity (initial)</td>
<td>75–90 Å/ps</td>
<td>Velocity vector of HOCl at (t = 0) ps</td>
</tr>
<tr>
<td>Number of QR cycles</td>
<td>5</td>
<td>Water optimization between each QR cycle</td>
</tr>
<tr>
<td>MD runtime regression coefficient</td>
<td>0.1</td>
<td>Successive QR cycle runtimes reduced by 0.1</td>
</tr>
<tr>
<td>MD time step regression coeff</td>
<td>0.1</td>
<td>Successive QR cycle time steps reduced by 0.1</td>
</tr>
<tr>
<td>Initial MD time step</td>
<td>0.5 fs</td>
<td>Floor = 0.25 fs</td>
</tr>
<tr>
<td>Initial MD runtime per QR</td>
<td>0.075 ps</td>
<td>Floor = 0.025 ps</td>
</tr>
<tr>
<td>Dynamics temperature cutoff</td>
<td>6000 K</td>
<td>Temperature above which Rxn is terminated</td>
</tr>
<tr>
<td>Include water in QRI cycle</td>
<td>Yes</td>
<td>Included water treated by QM (can enter Rxn)</td>
</tr>
<tr>
<td>Water distance cutoff</td>
<td>5 Å</td>
<td>Distance from HOCl and/or OxBZ</td>
</tr>
<tr>
<td>Pre-collision convergence</td>
<td>10 kcal/mol.Å</td>
<td>Rough RM1 optimization prior to each QR cycle</td>
</tr>
<tr>
<td>Maximum OxBZ rotation</td>
<td>72°</td>
<td>Random independent rotation about XYZ axes</td>
</tr>
<tr>
<td>Maximum HOCl rotation</td>
<td>360°</td>
<td>Random independent rotation about XYZ axes</td>
</tr>
<tr>
<td>OxBZ targeted atoms</td>
<td>1, 3, 10, and 15</td>
<td>Random selection of target atom</td>
</tr>
<tr>
<td>Target atoms: other molecules</td>
<td>COM</td>
<td>COM = center of mass</td>
</tr>
<tr>
<td>Max rotation: other targets</td>
<td>360°</td>
<td>Random rotation about XYZ axes</td>
</tr>
</tbody>
</table>
prior RM1 studies (data not presented). Initial QR velocities and reaction durations depended to some extent on the chemical nature of the reacting species; thus, numerous $N_{\text{FMD}}$ test runs were conducted for each reactant pair to establish the initial $N_{\text{FMD}}$ run parameters. Based on their mass, charge, elemental composition, atomic coordination, and other factors, the re-bonding script automatically scanned, enumerated, and categorized all reaction products regardless of their chemical stability, for example, hydrogen-abstracted, halogen substituted species, arenium ions, and so forth. The combined (i.e., the $N_{\text{FMD}}$ + re-bonding) algorithms were iterated over multiple independent collision events (typically >1,000 in this study) for each reactant pair to generate a frequency distribution of first-generation reaction products. Finally, because all parameters for each reaction are saved, it is possible to reproduce any given reaction for later detailed analysis.

3. Results and discussion

3.1. Model compound properties

Semiempirical (RM1 method) and ab initio (6-31G**) basis set) calculated ground-state orbital energies, atomic partial charges, and other structural properties for the HOCl and OxBZ models used in this study are summarized in Figs. 4 and 5, respectively. The HOCl (RM1/UHF) model used in this study was in partial agreement with experimental and calculated electronic and structural data for this compound, as cited in the Computational Chemistry Comparison and Benchmark DataBase (http://cccbdb.nist.gov/exp2x.asp?casno = 7790923) maintained by the US National Institute of Standards and Technology (NIST, https://www.nist.gov/). The RM1-computed bond lengths (O–H and O–Cl) and H–O–Cl angle were in good agreement with experimental values reported by NIST, although the ab-initio (6-31G**) calculated heat of formation (~20.98 kcal/mol) was substantially less than the reported experimental value of ~74.5 kcal/mol. The ab-initio computed partial atomic charge on the oxygen...
atom of \(-0.592 \text{e}\) was significantly less than the RM1 computed charge of \(-0.176 \text{e}\). There was a similar discrepancy with respect to the assigned charges on the chlorine atom (RM1 = \(-0.04 \text{e}\); ab initio = \(+0.212 \text{e}\)). Despite the generally “softer” RM1 partial charges, the semiempirical method yielded a dipole moment of 1.507D, which was in reasonable agreement with the experimental value of 1.631D.

RM1/UHF calculations of OxBZ indicated a strongly, electronically polarized molecule (2.169D, calculated by RM1) with substantially greater electron density associated with the HOMO orbital of the methoxy-substituted ring (Fig. 5). Calculation of the of the HOMO orbital energy yielded a ground-state energy of about \(-8.26428 \text{kcal}\) for the alpha-spin electron and \(-8.92662 \text{kcal}\) for the beta-occupation electron. Most of the HOMO electron density (Ѱ) was distributed over atoms 3 (ipso to the –OCH\(_3\) group), 4 (ortho), and 8 (ortho) in the methoxy-substituted aromatic ring. Strong HOMO orbital density was also associated with oxygen atom 12 comprising the carbonyl bond linking the two aromatic rings.

3.2. Regioselectivity of oxybenzone halogen addition/substitution in oxybenzone

About 25% of N\(_{\text{eq}}\)FMD collisions resulted in chlorine addition or substitution TPs, ostensibly resulting from electrophilic attack at preferred atom centers associated with the conjoined aromatic rings (Figs. 6 and 7). Halogen addition/substitution occurrence was highly asymmetrically distributed over the OxBZ target molecule with an approximate twofold greater frequency of halogen addition observed in the electron-rich methoxy ring. This observation is in accordance with the higher observed \Ѱ\(^2\) HOMO density of the methoxy ring compared with the unsubstituted benzene ring. Enhanced methoxy-ring reactivity also correlates favorably with the expected influences of the –OH and –OCH\(_3\) activating (i.e., electron-donating) groups, both of which are moderately strong ortho-/para-directing substituents.

In contrast to ring activation by the hydroxy and methoxy groups, the electron-withdrawing carbonyl bond (\(-\text{C}=\text{O}\)) adjoining the two aromatic rings is a moderate deactivateing group with a meta-directing influence, which probably contributed to the overall lower reactivity of the benzene ring. The elevated chlorine addition frequency noted at atom 14 in the benzene ring also appears to be consistent with the meta-directing influence of the carbonyl group. Based on these observations, the \(N_{\text{eq}}\)FMD method appears to be in reasonable qualitative agreement with what would be expected based on the known activating and deactivating influences of the OxBZ substituent groups.

Both the hydroxy and methoxy substituents would be expected to complement one another to possibly reinforce their ortho-/para-directing effects and increase electrophilic addition/substitution at methoxy-ring atoms 4, 6, and 8 (Fig. 6). In contrast, repressed halogen attack would be expected at methoxy-ring atoms 3, 5, and 7. While the expected enhancement of chlorine addition was indeed observed at atoms 4 and 8, little or no enhancement was recorded at atom 6 located para to the methoxy group and ortho to the –OH group. A plausible explanation for reduced halogen addition at atom 6 is that reactivity at this locus may have been partially curtailed by the meta-directing/deactivating influence of the adjacent carbonyl bond.

Interestingly, a heightened rate of chlorine addition was observed at atom 3 in the ipso position relative to the methoxy ring.
methoxy group, which controverted the positive partial atomic charge at this location (+0.248e). While it is possible that the enhanced ipso halogen addition was artefactual in nature (e.g., resulting from inadequate handling by RM1 of the oxygen lone pairs in the alkoxy –OCH₃ group), this result was nevertheless correlated with elevated HOMO ψ₂ electron density distribution at this location (inset). The carbonyl bond likely contributed to deactivation of: (1) the unsubstituted benzene ring and (2) methoxy carbon atom 6 located ortho to the –OH group and para to the methoxy group.

Ipso halogen addition may have also been positively influenced (i.e., promoted) by sterically induced underestimation of hyperconjugative effects involving the OxBZ methoxy group. Hyperconjugation is known to occur in methoxy groups when favorable alignment occurs between the π-orbital of an oxygen lone pair and one of the adjoining C–H σ* anti-bonding orbitals [39,40]. Under normal circumstances, hyperconjugation deferentially facilitates delocalization of electron density from the π-orbital of the oxygen lone pair into the methyl moiety, which effectively shunts charge density away from the aromatic ring rendering it less susceptible to electrophilic attack, particularly at the ipso position. However, hyperconjugation depends critically on the conformational space occupied by the methoxy group [39], and in the relatively high-velocity collision dynamics imposed on the reacting species in this study, the methoxy conformations were oftentimes highly strained sterically. It is speculated that such departure from an optimized –OCH₃ geometry over the collision trajectories may have led to severe misalignment of the –O–C–H– σ*–π bond. This, in turn, could lead to greater electron density being delocalized into the aromatic ring at the ipso carbon, which would ultimately be manifested as an enhancement of electrophilic attack at this location.

Ipso additions/substitutions typically occur under specialized circumstances, such as certain nitration reactions [41].

**Fig. 6.** Frequency of chlorine addition/substitution in the Nₑq FMD simulation over the 29 atoms comprising OxBZ. Note: the methoxy ring (blue bars) had about 2× more halogen additions compared with the benzene ring (green bars). Areas of blue circles approximate the degree of halogen addition at those atom locations. Although halogen addition at the methoxy ipso position (C3) was unexpectedly high, it was correlated with elevated HOMO ψ₂ electron density distribution at this position (inset). The carbonyl bond likely contributed to deactivation of: (1) the unsubstituted benzene ring and (2) methoxy carbon atom 6 located ortho to the –OH group and para to the methoxy group.

**Fig. 7.** Frequencies of chlorine addition to aromatic ring atoms in anisole (1,100 collisions), phenol (1,150 collisions) and aniline (1,523 collisions) in separate Nₑq FMD simulations. While there was a clear trend toward ortho-addition in all three compounds, halogen addition at the para-carbons was somewhat repressed. In addition, there was greater than expected chlorine addition at one of the meta-carbon atoms in phenol. Areas of circles over ring atoms approximate relative frequency of halogen addition at those locations.
or enzyme-catalyzed hydroxylation of para-cresol and related aromatic phenols by cytochrome P450 [42]. In the later situation, substituent elimination is accompanied with ipso substitution by an oxygen atom of the active species, and there is a requirement for a hydroxyl group at the para-position to the leaving group. Ipso-mediated hydroxylation reactions have also been reported in the enzyme-catalyzed biodegradation of BPA (which has a bi-phenolic structure resembling that of OxBZ) and nonylphenols by *Sphingomonas* [43]. The authors reported that subsequent to ipso hydroxylation, various molecular rearrangements took place consistent with the formation of transient pseudo-stabilized carbocation intermediates. A similar mechanism involving ipso ring hydroxylation at the site of the substituent has been described by Gabriel et al. [44]. Fischer and Henderson [45] reported bromination of para-Cresol via a mechanism involving halogen addition ipso to the 2-methyl group followed by migration of the bromine to the adjacent 3-position.

Similar ipso substitution mechanisms to those described above were observed in the N\textsubscript{FMD} simulations conducted in this study (see section 4.7). For example, reactions 115 and 885 resulted in the release of a formaldehyde di-radical (\(\text{CH}_3\text{O}\)) leaving group following ipso hydroxylation or chlorination, respectively. The former reaction yielded the stable product di-hydroxybenzophenone while the later resulted in formation of chloro-hydroxybenzophenone, neither of which has been previously reported in experimental studies. Similarly, at least five reactions (Nos. 95, 225, 577, 681, and 976) involved hydroxylation at the methoxy ipso carbon coupled with cleavage of the O–CH\(_2\) bond to yield a chloromethane product + di-hydroxybenzophenone.

Formation of ipso halogenated TPs in OxBZ chlorination reactions has not been reported by others [16–20], but this may have been due to the transient nature of such quasi-stable intermediates and the difficulty of their detection. Moreover, because only first-generation N\textsubscript{FMD} collision products were captured in this study, it is unknown if the carbocation ipso substitutents might have undergone subsequent structural rearrangements to yield energetically more favorable iso-meric forms via resonance delocalization processes.

### 3.3. Halogen addition/substitution in phenol, anisole, and aniline

To further evaluate the ability of the N\textsubscript{FMD} method to emulate the activating/directing influences of specific OxBZ substituent groups, the N\textsubscript{FMD} method was applied separately to phenol (1,100 collisions) and anisole (1,150 collisions), whose single hydroxy and methoxy substituents, respectively, are known ring-activating (electron-donating) and ortho-/para-directing groups. As an additional test of the ability of N\textsubscript{FMD} simulations to differentiate between the direct/resolving effects of certain substituent groups (namely, –NH\(_2\)), a third N\textsubscript{FMD} run was included for aniline (1,525 collisions).

A summary comparison of the results from these simulations is presented in Fig. 7. Inspection of the data suggest the following conclusions: (1) halogen addition was in general more likely (by about twofold) for aniline and anisole as it was for phenol under similar or identical N\textsubscript{FMD} simulation conditions, and (2) for all three compounds, ortho-addition of chlorine was mostly favored over addition at the ipso-, para-, and meta-positions. One exception to the later generalization was a comparatively high halogen addition frequency observed for the first meta-position of phenol (atom 1).

On strictly quantitative grounds, the later observation stands in contrast to the experimental findings of Ogata et al. [46] who reported that the ortho/para ratio in the chlorination of phenol and anisole with aqueous NaOCl was about 0.64 and 0.66, respectively, under acidic conditions. However, these authors also reported a strong preference for ortho-substitution of phenol at pH 10 (ortho/para ratio = 4.3), an observation that is clearly more closely aligned with results from the current N\textsubscript{FMD} simulations. Because the simulations did not explicitly account for pH effects and different forms of chlorine were used in the two studies, it is difficult to rationalize the current modeling results with the findings of Ogata et al. [46].

The observed higher reactivity of aniline compared with phenol and anisole in the N\textsubscript{FMD} simulations was in good qualitative agreement with the known experimental behavior of these compounds with regard to their halogen (electrophilic) reactivity. The finding that aniline and anisole exhibited a greater frequency of halogen addition than phenol in the N\textsubscript{FMD} simulations suggests the methoxy and primary amine groups, respectively, were more activating than the hydroxy group. In practice, however, phenol is usually found to be slightly more reactive with halogen species than is anisole. The reduced halogen reactivity of aniline (in practice) presumably results from stabilizing hyperconjugation between the methoxy oxygen π-lone pair(s) and the antibonding σ*–MO of the aligned –C–H bond [40,47]. In anisole a negative hyperconjugative effect tends to delocalize electron density away from an oxygen lone pair (and into the –C–H σ*–MO) that would otherwise be contributed to the aromatic ring through a π–π resonance mechanism. In phenol, hyperconjugation is absent leaving more electron density available at the oxygen lone pair(s) to be donated to the ring via resonance charge transfer.

It is concluded that some incongruity exists between the N\textsubscript{FMD} modeling results that indicate a somewhat reduced activating influence of the hydroxy substituent in phenol compared with the overall higher reactivities of anisole and aniline. Whereas the basis for this discrepancy is unclear at this time, it is probably not related to an inability of the RM1 method to model hyperconjugation effects in anisole, since these processes are computed with reasonable confidence by similar “neglect of diatomic differential overlap” (NDDO) semiempirical methods, including PM3 [48]. As described above for OxBZ, a likely source of error in anisole might be excessive (collision-induced) bond vibrational fluctuations during simulations causing poor MO alignments and reduced –OCH\(_3\) hyperconjugation, resulting in greater aromatic electron density and a corresponding rise in the probability of halogen attack [40].

The modeling data also indicated that halogen addition preferences, that is, substituent directing effects, were milder than what might be expected based on experimental reactions between halogens and substituted aromatic compounds [49]. Indeed some repression of halogen addition was evident in the para-positions of all three compounds, and addition at one of the phenol meta-positions (atom 1)
was inordinately high for unknown reasons. Thus, it may be concluded that overall the \( N_\text{FMD} \) algorithm, as currently implemented, is in partial qualitative accord with expectations based on experimental observations, but tends to discriminate less strongly than would be observed in practice between the directing effects of certain substituent groups, for example, the methoxy group.

The divergence from experimental observations may also be attributed in part to relatively “soft” partial atomic charges computed by the RM1/UHF method (e.g., compared with ab initio computed charges; see section 4.1), which might diminish polarization effects and dipole interactions with the oxidizing species resulting in weaker directing influences of some substituent groups. A second possibility is that the collision velocities used in the current study were sufficiently high as to partially dampen (i.e., swamp) the effects of higher activation energies associated with less preferred sites of halogen attack. Yet a third possibility could be related to the rather extreme conformational changes that occur in the oxidant and target species during the molecular collisions. Large conformational fluctuations resulting from kinetic transference from the oxidant species would tend to alter and destabilize ground-state MO arrangements in such a manner as to partially scramble and thus weaken the discriminatory influences of substituent groups. Additional studies will need to be undertaken to address these and some other possibilities.

### 3.4. Occurrence of OxBZ transformation products and intermediates

Despite the limitations described above in the ability of the \( N_\text{FMD} \) method to precisely emulate and strongly distinguish between the directing/activating influence of some substituent groups, the algorithm successfully predicted a range of experimentally verified first-generation TPs from a total of 1,150 reactive collisions between HOCl and OxBZ dissolved in a TIP3P water NSC. These results, which are summarized in Fig. 8, indicated that about 15.3% of the 1,150 collisions resulted in “failed reactions”, that is, no TPs were observed, and the parent reactants remained unchanged. Such failed reactions were principally caused by poor or unfavorable reactant geometries and orientations over the collision trajectories, which effectively nullified any MO rearrangements and subsequent TP formation. Nearly 4% of collisions resulted in “unstable reactions” in which the \( N_\text{FMD} \) calculation was automatically terminated when a user-adjustable thermal threshold criterion (Table 1) was exceeded due to an excessive energy gradient for the molecular system.

Among the largest product categories observed was the formation of free hydroxyl (‘\( \cdot \)OH; \( \approx 66\% \) of collisions) and chlorine (‘\( \cdot \)Cl; \( \approx 49\% \) of collisions) radicals resulting from homolytic dissociation of the HOCl molecule as it approached the OxBZ target compound. In some collisions, chlorine dissociation was observed to occur prior to the point of closest approach to the OxBZ molecule. In these instances, chlorine dissociation was typically initiated by interaction of HOCl with proximal water molecules lying in the OxBZ collision path that were also treated quantum mechanically in the QRI cycle.

Chlorine and ‘\( \cdot \)OH radical formation was not entirely unexpected since such species are generally produced by homolytic bond scission between two atoms of similar electronegativity (en), in this instance between oxygen (en = 3.44) and chlorine (en = 3.16). In addition the HO–\( \cdot \)Cl bond energy is known to be comparably labile (about 48 kcal/mol calculated by RM1 in this study), and it is therefore especially vulnerable to homolytic splitting under the collision conditions used in the \( N_\text{FMD} \) simulations (Table 1).

A range of first-generation ring-halogenated (~24.8% of collisions) and hydrogen-abstracted (~13.2% of collisions) TPs were observed, including a high frequency of Cl addition at the ortho- and ipso-positions relative to the –OCH\(_3\) methoxy group, as described above. Mono-halogenated single-ring products resulting from carbonyl bond cleavage, such as mono-chlorobenzene (rxn238) and para-chloromethoxyphenol (rxn2x), were also observed in about 3.1% of collisions. The formation of HCl was also frequently observed in the simulations and typically resulted from the reaction of a hydrogen atom abstracted from the OxBZ parent with a free chlorine radical following homolytic cleavage of the HO–Cl bond. Hydrochloric acid as well as similar or identical mono-halogenated TP categories and chlorinated ring-scission products have been observed in OxBZ freshwater chlorination experiments reported by Negreira et al. [20] and more recently by Zhang et al. [16], as well as in seawater experiments reported by Manasfi et al. [50].

The \( N_\text{FMD} \) simulations revealed a number of other TPs that have not been reported or only marginally documented
in earlier experimental chlorination studies of OxBZ. Some of these products included relatively rare ring opening reactions (~0.3%), hydroxylation (–OH) additions/substitutions (~8.5%), formaldehyde or formaldehyde di-radical precursor (•CH₂O•) formation (~0.3%), mono-chloromethane (CH₃Cl; ~3.1%), and formation of methyl (•CH₂; ~1.3%) and methoxide radicals (•CH₃O; ~0.3%; Fig. 9). The observation of early radical formation suggests that some second-generation TPs (with multiple halogen substitutions) could be readily formed by their further reaction with first-generation products and intermediates.

The significant chemical diversity of TPs and intermediates observed in the \( N_{eq}FMD \) simulations appears consistent with a scenario in which multiple potential HOCl oxidation routes lead to TP formation. Some of the key routes revealed by the \( N_{eq}FMD \) simulations involve: (1) carbonyl bond scission reactions that may or may not be combined with ring halogenation, (2) methoxy group elimination leading to formaldehyde or mono-chloromethane formation, (3) facile abstraction of hydrogen from the –OH group, (4) a variety of ring opening reactions, and (5) ring hydroxylation, for example, resulting in formation of 4-methoxycatechol (Fig. 9).

In contrast, experimental studies have generally described a somewhat more limited range of TP formation pathways based on the detection and quantification of a few stable or pseudo-stable reaction end products. However, the experimentally observable end products (and intermediates) may not adequately reflect the potential richness of TP formation mechanisms that is suggested by the present modeling study. A possible informative strategy for future experimental research would be to specifically analyze for some of the potential TPs that have been revealed in the current \( N_{eq}FMD \) simulations effort.

3.5. Detailed analysis of chlorine ortho-addition/carbocation formation

As indicated above, Zhang et al. [16] reported a comparatively short-lived (<60 min) semistable monochlorinated OxBZ product with a single halogen substitution in the para-position relative to the –OH side group (ortho relative to the –OCH₃ methoxy group). This same halogenated TP, in its meta-semistable carbocation intermediate form, was also observed in the \( N_{eq}FMD \) simulations. In an effort to shed light on the possible mechanism of this reaction outcome, a more detailed investigation was undertaken of one \( N_{eq}FMD \) simulation that resulted in electrophilic chlorine addition to the methoxy-ring C8 carbon atom + release of a free •OH radical. This particular reaction outcome was found to be reproducible across >10 independent \( N_{eq}FMD \) trials given the same starting positions and (rotational) orientations of the HOCl and OxBZ molecules (data not presented). An analysis of the time course of the reaction, which is depicted in Fig. 10, revealed high initial system strain energy (gradient) as the HOCl approached and underwent a reactive collision with the OxBZ target.
compound in the vicinity of the methoxy ring. As indicated in the right-hand panel of Fig. 10, elongation of the HO–Cl bond first became apparent at about 0.03 ps and continued for the duration of the reaction coordinate (tend = 0.3 ps).

Actual dissociation of the HO–Cl bond occurred between an Neq FMD elapsed time of about 0.04–0.05 ps (Figs. 10 and 11) and an HO–Cl bond distance of about 1.5–1.7 Å. This bond dissociation event was immediately preceded by a brief collision-induced compression (of about 0.2 Å) that occurred between 0.02 and 0.04 ps. The HO–Cl bond compression minimum at about 0.03 ps corresponded to the maximum system energy gradient and represents a significant portion of the activation energy for the C8–Cl σ–π conjugation leading to halogen addition to the aromatic ring. This strain energy was abruptly released between about 0.03–0.04 ps as the HO–Cl bond underwent dissociation. The HO–Cl bond dissociation also corresponded to the point of closest approach of the chlorine atom to atom C8 in the methoxy ring, which was located in the para-position relative to the –OH side group.

Dissociation of the HO–Cl bond was also accompanied by release of an •OH free radical between about 0.05–0.075 ps, after which time the system continued to undergo rapid but comparatively minor conformational changes (i.e., steric relaxation) due to prior conveyance of kinetic energy from the HOCl molecule to the OxBZ target compound. Conformational enthalpies (ΔH) calculated by RM1/UHF indicated net downhill thermodynamics for the formation of a stable halogenated ortho-substituted product, assuming eventual loss of the unstable arenium proton in the ortho-position to the water phase (Fig. 12).

3.6. Electron delocalization and resonance charge transfer effects during chlorine addition

In halogen-mediated electrophilic addition/substitution reactions, the partial atomic charge of the halogen species is expected to deepen (i.e., become more negative) as a result of resonance electron withdrawal from the donor atom(s). As shown in Fig. 13(A), such a charge deepening effect of the chlorine atom was in fact observed in the Neq FMD simulations, and this deepening effect was inversely correlated with the separation distance between the chlorine atom and carbon atom C8 in the ortho-position of the methoxy aromatic ring.

A similar inverse relationship was observed between the chlorine partial atomic charge and (1) the charge averaged over all of the methoxy-ring carbon atoms (Fig. 13(B)) and (2) the total charge of the OxBZ target molecule (Figs. 13(C) and (D)). These observations are consonant with a classical resonance charge transfer mechanism in which the approaching chlorine atom withdraws electron density primarily from the aromatic methoxy ring during collision leading to Cl–C8 covalent bond formation. The sudden increase in the total electronic charge of the OxBZ target molecule is tantamount to nascent carbocation intermediate formation as chlorine is added to the methoxy ring.

As indicated above, the HO–Cl homolytic bond cleavage in the HOCl approach region (Fig. 12) resulted in formation of •Cl and •OH radicals. However, as shown in Fig. 14, the former was exceedingly ephemeral in nature existing only in a “semi-bound” state around its neutral charge point between about 0.04 ps. It then underwent immediate hybridization with the methoxy-ring HOMO orbital at C8 to form the meta-stable ortho-addition carbocation intermediate.

Key aspects of this proposed mechanism are summarized in Fig. 15. Semiempirical RM1/UHF calculations revealed that the major proportion of charge resonance transfer (Δ376e) occurred at the C8 ortho-carbon atom leaving it with a positive partial charge (+0.104e) compared with its original negative charge of –0.272e in the unreacted OxBZ parent. A smaller but still significant charge transfer also took place at the C8 terminal hydrogen whose partial charge transitioned...
Fig. 11. Early part (<0.075 ps) of the reaction trajectory yielding the mono-chlorinated pseudo-stable OxBZ arenium intermediate. The system gradient (GRAD) increased then decreased during the initial 0.01 ps as HOCl approached OxBZ. HO–Cl bond compression between about 0.02 and 0.03 ps (green down arrows) resulted in a sharp spike in strain energy, which was followed by an abrupt decline as the HO–Cl bond underwent stretching and dissociation (red up arrows). This was coupled with nascent Cl–C8 covalent bond formation. Vertical blue bar indicates the point of maximum system strain (GRAD) occurring at the closest approach of the Cl atom to the C8 carbon.

Fig. 12. Energies calculated by RM1/UHF for the geometry-optimized parent OxBZ compound, the meta-stable carbocation intermediate, and the presumptive ortho-substituted product that has been experimentally verified by Negreira et al. [20]. Although included in the energy calculation, the HOCl molecule is hidden (for clarity) in the image of the parent compound. Similarly, the free *OH radical is not shown for the intermediate. The calculation for the ortho-product included a single water molecule (*H + *OH). In all calculations, the HOCL and *OH molecules were present in their original locations following the N_FMD simulation.
from +0.139e to +0.165e, a decrease of about 0.026e (Fig. 15). Though not large in magnitude, it is speculated that electron density was transferred from the terminal hydrogen to chlorine by a delocalization mechanism, possibly involving a direct “through-space” inductive (dipole) effect between the hydrogen and chlorine atom [51], or by a relatively weak positive σ–π hyperconjugation mechanism involving a –C–H– sigma-bond alignment with the C8 π bond [39]. Regardless of the delocalization mechanism involved, both C8 and its terminal hydrogen acquired enriched positive charge that would be expected to further destabilize the –C–H– bond, eventually resulting in hydrogen abstraction facilitated by Coulomb repulsion.

Inspection of the HOMO orbital maps presented in Fig. 16 indicates the dissociated chlorine atom underwent rapid hybridization with the OxBZ alpha HOMO orbital with concomitant release of the •OH radical. Moreover, the HOMO orbital energy of OxBZ over the simulation trajectory was directly correlated with a corresponding depression in the chlorine partial charge, an observation that is consistent with a thermodynamically favorable MO rearrangement associated with formation of a nascent meta-stable bond between chlorine and C8 in the methoxy ring (Fig. 16, graph inset).

Interatomic π-resonance charge transfer (i.e., electron density delocalization) effects were also recorded between neighboring sp² carbon atoms comprising the methoxy aromatic ring over the course of the $N_{eq}$FMD collision trajectory (Figs. 16 and 17). Neighboring ring-carbon atoms were observed to undergo opposing (i.e., inversely correlated) fluctuations in their RM1/UHF-computed partial electronic charges over the reaction coordinate. For example, an approximate linear direct correlation was observed between the C8 partial atomic charge and the C3–C8 conjugated-bond (stretch) distance (Fig. 17), with shorter C3–C8 bond distances being associated
Fig. 14. Charge transfer coupling during HO–Cl bond dissociation and Cl–C8 bond formation. Dissociation of the HO–Cl bond was accompanied by transitory existence of a “bound” •Cl radical (red circle) at its charge null point. The Cl and C8 atoms eventually assume equilibrium partial charges (negative for Cl; positive for C8) consistent with Cl–C8 bond covalency. Formation of the •Cl radical was accompanied by release of a free •OH radical. The •Cl radical rapidly undergoes hybridization with atom C8 and the region of the OxBZ (HOMO) orbital associated with the methoxy ring (Fig. 15). Note that the charge on the •OH species fluctuates briefly (hysteresis) and mirrors that of the Cl atom, then transitions to 0e by about 0.08 ps. Inset images show the reaction progress at specified times in ps. Oxygen = red; chlorine = violet; carbon = brown; and hydrogen = gray.

Fig. 15. Details of the proposed mechanisms of electrophilic attack at carbon C8 in the methoxy ring of OxBZ to form the meta-stable carbocation ortho-intermediate shown on the right. Atomic partial charges calculated by RM1/UHF infer charge resonance delocalization (eres) from C8 (thick green arrow) and minor transfer from the C8 terminal hydrogen (thin green and blue arrows), which also becomes more positive. It is unclear (?) if minor charge transfer from the terminal hydrogen could occur by direct inductive polarization (ein) with chlorine (thin green arrow) or by a –C–H– sigma-bond hyperconjugation (ehypr) mechanism (blue arrow). Evidence for hyperconjugation at the –OCH3 moiety is indicated by reduction of oxygen partial charge and gain by the ipso ring carbon (C3). Note that resonance donation of electron density from the hydroxy oxygen (O12) to the ring (C5), as well as from the ring (C6) to the carbonyl bond (C10) is also consonant with changes in neighbor-atom partial charges. Thick arrows indicate gain (red) or loss (green) of electron density at that locus.
with deeper (more negative) C8 partial charge. This relationship suggests that interatomic conjugated π-bond electron density delocalization was tightly coupled to conformational oscillations in the ring –C–C– bond lengths (and their accompanying MO distortions) arising from kinetic transfer into the OxBZ molecule by prior collision with HOCl. As indicated in Fig. 18, such coupled π-bond resonance oscillations were observed for all neighboring pairs of sp² aromatic carbon atoms comprising the methoxy ring. Partial charges associated with alternating atom pairs, for example, 3 and 5 or 4 and 6, were directly correlated, whereas charges associated with closest neighbor pairs were inversely correlated as a function of $N_{eq}$FMD elapsed time.

3.7. Formaldehyde formation by a di-radical mechanism involving ipso hydroxylation

As indicated above (Fig. 8), a formaldehyde “precursor” was formed in about 0.3% of the $N_{eq}$FMD simulations. This product was of special interest, not only because it is a known human carcinogen but also because its formation evidently involved cleavage of the methoxy group; and its mechanism of formation has not been previously described. Based on playback and analysis of Rxn115, which typified this pathway, the main steps involved in formaldehyde formation are proposed in Figs. 19(A)–(D). These steps may be summarized as follows:

- Dissociation of HOCl during its final approach to OxBZ in the vicinity of the methoxy group yielding transitory $\cdot$OH + $\cdot$Cl radicals. Reaction “success” was obligatorily correlated with simultaneous arrival of: (a) the chlorine atom proximal to the methyl hydrogen to be abstracted and (b) the $\cdot$OH radical proximal to the methoxy group ipso carbon atom (C8). In this orientation, the HOCl molecule was already largely, if not completely, dissociated and positioned with its major (HO–Cl) bond axis more-or-less parallel to that of the O–CH$_3$ methoxy bond (see inset, Fig. 19(B)).
Halogen-induced abstraction of one hydrogen atom from the terminal methyl group that rapidly combined with Cl to form one molecule of HCl.

Concomitant with the –CH$_3$ hydrogen abstraction, the OH radical hydroxylated the C8 ipso carbon atom (relative to the methoxy group) causing homolytic cleavage of the Caro–O bond and release of an unstable CH$_2$O di-radical species.

The di-radical undergoes resonance rearrangement as shown in Fig. 19(C) yielding the resonance-stabilized formaldehyde TP.

It is noteworthy that the reactive collisions resulting in formaldehyde formation were exceedingly sensitive to the reactant starting geometries and orientations, as well as the initial optimized positions of reactive TIP3P water molecules, which in many collisions tended to briefly chaperon (or otherwise interfere with) the HOCl oxidant along its trajectory. As indicated in Fig. 19(D), very small deviations (<1.5°) in the initial (t = 0 ps) XYZ axial rotations of the OxBZ (or HOCl) molecule resulted in unproductive reactions, that is, collisions yielding no formaldehyde formation. On the other hand, by constraining the initial HOCl and OxBZ rotational orientations to predetermined ideal values known to result in formaldehyde formation, it was possible to increase the success rate of formaldehyde formation from about 0.3% to nearly 100% (data not presented).

Finally, the velocity of collision between HOCl and OxBZ were also critical in determining the reaction outcomes and whether formaldehyde would be formed (Fig. 19(E)). Formaldehyde was only formed within a relatively narrow range of collision velocities (about 90–100 Å/ps). At low collision velocities (e.g., less than about 50 Å/ps), the trajectory of HOCl could be slightly altered due to non-bond interactions with proximal water molecules, which tended to exert a relatively greater influence on HOCl transit and orientation leading to unproductive collisions. Lower velocities also were generally unable to overcome non-bond repulsive forces between HOCl and OxBZ, which was a requirement for successful MO hybridizations to occur.

4. Concluding remarks

Results presented in this paper demonstrate that $N_{eq}$FMD reaction simulations can be applied to the detailed analysis of disinfection by-product (DBP) formation reactions using the sunscreen filter OxBZ as an example. The $N_{eq}$FMD simulation algorithm is distinguished from other similar approaches by the inclusion of a Newtonian background of explicit water (TIP3P models in this case) approximating the density of normal liquid water. Additionally, water molecules within a user-specified cutoff distance from the reacting species may be included in the reaction coordinate, a feature that allows water to undergo MO rearrangements and interactions with the other reacting species.

Another unique feature of the $N_{eq}$FMD algorithm is that reaction trajectories can be partitioned into a series of discrete steps or cycles (so-called QRs) that allow judicious management of the collision dynamics. For example, as the...
Reactant molecules undergo a forced collision, key parameters, for example, the MD time step, QR duration (i.e., the QR cycle time), velocity vector, number of QR cycles, and so forth, can be independently and automatically adjusted as deemed appropriate after each QR cycle to minimize excessive (repulsive) strain energies, thereby increasing the likelihood of reaction success and minimizing the possibility of calculation failure (by the MD time integrator) due to large energy gradients. Another distinctive aspect of the $N_{eq}$FMD algorithm is the ability to randomly or systematically modify reactant conformations and orientations at the beginning of each new reaction in a series. Finally, the algorithm is designed to permit the water field to semicontinuously adapt (via MM energy minimizations) to the updated nuclear positions of the reactants following each QR cycle, and vice versa.

One potential concern regarding the $N_{eq}$FMD algorithm is the use of valence-shell semiempirical QM methods (RM1 in this study) for computing MO reorganizations to describe bond dissociation and formation. However, the use of semiempirical methods is rationalized in this study on the basis of: (1) much greater computational efficiency compared with ab initio and DFT approaches; (2) reasonable accuracy for most organic compounds in terms of computed geometries, heats of formation, dipole moments, and so on, when compared with ab initio and DFT calculations [35,36,38,52,53]; and (3) a history of successful applications of semiempirical QM/MM hybrid methods for modeling enzyme catalysis [24] and many other chemical processes [22]. Significantly, the use of core-core interaction terms and other empirical parameters and formal approximations in RM1 and related NDDO methods partially compensates for the lack of explicit treatment of electron correlation [35].

In the current investigation, RM1/UHF calculations as implemented in the $N_{eq}$FMD algorithm qualitatively emulated certain key aspects of substituent directing and activating effects. In particular, the $N_{eq}$FMD algorithm forecasts the
activating influence of the electron donating –OH and –OCH$_3$ side groups on the OxBZ methoxy aromatic ring, as well as the deactivating influence of the central carbonyl linkage on the unsubstituted benzene ring. Furthermore, the expected ortho-directing effect of the –OH and –OCH$_3$ groups were clearly observed in phenol and anisole, respectively. Although para direction was weak for these test compounds, it was nevertheless evident to a minor degree (Fig. 7). The expected para-directing influence of the methoxy group at atom 6, as well as the expected ortho-directing effect of the –OH group at this same position, was likely suppressed to a significant degree by the deactivating and meta-directing behavior of the adjacent carbonyl bond. For the test compound aniline, the primary amine (–NH$_2$) exerted pronounced ortho-para-direction, as would be anticipated for this donating group.

Although not quantitatively evaluated in this study, elevated halogen addition at the methoxy group ipso position of OxBZ (atom C3) may have been caused by: (1) underestimation of –O–C–H hyperconjugation effects by RM1 at this position, which would effectively delocalize more electron density from oxygen lone pairs to the ring or (2) poor alignment of the –O–C–H π–σ* bond due to conformational fluctuations at or around this substituent group. Hyperconjugation is known to be extremely sensitive to steric effects [39,40]. Anomalous chlorine addition at the meta-position of phenol was unforeseen and cannot be easily reconciled without additional research, perhaps focused on the systematic evaluation of alternative semiempirical QM methods (or parameters) and MM force fields (e.g., OPLS for liquid solutions).

In this study, only first-generation TPs were analyzed, i.e., initial TPs originating from bi-molecular HOCl collisions with only the parent target compound. However, by cycling the $N_{\text{FMD}}$ algorithm over first-generation TPs of interest, it is possible to explore the formation of second- and later-generation TPs, as recently described by Ridgway et al. [30] for reactions of certain volatile gas-phase organics with ozone. Because collision durations were purposely restricted to <0.5 ps to maximize the number of reactions in a series, the $N_{\text{FMD}}$ algorithm as applied in this investigation did not allow sufficient time for potentially energetically favorable molecular (MO) rearrangements to take place, particularly for meta-stable (carbocation) intermediates, such as a meta α ortho migration of a halogen atom, or a hydrogen abstraction from a halogenated Wheland intermediate. It should be emphasized, however, that such rearrangements can indeed be explored for specific TPs of interest by subsequently carrying out conventional equilibrium MD using temperatures high enough to sample rearrangement barriers.

As applied to OxBZ, the $N_{\text{FMD}}$ algorithm revealed a range of TPs, including a number that have not yet been experimentally verified by other researchers [16,20]. TPs observed in the $N_{\text{FMD}}$ simulations as well as in experimental studies included mono-chlorinated OxBZ (especially, the ortho-addition product), mono-chlorobenzene, OH radicals, HCl, chloromethane, hydrogen abstracted TPs, and opening and carbonyl-bond scission products [16–20].

Several other compounds that have not yet been experimentally verified were also observed, including mono-chloroguaiacol (ortho-chloromethoxyphenol), mono-chlorohydroxybenzophenone, di-hydroxybenzophenone, 4-methoxycatechol, hydroxymethoxybenzaldehyde (precursor), benzaldehyde (precursor), and formaldehyde (di-radical precursor). At least some of these compounds (e.g., mono-chloroguaiacol and 4-methoxycatechol) have been demonstrated to exhibit human and/or environmental health hazards, including cytotoxicity, mutagenicity, and carcinogenic activity [54,55].

In addition to yielding a statistical distribution of expected TPs, as well as some compounds and intermediates that have not been previously reported, the $N_{\text{FMD}}$ algorithm was used to analyze the formation of an ortho-halogenated Wheland adduct of OxBZ. This meta-stable intermediate compound was of special interest since the stable hydrogen abstracted end-product was reported by Zhang et al. [16] and Negreira et al. [20]. Analysis of $N_{\text{FMD}}$ collision trajectories for this reaction revealed a pseudo-stable CI-adduct formation that occurred by electrophilic attack at methoxy ring atom C8 situated ortho to the –OH group and para to the –OH group. A critical step in the reaction path was homolytic cleavage of the relatively labile HO–Cl bond in HOCl followed by release of a free ‘OH radical and a transitory “semi-bound”’ CI radical. The later initiated electron delocalization by a π–resonance charge transfer mechanism in which electron density transmigrated from the methoxy ring to chlorine, resulting in rapid covalent bond formation (<50 fs). It was also noted that conjugated π-bond resonance fluctuations in the OxBZ methoxy ring were tightly coupled to kinetically invoked conformational oscillations stemming from the initial collision event with HOCl.

Collision orientation during $N_{\text{FMD}}$ simulations was critical to determining whether a reaction proceeded to completion (i.e., whether specific TP formation occurred). Very subtle alterations in the starting conformations and orientations of the reactants completely changed the reaction dynamics and the kinds of TPs generated (data not shown). Solvent effects in the form of Coulomb and Lennard–Jones interactions with the reactants, as well as random steric interference along the reaction trajectories, also influenced the $N_{\text{FMD}}$ collision orientations and reaction outcomes. These observations argue for the importance of reaction “preorganizational steps” as described by Rosokha and Kochi [56] in which ephemeral semistable pre-equilibrium (arene) complexes evolve through inductive and/or resonant charge transfer processes between non-bonded species prior to formation or dissociation of covalent bonds. Formation of the meta-stable halogen-addition OxBZ intermediate described in this report and the formaldehyde di-radical intermediate are two examples of pre-organizational states that would be expected to undergo MO rearrangements leading to stable TPs.

Recently Galabov et al. [49] challenged the traditional view of electrophilic aromatic addition/substitution reactions involving the standard SEAr (arenium ion) mechanism. Using computational and NMR approaches, they identified two alternative reaction pathways, an “addition–elimination” sequence and a “direct concerted mechanism”, with the former being more favorable with slightly lower activation barriers. According to the authors, no arenium σ-complex is found along the reaction path for direct substitution, yet
several new hybrid carbocation-HCl intermediates are proposed that respond to autocatalytic HCl elimination yielding the ortho-product. Either alternative explained the preference for ortho-/para-anisole derivatives, since these orientations represent a natural consequence of the favored stabilizing influence of the –OCH3 group.

Although HCl formation was observed in a significant fraction of the HOCI–OxOBZ and other reactions performed in this study (data not shown), no compelling evidence was observed for the formation of HCl-carbocation complexes. However, notwithstanding a more conventional interpretation, it would be possible to interpret the nascent H—Caro—Cl bond as the earliest stage of such a putative complex. Indeed, as suggested by data presented in Figs. 15–18, coupled changes in the atomic partial charges of the participating atoms (i.e., the terminal H—Caro— and Cl—Caro—) could have interacted through an inductive mechanism, not unlike that described by Galabov et al. [49].

Based on the findings reported in this investigation, it is concluded that N-MeFMD simulations can be successfully applied in combination with experimental approaches to help elucidate the roles of transitory or semistable intermediates that are sometimes challenging to detect and quantify in complex chemical reaction pathways. The modeling approach could be expected to have broad application in identifying potential reaction intermediates that cannot be readily experimentally observed, as well as in deciphering other details of proposed TP formation pathways.

Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BPA</td>
<td>Bis-phenol-A</td>
</tr>
<tr>
<td>DFT</td>
<td>Direct inversion of the iterative subspace</td>
</tr>
<tr>
<td>HOMO</td>
<td>Highest occupied molecular orbital</td>
</tr>
<tr>
<td>MD</td>
<td>Molecular dynamics</td>
</tr>
<tr>
<td>MM</td>
<td>Molecular mechanics</td>
</tr>
<tr>
<td>MO</td>
<td>Molecular orbital</td>
</tr>
<tr>
<td>NDDO</td>
<td>Neglect of diatomic differential overlap; category of semiempirical calculation</td>
</tr>
<tr>
<td>NSC</td>
<td>Nano-solvation cage</td>
</tr>
<tr>
<td>N-MeFMD</td>
<td>Non-equilibrium forced molecular dynamics</td>
</tr>
<tr>
<td>OxOBZ</td>
<td>Oxybenzone</td>
</tr>
<tr>
<td>QM</td>
<td>Quantum mechanics</td>
</tr>
<tr>
<td>QM/MM</td>
<td>Quantum/molecular mechanics</td>
</tr>
<tr>
<td>RM1</td>
<td>Semiempirical method similar to AM1 or PM3</td>
</tr>
<tr>
<td>SEAr</td>
<td>Electrophilic aromatic substitution reaction mechanism with arenium ion formation</td>
</tr>
<tr>
<td>SFC</td>
<td>Self-consistent field calculation</td>
</tr>
<tr>
<td>STP</td>
<td>Standard temperature and pressure; 25°C, 1 atmosphere</td>
</tr>
<tr>
<td>TIP3P</td>
<td>Water model of Jorgenson et al. [32]</td>
</tr>
<tr>
<td>TP</td>
<td>Transformation product</td>
</tr>
<tr>
<td>UHF</td>
<td>Unrestricted Hartree–Fock</td>
</tr>
<tr>
<td>VDW</td>
<td>Van der Waals</td>
</tr>
</tbody>
</table>

References
