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Molecular level insights into the direct health

impacts of some organic aerosol components?



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Fatemeh Keshavarz Dab

Organic aerosols (OAs) comprise 20-90 w% of atmospheric aerosols. While they have some positive impacts (e.g. acting as precipitation nuclei), they can pose some negative impacts on human health. The health impacts range from pulmonary inflammation to suppression of defense mechanisms, cancer and death. To understand how OA components can affect human body, this study employed a combination of in silico techniques: molecular docking, quantum chemical analysis of interactions, thermodynamics and kinetics of the reaction of OA components with DNA and oxidative stress mechanism, in addition to theoretical evaluation of their absorption, distribution, metabolism and excretion (ADME) properties. As OAs contain many components, this study just focused on benzo[a]pyrene (BaP), benzo[a]pyrene diol (BPDE), the cyclopentanoperhydrophenanthrene (CPPP) sterane, 2,6-dimethyl-1,4benzoguinone (DMBQ), hopane (HOP), 9,10-phenanthreneguinone (PQ), and an OA model (fulvic acid; FA), which are all known for their negative health impacts. According to the ADME predictions, the low bioavailability of FA would limit its negative impacts. However, the ADME estimations and docking simulations suggested that BPDE, DMBQ and PQ have significant impact in affecting human health negatively: they can be feasibly distributed throughout human body by human serum albumin, they can likely permeate cell membranes, the blood-brain barrier and the gastrointestinal tract, and they show higher DNA binding efficiencies compared to the other OA components. Based on the quantum chemical calculations, BPDE, DMBQ and PQ can interact with the adenosine nucleobases of DNA, but they cannot directly form DNA adducts because of the nonspontaneous and kinetically infeasible nature of the corresponding reaction routes. Similarly, they cannot directly cause oxidative stress. However, in reference to the experimental evidence offered by earlier studies, their metabolites can result in both oxidative stress and carcinogenicity.

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Introduction

Organic aerosols (OAs) comprise a large proportion of atmospheric aerosols¹ ranging from 20 to 60 w% in continental mid-latitudes to about 90 w% in tropical forested areas.² They not only control the energy budget, biogeochemical cycles (including precipitation patterns) and climate of Earth, but also pose adverse health effects. 3-6 Particularly, as a significant portion of particulate matter (PM), OAs can be responsible for a portion of the 3 million deaths per year caused by PM.⁷

An important point is that the health impacts of aerosols depend on many parameters such as their particle size, organic compounds, transition metals and specific surface.8,9 Therefore, their health effects are highly sensitive to their physiochemical properties. However, generally speaking, exposure to aerosol particles can induce pulmonary inflammation, lung cancer, oxidative stress caused by organic compounds or transition metals, inflammation and immune impacts initiated by biological aerosol components, covalent modification of enzymes and other key intracellular proteins, regulation of airway reactivity and heart rate variability, stimulation of autonomic nervous system activity, enhanced response to environmental allergens, and suppression of defense mechanisms.9-11 Some other health implications of PM are related to reproductive systems and include low birth weight, preterm labor, growth retardation and female infertility. 12 All these health risks increase on prolonged exposure to PM. 13

While most studies have considered the health impacts of PM rather than focusing on OAs, the hazardous nature of some OA components has been the subject of ongoing

^a Institute for Atmospheric and Earth System Research, Faculty of Science, University of Helsinki, FI-00014 Helsinki, Finland. E-mail: fatemeh.keshavarz@helsinki.fi

^b Department of Physics, School of Engineering Science, LUT University, FI-53851 Lappeenranta, Finland. E-mail: fatemeh.keshavarz@lut.fi

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research. The most outstanding examples are quinones (such as 9,10-phenanthrenequinone (PQ) and 2,6-dimethyl-1,4benzoquinone (DMBQ)), polycyclic aromatic hydrocarbons (PAHs; e.g. benzo[a]pyrene (BaP)), triterpenes (e.g. hopanes (HOP)) and phenanthrenes (like the cyclopentanoperhydrophenanthrene (CPPP) sterane).

PQ represents the quinones found in diesel exhaust particles, and it is believed to induce cytotoxic effects through the generation of reactive oxygen species (ROSs; i.e. oxidative stress). 14 Similarly, DMBQ in ambient particles leads to oxidative stress. However, the contribution of DMBQ to the toxicity of PM should be lower than that of PQ. 15 BaP has been assessed as the most genotoxic, mutagenic and carcinogenic component of PM. 16-18 These properties are owed to its ability to form adducts with DNA. 19 BaP is mainly found in its oxygenated forms (e.g. (+)-benzo[a] pyrene-7,8-dihydrodiol-9,10-epoxide (BPDE)), and its oxygenated forms might be responsible for its genotoxicity.²⁰ Finally, hopanes and steranes, which are fossil fuel markers, 21 are linked to toxicity but no plausible mechanism has been proposed for them.²²

To the best of our knowledge, no study has evaluated the molecule-level impacts of OAs. However, some details about the molecule-level effect of several quinones on oxidative stress (e.g. the reaction of tetrachloro-o-benzoquinone with H₂O₂²³ and the redox mechanism of several quinones²⁴) and DNA (see ref. 25-27) have been studied. To this end, we consider an OA model, PQ, DMBQ, BaP, BPDE, the simplest HOP and CPPP, and analyze their corresponding absorption, distribution, metabolism and excretion (ADME) properties. While the metabolic response to OAs is completely complicated, we assess their direct impacts and potential toxicity by analyzing their interaction with human serum albumin (HSA) and DNA through molecular docking, in addition to concerning their mechanistic role in oxidative stress and DNA alkylation using quantum chemical approaches. The docking and quantum chemical investigations are devised to focus on an isolated process. However, the ADME properties are used to study the health impacts of the OA components at a broader scope. The ADME results can offer pharmacokinetic/physiological details beyond extensive quantum chemical or molecular docking simulations because they can follow metabolic networks rather than a single isolated reaction.

The main working hypothesis of the study is that the applied molecular modeling tools can qualitatively describe what will happen to the OA components at their first encounter with important biomolecules. However, because of the complexity and vast range of bioprocesses, all plausible bioprocesses are not studied. Particularly, the metabolic products of the OA components and their reaction/interaction with the evaluated biomolecular targets are neglected, and only the direct impacts of the OA components have been concerned. Furthermore, although all the adopted techniques are strong and advantageous to some extent, each has some shortcomings which are addressed along with the results. Therefore, we suppose that the results are qualitatively reliable, but the quantitative results might suffer from some level of inaccuracy. In addition, we encourage the use of the findings of this study as a starting

point for a more comprehensive experimental/computational study of the health impacts.

HSA is eventually selected because it is the transportation vehicle in human body, and it controls the distribution and metabolism of different organic compounds (like drugs). 28,29 DNA is chosen as the principal target of genotoxicity. For comparison purposes, the same analyses are done on cis-platin (CPT) and ThioTEPA (N,N',N''-triethylenephosphoramide) (globally known anticancer drugs both forming adducts with DNA)30,31 and ibuprofenate (IBPPA; isobutyl phenyl propionic acid or (RS)-2-(4-(2-methylpropyl)phenyl)propanoic acid) (a famous antiinflammatory drug with no impact on DNA)32 as reference compounds. Fig. 1 shows the chemical structure of the evaluated compounds.

Results and discussion

ADME properties

Table 1 reports the ADME properties of the aerosol components and the reference drugs. It should be noted that these results are obtained through machine learning based protocols, and they are not completely reliable without further experimental verification. According to the results, just FA is slightly hydrophilic (negative $\log P_{\text{o/w}}$), while the other compounds are lipophilic (positive $\log P_{\text{o/w}}$). As cell membranes are made of lipid-type structures (i.e. phospholipid bilayers), all the evaluated compounds, except FA, should be able to pass through cell membranes passively (rather than diffusing through membrane proteins). Also, just FA and ThioTEPA have substantial water solubility (WS). Regardless of solubility and hydrophilicity, the ADME results predict BPDE, DMBQ, IBPPA, PQ and ThioTEPA as the compounds that can be absorbed through the gastrointestinal system. ThioTEPA has been approved as a drug that can significantly affect (toxify) the gastrointestinal tract.³³ Also, the long-term use of IBPPA has been associated with gastrointestinal ulceration.34 Therefore, BPDE, DMBQ and PQ are likely to damage the gastrointestinal tract in a similar way. In addition, the blood-brain barrier (BBB) permeation results outline that BPDE, DMBQ and PQ can permeate the BBB like the IBPPA anti-inflammatory/painkiller agent. As these compounds are lipophilic and have the potential of crossing the BBB, they might affect the central nervous system.³⁵ However, both DMBQ and PQ are quinones, and some quinones, e.g. pyrroloquinoline quinone,³⁶ have neuroprotective effects on the nervous system, meaning that they can prevent/reduce damages to the nervous system. Consequently, the predicted BBB permeation should not be considered as a negative health impact by organic aerosols without experimental evidence. None of the compounds can pass through skin (negative $\log K_p$) as expected. However, there is experimental evidence for the entrance of OAs and their components through lungs.³⁷ Also, as BPDE, DMBQ, IBPPA, PQ and ThioTEPA are predicted to pass through the gastrointestinal tract, processing of OA contaminated drinks and foods by the gastric system can be considered as a potential route for their entrance to cellular areas.

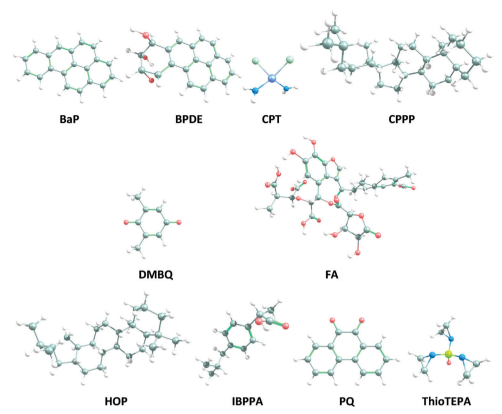


Fig. 1 Geometries of the model OA (FA), PQ, DMBQ, BaP, BPDE, HOP, CPPP, CPT, ThioTEPA and IBPPA optimized at the ωB97X-D/def2-TZVP level of theory, in water (see Fig. S1, ESI,† for the 2D structures). The carbon, hydrogen, oxygen, nitrogen, chloride, sulfur and platinum atoms are represented as grey, white, red, blue, light green, yellowish and light purple balls, respectively

Among the tested compounds, BPDE, FA and ThioTEPA are proposed as P-glycoprotein (a drug transporter) substrates by theoretical ADME estimations, which means that their pharmacokinetic properties would likely change in human body. However, the bioavailability of FA is estimated to be low,

meaning that just a small fraction of it can potentially circulate among cells and organs systematically. Therefore, the metabolism of FA and its interaction with other metabolites should be limited. In the case of ThioTEPA, being a Pglycoprotein substrate can be related to tumor chemoresistance.³⁸

Table 1 ADME properties of the evaluated compounds^a

Molecule	BaP	BPDE	CPT*	DMBQ	FA	НОР	IBPPA*	PQ	CPPP	ThioTEPA*
MW	252.31	303.33	298.03	136.15	774.63	370.65	206.28	208.21	372.67	201.21
NRB	1	0	0	0	14	1	4	0	5	3
MR	87.65	88.34	19.23	37.91	176.76	120.81	62.18	60.54	122.67	64.96
TPSA	0.00	52.99	52.04	34.14	339.23	0.00	37.30	34.14	0.00	35.91
$\log P_{\mathrm{o/w}}$	5.34	1.86	_	1.21	-0.26	7.56	3.01	2.46	8.21	0.51
WS	1.38×10^{-6}	1.08×10^{-2}	_	3.66	3.08×10^{2}	6.34×10^{-4}	7.49×10^{-2}	1.17×10^{-3}	2.43×10^{-5}	5.59×10^{2}
GIA	Low	High	_	High	Low	Low	High	High	Low	High
BBB	No	Yes	_	Yes	No	No	Yes	Yes	No	No
Pgp	No	Yes	_	No	Yes	No	No	No	No	Yes
CYP1A2	Yes	Yes	_	No	No	Yes	No	Yes	No	No
CYP2C19	No	No	_	No	No	No	No	Yes	No	No
CYP2C9	No	No	_	No	No	Yes	No	No	No	No
CYP2D6	No	Yes		No	No	No	No	No	No	No
CYP3A4	No	No	_	No	No	No	No	No	No	No
$\log K_{\rm p}$	-3.6	-6.74		-6.26	-10.95	-1.26	-5.07	-5.78	-0.61	-7.12
BS	0.55	0.55	_	0.55	0.11	0.55	0.56	0.55	0.55	0.55

^a MW: molecular weight in g mol⁻¹; NRB: number of rotatable bonds; MR: molar refractivity as a measure of polarizability; TPSA: total polar surface area in \mathring{A}^2 ; $\log P_{0/w}$: octanol-water partition as a lipophilicity measure; WS: water solubility in g L⁻¹; GIA: gastrointestinal absorption; BBB: blood-brain barrier permeant; Pgp: P-glycoprotein substrate; CYP1A2: cytochrome P450 1A2 inhibitor; CYP2C19: cytochrome P450 2C19 inhibitor; CYP2C9: cytochrome P450 2C9 inhibitor; CYP2D6: cytochrome P450 2D6 inhibitor; CYP3A4: cytochrome P450 3A4 inhibitor; $\log K_{\rm p}$: skin permeation in cm s⁻¹; and BS: bioavailability score. The reference drugs are distinguished by an asterisk.

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Therefore, this result can imply that P-glycoprotein can likely alter the health impacts of BPDE including the inhibition of the cytochrome P450 1A2 and 2D6 enzymes. The negative or positive effect of P-glycoprotein on the health impacts of BPDE requires experimental assessment. Also, BaP, HOP and PQ are predicted to inhibit cytochrome P450 1A2. Similarly, HOP and PQ are predicted to respectively inhibit cytochromes P450 2C9 and P450 2C19. All these enzymes are involved in the metabolism of drugs and physiologically important compounds.39-41 Therefore, BaP, BPDE, HOP and PQ seem to have the potential to negatively affect the metabolism of some pharmaceutics and metabolic agents.

DNA and HSA docking

The docking results are summarized in Table 2 and Tables S1, S2 and depicted in Fig. S2-S4 (ESI†). According to Table S1 (ESI†), the strongest interaction (i.e. the lowest interaction energy) with HSA refers to FA (the largest compound) and the weakest interaction refers to CPT (the smallest compound). None of the tested compounds bind to HSA through electrostatic interactions, while hydrogen bonding plays a significant role in the binding of BPDE, CPT, DMBQ, FA and IBPPA and steric (van der Waals) interactions are the main drivers of the compounds binding to HSA. The dominance of steric interactions can be attributed to the hydrophobicity (lipophilicity) of most ligands (see the positive values of $\log P_{\text{o/w}}$ in Table 1) and the hydrophobicity of most HSA domains.

In physiological conditions, ligand binding is not just governed by the sum of molecular interactions. In competition with other ligands, the ligand with the highest diffusion rate and the highest concentration would bind to the target more efficiently. Larger molecules diffuse slower toward the target, and the ligand with the highest concentration in the microenvironment has the highest potential of binding. To account for diffusion contribution and eliminate the effect of molecular size on the binding of the ligands, ligand efficiency (LE) was calculated as the interaction energy divided by the number of non-hydrogen atoms of the compound. 42 The LE values showed that CPPP, HOP and FA have the lowest HSA binding efficiencies, whereas BaP, BPDE, CPT, DMBQ, IBPPA and PQ have noticeable

Table 2 Summary of the 3USO docking results^a

Ligand	Interaction energy	Electrostatic interaction	Hydrogen bonding	LE	Binding site
BaP	-156.9	0.0	0.0	-6.96	CAA/GTT
BPDE	-160.0	0.0	0.0	-7.84	ATT/TAA
CPPP	-102.9	0.0	0.0	-3.81	AAT/TTA
CPT	-52.2	0.0	0.0	-10.45	CA/GT
DMBQ	-89.9	0.0	0.0	-8.99	TTT/AAA
FA	-231.5	0.0	0.0	-4.21	AATA/TTAT
HOP	-110.4	0.0	0.0	-4.09	TAA/ATT
IBPPA	-114.8	0.0	0.0	-7.65	AAA/TTT
PQ	-132.8	0.0	0.0	-8.30	TTT/AAA
ThioTEPA	-74.0	0.0	0.0	-6.73	TA/AT

^a The energy and ligand efficiency (LE = interaction energy/number of non-hydrogen atoms) values are unitless.

LE values (LE < -5.00) and a high potential of transportation to all cellular targets by HSA.

Similar to HSA binding, Table 2 and Table S2 (ESI†) indicate the lowest (strongest interaction) and highest DNA binding energies, respectively, for FA and CPT, and the involvement of no electrostatic interaction. Here, no hydrogen bonding interaction can be observed. Therefore, the docking results indicate that the DNA-ligand binding of the studied compounds should be mainly driven by steric interactions. However, one should note that the employed docking simulations are based on semi-flexible treatment of ligand binding to the targets, and the calculated interaction energies/types are based on classical force field parameters. Therefore, the docking results might be biased with inaccurate estimation of atomic parameters and intermolecular interactions, leading to underestimation of electrostatic and hydrogen-bonding interactions.

For both DNA strands, the best LE values (LE < -6.00) are obtained for BaP, BPDE, CPT, DMBQ, IBPPA, PQ and ThioTEPA, which are the compounds giving HSA LE values lower than -5.00. Particularly, the CPT anticancer drug gives the best ligand efficiency, and the ThioTEPA alkylating agent shows a fairly noticeable LE value. These two reference compounds were neglected in the quantum mechanical calculations because their DNA cleavage mechanisms are reported elsewhere, 31,43 and just BaP, BPDE, DMBQ, IBPPA and PQ were concerned in further evaluation of the OA-DNA interaction and oxidative stress.

Mutagenicity potential/DNA interaction

To increase the accuracy of the DNA/ligand interaction analysis and to identify any missing interaction type/pairs, NBO analysis was performed. NBO analysis goes beyond classical force fields and atom pairs, and it utilizes wave functions to calculate orbital properties and follow the trends of charge transfer between them. In this way, it can specifically distinguish hydrogen bonds. The results of Natural Bond-Orbital (NBO) analysis on the interactions of BaP, BPDE, DMBQ, IBPPA and PQ with 3US0 are outlined in Table S3 (ESI†), where higher $E^{(2)}$ stabilization energy values are indicative of stronger interactions. 3US0 is selected as the DNA target because it interacts with the case compounds more strongly, relative to 4AWL (see Table 2 and Table S2, ESI†). Although the docking results of none of the compounds predict hydrogen bonding interactions, the quantum chemistry based NBO results identify very weak to strong hydrogen bonding interactions in the form of charge transfer from the lone-pair (LP) orbital of proton donors (such as O and N atoms) to the antibonding (σ^*) orbital of proton acceptors (e.g. OH and CH). The BaP interaction with the T nucleobase of 3US0 involves a very weak hydrogen bond. Also, a weak hydrogen bond is observed between PQ and an A nucleobase. DMBQ and IBPPA do not form any hydrogen bonds with 3USO. However, a strong hydrogen bonding network helps BPDE stabilize along the A and T nucleotides. On the other hand, both docking-based interaction energies and $E^{(2)}$ results show that BPDE and BaP can interact with 3US0 the strongest, whereas

DMBQ can interact with DNA weakly, relative to the other tested compounds. This increases the DNA adduct formation potential of BPDE (and its BaP non-oxidized form), but it cannot be directly interpreted as the experimentally observed BaP carcinogenicity¹⁹ without further theoretical and experimental evaluations. In addition, the results suggest that BaP, BPDE, DMBQ and PQ mainly interact with the A nucleobases of DNA, while IBPPA just interacts significantly with the deoxyribose moiety of the A, C and T nucleotides. This is in line with the experimental findings that report that IBPPA has no significant negative impact on DNA,44 or the studies that show it prevents DNA damage. 45 Also, it should be noted that A nucleobases are more susceptible to mutagenic lesion formation when interacting with some carbonyl containing compounds (such as formaldehyde⁴⁶), and BaP, BPDE, DMBQ and PQ interact efficiently with the A nucleobases. This opens up a question about the possible contribution of such interaction mode to their mutagenicity, which should be answered through more accurate calculations (such as QM/MM molecular dynamics simulations or quantum chemical modeling) or experimental evidences. To answer this question, we used quantum chemical calculations to re-evaluate the binding of BaP, BPDE, DMBQ and PQ to their corresponding DNA binding site (according to the docking and NBO results) by full geometry optimization of the binding pairs and re-calculation of their binding energies under cell-like conditions. Also, we conducted quantum chemistry level mechanism exploration to assess any DNA alkylating/non-alkylating reaction between these OA components and DNA.

Based on the NBO results, more accurate interaction energies were calculated only for BaP, BPDE, DMBQ and PQ by cropping the effective DNA binding sites (i.e. the main interacting nucleotides). The ZPE-corrected and Gibbs free interaction energies, along with the optimized structures, are shown in Fig. 2. According to the obtained values, the interactions of BPDE, DMBQ and PQ with the nucleotides are slightly nonspontaneous (positive Gibbs free energies), but the BaP interaction is slightly spontaneous. Therefore, the binding of these compounds to DNA cannot be completely neglected, while it cannot be thermodynamically highly plausible. Nevertheless, the potential mutagenicity and carcinogenicity of the OA components

were mechanistically assessed by considering their roles in DNA alkylation (adduct formation), DNA cleavage or any other structural changes in DNA. However, exhaustive reaction path scanning and TS predictions failed to propose any mechanism.

Because many experimental studies have associated BPDE with genotoxicity, mutagenicity and carcinogenicity, 16-18,20 two common mechanisms were further studied for adduct formation between DNA and BPDE, DMBQ and PQ; type-2 nucleophilic substitution (SN2; formation of one bond between two reactants simultaneously with the cleavage of a bond in one of the reactants) similar to the alkylation mode of the ThioTEPA anticancer agent; ³¹ and Michael addition (MA; nucleophilic addition of a nucleophile to a carbonyl compound) between the electron deficient carbon of quinones and the exocyclic NH2 moiety of nucleobases.47 Here, just an A nucleobase was targeted to exclude steric restrictions and simplify the calculations. The SN2 scenario (bond formation between the OA components and the A nucleobase simultaneously with opening of the epoxy functional group in BPDE, or conversion of the C=O double bond to a C-O⁻ single bond) was rejected by the scanning and TS calculations, and every time the predicted TSs converted into the individual reactants.

The MA scenario was more successful, and a reaction path was approved for DMBQ (R1) and PQ (R2) which is shown in Fig. 3. Also, a TS was found for adduct formation between BPDE and the A nucleobase at the ωB97X-D/3-21g level and verified using IRC calculations ((R3); see Fig. S5, ESI,† for the TS, and the connecting pre-reaction complex and product). But the TS could not be validated at the ωB97X-D/def2-TZVP level. Notably, in reactions (R1)-(R3), "A" stands for the adenine (A) nucleobase. To assure that the reaction path observed for BPDE at the ωB97X-D/3-21g level is an artifact of low-level calculations, the TS was reoptimized at the BVP86/def2-TZVP, M06-2X/def2-TZVP and B3LYP/def2-TZVP levels. All alternative level calculations failed, asserting the absence of any DNA adduct formation route for BPDE. However, there might be still some reaction paths involved in mutagenicity induction but missing from our calculations. Some experiments have stated that it is the metabolites of BaP and BPDE (like phenols and quinines) that can induce mutagenic, teratogenic and cytotoxic effects⁴⁸⁻⁵¹ through ROS

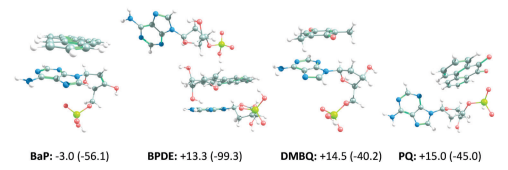


Fig. 2 Geometry and binding Gibbs free energy (and ZPE-corrected interaction energy) of the ligand-nucleotide complexes retrieved from NBO analysis. The energies are in kJ mol $^{-1}$. The complexes, nucleotides and ligands are optimized and energy analyzed under cellular conditions while conserving their conformational state in their corresponding ligand-nucleotide complexes.

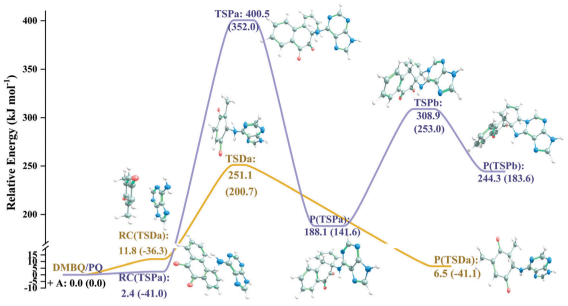


Fig. 3 Mechanism of DNA adduct formation by DMBQ and PQ. The Gibbs free energies (and ZPE-corrected energy values) are calculated relative to the corresponding reactants in kJ mol⁻¹. The carbon, hydrogen, oxygen, and nitrogen atoms are represented as grey, white, red, and blue balls, respectively.

generation.52

$$A + DMBQ \rightarrow RC(TSDa) \xrightarrow{TSDa} P(TSDa)$$
 (R1)

$$A + PQ \rightarrow RC(TSPa) \xrightarrow{TSPa} P(TSPa) \xrightarrow{TSPb} P(TSPb) \qquad \text{(R2)}$$

$$\begin{array}{c} A + BPDE \rightarrow RC(TSBa) \xrightarrow{TSBa} P(TSBa) \\ \\ (R3: unapproved \ reaction) \end{array}$$

As seen in Fig. 3, there are transition states found along the DNA adduct formation paths of DMBQ and PQ through the MA reaction scheme. However, the related paths are not kinetically feasible. When DMBQ and PQ form a pre-reaction complex (PC) with A, they, respectively, need 251.1 and 400.5 kJ mol⁻¹ energy to surpass the TSDa and TSPa barriers and form the corresponding products (Ps), namely, P(TSDa) and P(TSPa). For DMBQ, the reaction is single-step and TSDa is a result of concerted NH2 attack on the electron deficient carbon of DMBQ and hydrogen transfer from NH2 to the neighboring carbon atom. The resultant P(TSDa) product is more stable than the reactants based on its electronic energy, but the reaction is slightly nonspontaneous. For PQ, the reaction is two-step: the first step being nonspontaneous and mimicking TSDa but giving a highly unstable P(TSPa) product; in the second step, the endocycle pyrimidinimine nitrogen attacks the carbon atom neighboring the newly NH-bound carbon by surmounting the 308.9 kJ mol⁻¹ free energy barrier of TSPb, yielding the unstable adenosine-bound PQ structure of P(TSb).

As all our quantum chemical evaluations suggest, the tested OA components are incapable of causing mutagenicity directly through the evaluated mechanisms under cellular conditions. However, some mechanistic schemes might be missing from our calculations. Particularly, as the ADME results predicted, binding to P-glycoprotein might convert BPDE to secondary

metabolites, which can end up in carcinogenic effects. Also, experimental studies on some quinones47,53,54 have attributed DNA adduct formation to quinone metabolites. Therefore, while our results decline a direct DNA effect by the OA components, the possibility of un-examined reaction mechanisms and the indirect effect of OA metabolites on DNA should not be ignored without sufficient computational and/or experimental evidence.

Oxidative stress

According to Squadrito et al.,55 the first step in oxidative stress induced by different organic compounds is hydride transfer from NADPH to the organic compound in a nucleophilic substitution reaction (simplified as (R4) with OC being the organic compound and the transferred hydride being represented by h⁻). Among the studied OA components, the quinones (PQ and DMBQ), BPDE and FA can offer an electrophilic site for acceptance of hydrides, but FA has low bioavailability based on the HSA docking and ADME estimations. Therefore, the oxidative stress mechanism was just followed for PQ, DMBQ and BPDE. Fig. 4, 5 and Fig. S5 (ESI†) depict the minimum energy reaction paths of the observed mechanism, which takes place in five individual steps. In the first step (Fig. 4; (R5)-(R7)), NADPH and DMBQ/ BPDE diffuse toward each other to form RC complexes, respectively, named RC(TSD1) and RC(TSB1). For PQ, no stable RC forms. Then, a hydride transfers from NADPH to the carbon in the carbonyl groups of PQ and DMBQ, or the oxirane carbon of BPDE. Simultaneously, the attacked bond cleaves, forming a single bond between the attacked carbon and a (now) negatively charged oxygen atom along with NADP+ in the form of a post-reaction complex (PC). The products then diffuse apart. Under cell-like conditions, the Gibbs free energy barriers of this step for BPDE (TSB1), DMBQ (TSD1) and PQ (TSP1) are, respectively, 118.7, 109.7 and 98.3 kJ mol⁻¹ high, meaning that this reaction is the most

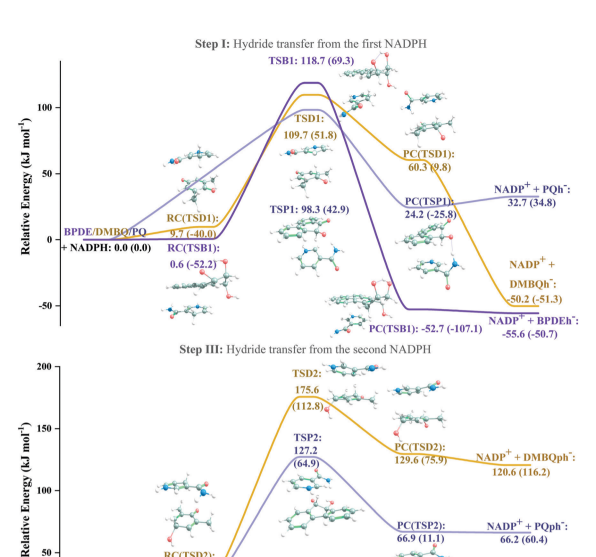


Fig. 4 Steps I and III of the oxidative stress mechanism initiated by the organic aerosol components. The Gibbs free energies (and ZPE-corrected energy values) of the BPDE, DMBQ and PQ reaction sets are calculated relative to the corresponding reactants in kJ mol⁻¹. The carbon, hydrogen, oxygen, and nitrogen atoms are represented as grey, white, red, and blue balls, respectively.

feasible for PQ. However, in general, these reaction steps are not kinetically feasible, unless side metabolic reactions provide sufficient energy for surpassing the corresponding barriers.

DMBQp/PQp NADPH: 0.0 (0.0)

100

50

$$NADPH + OC \rightarrow NADP^{+} + (OC)h^{-}$$
 (R4)

RC(TSP2):

$$NADPH + BPDE \rightarrow RC(TSB1) \xrightarrow{TSB1} PC(TSB1)$$

$$\rightarrow NADP^{+} + BPDEh^{-}$$
(R5)

$$NADPH + DMBQ \rightarrow RC(TSD1) \xrightarrow{TSD1} PC(TSD1)$$

$$\rightarrow NADP^{+} + DMBQh^{-}$$
(R6)

$$NADPH + PQ \xrightarrow{TSP1} PC(TSP1) \rightarrow NADP^{+} + PQh^{-} \quad \mbox{(R7)}$$

In the second step (Fig. S6, ESI†), the negatively charged hydride containing BPDE, DMBQ and PQ compounds (i.e. BPDEh⁻, DMBQh⁻ and PQh⁻) are, respectively, protonated to BPDEp, DMBQp and PQp through the general reaction of (R8). This reaction step is thermodynamically spontaneous and highly favorable with protonation Gibbs free energies being lower than $-690.0 \text{ kJ} \text{ mol}^{-1}$. In the third step (Fig. 4; (R9) and (R10)), a second hydride transfer reaction occurs similar to the first step. Here, just PQ and DMBQ are attacked as they have another electrophilic center (carbonyl group) available. The barriers of the second hydride transfer step are higher in Gibbs free energy mainly because of the lower electron deficiency of the attacked centers: 175.6 kJ mol⁻¹ (TSD2) vs. 109.7 kJ mol⁻¹ (TSD1) for DMBQ, and 127.2 kJ mol⁻¹ (TSP2) vs. 98.3 kJ mol⁻¹

120.6 (116.2)

NADP⁺ + PQph⁻:

66.2 (60.4)

PC(TSP2):

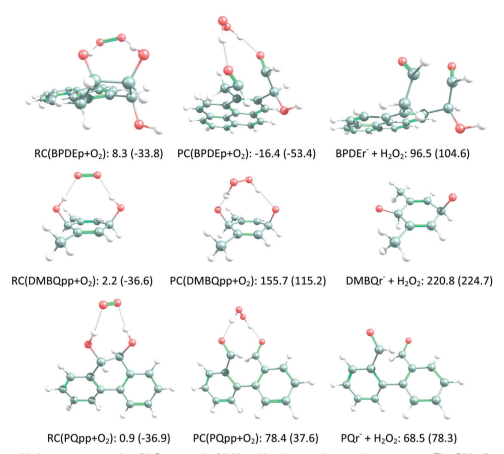


Fig. 5 Step V of the oxidative stress mechanism (H₂O₂ generation) initiated by the organic aerosol components. The Gibbs free energies (and ZPEcorrected energy values) of the BPDE, DMBQ and PQ reaction sets are calculated relative to the corresponding reactants in kJ mol⁻¹. The carbon, hydrogen, oxygen, and nitrogen atoms are represented as grey, white, red, and blue balls, respectively.

(TSP1) for PQ. The third step is followed by another highly favorable protonation step (see Fig. S6, ESI,† and the general reaction (R11)) with protonation Gibbs free energies lower than $-670.0 \text{ kJ mol}^{-1}$.

$$H^+ + (OC)h^- \rightarrow (OC)p$$
 (R8)

$$\begin{split} NADPH + DMBQp &\rightarrow RC(TSD2) \xrightarrow{TSD2} PC(TSD2) \\ &\rightarrow NADP^+ + DMBQph^- \end{split} \tag{R9}$$

$$NADPH + PQp \rightarrow RC(TSP2) \xrightarrow{TSP2} PC(TSP2)$$

$$\rightarrow NADP^{+} + POph^{-}$$
(R10)

$$H^+ + (OC)ph^- \rightarrow (OC)pp$$
 (R11)

While steps I to IV are responsible for hydrogenation of the OA components, step V is the key step producing ROSs. In the fifth step (Fig. 5; reactions (R12)-(R14)), an O₂ radical reacts with the BPDEp, DMBQpp and PQpp compounds produced in the earlier steps, yielding H₂O₂ along with the corresponding organic compounds. We were not able to find any TSs for the associated reaction. Therefore, this reaction should have a bottleneck rather than a TS. Based on the energy profiles, the

reaction is thermodynamically nonspontaneous. In the end, the generated H₂O₂ molecules would give the OH ROS via Fenton reaction, 56 Fe²⁺ + H₂O₂ to Fe³⁺ + OH + OH⁻.

$$O_2^- + BPDEp \rightarrow RC(BPDEp + O_2) \rightarrow PC(BPDEp + O_2)$$

 $\rightarrow BPDEr^- + H_2O_2$ (R12)
 $O_2^- + DMBQpp \rightarrow RC(DMBQpp + O_2) \rightarrow PC(DMBQpp + O_2)$
 $\rightarrow DMBQr^- + H_2O_2$ (R13)
 $O_2^- + PQpp \rightarrow RC(PQpp + O_2) \rightarrow PC(PQpp + O_2)$
 $\rightarrow PQr^- + H_2O_2$ (R14)

Although the obtained reaction barriers are high and step V is nonspontaneous, suggestive of negligible oxidative stress induced directly by the OA components, the roles of BPDE, DMBO and PO in oxidative stress should not be neglected. In the case of BPDE, experiments have suggested that it is the BaP and BPDE metabolites that lead to ROS generation. 51,52 Also, PQ is known for enhancing oxidative stress¹⁴ and consequently causing lung inflammation.⁵⁷ Oginuma et al.⁵⁸ have stated that PQ might pose oxidative stress through inhibition of the carbonyl reductase enzyme, but there are also studies that have proposed redox cycling (the mechanism outlined in Fig. 4, 5 and Fig. S6, ESI†) as the route of oxidative stress caused by

PQ.14 In general, many quinones are known for their involvement in oxidative stress.⁵⁹ The point here is that we have

assessed the direct impact of the OA components on oxidative stress, ignoring the metabolic changes and the enzymatic networks affected by or affecting OAs. Moreover, one of the crucial components in oxidative stress is NADPH whose depletion causes oxidative stress as an outcome of a decreased level of intracellular glutathione.60

Conclusion

NJC

The theoretically examined reaction schemes indicated that the studied OA components are unable to have any direct negative impacts on human health through oxidative stress or DNA alkylation. However, there might be some potential direct/nondirect reaction mechanisms that have not been considered in this study. Particularly, as suggested by earlier experimental studies, the OA metabolites can pose some inevitable health impacts. Based on the ADME predictions, the OA components cannot enter human body through skin. However, as BPDE, DMBQ and PQ enter the system of human body, they can likely permeate cellular membranes, the gastrointestinal tract and the blood-brain barrier, affecting both the gastric and nervous systems. The consequence of their permeation is not necessarily negative, and they should be studied in detail using experimental techniques. Also, as BPDE and PQ (along with BaP and HOP) were predicted to have the potential of inhibiting some cytochrome enzymes, and they can be easily transported by HSA to all metabolism centers, their plausible negative effects on the metabolism of pharmaceutics and metabolic agents should be addressed by further experimental or computational research. Lastly, this study serves as a first step toward understanding the health impacts of organic aerosols at the molecular level, and further in vitro and in vivo experiments are recommended to complement the results and enhance our understanding of the impacts.

Computational details

OA model

Fulvic acid (FA) was selected to represent OAs because (1) it mimics both the aliphatic and aromatic structures in the polycarboxylic acid fraction of OAs, 61 (2) it displays the functionality and behavior of organic PM fragments, 62 (3) 6-11% of the total organic mass extracted from dust aerosols is composed of humic and fulvic acids, 63 and (4) it has been accepted as a suitable surrogate for highly oxidized OAs⁶⁴ and a proxy for atmospheric PM. 65,66 Among various FA chemical structures, the model structure selected and geometry optimized by Sadhu et al. 67 were adopted. This OA model (Fig. 1) is composed of aromatic and non-aromatic moieties, a quinone group, phenolic and heterocyclic hydroxyl functionalities, and multiple carboxylate groups.

ADME properties

The ADME properties of the selected OA components (PQ, DMBQ, BaP, BPDE, HOP and CPPP), the reference drugs (CPT, ThioTEPA and IBPPA) and the OA model were evaluated using SwissADME.68 SwissADME is a web tool that applies machine learning principles, and offers fast and robust models for prediction of the pharmacokinetic, physiochemical and medicinal properties of various molecules. In other words, it describes any input molecule (read as a 2D structure or SMILES notation) by a set of molecular descriptors, and then it crossmatches the descriptors with those of molecules with known properties to predict their ADME performance.

Molecular docking

For analysis of the interaction of PQ, DMBQ, BaP, BPDE, HOP, CPPP, CPT, ThioTEPA, IBPPA and FA with DNA and HSA, molecular docking simulations were performed. The target DNA structures included a DNA structure complexed with the NF-Y transcription factor (pdb (protein data bank)⁶⁹ code: 4AWL;⁷⁰ 25 nucleobase pairs, sequences: TTCTGAGCCAAT-CACCGAGCTCGAT and ATCGAGCTCGGTGATTGGCTCAGAA) and a free DNA domain with the pdb code 3US0⁷¹ (22 nucleobase pairs with the AAACATGTTTATAAACATGTTT and AAA-CATGTTTATAAACATGTTT sequences). For HSA, chain A of the 2BXD⁷² protein structure was taken. The structures of the DNA strands and HSA (with the warfarin ligand removed from the 2BXD crystal) equilibrated in cell-like conditions (using molecular dynamics simulations with identical procedures and similar parameters) were, respectively, adopted from the studies of Keshavarz and Mohammad-Aghaei⁷³ and Alavianmehr et al.74

The docking simulations were performed using Molegro Virtual Docker (MVD) 5.0 software, 75,76 adding the optimized structures (aqueous-phase; see Quantum chemical calculations) of FA, the OA components and the reference drugs as ligands, and considering the ligand-free DNA and HSA structures. Each ligand/target docking simulation was performed by ignoring the possibility of the presence of any other ligand in the binding site. During the simulations, the ligand structures, the DNA strands, and the side chain of HSA were structurally flexible. First, the top ten sterically available regions (probable binding sites; called cavities) of the targets were identified. As MVD basically detects proteins as the docking target and identifies DNA as a cofactor, the cavity searching was manually directed toward the DNA strands by adjusting the cavity search space on them. After that, a spherical search grid with 0.30 Å resolution was adjusted on all identified cavities, and the docking search was restrained to the search grid. The search grid covered almost all parts of the DNA and HSA targets. For each ligand, the docking run was repeated 10 times. At each run, 1500 pose iterations were applied to a maximum of 80 poses (max size) and stopped at the energy threshold of 100 (unitless). One top molecular pose was returned from each run for further analysis. At the end of pose generation, the ligand structure was energy-minimized in the binding site to enhance the energy results. This resulted in 10 of the most energetically feasible ligand/target binding configurations for each ligand and target. After completion of the docking runs, the 10 poses of each ligand corresponding to the 10 runs (ligand/target binding configurations) were inspected visually, the

binding site hosting the highest number of strongly binding poses (i.e. the poses giving the lowest MolDock and Rerank scores) was distinguished as the main binding site, and the interaction of the pose giving the lowest binding energy (the overall interaction energy, in addition to the MolDock and Rerank energies) was analyzed with respect to the strength of interaction and the factors affecting the interaction. A description of the docking algorithm (the hybrid guided differential evolution algorithm) and the scoring functions (which are based on piecewise linear potentials) can be found in ref. 75.

Quantum chemical calculations

All quantum chemical calculations were carried out using Gaussian 16 revision A.03.⁷⁷ For all computations, the ωB97X-D density functional was used in combination with the def2-TZVP^{78,79} basis set to provide long-range exchange-correlation correction, high accuracy of thermodynamics and chemical kinetics analyses, 80-82 and cost-effectiveness. 83 The application of the ωB97X-D/def2-TZVP level of theory was justified by taking the experimentally resolved structures of PQ,84 DMBQ,85 HOP, 86 CPT, 87 and IBPPA, 88 optimizing them and comparing the optimized structures to the experimental geometries. As seen in Fig. S7 (ESI†), the selected computational level gives bond lengths as accurate as 0.04 Å for the organic compounds. However, its accuracy error for reproducing the Pt-N bond length of CPT is 0.23 Å, and it predicts the N-Pt-N and Cl-Pt-Cl bond angles with, respectively, 17.1° and 6.8° error. This can be a result of the inefficiency of the ωB97X-D/def2-TZVP level in precise treatment of the Pt transition metal. Therefore, the performance of the selected computational level is not completely acceptable for quantum chemical simulations of CPT, but it can be reliably used for modeling of the studied organic compounds. Notably, experimental structures were not available for all studied compounds. In the case of ThioTEPA, the starting structure was taken from former theoretical studies.31,74 For BaP and BPDE, the overall structure has no specific conformational flexibility, and no reference starting structure was needed. However, CPPP could pose various conformers with close energy values. Because of the nature of its structure (having multiple alkyl and carbocycle groups), we supposed the barrier toward conformational changes to be low in energy and focused on just one conformer (shown in Fig. 1) for simplicity of the calculations.

To simulate cellular conditions, cellular ionic strength should be considered. However, as cellular ionic strength varies with extracellular environment, 89 all geometry optimizations, (harmonic) frequency and energy calculations (at 310 K and 1 atm), and mechanism explorations were performed in the pure water solvent. We suppose that this approximation is adequate for description of the microenvironment of the evaluated targets, and it does not introduce a large source of error in the energy profiles considering the non-ionic nature of the studied OA components. The aqueous phase was modeled using the SMD polarizable continuum model, 90 which has been commonly used as the best solvent model in many recent studies, e.g. see ref. 91-95. This implicit model can outperform

explicit water treatments⁹⁶ and accurately predict reaction free energy barriers. 97,98 Accordingly, all energy values were directly extracted from the aqueous-phase properties, with the Gibbs free energies and zero-pint energy (ZPE) corrections being calculated using harmonic oscillator and rigid-rotor partition functions.

For analysis of the OA interaction with DNA, a subset of the studied compounds was selected based on their ligand efficiencies (interactions with DNA and HSA during the docking simulations) and their mutagenicity potential (i.e. BaP and BPDE¹⁶⁻¹⁸). Then, the ligand/DNA interactions were analyzed through natural bond-orbital (NBO) analysis on the ligand/DNA complexes obtained from molecular docking (just the binding site and a few additional nucleotides). After the analysis of the NBO results, the main interacting nucleobases were identified, and the ligand/nucleobase complexes were used for mechanism exploration.

To study oxidative stress, we focused on the mechanism suggested by Squadrito et al., 55 which leads to the generation of the O2- radical (and OH; through H2O2 formation) by the intervention of nicotinamide adenine dinucleotide phosphate (NADPH; H - transfer to the organic compounds). The most stable conformers of NADPH and NADP+ were taken from the study of Cao et al. 99 and re-optimized under cell-like conditions at the ωB97X-D/def2-TZVP level (see Fig. S8, ESI,† for their optimized structures). To reduce computational costs, we replaced the full NADPH and NADP+ structures with simple models. Fig. S9 (ESI†) compares the geometries of the models with the full experimental and theoretical NADPH and NADP+ structures and confirms that the models resemble the structural features of NADPH and NADP⁺. Also, the cellular activity of the models was found fairly acceptable by calculating the Gibbs free energy (G) of H⁻ loss from NADPH (NADPH \rightarrow NADP⁺ + H⁻) in cell-like conditions (i.e. aqueous phase, 310 K and 1 atm; 167.9 kJ mol^{-1} for the model vs. 152.2 kJ mol^{-1} for NADPH).

After NADPH/NADP+ model approval, configurational sampling¹⁰⁰ was used to obtain the lowest energy NADPH/ organic compound/water pre-reaction complexes. For this purpose, the aqueous-phase NADPH model, water and organic compounds were imported to the ABCluster 1.4 program^{101,102} and 200 local minima complexes were created using the artificial bee colony (ABC) algorithm, 700 initial guesses, 200 generation rounds and 4 scout bees. To sort the generated complexes and determine the local minima (LM) structures, the energy of the complexes was computed as the sum of coulombic interactions (using atomic polar tensor (APT) atomic charges; ωB97X-D/def2-TZVP) and Lennard-Jones interactions (CHARMM36 force field 103,104). Next, all LM structures were re-optimized using the XTB 6.0.1105 program at the GFNxTB¹⁰⁶ semi-empirical level. Then, a maximum of 35 nonidentical structures meeting the 100 kJ mol⁻¹ energy threshold relative to the most stable complex were filtered and reoptimized at the ωB97X-D/def2-TZVP level. The complex giving the lowest G value was chosen as the global minima (GM) structure and processed to the reaction mechanism explorations, along with three to five low-lying LM structures.

For mechanism investigation, various reaction coordinates of the ligand/nucleobase complexes or organic compound/ NADPH/water systems were scanned. When no transition state (TS) or reaction intermediate was found, several TSs were guessed. In extreme cases, when a TS was suspected but no TS could be found by the two other approaches, the Synchronous Transit-Guided Quasi-Newton (STQN) method 107 (i.e. the OST2 feature of the Gaussian package) was applied to search for the "missing" TSs. Regardless of the approach, all structures were optimized and their harmonic frequencies were computed to label them as intermediates/products/reactions (no imaginary frequencies) or TSs (with imaginary frequency). After that, intrinsic reaction coordinate (IRC)^{108,109} analysis was carried out on the TSs to follow their corresponding reaction paths and connect them to their associated reactant and product sets.

Conflicts of interest

There are no conflicts to declare.

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