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QSAR models for the ozonation of diverse volatile organic compounds at different temperatures†

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In order to assess the fate and persistence of volatile organic compounds (VOCs) in the atmosphere, it is necessary to determine their oxidation rate constants for their reaction with ozone (k_{O_2}). However, given that experimental values of k_{O_3} are only available for a few hundred compounds and their determination is expensive and time-consuming, developing predictive models for k_{O_3} is of great importance. Thus, this study aimed to develop reliable quantitative structure-activity relationship (QSAR) models for 302 values of 149 VOCs across a broad temperature range (178-409 K). The model was constructed based on the combination of a simplified molecular-input line-entry system (SMILES) and temperature as an experimental condition, namely quasi-SMILES. In this study, temperature was incorporated in the models as an independent feature. The hybrid optimal descriptor generated from the combination of quasi-SMILES and HFG (hydrogen-filled graph) was used to develop reliable, accurate, and predictive QSAR models employing the CORAL software. The balance between the correlation method and four different target functions (target function without considering IIC or CII, target function using each IIC or CII, and target function based on the combination of IIC and CII) was used to improve the predictability of the QSAR models. The performance of the developed models based on different target functions was compared. The correlation intensity index (CII) significantly enhanced the predictability of the model. The best model was selected based on the numerical value of $R_{
m m}^2$ of the calibration set (split #1, $R_{
m train}^2$ 0.9834, $R_{\text{calibration}}^2 = 0.9276$, $R_{\text{validation}}^2 = 0.9136$, and $\overline{R_{\text{m}}^2}$ calibration = 0.8770). The promoters of increase/decrease for $\log k_{\mathrm{O}_{\mathrm{x}}}$ were also computed based on the best model. The presence of a double bond (BOND10000000 and \$10 000 000 000), absence of halogen (HALO00000000), and the nearest neighbor codes for carbon equal to 321 (NNC-C...321) are some significant promoters of endpoint increase.

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1. Introduction

Organic structures with high vapor pressure, low boiling point, and low water solubility at room temperature and pressure (293.15 K and 101.325 kPa, respectively) are known as volatile organic compounds (VOCs).¹ VOCs come from two primary sources, namely, anthropogenic VOCs (AVOCs) released from humans and biogenic VOCs (BVOCs) from soil ecosystems. It should be noted that AVOCs are hydrocarbons released by human activities. These compounds are emitted from various daily activities such as industrial processes, traffic, energy production, and the use of solvents, paints, adhesives, lubricants, wear-reducing products, cosmetics, and personal care

items.² Alternatively, BVOCs are mostly derived from microorganisms, plants, and animals.³

Typical VOCs are halogenated compounds, aromatic compounds, aldehydes, ketones, alcohols, and ethers. High concentrations of these VOCs can lead to headaches, nausea, dizziness, and irritation. Unfortunately, significant amounts of VOCs are being emitted into the environment, posing a potentially significant threat to both climate and life.⁴ Also, they secondarily act as ozone/smog precursors and directly as poisonous materials in the environment. Inferior indoor air quality can lead to various short-term and long-term harmful health effects.⁵ In this case, reaction with ozone is a meaningful way to remove most VOCs in the atmosphere.⁶ The kinetic rate constant for the degradation of VOCs is a crucial parameter that must be considered to assess their removal efficiency and the ecological risk of contaminants.⁷

Ozonolysis is a chemical reaction involving the breakdown of organic compounds in the presence of ozone (O_3) . This process plays a central role in atmospheric chemistry, contributing to the formation of secondary organic aerosols and the

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degradation of VOCs emitted by diverse sources. The chemical oxidation process in the atmosphere plays a primary role in the composition of the atmosphere, resulting in the elimination of initially released species and the production of secondary products. In many instances, emitted species or their oxidation products adversely affect the air and climate quality. Among the many ingredients of atmospheric aerosol fragments, organic aerosol particles are less well-known. Secondary organic aerosol (SOA) is a significant component of organic aerosols. Thus, identifying the chemical pathways of compressible products is essential for predicting the formation of SOA. 10-13

Quantitative structure–property relationship (QSAR) is a computational tool for building models to predict various activities. ^{14,15} In this case, different machine learning packages are available to build reliable models. Among them, CORAL is one of the user-friendly packages for building valid QSAR models based on the simplified molecular-input line-entry system (SMILES) notation. ^{16,17} One of the excellent applications of CORAL software is entering the experimental condition into SMILES of a molecule, namely as quasi-SMILES. ^{18–21}

To date, researchers have developed various QSAR models for predicting the reaction rate constants of organic compounds in ozonation reactions. Zhu et al. (2014 and 2015) constructed two optimized QSAR models to estimate the reaction rate constants in ozonation reactions under acidic and neutral conditions at room temperature. These models successfully predicted the reaction rates of diverse organic compounds, yielding the determination coefficients of $R^2 = 0.802$ and 0.723, respectively. In both models, the Fukui indices of a molecule had a notable impact on the reaction rate constants. 22,23 Sudhakaran et al. (2013) developed a QSAR model for the ozone oxidation of organic micropollutants. This model incorporated parameters such as double bond equivalence, solvent accessible surface area, and ionization potential, achieving a notable determination coefficient of 0.832.24 In a separate study, McGillen et al. (2008) employed an SAR model to predict the rates of alkyl substituents. The results indicated a strong agreement between the experimental and predicted values.25

Due to the significant impact of temperature on degradation behavior, it is imperative to incorporate this variable as an independent factor in QSAR models for accurately predicting the reaction rate constants at various temperatures. Recently, several temperature-dependent QSAR models have been developed. For example, Li et al. (2014) devised a QSAR model for room temperature and a temperature-dependent model for the hydroxyl radical oxidation process, demonstrating high goodness-of-fit and robustness measures.²⁶ Similarly, Gupta et al. (2016) established QSAR models for nitrate radical oxidation at room temperature and under temperaturedependent conditions. In a recent study, our group investigated the quantitative relationship between the rate of Fenton oxidation and various parameters, including temperature and quantum chemical and physical-chemical properties of molecules. The findings indicated that temperature exerted the most significant influence on the reaction rate constants.27

Li *et al.* (2013) constructed a QSAR model for predicting ozonation reaction rates at different temperatures, displaying robust predictive capability for 379 reaction rate values, ²⁸ despite the limitation that the molecular weights (MWs) of the studied organics were 200.03 (linalool) or smaller.

Liu et al. (2021) developed QSAR models to predict the rate constant of VOC degradation by O₃. The models were developed based on factors such as bond order, Fukui indices, and other relevant descriptors, in addition to considerations related to temperature. The utilized dataset consisted of $302 \log k_{\Omega}$ values, ranging from 178 to 409 K. This dataset was partitioned into training and test sets for the development and evaluation of the model. The optimized QSAR model demonstrated a favorable determination coefficient for both the training and test sets, achieving R^2 and Q^2 values of 0.83 and 0.72, respectively. These temperature-dependent QSAR models have expanded the applicability domain of traditional OSAR models. However, it is crucial to acknowledge that measured data are subject to errors, impacting the reliability of the models. In this case, utilizing data obtained within the same laboratory can mitigate these errors and enhance the accuracy of the models.

This study aimed to develop a simple and reliable model to predict the rate constants of VOC reaction with ozone at different temperatures based on the Monte Carlo technique. To identify the optimal model, various target functions were assessed through the utilization of the correlation intensity index (CII) and the index of ideality correlation (IIC) employing the CORAL software.

2. Materials and methods

2.1. Dataset

The data set included diverse organic compounds such as alkanes, alkenes, alkynes, and aldehydes. It also included aromatic compounds containing nitrogen, oxygen, and fluorine. Here, $302 \log k_{\rm O_3}$ values in a broad temperature range (178–409 K) for 149 VOCs were obtained from the literature. Leg $k_{\rm O_3}$ was selected as the dependent variable for QSAR modeling, which ranged from -25.3 to -13.92. All QSAR models were constructed using the latest version of the CORAL free software (https://www.insilico.eu/coral).

2.2. Optimal quasi-SMILES descriptors

In the CORAL software, three types of optimal descriptors are available, *i.e.*, SMILES-based, graph-based, and hybrid descriptors (a combination of SMILES and graph) for the creation of QSAR models.^{30,31}

One of the excellent features of the CORAL software is entering the experimental condition with SMILES of the compounds. Here, the experimental temperature was entered as quasi-SMILES. The temperature with a 5° increment was divided, and each increment was defined as [T0], [T1], [T2], etc., as shown in Table 1.

Each quasi-SMILES for each data point was obtained by combining the SMILES with code for temperature [Tx]. Some examples of the created quasi-SMILES and the relevant

Table 1 Defined codes for different temperature ranges to convert the temperature range of experimental data to quasi-SMILES

T(K) range	Code	T(K) range	Code	T(K) range	Code	T(K) range	Code
$T \le 178$	[T0]	$233 < T \le 238$	[T12]	$293 < T \le 298$	[T24]	$353 < T \le 358$	[T36]
$178 < T \le 183$	[T1]	$238 < T \le 243$	[T13]	$298 < T \le 303$	[T25]	$358 < T \le 363$	[T37]
$183 < T \le 188$	[T2]	$243 < T \le 248$	[T14]	$303 < T \le 308$	[T26]	$363 < T \le 368$	[T38]
$188 < T \le 193$	[T3]	$248 < T \le 253$	[T15]	$308 < T \le 313$	[T27]	$368 < T \le 373$	[T39]
$193 < T \le 198$	[T4]	$253 < T \le 258$	[T16]	$313 < T \le 318$	[T28]	$373 < T \le 378$	[T40]
$198 < T \le 203$	[T5]	$258 < T \le 263$	[T17]	$318 < T \le 323$	[T29]	$378 < T \le 383$	[T41]
$203 < T \le 208$	[T6]	$263 < T \le 268$	[T18]	$323 < T \le 228$	[T30]	$383 < T \le 388$	[T42]
$208 < T \le 213$	[T7]	$268 < T \le 273$	[T19]	$328 < T \le 333$	[T31]	$3888 < T \le 393$	[T43]
$213 < T \le 218$	[T8]	$273 < T \le 278$	[T20]	$333 < T \le 338$	[T32]	$393 < T \le 398$	[T44]
$218 < T \le 223$	[T9]	$278 < T \le 283$	[T21]	$338 < T \le 343$	[T33]	$398 < T \le 403$	[T45]
$223 < T \le 228$	[T10]	$283 < T \le 288$	[T22]	$343 < T \le 348$	[T34]	$403 < T \le 408$	[T46]
$228 < T \le 233$	[T11]	$288 < T \le 293$	[T23]	$348 < T \le 353$	[T35]	>408	[T47]

experimental $\log k_{\rm O_3}$ of the VOCs are presented in Table 2. The corresponding quasi-SMILES for the total dataset are presented in Table S1.†

Following the generation of quasi-SMILES, the dataset was divided nine times. Subsequently, each VOC within each split was randomly allocated to the active training (ATRN, 25%), passive training (PTRN, 25%), calibration (CAL, 20%), and validation (VAL, 30%) sets. The quasi-SMILES symbol, split distribution, observed $\log k_{\rm O_3}$ and calculated $\log k_{\rm O_3}$ are presented in Table S1.† The role of each set in the developing QSAR models was previously described in the literature. 32,33

The one variable model used in this study is based on the "descriptors of correlation weights" (DCWs). In the CORAL software, the DCWs for each feature are optimized by the Monte Carlo algorithm. The final QSAR equation is a univariate equation based on the summation of DCWs. Here, the hybrid descriptor was used to build the QSAR models. 34,35 The following equations were used based on optimal descriptors for $\log k_{\rm O_3}$ modeling:

$$DCW(T^*, N^*) = {}^{SMILES}DCW(T^*, N^*) + {}^{Graph}DCW(T^*, N^*) (1)$$

$$SMILES DCW(T^*, N^*) = \sum CW(SSS_k) + CW(BOND) + CW(NOSP) + CW(HALO) + CW(HARD) (2)$$

$$G^{\text{Graph}}DCW(T^*, N^*) = \sum CW(EC2_k) + \sum CW(pt2_k) + \sum CW(pt3_k) + \sum CW(VS2_k) + \sum CW(nn_k) + \sum CW(APP_k)$$
(3)

where T is the threshold and N indicates the number of epochs. T is an integer that divides the SMILES features into active and rare classes. If a molecular feature, F, occurs less than T times, this molecular feature should be removed from the model building (the molecular feature is calculated from SMILES in the training set); therefore, the correlation weight F, CW(F) = 0. Consequently, this molecular feature is known as rare. T^* and N^* are the optimal values of T and N that give the best statistical result for the calibration set. The details of the notation given in eqn (2) are as follows:

The notation details presented in eqn (2) are as follows: SSS_k is fragments of SMILES containing one symbol; the presence/ absence of double ('='), triple ('#'), and stereochemical ('@' or '@@') bonds are indicated by BOND; the presence/absence of nitrogen (N), oxygen (O), sulfur (S), and phosphorus (P) is displayed by NOSP; HALO is the presence of fluorine, chlorine, and bromine; and HARD implies the combination of BOND, NOSP, and HALO. CW(F) demonstrates the correlation weight for the SMILES features, *e.g.*, SSS_k , BOND, NOSP, HALO, and HARD.³⁶

Moreover, in eqn (3), the attribute EC2 is the extended Morgan's connectivity of second order; $\operatorname{pt2}_k$ and $\operatorname{pt3}_k$ are the number of path lengths 2 and 3, which start from the k^{th} vertex of the molecular graph, respectively; VS2 is the valence shells of radius 2 in the hydrogen field graph (HFG); and nn_k is the nearest neighbor code for the k^{th} vertex of the molecular graph. The correlation weights (CWs) were calculated using Monte Carlo optimization.³⁷⁻⁴¹

Table 2 Some examples of the name, temperature reaction, SMILES, code for temperature, quasi-SMILES, and the relevant experimental $\log k_{\rm O_3}$ of VOCs

No.	Name	T(K)	SMILES	Code for $T(K)$	Quasi-SMILES	$\text{Log } k_{\text{O}_3} \text{ (exp.)}$
1 10	Alpha-phellandrene 2,3-Dimethyl-2-butene	295 227	CC(C)C1CC=C(C)C=C1 CC(=C(C)C)C	[T24] [T10]	CC(C)C1CC=C(C)C=C1[T24] CC(=C(C)C)C[T10]	-13.92 -15.05
61 128 242 183 185 251	trans-4-Octene Trimethylamine 1,1,1-Trifluoroethane Tetrachloroethene trans-1,2-Dichloroethene cis-2-Butene	290 296 298 409 380 336	CCC\C=C\CCC CN(C)C CC(F)(F)F ClC(Cl)=C(Cl)Cl Cl\C=C\Cl C\C=C\C	[T22] [T24] [T24] [T47] [T40] [T31]	CCC\C=C\CCC[T22] CN(C)C[T24] CC(F)(F)F[T24] ClC(Cl)=C(Cl)Cl[T47] Cl\C=C\Cl[T40] C\C=C\C[T31]	-16.00 -17.01 -25.30 -18.23 -18.25 -15.71
300	Ethene	193	C=C	[T3]	C=C[T3]	-19.83

Using the APP_k features in the CORAL software is another new conceptual method to improve the predictability of models. APP_k is the vector of the atom pair proportions³⁵ related to fluorine ('F'), chlorine ('Cl'), bromine ('Br'), nitrogen ('N'), oxygen ('O'), double bonds ('='), and triple bond ('#') proportions. APP_k indicates that the compound contains atoms Atom1 and Atom2 and the ratio of Atom1 and Atom2 in the molecule, e.g., 2:1, 1:3, 2:3, and 3:1.

The correlation weights for these events (positions in compounds) can be derived through the Monte Carlo approach. Finally, by calculating the numerical data of DCW (algebraic sum of weights for all features included in the model), the prediction of $\log k_{\rm O_3}$ of VOCs by the least square method is obtained based on the following equation:

$$\text{Log } k_{\text{O}_3} = C_0 + C_1 \times \text{DCW}(T^*, N^*)$$
 (4)

2.3. Monte Carlo optimization

In this study, four distinct types of target functions, namely TF₀, TF₁, TF₂, and TF₃, were employed for the development of robust QSAR models. Subsequently, the resultant statistical outcomes were compared for evaluation.⁴²

The following equations are the mathematical relationship for each target function:

$$TF_0 = R_{ATRN} + R_{PTRN} - |R_{ATRN} - R_{PTRN}| \times dr_{weight}$$
 (5)

$$TF_1 = TF_0 + IIC_{CAL} \times weight for IIC (IIC_{weight})$$
 (6)

$$TF_2 = TF_0 + CII_{CAL} \times weight for CII (CII_{weight})$$
 (7)

$$TF_3 = TF_0 + IIC_{CAL} \times IIC_{weight} + CII_{CAL} \times CII_{weight}$$
 (8)

where the correlation coefficients between the experimental and predicted $\log k_{\rm O_3}$ for the active and passive training sets were denoted by $R_{\rm ATRN}$ and $R_{\rm PTRN}$, respectively. The parameters ${\rm dr_{weight}}$, ${\rm IIC_{weight}}$, and ${\rm CII_{weight}}$ represent the weights assigned to IIC and CII, and they are constant throughout the analysis. Here, the numerical values assigned to the parameters ${\rm dr_{weight}}$, ${\rm IIC_{weight}}$, and ${\rm CII_{weight}}$ were 0.1, 0.5, and 0.3, respectively.

 IIC_{CAL} and CII_{CAL} were computed for the calibration set using eqn (9).

$$IIC_{CAL} = R_{CAL} \times \frac{\min(-MAE_{CAL}, +MAE_{CAL})}{\max(-MAE_{CAL}, +MAE_{CAL})}$$
(9)

positive errors, which were calculated using the following equations:

$$^{-}MAE_{CAL} = -\frac{1}{N} \sum_{y=1}^{N^{-}} |\Delta_{k}| \quad \Delta_{k} < 0, \, ^{-}N \text{ is no. of } \Delta_{k} < 0 \quad (10)$$

$$^{+}\text{MAE}_{\text{CAL}} = +\frac{1}{N} \sum_{\nu=1}^{N^{+}} |\Delta_{k}| \quad \Delta_{k} \ge 0, \, ^{+}N \text{ is no. of } \Delta_{k} \ge 0 \quad (11)$$

$$\Delta_k = \operatorname{Exp}_k - \operatorname{Prd}_k \tag{12}$$

where Exp_k and Prd_k are the experimental and predicted endpoint values, and 'k' ranges from 0 to N.

$$CII_{CAL} = 1 - \sum Protest_k$$

$$Protest_k = \begin{cases} R_k^2 - R^2, & \text{if } R_k^2 - R^2 > 0 \\ 0 & \text{otherwise} \end{cases}$$
(13)

 R^2 is the correlation coefficient for a set with n samples. R_k^2 is the correlation coefficient for n-1 samples of a set after removing the k^{th} sample. Therefore, if $(R_k^2 - R^2) > 0$, the k^{th} substance is an "oppositionist" for the correlation between the observed and predicted values of the set. The more "intensive" correlation appears with the small sum of "protest".

2.4. Domain of applicability

Applicability domain (AD) analysis indicates whether the developed QSAR model can be applied to any set of chemicals. AD is defined based on the theoretical region in the chemical space of molecular descriptors and the activity region modeled by the training dataset. In the CORAL software, AD assessment is done through the probability density distribution. The distribution of the quasi-SMILES features in the ATRN, PTRN, and CAL sets defines AD. Thus, the AD of the model built by Monte Carlo optimization varies depending on the distribution of the datasets in the training and calibration sets. In the CORAL software, the statistical defects of quasi-SMILES are used to define AD. The "statistical defect," d(A) is obtained by the following equation:⁴³

$$Defect_{F_{k}} = \frac{|P_{ATRN}(F_{k}) - P_{PTRN}(F_{k})|}{N_{ATRN}(F_{k}) + N_{PTRN}(F_{k})} + \frac{|P_{ATRN}(F_{k}) - P_{CAL}(F_{k})|}{N_{ATRN}(F_{k}) + N_{CAL}(F_{k})} + \frac{|P_{PTRN}(F_{k}) - P_{CAL}(F_{k})|}{N_{PTRN}(F_{k}) + N_{CAL}(F_{k})} \quad \text{if } F_{k} > 0$$

$$Defect_{F_{k}} = 1 \quad \text{if } F_{k} = 0$$
(14)

The correlation coefficient between the observed and predicted values of $\log k_{\mathrm{O_3}}$ for the calibration set is indicated by R_{CAL} . $^-\mathrm{MAE}$ and $^+\mathrm{MAE}$ are the mean absolute of negative and

where, $P_{\text{ATRN}(F_k)}$, $P_{\text{PTRN}(F_k)}$, and $P_{\text{CAL}(F_k)}$ are the probability of features in the ATRN, PTRN, and CAL sets, and $N_{\text{ATRN}(F_k)}$, $N_{\text{PTRN}(F_k)}$, and $N_{\text{CAL}(F_k)}$ are the frequencies of the features in the ATRN, PTRN and CAL sets, respectively.

The statistical defect of quasi-SMILES was obtained from the sum of the statistical defects of all the features.

$$Defect_{quasi-SMILES} = \sum_{k=1}^{N_{F}} Defect_{F_{k}}$$
 (15)

where $N_{\rm F}$ denotes the number of active quasi-SMILES features for the specified data.

A quasi-SMILES is considered an outlier if:

Table 3 Mathematical equations of goodness-of-fit criteria for QSAR models built using the CORAL software

Type of validation	Criterion of the predictive potential	Ref
Internal	$R^2 = 1 - \frac{\sum (Y_{\rm obs} - Y_{\rm prd})^2}{\sum (Y_{\rm obs} - \overline{Y})^2}$	44
	$Q^{2} = 1 - \frac{\sum (Y_{\text{prd}} - Y_{\text{obs}})^{2}}{\sum (Y_{\text{obs}} - \overline{Y}_{\text{train}})^{2}}$	
External	$Q_{F_1}^2 = 1 - \frac{\sum (Y_{\text{per(test)}} - Y_{\text{obs(test)}})^2}{\sum (Y_{\text{obs(test)}} - \overline{Y}_{\text{train}})^2}$	45
	${Q_{F_2}}^2 = 1 - \frac{\sum (Y_{\mathrm{prd(test)}} - Y_{\mathrm{obs(test)}})^2}{\sum (Y_{\mathrm{obs(test)}} - \overline{Y}_{\mathrm{ext}})^2}$	44 45 46
	$Q_{F_2}^2 = 1 - \frac{\sum (Y_{\text{prd(test)}} - Y_{\text{obs(test)}})^2 / n_{\text{ext}}}{\sum (Y_{\text{obs(test)}} - \overline{Y}_{\text{train}})^2 / n_{\text{train}}}$	
	$R_{\rm m}^2 = R^2 \times (1 - \sqrt{R^2 - R_0^2})$	46
	$CCC = \frac{2\sum(X - \overline{X})(Y - \overline{Y})}{\sum(X - \overline{X})^2\sum(Y - \overline{Y})^2 + n(\overline{X} - \overline{Y})^2}$	47
	$\text{MAE} = \frac{1}{n} \times \sum Y_{\text{obs}} - Y_{\text{prd}} $	
Y-randomization	$C_{R_{\rm p}^2} = R\sqrt{(R^2 - R_{\rm r}^2)}$	48

$$Defect_{quasi-SMILES} > 2 \times \overline{Defect}_{ATRN}$$
 (16)

Defect_{ATRN} represents the average statistical defects for the active training set.

2.5. QSAR model validation

The goodness-of-fit of the generated QSAR models for $\log k_{\rm O_3}$ of VOCs was assessed based on three methods, as follows: (i) internal validation by measuring R^2 , IIC, CCC, Q^2 , and F test in the training set; (ii) external validation by measuring Q^2F_1 , Q^2F_2 , Q^2F_3 , $C_{R_p}^2$, RMSD, MAE, $\bar{R}_{\rm m}^2$, and $\Delta R_{\rm m}^2$ using the test set materials and (iii) data randomization or Y-scrambling (Table 3).

In Table 3, $Y_{\rm obs}$ is the experimental activity; $Y_{\rm prd}$ is the calculated activity; R^2 and R_0^2 are the squared correlation coefficient values between the experimental and predicted property/activity with intercept and without intercept, respectively; and $R_{\rm r}^2$ is R^2 for the randomized models.

Results and discussion

3.1. QSAR models

To obtain predictive and reliable models, nine different QSAR models were constructed for each type of objective function (TF₀, TF₁, TF₂, and TF₃) using hybrid optimal descriptors. Fig. 1 depicts a graphical representation of the attributes and various goodness-of-fit criteria for split #1, as determined by TF₂ using the CORAL software. This figure shows the graphics of the software. The descriptors derived from SMILES and GRAPH are

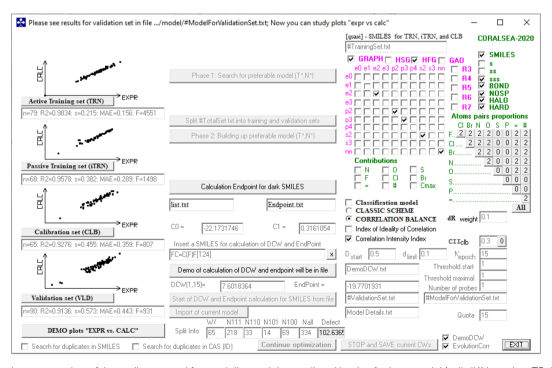


Fig. 1 Graphical representation of the attributes used for modeling and the predicted log k_{O_3} for best model (split #1) based on TF₂ by the CORAL software.

Table 4 Goodness-of-fit criteria for QSAR models developed based on TF₂ for $\log k_{O_2}$ of VOCs

Split	Set	n	R^2	CCC	IIC	CII	Q^2	${Q_{\mathrm{F1}}}^2$	${Q_{\mathrm{F2}}}^2$	${Q_{\mathrm{F3}}}^2$	RMSE	MAE	F	$\overline{{R_{\rm m}}^2}$	$\overline{\Delta R_{\mathrm{m}}^{2}}$	Y-test	$C_{R_{ m p}}{}^{^2}$
1	ATRN	79	0.9834	0.9916	0.7888	0.9882	0.9825				0.215	0.156	4551				0.9768
	PTRN	68	0.9578	0.9710	0.8675	0.9682	0.9554				0.382	0.289	1498				0.9532
	CAL	65	0.9276	0.9592	0.7878	0.9615	0.9230	0.9129	0.9129	0.9224	0.455	0.359	807	0.8770	0.0709		0.9218
	VAL	90	0.9136	0.9464	0.5804	0.9410	0.9086				0.5730	0.4433	937	0.8698	0.0824	0.0141	
2	ATRN	79	0.9650	0.9822	0.9578	0.9749	0.9630				0.308	0.215	2125				0.9568
	PTRN	79	0.9446	0.9662	0.8630	0.9617	0.9416				0.442	0.321	1313				0.9383
	CAL	54	0.8982	0.9416	0.6120	0.9563	0.8893	0.8894	0.8894	0.9200	0.462	0.367	459	0.8364	0.0978		0.8932
	VAL	90	0.9037	0.9501	0.8266	0.9324	0.8998				0.5670	0.4319	823	0.8589	0.0555	0.0093	
3	ATRN	88	0.9866	0.9932	0.8665	0.9901	0.9860				0.191	0.132	6325				0.9832
	PTRN	87	0.9574	0.9777	0.8604	0.9695	0.9556				0.411	0.297	1912				0.9532
	CAL	42	0.9361	0.9231	0.6887	0.9850	0.9268	0.8914	0.7953	0.9450	0.425	0.324	586	0.7224	0.1087		0.9236
	VAL	85	0.8955	0.9368	0.7375	0.9386	0.8897				0.5178	0.4113	712	0.8149	0.1030	0.0148	
4	ATRN	84	0.9707	0.9851	0.8956	0.9815	0.9691				0.229	0.166	2713				0.9648
	PTRN	70	0.9509	0.9736	0.9504	0.9630	0.9481				0.431	0.303	1318				0.9487
	CAL	61	0.9495	0.9641	0.6687	0.9769	0.9412	0.9177	0.9159	0.8759	0.567	0.390	1109	0.8105	0.0684		0.9445
	VAL	87	0.8952	0.9334	0.6880	0.9443	0.8872				0.5348	0.4116	747	0.8102	0.1041	0.0138	
5	ATRN	75	0.9739	0.9868	0.9110	0.9808	0.9725				0.239	0.163	2726				0.9689
	PTRN	80	0.9460	0.9714	0.9139	0.9654	0.9421				0.327	0.223	1365				0.9386
	CAL	61	0.9419	0.9686	0.8129	0.9688	0.9327	0.9415	0.9409	0.8781	0.504	0.361	956	0.8846	0.0587		0.9337
	VAL	86	0.8910	0.9434	0.7194	0.9266	0.8852				0.5330	0.4071	687	0.8412	0.0156	0.0134	
6	ATRN	84	0.9810	0.9904	0.8584	0.9857	0.9801				0.228	0.155	4227				0.9783
	PTRN	71	0.9569	0.9723	0.7565	0.9677	0.9543				0.391	0.282	1532				0.9483
	CAL	56	0.9097	0.9422	0.8120	0.9543	0.9006	0.8668	0.8639	0.8902	0.546	0.432	544	0.7757	0.1033		0.8990
	VAL	91	0.9126	0.9471	0.6621	0.9469	0.9073				0.5758	0.4389	929	0.8488	0.0856	0.0105	
7	ATRN	86	0.9786	0.9892	0.7470	0.9844	0.9776				0.238	0.156	3842				0.9730
	PTRN	80	0.9546	0.9761	0.8704	0.9667	0.9526				0.370	0.256	1641				0.9529
	CAL	43	0.9124	0.9289	0.5774	0.9727	0.9009	0.9098	0.8295	0.9362	0.421	0.317	427	0.7824	0.1000		0.8968
	VAL	93	0.9085	0.9432	0.5673	0.9365	0.9046				0.5763	0.4287	902	0.8661	0.0266	0.0124	
8	ATRN	83	0.9817	0.9907	0.9672	0.9853	0.9808				0.231	0.156	4336				0.9762
	PTRN	73	0.9509	0.9713	0.5904	0.9637	0.9485				0.450	0.319	1375				0.9433
	CAL	59	0.9080	0.9509	0.8728	0.9655	0.8972	0.8983	0.8968	0.9198	0.498	0.408	562	0.8609	0.0876		0.9012
	VAL	87	0.9031	0.9478	0.5651	0.9485	0.8953				0.4817	0.3752	812	0.8387	0.0940	0.0124	
9	ATRN	91	0.9828	0.9913	0.7435	0.9873	0.9821				0.236	0.163	5099				0.9711
	PTRN	71	0.9830	0.9797	0.4955	0.9889	0.9812				0.313	0.264	3999				0.9778
	CAL	52	0.8898	0.9395	0.9119	0.9602	0.8794	0.8696	0.8693	0.9237	0.467	0.392	404	0.8274	0.1050		0.8813
	VAL	88	0.9173	0.9501	0.6652	0.9431	0.9142				0.5463	0.4355	954	0.8787	0.0202	0.0097	

marked in green and pink, respectively. The different types of descriptors selected are marked with a tick mark. Also, the type of the target function and the corresponding coefficients can be seen. In addition, a plot of the predicted values according to the experimental values of $\log k_{\rm O_3}$ can be seen on the left side of the graph.

The goodness-of-fit criteria for all the models obtained by TF_2 are shown in Table 4. The goodness-of-fit criteria for all splits obtained by TF_0 , TF_1 , TF_2 , and TF_3 are indicated in Table S2.†

The comparison of the fit criteria of the models shows that for all models, the \mathbb{R}^2 of the validation set based on TF_2 (eqn (7)) is higher than that of the other target functions. Fig. 2 compares the \mathbb{R}^2 for the validation set across all models obtained based on the four target functions. The \mathbb{R}^2 of the validation set for split 1 (0.9136) calculated based on TF_2 is the highest, and thus this split was selected as the best model.

In the validation of models, apart from evaluating R^2 , it is essential to check the value of MAE. Based on the comparison of this parameter in all the models, it can be concluded that split 1 exhibits the lowest value of MAE (Fig. 3). Therefore, in this

study, TF_2 was chosen as the best target function and split #1 as the best split.

The observed *versus* predicted graph is a valuable tool in modeling to evaluate the performance of a forecasting model.

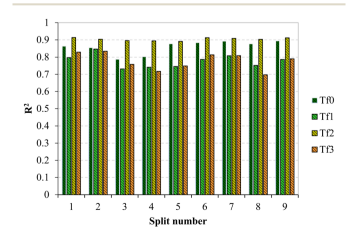


Fig. 2 Comparison of determination coefficients of models constructed based on TF_0 , TF_1 , TF_2 , and TF_3 of all nine splits.

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Fig. 3 Comparison of mean absolute error of models constructed based on TF_0 , TF_1 , TF_2 , and TF_3 of all nine splits.

Split number

Model evaluation, accuracy evaluation, pattern recognition, outlier detection, heterogeneity analysis, and model refinement are several methods in which this chart is helpful. Fig. 4 presents a direct comparison between the experimental values of $\log k_{\Omega_0}$ and the corresponding predictions generated by the model. This visual inspection helps to understand how well the model captures the underlying patterns in the data. By evaluating the proximity of points to the diagonal line (y = x), one can gauge the accuracy of the model. The points near the diagonal line indicate accurate predictions, while deviations from the line suggest discrepancies between the predicted and observed values. Also, the plot helps identify systematic patterns or trends in the predictions by the model. Detecting any consistent overestimation or underestimation can provide insights into potential biases in the model. Outliers, or data points that deviate significantly from the general trend, are shown on the graph. Recognizing and understanding these outliers are crucial for improving the robustness of the model. Heteroscedasticity, which is the presence of non-constant variability in the errors across predicted values, can be observed in the plot. Uneven spreads of points around the diagonal line may indicate varying levels of uncertainty in the model predictions. The insights gained from the graph can guide model refinement. Adjustments, such as feature engineering or modifying the model structure, can be informed by the observed patterns to enhance the predictive accuracy. In essence, the observed vs. predicted plot serves as a diagnostic tool, offering a visual representation of how well the model aligns with actual data. It helps modelers understand the strengths and weaknesses of the model, facilitating informed decisions for model improvement.

As shown in Fig. 4, there are no outliers, and the points near the diagonal line indicate accurate prediction. Furthermore, there is no bias and non-linearity in the reported models.

The following equations represent the QSAR models for predicting the $\log k_{\rm O_3}$ of VOCs from 9 splits by TF₂: Split 1

$$Log k_{O_3} = -22.1732(\pm 0.0087) + 0.3161(\pm 0.0005) \times DCW(1, 15)$$
 (17)

Split 2

$$\text{Log } k_{\text{O}_3} = -22.0600(\pm 0.0151) + 0.2551(\pm 0.0007) \times \text{DCW}(1, 15)$$
 (18)

Split 3

$$\text{Log } k_{\text{O}_3} = -21.9851(\pm 0.0063) + 0.1813(\pm 0.0002) \times \text{DCW}(1, 15)$$
 (19)

Split 4

$$\text{Log } k_{\text{O}_3} = -21.9109(\pm 0.0124) + 0.2606(\pm 0.0006) \times \text{DCW}(1, 15)$$
 (20)

Split 5

$$\text{Log } k_{\text{O}_3} = -21.7750(\pm 0.0115) + 0.2765(\pm 0.0006) \times \text{DCW}(1, 15)$$
 (21)

Split 6

$$\text{Log } k_{\text{O}_3} = -23.1789(\pm 0.0103) + 0.2412(\pm 0.0003) \times \text{DCW}(1, 15)$$
 (22)

Split 7

$$\text{Log } k_{\text{O}_3} = -21.7489(\pm 0.0076) + 0.2546(\pm 0.0004) \times \text{DCW}(1, 15)$$
 (23)

Split 8

$$\text{Log } k_{\text{O}_3} = -22.3377(\pm 0.0088) + 0.2845(\pm 0.0004) \times \text{DCW}(1, 15)$$
 (24)

Split 9

$$\text{Log } k_{\text{O}_3} = -22.5932(\pm 0.0082) + 0.2430(\pm 0.0004) \times \text{DCW}(1, 15)$$
 (25)

Ojha *et al.* (2010) proposed $R_{\rm m}^2$ as a reliable criterion for determining the optimal model.⁴⁹ The best split is split #1, with the maximum average $R_{\rm m}^2$ for the CAL and VAL sets. According to the AD results for the models in Table S3,† 86%, 88%, 85%, 90%, 91%, 91%, 91, 90%, and 86% of the dataset are in the AD models for splits 1–9, respectively. This shows that nine reliable and robust QSAR models can predict more than 85% of the new data.

3.2. Model interpretation

Mechanistic interpretation is one of the basic steps in QSAR modeling. In the CORAL software, the procedure is carried out relying on the structural features extracted from SMILES or HFG, which are responsible for the enhancement or reduction of the targeted activity. If the correlation weight of these structural features is negative in at least three Monte Carlo optimization runs, then these structural attributes are considered activity reduction drivers. Otherwise, if the correlation weights of these structural attributes are positive in at

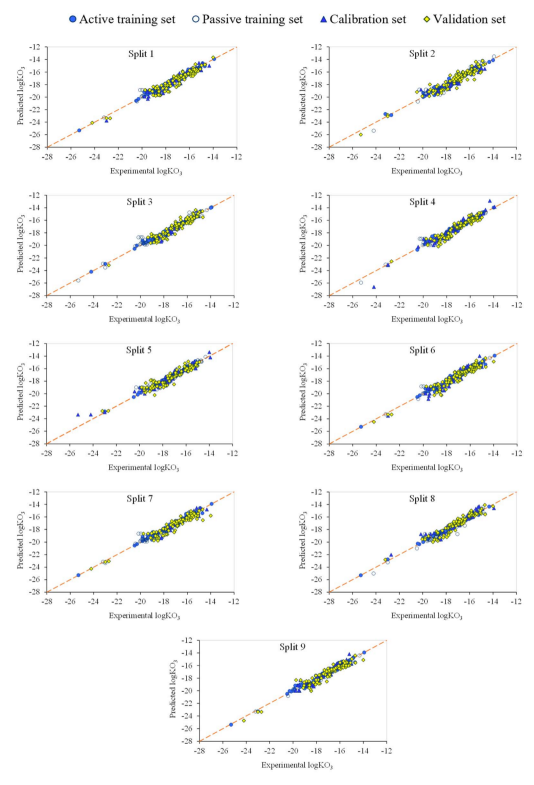


Fig. 4 Plot of the experimental versus predicted $\log k_{O_3}$ of splits 1 to 9 for VOCs based on TF₂.

least three runs, these structural attributes are considered triggers for increasing the activity. However, if the correlation weights of the structural features are positive in some optimization runs and negative in others, the structural features are not considered.

The promoters responsible for an increase/decrease in log $k_{\rm O_3}$ were calculated from the best model (split 1) and are shown in Table 5. The presence of a double bond (BOND10000000 and \$10 000 000 000), absence of a halogen atom (HALO00000000), the number of paths of length two, which started from a carbon

Table 5 The promoters responsible for an increase/decrease in $\log k_{\rm O_3}$ for the best model based on TF₂

No.	Structural attributes	CWs probe 1	CWs probe 2	CWs probe 3	$N_{ m ATRN}{}^a{ m a}$	$N_{\mathrm{PTRN}}^{b}\mathrm{b}$	$N_{\mathrm{CAL}}{}^{c}\mathrm{c}$	Defect	Description
The	The promoters of $\log k_{ m O_i}$ increase	ıcrease							
1	BOND10000000	2.67165	2.51987	3.83649	73	62	61	0.0001	Presence of double bond
2	HALO000000000	1.07055	0.84955	1.74407	89	55	52	0.0005	Absence of halogen
3	PT2-C···5	0.17027	0.12434	0.99446	89	61	09	0.0005	The no. of paths of length 2, which started from a carbon atom, is equal to 5
4	NNC-C…321	0.87169	0.98392	0.73303	61	49	47	0.0005	The nearest neighbor codes for carbon equal to 321
5	PT2-C…2	0.08082	0.02056	0.49237	09	52	46	0.0005	The no. of paths of length 2, which started from a carbon atom, is equal to 2
9	VS2-H···5	0.47711	1.07817	0.36872	58	53	49	0.0002	Valence shell of second order for hydrogen atom equal to 5
^	PT2-C…3	0.22796	0.97566	0.40759	49	35	30	0.002	The no. of paths of length 2, which started from a carbon atom, is equal to 3
8	\$10 000 000 000	1.56878	2.33819	2.97527	48	39	42	0.0004	Presence of a double bond
6	PT3-C···6	0.83176	0.09089	0.24243	48	38	37	0.0005	The no. of paths of length 3 which started from a carbon atom is equal to 6
10	EC2-H…9	0.71934	0.08452	0.1885	44	39	34	0.0004	Morgan extended connectivity of second-order for hydrogen atom equal to 9
11	[T24]	1.62695	1.40185	1.67637	44	32	36	0	Temperature between 124 and 298 K
12	NNC-C…312	0.55561	0.81517	0.50168	28	26	13	0.0038	The nearest neighbor codes for carbon equal to 312
13	VS2-C···6	0.44593	0.00567	0.79277	28	10	15	0.0029	Valence shell of second order for carbon atom equal to 6
14	C···C···=···	0.51624	0.59934	0.09671	27	15	21	0.0004	Two successive aliphatic carbon with double bond
15	$=\cdots C\cdots (\cdots)$	0.40825	0.1176	0.13709	24	13	14	0.0023	Carbon-bonded double bond with branching
1	;								
The	The promoters of $\log k_{O_3}$ decrease	ecrease							
Η	NNC-H…110	-0.08218	-0.20453	-0.02848	71	57	53	0.0007	The nearest neighbors code for hydrogen equal to 110
2	EC2-H···7	-0.21406	-0.25547	-0.08738	63	46	45	0.001	Morgan extended connectivity of second-order for hydrogen atom equal to 7
3	C···=···C	-0.75092	-0.15264	-0.72575	28	46	48	0	Two aliphatic carbons joined by double bond
4	PT3-H…3	-0.03253	-0.00237	-0.24255	53	41	38	0.0009	The no. of paths of length 3, which started from a hydrogen atom, is equal to 3
2	PT3-C…3	-0.28429	-0.0233	-0.53459	39	40	42	0.0019	The no. of paths of length 3, which started from a carbon atom, is equal to 3
9	EC2-C···26	-0.14966	-0.33885	-0.48099	28	15	21	900000	Morgan extended connectivity of second-order for carbon atom equal to 26
7	EC2-H···5	-0.43867	-0.24929	-0.21481	28	22	14	0.0033	Morgan extended connectivity of second-order for hydrogen atom equal to 5
∞	PT2-C···4	-0.52364	-0.36281	-0.64257	56	15	8	0.0061	The no. of paths of length 2, which started from a carbon atom, is equal to 4
6	PT2-C…6	-0.08319	-0.10097	-0.35475	26	13	23	0.0005	The no. of paths of length 2, which started from a carbon atom, is equal to 6
10	$C\cdots C\cdots (\cdots)$	-0.85339	-1.02505	-1.27507	24	13	19	0.0003	Two successive aliphatic carbons with branching
11	EC2-C…22	-0.4121	-0.20762	-0.92869	21	11	16	0.0005	Morgan extended connectivity of second-order for carbon atom equal to 22
12	$C \cdots = \dots (\cdots)$	-0.78929	-1.29912	-1.39466	17	15	12	0.0011	Carbon-bonded double bond with branching
13	NOSP01000000	-0.7395	-0.10479	-0.05487	16	8	7	0.0041	Presence of oxygen
14	VS2-C···13	-0.79019	-0.16384	-1.07209	12	10	14	0.0024	Valence shell of second order for carbon atom equal to 13
15	EC2-C…19	-0.00369	-0.47041	-0.23522	11	7	2	0.0083	Morgan extended connectivity of second-order for carbon atom equal to 19

a Frequencies of SMILES feature in the active training. b Frequencies of SMILES feature in the passive training. Frequencies of SMILES feature in the calibration sets.

atom, is equal to 2, 3, or 5 (PT2-C···2..., PT2-C···3..., and PT2-C···5...), the number of paths of length three, which started from a carbon atom, is equal to 6 (PT3-C···6...), valence shell of second order for hydrogen atom equal to 5 (VS2-H···5...), valence shell of second order for carbon atom equal to 6 (VS2-C···6...), Morgan extended connectivity of second-order for hydrogen atom equal to 9 (EC2-H···9...), two successive aliphatic carbon with a double bond ($C \cdot \cdot \cdot C \cdot \cdot \cdot = = ...$), carbonbonded double bond with branching ($=\cdots C\cdots (\cdots)$, the nearest neighbor codes for carbon equal to 312 (NNC-C···312), and temperature between 353 and 358 K ([T24]...) were some significant promoters of a $\log k_{O_2}$ increase. The nearest neighbor code for hydrogen is equal to 110 (NNC-H···110), Morgan extended connectivity of second-order for hydrogen atoms equal to 5 and 7 (EC2-H···5... and EC2-H···7...), Morgan extended connectivity of second-order for carbon atoms equal to 19, 22, and 26 (EC2-C···19..., EC2-C···22..., and EC2-C··· 26...), the number of paths of length three, which started from a hydrogen atom, is equal to three (PT3-H···3...), the number of paths of length three, which started from a carbon atom, is equal to three (PT3-C···3...), the number of paths of length two, which started from a carbon atom, is equal to four and six (PT2-C···4..., and PT2-C···6...), valence shell of second order for a carbon atom equal to 13 (VS2-C···13...), two aliphatic carbons joined by a double bond (C = C = C), two successive aliphatic carbons with branching (C···C···(···)), carbon-bonded double bond with branching (C...=...(...)), and presence of oxygen (NOSP01000000) were some significant promoters of a $\log k_{\Omega}$ decrease.

Table S4 \dagger presents the correlation weights assigned to each attribute incorporated in the model for split #1 based on TF₂. Another noteworthy observation is that despite the evident

impact of temperature on VOC degradation, as indicated in Table S4,† the correlation weights for temperature (CW(SAK)) are predominantly positive, with the exception of some lower temperatures, where they exhibit a negative trend. Furthermore, a positive coefficient of temperature is also found in increasing descriptors ([T24], temperature between 353 and 358 K), also explaining the positive effect of high temperature on the degradation of VOCs. This conclusion is consistent with the results of the latest OSAR model for this data set.²⁹

3.3. Reliability of QSAR models compared to the best available predictive methods

The literature review shows that only one QSAR model has been reported to predict the rate constants of 302 VOCs with ozone reaction.50 Table 6 compares the goodness-of-fit criteria of the current QSAR model with previous QSAR models. Based on statistical criteria, all the proposed models show a good performance. The datasets for models no. 1, 2, 3, 4, 5, and 7 (Table 6) are relatively small, and the influence of temperature was not considered. Moreover, the previous model was performed with only one partition, but in the current QSAR models, nine partitions were produced to design 36 QSAR models using four objective functions (TF_0 , TF_1 , TF_2 , and TF_3). In this study, two crucial criteria, namely the ideal correlation index (IIC) and the correlation intensity index (CII), were explored. These criteria have not been examined in previous studies. The numerical value of the coefficient of determination for the validation set $(R_{\rm val}^2)$ of the QSAR model obtained by TF $_2$ for split 1 is 0.914, which is better than the proposed model based on the same data.29 Thus, the current QSAR model is more accurate and robust.

Table 6 Comparison of the goodness-of-fit of the developed QSAR model with other reported models

No.	Set	n	T (K)	Descriptor generator package	Regression method	R^2	RMSD	Ref.
1	Total set	117	298	MOPAC and CODESSA	MLR	0.83	0.99	51
2	Training	83	298	DRAGON	MLR	0.88	0.73	52
	Test	42				_	_	
3	Training	103	298	CODESSA	ANN	0.99	0.36	53
	Test	17				0.98	0.46	
	Validation	17				0.98	0.48	
4	Training	93	298	CODESSA	Projection pursuit regression	0.92	0.66	54
	Test	23				0.91	1.04	
5	Training	68	298	Gaussian	Support vector machine	0.86	0.68	55
	Validation	36				0.77	0.77	
	Test	35				_	0.71	
6	Training	306	178-409	MOPAC and DRAGON	PLS	0.840	0.551	28
	Test	73				0.813	0.612	
7	Training	109	295	DRAGON and Gaussian	MLR	0.734	1.05	56
	Validation	27				0.797	0.858	
	Training	109			SVM	0.862	0.801	
	Validation	27				0.782	0.970	
8	Training	242	178-409	Gaussian, Material Studio	MLR	0.83	0.48	29
	Test	60				0.72		
9	ATRN	79	178-409	CORAL package	LR	0.983	0.215	Present work (split 1)
	PTRN	68				0.958	0.382	,
	CAL	65				0.928	0.455	
	VAL	90				0.914	0.573	

Conclusion 4.

Paper

In this study, 36 QSAR models were developed to predict 302 log $k_{\rm O}$ values from 149 VOCs across a broad temperature range (178-409 K). These models were derived from nine random splits of the dataset. The QSAR modeling was done using the CORAL software based on the Monte Carlo approach. The different temperature feature was incorporated in models by considering the quasi-SMILES of compounds instead of SMILES. To investigate the importance of different target functions for the optimization weights of descriptors, four different target functions were used based on IIC and CII or without using these objective functions. The OSAR models using CII (TF₂) produce more predictive and reliable models. All the proposed models provided satisfactory fit criteria for predicting the $\log k_{\mathrm{O_3}}$ of VOCs. However, TF₂ for split #1 was identified as the best model. Various goodness-of-fit criteria such as R^2 , IIC, CII, CCC, Q^2 , Q_{F1}^2 , Q_{F2}^2 , Q_{F3}^2 , s, MAE, F, RMSE, $\overline{R_m^2}$, $\overline{\Delta R_m^2}$, $C_{R_m^2}$ and Y-test were used to assess the reliability and predictive ability of all the proposed models. The applicability domain of the models is defined based on "statistical defect" d(A). Structural features based on both graphs and SMILES were generated from split #1 (considered the best model) and employed to identify the factors promoting either an increase or decrease in $\log k_{O_3}$. The presence of a double bond (BOND10000000 and \$10 000 000 000), absence of halogen (HALO00000000), and the nearest neighbor codes for carbon equal to 321 (NNC-C···321) are some of the significant promoters of endpoint increase. Alternatively, two successive aliphatic carbons with branching $(C \cdots C \cdots (\cdots))$, valence shell of second order for carbon atom equal to 13 (VS2-C···13...), and two aliphatic carbons joined by a double bond (C···=···C...) are some significant promoters of endpoint decrease.

Author contributions

S. Ahmadi designed the study. A. Azimi performed data processing and building the QSAR models. S. Ahmadi and M. Jebeli Javan and wrote the manuscript and performed the interpretation of models. M. Rouhani and Zohreh Mirjafary participated in editing the manuscript.

Conflicts of interest

The authors declare no competing interests.

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