# Machine learning guided design of covalent organic frameworks for CO2 capture in wet flue gas

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#### Abstract

Discovery of remarkable porous materials for CO2 capture from wet flue gas is of great significance to reduce the CO2 emissions, but elucidating the most critical structure features for boosting CO2 capture capabilities remains a great challenge. Here, machine learning assisted computational screening on 516 experimental covalent organic frameworks (COFs) identify the superior secondary building units (SBUs) for wet flue gas separation, which are tetraphenylporphyrin unit in sql-type COFs and functional groups. Accordingly, 1233 COFs are assembled using the superior SBUs. Combined with DFT calculations, the "electron-donating induced vdW interaction" effect is discovered to design better-performing COFs with superior CO2 uptake, which can achieve 4.4 mmol·g-1 with CO2/N2 selectivity of 104.8; while the "electron-withdrawing induced vdW+electrostatic coupling interaction" effect is proposed to design better-performing COFs with superior CO2/N2 selectivity, which can arrive 277.6 with CO2 uptake of 2.2 mmol·g-1, in this case, H2O contributes to improving the CO2/N2 selectivity.

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**Graphic Abstract** 



# Abstract

Discovery of remarkable porous materials for CO<sub>2</sub> capture from wet flue gas is of great significance to reduce the CO<sub>2</sub> emissions, but elucidating the most critical structure features for boosting CO<sub>2</sub> capture capabilities remains a great challenge. Here, machine learning assisted computational screening on 516 experimental covalent organic frameworks (COFs) identify the superior secondary building units (SBUs) for wet flue gas separation, which are tetraphenylporphyrin unit in **sql**-type COFs and functional groups. Accordingly, 1233 COFs are assembled using the superior SBUs. Combined with DFT calculations, the "electron-donating induced vdW interaction" effect is discovered to design better-performing COFs with superior CO<sub>2</sub> uptake, which can achieve 4.4 mmol  $g^{-1}$  with CO<sub>2</sub>/N<sub>2</sub> selectivity of 104.8; while the "electron-withdrawing induced vdW+electrostatic coupling interaction" effect is proposed to design better-performing COFs with superior CO<sub>2</sub> uptake of 2.2 mmol  $g^{-1}$ , in this case, H<sub>2</sub>O contributes to improving the CO<sub>2</sub>/N<sub>2</sub> selectivity.

Keywords: covalent organic framework, wet flue gas separation, machine learning, material design

# **1.Introduction**

In recent decades, the rapidly rising levels of  $CO_2$  in the atmosphere have sparked public concerns around the world<sup>1-3</sup>. Many governments have formulated policies to reduce  $CO_2$  emissions in the midterm and longterm plan including carbon capture and storage (CCS)<sup>4</sup>. However, the current CCS technology relies on high  $cost^{5,6}$ , energy inefficient and corrosive aqueous amine solutions<sup>7</sup>. Alternatively, porous adsorbent materials can be treated as a more effective  $CO_2$  capture process<sup>8-11</sup>. A major source of  $CO_2$  emissions comes from the generated flue gas after burning fossil fuels to produce electricity in power plants. The flue gas mainly contains N<sub>2</sub>(70-75%) ,  $CO_2(15-16\%)$  and H<sub>2</sub>O(5-7%)<sup>11</sup>. Although the H<sub>2</sub>O in flue gas stream is of trace amount, the tolerance of adsorbent to H<sub>2</sub>O is quite important to its actual use<sup>12</sup>. Most conventional adsorbent materials (e.g. zeolites and porous carbon) are subjected to water poisoning<sup>13-15</sup>. Hence, the development of new adsorbents represents a worth pursuing target<sup>16</sup>.

Covalent organic framework (COF) represents a type of crystalline porous material combining organic linkers by strong covalent bonds. The covalent linkage in COFs ensures their application even in some harsh conditions<sup>17</sup>. The advantages like high specific surface area, high porosity and framework tenability endow COFs potentials as promising adsorbents. Li et al.<sup>18</sup> studied water adsorption and the impact on CO<sub>2</sub> capture by GCMC simulations of 18 chemically stable COFs, in which COF-300 exhibit excellent resistance to high relative humidity (p/p0 = 0.8) when used to capture CO<sub>2</sub>. Zhao et al.<sup>19</sup> estimated a series of Schiff-base COFs for wet flue gas separation. It is indicated that keto-COFs of NUS-2 and TpPa-1 outperform the imine-COFs; moreover, they exhibit excellent hydrothermal and chemical stability. Keskin et al.<sup>20</sup> used high throughput computational screening to study the adsorption based flue gas separation performance of 295 COFs. It is found that COF adsorbents could compete with metal-organic frameworks (MOFs) in capturing CO<sub>2</sub> in flue gas, especially most COFs maintain high CO<sub>2</sub> selectivity in the presence of water.

In published works, hydrophobic framework is usually the most recommended structure feature as it is water-resistant. However, hydrophobic pores without sufficient polarity usually show poor separation performance on  $CO_2/N_2$ . Zhao et al.<sup>19</sup> suggested that COFs with

moderate hydrophobicity would be promising adsorbent candidates for practical post combustion  $CO_2$  capture. An in depth and meticulous scientific issue is that what structural unit can commonly make COFs show both high  $CO_2/N_2$  selectivity and  $CO_2$  uptake in humid environment. To answer this question, we need to do two sub-works: 1. a thorough performance evaluation on the present synthesized COFs; 2. a deep mining of the key structure feature that can universally facilitates  $CO_2/N_2$  separation.

Machine learning is an increasingly popular tool in material science due to its powerful ability in analyzing and extracting important feature from big data<sup>21-30</sup>. The machine learning obtained useful information can contribute to design materials in a more effective way. For example, Qiao et al.<sup>28</sup> combined machine learning and molecular fingerprint to identify the excellent bits (aromatic rings, double bonds, transition metals, halogens and oxygen heterocycles) that could promote the non-methane hydrocarbons capture performance of MOFs. Instead of counting the common characteristics that may benefit the property, Boyd et al.<sup>12</sup> used data mining to directly search for the water-resistant CO<sub>2</sub> adsorption sites, named adsorbaphores, from the top-ranked 8325 MOFs. Then, they generated a new set of hypothetical structures that contain the adsorbaphores and synthesized two hypothetical structures with expectant good performance. As is shown, the machine learning techniques develop fast in MOFs that specific substructure features are defined for guiding design of MOFs. By contrast, the application of machine learning in COFs is lately started and is mainly focused on establishing precise correlation model to forecast material properties<sup>31,32</sup>. Machine learning based mining of the excellent substructure features so as to design COFs with optimized performance are not yet reported as far as we know.

In this work, the superior secondary building units (SBUs) of COFs that can promote the  $CO_2$  capture in wet flue gas are identified through machine learning analysis on the separation performance data of 516 computational ready experimental (CoRE) COFs. Then, 1233 potential COFs are constructed based on the superior SBUs. DFT calculations are applied to clarify the separation mechanism of the superior SBUs in facilitating  $CO_2/N_2/H_2O$  separation on the electronic level. By accomplishing the research pipeline of superior substructure mining, material construction and mechanism analysis, we propose the "electron-donating induced vdW interaction" strategy to obtain the better-performing COFs with superior  $CO_2$ 

uptake, and the "electron-withdrawing induced vdW+electrostatic coupling interaction" strategy to obtain the better-performing COFs with superior  $CO_2/N_2$  selectivity. Through combining machine learning, materials genomics based material construction and multi-scale simulation, this work provide specific and in-depth theoretical guidance on designing COFs with both high  $CO_2/N_2$  selectivity and high  $CO_2$  uptake in wet environment.

# 2.Materials and Methods

#### 2.1 CoRE COF

The number of experimentally synthesized COFs in our built CoRE COF database<sup>33</sup> is enlarged to 613. Among them, 97 boric-based COFs in the collected COFs are not calculated in this work considering their instability in the presence of water. Hence, 516 experimentally synthesized COFs (58 3D-COFs and 458 2D-COFs) are estimated for CO<sub>2</sub> capture in wet flue gas. Zeo++ software package<sup>34,35</sup> is used to calculate the structural properties of COFs materials, such as void fraction ( $V_f$ ), surface area ( $S_{acc}$ ), pore limiting diameter (PLD), largest cavity diameter (LCD) for each material,  $S_{acc}$  is calculated using a probe molecule with size equal to the kinetic diameter of N<sub>2</sub> (3.68 Å).  $V_{free}$  is computed with a probe size of 0.0 Å, which is the absolute amount of volume unoccupied by the framework atoms.

#### 2.2 Force fields

The Lennard-Jones (LJ) interaction of COF framework atoms are taken from DREIDING force field<sup>36</sup> as it can accurately describe the gas adsorption behavior of COFs<sup>37</sup>. Partial point charges are assigned to frameworks using charge equilibration method (QEq)<sup>38</sup>. The LJ interaction and atomic charges for CO<sub>2</sub> and N<sub>2</sub> are taken from the TraPPE force field<sup>39</sup>. Previous studies have validated the reliability of the above force filed set in accurately describing CO<sub>2</sub> and N<sub>2</sub> adsorption behavior in COFs<sup>37,40,41</sup>. A set of four commonly used water models<sup>18,42,43</sup>, including TIP4P, TIP4P\_Ew, SPC and SPC/E, are evaluated in our study. The SPC/E model is chosen in our studies due to its computational efficiency and good accuracy to reproduce the experimental absorption performance of COFs (Figure S1).

#### 2.3 GCMC simulation

The separation performances of CO<sub>2</sub>/N<sub>2</sub> (0.1: 0.9) mixture under dry and wet condition

(relative humidity is ~90%) at 1 bar and 298K are studied by Grand canonical Monte Carlo (GCMC) simulations. Leonard Jones (LJ) 12-6 is used to calculate the van der Waals interaction (vdW) parameters between framework atoms and adsorbents. The Coulomb potential is used to consider the electrostatic interactions of N<sub>2</sub>, H<sub>2</sub>O, and CO<sub>2</sub> molecules. A cutoff radius is set to 14.0 Å for the simulation system. The long range interactions are handled by the Ewald summation technique. The fugacity of GCMC simulations are calculated by Peng-Robinson equation of state. For each state point, GCMC simulations consisted of  $1 \times 10^7$  steps to confirm the equilibration, and the following  $1 \times 10^7$  steps to sample the desired properties. The adsorption selectivity ( $S_{CO_2/N_2}$ ) of adsorbent performance evaluation metrics from molecular simulation is calculated from the following definition, where *x* and *y* are the mole fractions of the gases in the adsorbed and bulk phases, respectively.

$$S_{\text{CO}_2/\text{N}_2} = (x_{\text{CO}_2} / x_{\text{N}_2})(y_{\text{N}_2} / y_{\text{CO}_2})$$
(1)

#### **2.4 DFT simulation**

DFT calculations are performed by using  $D3^{44}$  dispersion corrected  $M062x^{45}$ /6-31+g(d)<sup>46,47</sup> as implemented in Gaussian 09<sup>48</sup>. The geometry and electronic structure optimizations of COF SBUs are carried out in gas phase with no constraints imposed beyond the multiplicity of the electronic state, but partial atoms are fixed to keep stacking modes of SBUs unchanged during optimizations. The stability of molecular structure of validated by frequency analysis, and the highest occupied molecular orbital (HOMO) and the electrostatic potential (ESP) of the SBUs are analyzed. Binding energy (BE) between CO<sub>2</sub> and the SBUs are evaluated with basis set superposition error (BSSE) correction by following equation:

$$BE = E_{\text{SBU+CO},} - E_{\text{SBU}} - E_{\text{CO},} - E_{\text{BSSE}}$$
(2)

#### 2.5 Machine learning

In the machine learning process of this work, four algorithms are used to analyze the quantitative relationship between input descriptors and output separation performance of COFs. The algorithms are decision tree (DT), random forest (RF), Xtreme Gradient Boosting (XGBoost), and Categorical Boosting (CatBoost) in scikit-learn library function coded in Python<sup>49</sup>. Here, 90% of CoRE COFs are randomly selected as the training set, and the

remaining 10% are used as the test set. During the training process, each model is trained with Bayesian optimization algorithm to determine the hyper-parameters that could yield the best accuracy according to a 5-fold cross validation. After obtaining the optimal hyper-parameters, each model is retrained on the entire training dataset, and the final prediction performance evaluation is performed on the test dataset. The accuracy of the model is judged by the mean absolute error (MAE), root-mean-square error (RMSE) and R-squared (R 3, as shown in equations  $(3-5)^{32,50,51}$ .

$$R^{2}(x, y) = 1 - \frac{\sum_{i=1}^{n_{samples}} (x_{i} - y_{i})^{2}}{\sum_{i=1}^{n_{samples}} (y_{i} - \overline{y})^{2}}$$
(3)

$$MAE(x, y) = \frac{1}{n_{samples}} \sum_{i=1}^{n_{samples}} |x_i - y_i|$$
(4)

$$RMSE(x, y) = \sqrt{\frac{1}{n_{samples}} \sum_{i=1}^{n_{samples}} (x_i - y_i)^2}$$
(5)

where  $n_{\text{samples}}$ ,  $x_i$ ,  $y_i$  and  $\bar{y}$  denote the number of structures, simulated values from GCMC, machine learning predicted values, and average machine learning predicted values, respectively.

# 3. Results and Discussion

#### 3.1 CO<sub>2</sub> capture from wet flue gas using CoRE COFs

Two key indicators of separation performance in wet flue gas, CO<sub>2</sub> adsorption uptake  $(N_{CO_2})$  and CO<sub>2</sub>/N<sub>2</sub> selectivity  $(S_{CO_2/N_2})$ , are calculated for the 516 COFs under the operating conditions of 298K and 1bar, as shown in Figure 1a. Among them,  $N_{CO_2}$  of 47 COFs exceeds 0.5 mmol g<sup>-1</sup>, and  $S_{CO_2/N_2}$  of 38 COFs exceeds 50. 2D-COFs show better separation performance than 3D-COFs do. The highest  $N_{CO_2}$  and  $S_{CO_2/N_2}$  can achieve 2.87 mmol g<sup>-1</sup> and 360, which are quite outstanding performance in porous adsorbents<sup>52</sup>. However, only 22 COFs outperform when using comprehensive indicators ( $N_{CO_2} > 0.5 \text{ mmol g}^{-1}$  and  $S_{CO_2/N_2} > 50$ ), due to the trade-off effect between  $N_{CO_2}$  and  $S_{CO_2/N_2}$ . To analyze the influence of structure features on CO<sub>2</sub> capture performance, four decision tree based machine learning algorithms (DT, RF, XGBoost, CatBoost) are used. We select seven structural parameters as descriptors that are functional group (FG), interlayer spacing (IS),  $V_{\rm f}$ ,  $S_{\rm acc}$ , PLD, LCD and

density of framework (DF). FG is a category feature described by 1 or 0, representing the COF containing or not containing functional groups. Except for FG, other descriptors are numeric parameters. The energetic parameters (e.g. adsorption heat) are not used because they are the output parameters of GCMC simulations. Figure 1b,c present the prediction results of  $N_{\rm CO_2}$  and  $S_{\rm CO_2/N_2}$  obtained by machine learning and GCMC simulation. DT algorithm uses only one decision tree while other three algorithms use multiple trees, resulting the better fitting and generalization capability of the latter three algorithms. XGBoost and CatBoost are representatives of gradient boosting decision tree algorithms<sup>53</sup>. XGBoost is composed of multiple lift trees and develops trees in a level-wise way, while CatBoost uses oblivious trees as base predictor. Thus, CatBoost is good at dealing with category features, and outperform in this work (CatBoost has the highest R<sup>2</sup> for  $N_{\rm CO_2}$  and  $S_{\rm CO_2/N_2}$  during fitting).



Figure 1. (a) Distribution diagram of the relationship between  $N_{\text{CO}_2}$  and  $S_{\text{CO}_2/N_2}$  of the 516 CoRE COFs under wet condition. (b) Accuracy comparison of four machine learning algorithm by using  $N_{\text{CO}_2}$  and (c)  $S_{\text{CO}_2/N_2}$  as separation performance indicators. DT, RF, XGBoost and CatBoost denote for decision tree, random forest, xtreme gradient boosting, and categorical boosting, respectively.

By using the Catboost model, the contribution (relative importance) of each descriptor in predicting the  $N_{\text{CO}_2}$  and  $S_{\text{CO}_2/N_2}$  can be determined. Here, we use the Shapley's additive interpretation method<sup>54</sup> to evaluate the feature importance values (Figure 2a,b). The higher the importance of a feature, the more it is used in building the boosting tree when predicting the target property. For  $N_{\text{CO}_2}$ , the descriptor importance follows the rank of IS >  $S_{\text{acc}}$  > PLD > LCD > DF >  $V_{\text{f}}$  > FG. Interlayer spacing is a structure parameter of 2D-COFs, a less studied

descriptor compared with the textural properties like  $S_{acc}$ , LCD and so on. A step-further analysis is performed to study the relationship between the adsorption capacity of materials and the interlayer spacing, as shown in Figure 2c. The COFs with high  $N_{CO_2}$  (> 1 mmol g<sup>-1</sup>) are mainly in two types of structures: 1. AB-stacking structures; 2. tetraphenylporphyrin (TPP) contained COFs in **sql** topology with large interlayer spacing of 6.4~8.0 Å, which in turn explain the high contribution of large interlayer spacing in prediction of CO<sub>2</sub> uptake. For  $S_{CO_2/N_2}$ , the descriptor importance follows the rank of IS > DF > FG >  $S_{acc}$  > PLD > LCD >  $V_f$ . By checking the COFs with top 20 high  $S_{CO_2/N_2}$ , we find that fifteen of them are functionalized with polar functional groups like -C=O, -F, -Cl and  $-NO_2$  (shown in Figure S4). The effectiveness of functionalization in improving  $S_{CO_2/N_2}$  is easy to understand because CO<sub>2</sub> is sensitive to electrostatic interaction<sup>55</sup>.



Figure 2. The relative importance of material influencing factors obtained from machine training on (a)  $N_{\text{CO}_2}$  and (b)  $S_{\text{CO}_2/N_2}$  under wet condition. (c) The relationship between the

interlayer spacing of the COFs and  $N_{CO_2}$ . (d) Humid CO<sub>2</sub>/N<sub>2</sub> adsorption configuration of COF-367 containing TPP SBU. (Framework: N, blue; C, gray and H, white; gas molecules: CO<sub>2</sub>, pink; N<sub>2</sub>, brown; H<sub>2</sub>O, green).

#### 3.2 Machine learning guided design of TPP COFs

Machine learning assisted analysis indicates the TPP contained COFs in sql topology with interlayer spacing of 6.4~8.0 Å can bring high CO<sub>2</sub> uptake, while functionalization benefits the CO<sub>2</sub>/N<sub>2</sub> separation. To solve the selectivity-uptake trade-off dilemma, the two favorable features for  $N_{\rm CO_2}$  and  $S_{\rm CO_2/N_2}$  are utilized to build COF adsorbents for wet flue gas separation, as shown in Figure 3a. First, in the parent TPP COFs construction step, TPP with 4-connection sites is used as the center unit, and 21 monomers with 2-connection sites are one by one used as the ligand unit to link with the TPP unit to build the sql network. We get 21 TPP COFs as the parent materials in this step. The structures of 21 parent TPP COFs are shown in Figure S5. Next, in the modified TPP COFs construction step, six common functional groups (-CH<sub>2</sub>CH<sub>3</sub>, -CH<sub>3</sub>, -OCH<sub>3</sub>, -OH, -Cl, -NO<sub>2</sub>) are applied to decorate the 21 parent TPP COFs. Every parent COF is decorated by one type of functional group each time. The decoration site and number depend on the structure of the ligand unit. If the functional group has close contact with the framework atom or adjacent functional group, this modified structure is discarded. Finally, we get 1212 modified TPP COFs. The performance change of the TPP COFs is recorded in Figure 3b. In the parent COF construction step, the highest  $N_{CO_2}$ is increased to 3.4 mmol g<sup>-1</sup> compared with the highest 2.9 mmol g<sup>-1</sup> of the TPP contained CoRE COFs (labeled as CoRE TPP COFs in Figure 3b). In the functionalization step, the highest  $S_{CO_2/N_2}$  is improved to 277.6 (the  $N_{CO_2}$  of this modified TPP COF is 2.2 mmol g<sup>-1</sup>); moreover, the highest  $N_{\rm CO_2}$  is increased to 4.4 mmol g<sup>-1</sup> (the  $S_{\rm CO_2/N_2}$  of this modified TPP COF is 104.8), demonstrating the synergistic effect of the two designing steps in boosting CO<sub>2</sub>/N<sub>2</sub> selectivity and CO<sub>2</sub> uptake. Among the six functional groups, -Cl and -NO<sub>2</sub> decorations contribute to build COFs with the highest CO<sub>2</sub>/N<sub>2</sub> selectivity, while -CH<sub>2</sub>CH<sub>3</sub>, -CH<sub>3</sub>, –OCH<sub>3</sub> and –OH are better at steadily increasing the CO<sub>2</sub> uptake, as shown in Figure 3c.



Figure 3. (a) The assembly scheme of the TPP COFs. TPP is the center unit, and 21 monomers are one by one used as the ligand units to build the parent TPP COFs. Six kinds of functional groups are postly modified to the 21 parent TPP COFs. (b)  $S_{CO_2/N_2}$  and  $N_{CO_2}$  of the 11 CoRE TPP COFs, 21 parent TPP COF and 1212 modified TPP COFs. (c)  $S_{CO_2/N_2}$  and  $N_{CO_2}$  of the CoRE COFs and designed TPP COFs. The performance of designed TPP COFs is classified by functional groups.

To have detailed knowledge of the improvement on both  $CO_2/N_2$  selectivity and  $CO_2$  uptake by functionalization on 21 parent TPP COFs, we calculate the proportion of these modified materials which outperform or underperform their parent COFs. The parameters are described as "proportion of outperformed COFs" (POC) and "proportion of underperformed

COFs" (PUC). The parameters can describe the universality of the functional group in decorating parent COFs with better performance. The corresponding calculation equations are shown as below,

$$POC = \frac{\sum_{i=1}^{21} g(y_i \ge x_i)}{\sum_{i=1}^{21} g(y_i)} \qquad PUC = \frac{\sum_{i=1}^{21} g(y_i < x_i)}{\sum_{i=1}^{21} g(y_i)}$$

where *i* denotes the 21 parent TPP COFs,  $x_i$ ,  $y_i$  denote the target index ( $N_{CO_2}$  or  $S_{CO_2/N_2}$ ) of the parent TPP COFs and the modified TPP COFs, g(...) is the number of the modified TPP COFs.

Here, the dry flue gas separation performance is also estimated to clarify the effect of water on CO<sub>2</sub> capture in TPP COFs. The data classified by functional groups are present in Figure 4a,b. For CO<sub>2</sub> uptake (Figure 4a), in dry flue gas, -CH<sub>2</sub>CH<sub>3</sub>, -CH<sub>3</sub>, and -OCH<sub>3</sub> can make 95%~99% decorated TPP COFs outperform the parent TPP COFs in CO<sub>2</sub> uptake. The same parameter for -OH, -Cl and -NO2 is only 34%~49%. In wet flue gas, -CH3 and -CH<sub>2</sub>CH<sub>3</sub> basically maintain the POC as they do in dry flue gas; -OH, -OCH<sub>3</sub>, -Cl and -NO<sub>2</sub> show obvious decline on POC. The H<sub>2</sub>O uptake shown in Figure 4c can explain the sharp decrease of CO<sub>2</sub> uptake POC to almost zero in -Cl, -NO<sub>2</sub> decorated TPP COFs, because the largely adsorbed H<sub>2</sub>O molecules will expel the CO<sub>2</sub> molecules. For CO<sub>2</sub>/N<sub>2</sub> selectivity (Figure 4b), the presence of H<sub>2</sub>O can promote the positive role of all functional groups on improving the  $CO_2/N_2$  selectivity, especially for -Cl and  $-NO_2$ . The above data imply the following information: 1. enhancing CO<sub>2</sub> uptake is more difficult than enhancing CO<sub>2</sub>/N<sub>2</sub> selectivity in wet flue gas; 2. -CH<sub>3</sub>, -CH<sub>2</sub>CH<sub>3</sub> and -OCH<sub>3</sub> are more universal than -OH, -Cl and -NO<sub>2</sub> in decorating different COFs to have higher CO<sub>2</sub> uptake in both dry and wet flue gas; 3. the performance enhancement mechanisms are quite different in -CH<sub>3</sub>, -CH<sub>2</sub>CH<sub>3</sub> and -Cl, -NO<sub>2</sub> decorated TPP COFs.



Figure 4. The proportion of the modified COFs which outperform (POC) or underperform (PUC) their parent COFs in (a)  $N_{\text{CO}_2}$  and (b)  $S_{\text{CO}_2/N_2}$ . (c) H<sub>2</sub>O adsorption capacity of 1212 functionalized COFs classified by functional groups.

#### 3.3 The role of superior SBUs on boosting separation performance

To clarify these issues, we further analyze the center of mass (COM) probability distributions of the adsorbed CO<sub>2</sub> molecules in the TPP COFs. The parent TPP-No.21 COF is used as an example. As shown in Figure 5a, the CO<sub>2</sub> molecules are mainly adsorbed in the interlayers of the TPP unit and the No.21 unit. This is the common feature of all TPP COFs. Specific to the No.21 unit, CO<sub>2</sub> distributions show differences according to the functional groups. In the  $-CH_3$ ,  $-CH_2CH_3$  and  $-OCH_3$  decorated No.21 units, the CO<sub>2</sub> molecules are evenly and densely distributed (additional figures are shown in Figure S6). It is worth noting that  $-CH_3$  and  $-CH_2CH_3$  themselves are not the adsorption sites but the adsorbed CO<sub>2</sub> molecules in the interlayered multi-phenyl unit are increased.  $-OCH_3$  can served as the weak adsorption sites as we can see CO<sub>2</sub> molecules around the  $-OCH_3$ . For -OH, -Cl and  $-NO_2$  decorated No.21 unit, the functional groups themselves are strong CO<sub>2</sub> adsorption sites, while the density of CO<sub>2</sub> in the interlayered multi-phenyl unit is decreased.

How the functional groups affect the multi-phenyl unit adsorb CO<sub>2</sub> molecules? Density functional theory (DFT) calculations are performed to get the highest occupied molecular orbital (HOMO) to answer the question. No.21 unit with para modification by one type functional group is taken as an example to carry out the HOMO analysis. The HOMO energy levels of the functionalized No.21 unit follow the order of  $-CH_2CH_3 > -CH_3 > -OCH_3 > H > -OH > -Cl > -NO_2$ , as shown in Figure 5b. The less negative HOMO value, the stronger quadrupole- $\pi$  electron interaction between the No.21 unit and CO<sub>2</sub> molecule. Therefore, the

HOMO analysis suggests the electron-donating effect of  $-CH_2CH_3$ ,  $-CH_3$ ,  $-OCH_3$  and the electron-withdrawing effect of -OH, -Cl, -NO<sub>2</sub> on the No.21 unit. The adsorption of CO<sub>2</sub> molecules in the interlayer of No.21 unit relies on the vdW force interaction between molecular quadrupole of CO<sub>2</sub> and an aromatic system. The more the electron on the multi-phenyl unit, the stronger it is to interact with the CO<sub>2</sub> through vdW force. An intuitive observation on the electrostatic potential of the functionalized No.21 units clearly proves the speculation. -CH<sub>2</sub>CH<sub>3</sub>/-NO<sub>2</sub> functionalized No.21 units exhibit the most negative/positive electrostatic potential, indicating the most abundant/insufficient electron of the  $\pi$  aromatic system that can interact with  $CO_2$  molecule (Figure 5c). The electrostatic potential of the – CH<sub>3</sub>, -OCH<sub>3</sub>, -OH, -Cl functionalized No.21 units are shown in Figure S7, which is in accordance with the order of HOMO for the functional groups. Then, the binding energy of  $CO_2$  in the interlayer of the No.21 units is calculated to explore the effect of electron aggregation or loss in the No.21 units on CO<sub>2</sub> adsorption. Figure 5d presents the binding energy follows the order of  $-CH_3 > -OCH_3 > -CH_2CH_3 > H > -OH > -Cl > -NO_2$ . The results declare that utilizing electron-donating group to modify the aromatic unit facilitates the  $CO_2$  adsorption through the quadrupole- $\pi$  interaction.



Figure 5. (a) Contour plots of the COM probability densities of  $CO_2$  adsorbed in  $-CH_2CH_3$ and  $-NO_2$  modified TPP COF using No.21 unit (N, blue; O, red; C, gray and H, violet). Conditions of other functional groups are shown in Figure S6. (b) HOMO values of the

modified No.21 units (c) Electrostatic potential of the  $-CH_2CH_3$  and  $-NO_2$  modified No.21 unit. Conditions of other functional groups are shown in Figure S7. (d) Binding energy of  $CO_2$  in the interlayer of the modified No.21 units (R =  $-CH_2CH_3$ ,  $-CH_3$ ,  $-OCH_3$ , -OH, -Cl,  $-NO_2$ ).

Besides the interlayers of No.21 units, -OH, -Cl and  $-NO_2$  themselves are strong adsorption sites for CO<sub>2</sub> relied on the polar electrostatic interaction (Figure 5a, Figure S6). Figure 6 depicts the CO<sub>2</sub>/N<sub>2</sub> separation performance under dry and wet environment classified by functional groups. In dry flue gas, a certain part of the -Cl and  $-NO_2$  functionalized TPP-COFs can achieve superior high CO<sub>2</sub> uptake of 4~6.3 mmol g<sup>-1</sup>. In fact, -Cl and  $-NO_2$ surpass  $-NH_2$  and  $-CH_3$  in CO<sub>2</sub> uptake under dry environment. In wet flue gas, H<sub>2</sub>O strongly competes with CO<sub>2</sub> for the polar electrostatic interaction sites of -Cl and  $-NO_2$  (Figure 4c), bringing the sharp decrease in CO<sub>2</sub> uptake. Even so, we should also note that the -Cl and -NO<sub>2</sub> functionalization can keep a high CO<sub>2</sub> uptake of 2~3 mmol g<sup>-1</sup> and realize a much higher CO<sub>2</sub>/N<sub>2</sub> selectivity between 180~300 in wet flue gas, because the largely adsorbed H<sub>2</sub>O also impair the adsorption of N<sub>2</sub>.



Figure 6.  $N_{\text{CO}_2}$  and  $S_{\text{CO}_2/N_2}$  of modified TPP COFs in dry and wet flue gas classified by functional groups.

That is to say, the introduction of functional groups initiates the electron transfer in the

aromatic unit, changing the way of CO<sub>2</sub> adsorption. As shown in Figure 7a, in the assembly of "electron-donating group + delocalized  $\pi$  aromatic system", delocalized  $\pi$  aromatic system is the CO<sub>2</sub> adsorption site, and the electron-donating group intensifies the  $\pi$ -quadrupole vdW interaction between the aromatic unit and CO<sub>2</sub> molecules, which is a water resistant adsorption site that can keep superior high CO<sub>2</sub> uptake under wet environment. In the assembly of "electron-withdrawing group + delocalized  $\pi$  aromatic system" shown in Figure 7b, the delocalized  $\pi$  aromatic system is still CO<sub>2</sub> adsorption sites. Under wet environment, the delocalized  $\pi$  aromatic system is kept as the adsorption site for CO<sub>2</sub>, while the electron-withdrawing group turns into H<sub>2</sub>O adsorption site which may also block the N<sub>2</sub> adsorption, and thus high CO<sub>2</sub> uptake and superior high CO<sub>2</sub>/N<sub>2</sub> selectivity can be achieved.



Figure 7.  $CO_2/N_2/H_2O$  separation mechanism in modified TPP COFs (a) assembly of "electron-donating group + delocalized  $\pi$  aromatic system", (b) assembly of "electron-withdrawing group + delocalized  $\pi$  aromatic system". Surface of the functionalized No.21 units are mapped with the  $2 \times 10^{-5}$  electron density isosurface. Color of atom in gas molecules : C, gray; N, blue; O, red; H, white.

### 4. Conclusion

In summary, we use high throughput computational screening combining with machine learning analysis to identify the superior COF SBUs for facilitating  $CO_2/N_2$  separation under wet environment. These superior SBUs, TPP center unit, 21 ligand units and 6 functional groups, are further adopted to construct 21 parent- and 1212 modified-TPP COFs with

potentially high  $N_{\rm CO_2}$  and high  $S_{\rm CO_2/N_2}$ . DFT calculation analysis on HOMO, electrostatic potential and binding energy reveal the influencing mechanism of functionalization on the wet flue gas separation performance of TPP COFs. The interlayers of the aromatic units in the TPP COFs are favorable and hydrophobic vdW adsorption sites for CO<sub>2</sub>. Electron-donating functional groups (-CH<sub>3</sub>, -CH<sub>2</sub>CH<sub>3</sub> and -OCH<sub>3</sub>) reinforce the vdW interaction between the CO<sub>2</sub> and the interlayered aromatic units, enabling the TPP COFs to show high  $S_{\rm CO_2/N_2}$ (70~180) and very high  $N_{\rm CO_2}$  (3~4.5 mmol g<sup>-1</sup>) under wet environment. Electron-withdrawing functional groups (-OH, -Cl, -NO<sub>2</sub>) reinforce the electrostatic interaction in the TPP COFs, which can bring superior high  $S_{\rm CO_2/N_2}$ (150~280) and moderate high  $N_{\rm CO_2}$  (2~3 mmol g<sup>-1</sup>). Our work represents a multi-scale computational study on revealing and then utilizing the critical structure features for boosting CO<sub>2</sub> capture capabilities of COFs, and may provide useful information for designing optimized 2D-COFs adsorbents for wet flue gas separation.

#### **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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