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1. Introduction

To slow the pace of global warming, considerable efforts have been made towards the development of novel strategies for mitigating $CO₂$ emission, particularly for the $CO₂$ capture and sequestration (CCS) in the past two decades.^{1,2} Among numerous materials under study, ionic liquids (ILs) have received remarkable attention from both industry and academia since the first report of using ILs in CCS.³ The outstanding capability of ILs to attract $CO₂$ lies in their unique physical and chemical properties including the high affinity towards $CO₂$, the vanishingly low vapor pressure, the high thermal and chemical stability and the low energy consumption for regeneration.⁴⁻⁶ Benefitting from these distinguished

Interface-enhanced $CO₂$ capture via the synthetic effects of a nanomaterial-supported ionic liquid thin film†

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Ionic liquids (ILs) are effective CO₂ capture media and recent experimental evidence has demonstrated that the addition of two-dimensional (2D) nanomaterials into ILs can effectively improve their $CO₂$ capturing capability. However, an in-depth mechanism on how 2D nanomaterials enhance $CO₂$ absorption is poorly documented. In this study, the adsorption of $CO₂$ by a representative IL, namely 1-ethyl-3methyl-imidazole-tetrafluoroborate ([EMIM][BF4]), coated on graphene (GRA, the prototype 2D nanomaterial) and nitrogenized graphene (C_3N) was investigated by molecular dynamics simulations. The influence of the IL film thickness on the amount of $CO₂$ adsorption was systematically analyzed. Our data clearly indicate that at the IL-gas interface the $CO₂$ accumulation is significantly enhanced. In contrast, at the IL-GRA and IL-C₃N interfaces, only slight enhancement was observed for $CO₂$ accumulation. Quantitative calculations of the adsorption-free energy for CO₂ inside the IL film further support the simulation results. Our present results also reveal that the sub-nanometer IL film possesses a considerably high $CO₂$ capture efficiency because of the formation of the reduced bulk IL region. Moreover, the nanomaterial substrate surfaces can effectively accelerate the diffusion of CO₂, which is beneficial for the $CO₂$ mass transfer. In general, our theoretical study provides a deep microscopic understanding of the $CO₂$ capture by nanomaterials and IL composites. These results could benefit the design and fabrication of a high-performance $CO₂$ capture and storage medium through the synthetic effects of ILs and nanomaterials.

> features, ILs are widely accepted as promising candidates for being economic and environmental-friendly capture-medium for $CO₂$. More importantly, the advantages of ILs are reflected in the abundant types of the composed cations and anions in ILs. Thus, the properties of ILs can be customized to match different CCS requirements.⁷

> Despite the high performance of ILs for $CO₂$ capture, severe drawbacks still exist, which hinder their industrial usage. In particular, the high viscosity induced by the hydrogen-bond network inside ILs unavoidably slows down the kinetic speed for $CO₂$ uptake and separation.⁸ In addition, it is also reported that the absorption of $CO₂$ can in turn increase the viscosity of ILs.⁹ To overcome this drawback, one solution is building supported IL membranes (SILMs) by coating ILs on the surface of solid materials.^{10,11} In previous studies, researchers found that the thickness of IL films is a key factor to attract $CO₂$. Earlier experiments have shown that when the thickness of an IL membrane decreases from the microscale to nanoscale, the $CO₂$ mass transfer rate has a hundred-fold increase.¹² This, according to the diffusion-reaction theory, 13 is because the diffusioncontrolled process in the thick layer is switched to the surface-reaction controlled process when IL thickness decreases. It is also reported that the thin IL film on the

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substrate can enhance the confinement effect, which accelerates its diffusion and the adsorption of $CO₂$,^{14,15} Despite these previous studies, the molecular mechanism of how IL molecules interact with the $CO₂$ gas in the confined space is still poorly documented, which becomes the main hindrance for further experimental research.

Here, molecular dynamics (MD) simulations have been conducted to explore the molecular mechanism of SILM for $CO₂$ capture. The commonly used 1-ethyl-3-methyl-imidazoletetrafluoroborate $[EMIM][BF₄]$ was used as the representing IL model. However, for nanomaterial substrates, we choose graphene (GRA, a prototype 2D nanomaterial) and C_3N (a recently synthesized 2D structure,¹⁶ which was used for controllable $CO₂$ capture and release $17-19$), as representative models. The influence of the IL film of various thicknesses in $CO₂$ capturing was systematically addressed.

Our results clearly indicate that the $GRA/C₃N$ substrates introduced two interfacial faces in the IL film, the substrate-IL interface and IL-gas interface, with distinct IL ordering structures compared to that in the bulk IL. The $CO₂$ capturing efficiency of the supported IL increases with a decrease in the IL film thickness. Among two interfacial regions, the clear accumulation of $CO₂$ is found at the IL-gas interface region, which plays a key role in enhancing the $CO₂$ capturing capability than that with bulk IL. In contrast, the substrate-IL interfacial region plays a minor role in $CO₂$ capture. However, the substrate-IL interface is beneficial for the diffusion of IL and $CO₂$, particularly for graphene, which is beneficial for the $CO₂$ mass transfer. More importantly, our results also reveal that the subnanometer IL film possesses a significantly higher $CO₂$ capture efficiency because of the formation of a reduced bulk IL region. Our present findings provide a deep understanding of the interactions between the SILM components with $CO₂$, which are important to guide the design and optimization of SILMs for the CCS usage.

2. Simulation methods

2.1 Model setup

The GRA nanosheet has a dimension of 2.993 \times 3.455 nm² and is composed of 392 carbon atoms. The C_3N nanosheet has a dimension of 3.104 \times 3.584 nm² and is composed of 96 nitrogen and 288 carbon atoms. Previous studies have shown that compared to the monolayer substrate the multilayer substrates can affect the adsorption of molecules in a limited manner.²⁰ Thus, in our simulation, we only focused on the monolayer substrate. The nanosheet is placed parallel to the $x-y$ plane of the simulation box and the cross-section of the simulation box is equal to the size of the nanosheet. A space of 10 nm is placed along the z direction. Five systems with different numbers of [EMIM][BF₄], which were 20, 50, 80, 110 and 140, coated on the surfaces of GRA and C_3N were prepared, respectively. The thickness of the IL film was estimated from the 2D layer to the IL-gas interface with the IL density being half of the bulk value. After equilibration of 10 ns, the IL film thicknesses were estimated to be 0.57 nm, 1.57 nm, 2.41 nm, 3.26 nm and 4.09 nm for the five GRA systems, and 0.57 nm, 1.43 nm,

2.21 nm, 3.00 nm and 3.76 nm for the five C_3N systems, as demonstrated in Fig. S1 in the ESI.[†] Then, 200 $CO₂$ molecules were randomly added to the vacuum region of each system, as depicted in Fig. 1c and d.

2.2 Simulation parameters

All MD simulations were performed using the GROMACS 5.1.3 package.²¹ The force field parameters for $[EMIM][BF₄]²² CO₂¹³$ and C_3N^{23} were adopted from previous studies in which they were optimized by precise quantum chemistry calculations. During the simulation, a leap-frog algorithm was used to integrate the Newton equations of motion, and the time step was set to 1 fs. All bonds involving hydrogen atoms were constrained using the LINCS algorithm.²⁴ All the atoms in $GRA/C₃N$ were position-restrained to maintain a planar surface during the whole simulation. The cutoff for both the electrostatic and van der Waals (vdW) interactions was 1.0 nm. Long-range electrostatic interactions were calculated using the particle mesh Ewald (PME) method.²⁵ All the simulations were performed under the NVT ensemble, where the V-rescale algorithm²⁶ was applied to the system to maintain a constant temperature of 298 K. Totally 200 ns production trajectory for each system was collected for $CO₂$ molecules to dissolve into the IL film, and the last 30 ns trajectories were used for the data analysis.

2.3 Quantitative interpretation of the $CO₂$ -binding affinity by umbrella sampling

To quantitatively evaluate the binding affinity of $CO₂$ by the IL film, the potential of mean force (PMF) of a single $CO₂$ molecule diffusing from the IL-gas interface into the substrate-IL interface was quantitatively calculated via the umbrella-sampling method.²⁷⁻²⁹ The separation between $CO₂$ and the nanosheet is selected as a collective variable (CV). Totally 20 points were sampled along with the CV. The harmonic force with a constant of 1000 kJ mol $^{-1} \cdot$ nm $^{-2}$ was applied to the CO₂ molecule. Each point was sampled for 10 ns, and the PMF profile was generated using the Weighted Histogram Analysis Method.²⁸

3. Results and discussion

3.1 IL distributions on the substrate

To investigate the role of GRA and C_3N substrates on the IL film, we first characterized the structural features of the IL film on the two substrates. Fig. 2a-d depict the number density profiles of cation $[EMIM]^+$ and ion $[BF_4]^-$ along the normal direction of GRA and C_3N , respectively. Except for the system with an IL thickness of 0.57 nm, two clear peaks, which represented the first and second binding shells of cations (Fig. 2a and b) and anions (Fig. 2c and d) around the substrate, were detected. The high value indicates a strong attraction of the substrate towards IL molecules. Such phenomenon agrees well with the previous reports about the 2D material and IL interaction.14,30 Beyond these two peaks, the density profiles of the ions reach a plateau with only slight fluctuation, until they drop to zero when being far away from the nanosheets. Based on the variation of the density profiles of ILs, the IL film is divided into three regions:

Fig. 1 Structures of cation [EMIM]⁺ (a) and anion [BF₄]⁻ (b), and schematic representations of the simulation box for IL coated on GRA (c) and C₃N (d).

 (1) the substrate-IL interfacial region that covers the first two binding shell of IL, (2) the bulk IL region where IL is evenly distributed, and (3) the IL-gas interfacial region where the IL density starts to diminish. For the three regions, the thicknesses of the two interfacial regions are relatively constant, except that the bulk IL region varies in different systems. In the extreme case of the IL film with a thickness of only 0.57 nm (the black lines in Fig. 2a–d), the substrate-IL and IL-gas interfacial regions merge without the bulk IL region.

3.2 The $CO₂$ assembly in the IL film

The $CO₂$ molecules were initially placed outside the IL film. During the simulations, varying number of $CO₂$ molecules adsorbed and intruded into the IL film. Fig. 2e and f depict the number density profiles of $CO₂$ along the normal direction of

GRA and C_3N . For each system, there are two detectable CO_2 peaks at the substrate-IL interface, which represent the direct attraction of GRA and C_3N towards CO_2 . In addition, one dominant peak can be detected for the density profile, which was located at the edges of the IL film, particularly, at the IL-gas interface. When the thickness of the IL film decreases, this peak shifts according to the IL-gas interface. For the thinnest IL film with a thickness of 0.57 nm, the two accumulative regions of $CO₂$ merge into one. More importantly, $CO₂$ accumulation inside a 0.57 nm IL film is significantly enhanced than that in other systems, as indicated by the highest peak among the five systems. This reflects synergistic effects from both the substrate-IL and IL-gas interfaces in the thinnest IL film. Moreover, the $CO₂$ densities in the bulk IL regions are found to be constant, which are considerably lower than those found in

Fig. 2 Number density profiles of different compositions. (a and b) Represent the number density profiles of cations; (c and d) represent the number density profiles of anions; (e and f) represent the number density profiles of CO_2 molecules in the C_3N and GRA systems.

the two interfacial regions, independent of the variation of the IL film thickness. Fig. 3a depicts the total number of captured $CO₂$ with respect to the thickness of IL film. In the studied thickness region, roughly a linear dependency is found for both the GRA and C_3N systems. This is because when the IL film thickness increases, more $CO₂$ can be stored inside. However, when the $CO₂$ storage is normalized by the number of IL molecules (storage efficiency of the IL), which is depicted in Fig. 3b, the $CO₂$ storage efficiency increases quickly with the decrease in IL thickness, which is well consistent with SILM experimental reports.³¹ These results are expected because a high-density layer of $CO₂$ is formed at the 0.7-1.1 nm from the substrates (Fig. 2e and f) where the IL density becomes considerably low (Fig. 2a–d). From these results, it is also deduced that the bulk IL region has lower $CO₂$ storage capability than two interfacial regions.

From the above-mentioned results, the role of two substrates is highlighted in the introduction of the two interfacial regions, which are rather attractive to $CO₂$ than to bulk IL. To quantitatively distinguish the roles of interfacial regions in attracting $CO₂$, we classified the captured $CO₂$ according to their locations, in detail: the substrate-IL interfacial region, the IL bulk and the IL-gas interfacial regions. As summarized in Table S1† (GRA systems) and Table S2† (C₃N systems), $CO₂$ residing inside the two interfacial regions are relatively constant despite the IL film thickness. More importantly, the IL-gas interface plays a more dominating role in the $CO₂$ storage than does the solid-IL interface. For the bulk IL region, the $CO₂$ storage efficiency is also considerably lower than that of the IL-gas interface.

In addition, no significant differences were detected between the GRA and C_3N systems. This is because the regulation of the substrate to the IL structure is short-ranged, which is limited to the solid-IL interface, and the contribution of the solid-IL interface in $CO₂$ storage is rather small. This is consistent with the previous report on using hydrophilic TiO₂ as $SIM¹⁴$ where the $CO₂$ storage in the substrate-IL interfacial region was found to be small. Taken together, these results suggest that the capability of SILM is not sensitive to the specific type of supporting materials, which are mainly treated as substrates in

3.3 Thin IL film facilitates quick $CO₂$ diffusion

the SILMs instead of amplifiers for the $CO₂$ capture.

Except for the gas capture process, the $CO₂$ mass transfer in the IL film is another determinative factor to evaluate the performance of the medium because a high rate of mass transfer is typically required. In addition, we studied the diffusion character of $CO₂$ molecules in the IL film. Technically, the mean square displacement (MSD) of $CO₂$ molecules along the transverse direction of the IL film was calculated. Then, the lateral diffusion coefficient (D) was fitted based on Einstein's relation $D = \lim_{t \to \infty} \frac{1}{4i}$ $\frac{1}{4t} \langle [r_i(t) - r_i(0)]^2 \rangle$, where t is the time, and $r_i(t)$ and $r_i(0)$

are the positions of $CO₂ i$ at time t and time 0, respectively.

Fig. 4a presents the specific values of D of the dissolved $CO₂$ in the GRA and C_3N systems with different IL film thicknesses. It is found that the diffusion of the dissolved $CO₂$ in both systems shows a decreasing trend with the increase in the IL thickness. According to the diffusion-reaction theory,¹³ the bulk IL region was deduced to act as major resistance for the $CO₂$ diffusion. A thinner IL film can enhance the $CO₂$ diffusion. Accordingly, it is notable that there is a clear increase in the $CO₂$ diffusion at the IL thickness of 0.57 nm where the bulk IL region totally vanishes. These results suggest that the introduction of the interfacial IL regions by substrates greatly facilities the $CO₂$ diffusion. This is consistent with previous experimental reports that $CO₂$ exhibits faster kinetics in IL with substrates than that in pure IL.^{11,32,33}

As in the IL medium, the diffusion of $CO₂$ is highly related to the cations and anions. We calculated the lateral diffusion coefficients of the cations and anions in both the GRA and C_3N systems (Fig. 4b) and found a similar vertiginous decrease tendency with the IL film thickness. Shi et al. reported that ions in IL diffuse faster with confinement from nano-materials at room temperature.¹⁵ In our simulations, the confinement rate of IL by nano-sheets decreases with the IL film thickness, thus

Fig. 3 (a) Total number of captured CO_2 with respect to IL thickness; (b) the CO_2 capture efficiency of the IL films with different thicknesses calculated by dividing the number of captured $CO₂$ by the total number of IL molecules.

Fig. 4 Self-diffusion coefficients of the dissolved $CO₂$ molecules (a) and IL ions (b).

the ion diffusion is the fastest in the systems with the smallest IL film thickness. Additionally, the diffusion coefficients of ions in the C_3N systems are found comprehensively lower than those in the GRA systems. This is attributed to the stronger surface-IL interactions, which slow down the self-diffusion of ions, as reported by Shi et al.¹⁵ We calculated the interaction energy of one ion binding onto GRA (-48.34 \pm 11.24 kJ mol⁻¹ for [EMIM]⁺ and -13.53 ± 3.82 kJ mol⁻¹ for $[BF_4]$ ⁻) and C_3N (-49.02 \pm 10.55 kJ mol⁻¹ for $[EMIM]^+$ and -14.76 ± 3.32 kJ mol⁻¹ for $[\text{BF}_4]^-$). For both the cation and anion, their interactions with C_3 N are slightly higher than those with GRA. This explains the slower ion diffusion in the C_3N systems than that in the GRA systems.

3.4 Quantitative interpreting $CO₂$ -binding affinity and structural mechanisms

To gain a deep understanding of the binding behavior of $CO₂$ inside the IL film, the binding affinity changes of a $CO₂$ molecule translocating from the gas region into the IL interior is studied by calculating the potential of mean force (PMF) by the umbrella-sampling method. As summarized in Fig. 5, the PMF profiles illustrate that the global minimum, around -6 kJ mol⁻¹, is located at the IL-gas interface in both GRA and C_3N systems, which is well consistent with the CO_2 distribution shown in Fig. 2c and f. In addition, there are two local minima near the two nanosheets, corresponding to the two density peaks of $CO₂$ at the solid-IL interface in Fig. 2c and f. These minima, particularly those at the IL-gas interface, reflect the strong attraction of interfacial regions to $CO₂$ molecules and the accumulation of $CO₂$. In the bulk IL, the PMF profiles are relatively flat and generally higher than those at the IL-gas interface, corresponding to the overall $CO₂$ storage capability of IL itself.

To probe the atomic mechanism of such patterned $CO₂$ distribution inside the IL film, we further accessed the microscopic structure of the molecule assembly in the IL film by calculating the orientations of the $[EMIM]^+$ cations and CO_2 . The $[BF_4]$ ⁻ anions were ignored because of their tetrahedron shapes. We defined two tilt angles for $[EMIM]$ ⁺ to describe the orientation of the cation: one is the angle between the ethyl

Fig. 5 Potential of mean force (PMF) for CO₂ diffusion from the IL-gas interface to the substrate-IL interface. The dashed lines represent the position of the two interfaces.

chain and the normal direction of the substrate (Fig. 6a); the other is the dihedral angle between the imidazole ring plane and the substrate (Fig. 6e). The orientation of the $CO₂$ molecule is described by the angle between the linear $CO₂$ and the normal direction of the substrate (Fig. 6i).

We used the GRA system as a representative for discussions, and three IL films with thicknesses of 0.57, 2.41 and 4.09 nm were compared, as shown in Fig. 6. The corresponding results for the C_3N system are generally similar, as shown in Fig. S2 in ESI. \dagger In a thin IL film, orientations of the cations are strongly influenced by the nanosurface. The orientations of the IL cations and $CO₂$ are highly biased at the substrate-IL and IL-gas regions. From the angle distributions, we observed two distinguishable spots for the ethyl chain tilt angle (Fig. 6b), which are labelled as s1 (around 50°) and s2 (90[°]), respectively. Two indistinguishable spots for the imidazole ring tilt angle (Fig. 6f), which are both labelled s3 (at equivalent 0° and 180°), can also be detected. From this information, it is seen that the cations stack to the substrate with the imidazole ring parallels to the substrate surface. However, for the ethyl chains, they are flexible enough to be alternatively attached to or detached from the surface, as illustrated by the two representative structures embedded in Fig. 6b. In addition, the $CO₂$ molecules at the substrate-IL interfacial region adopt a parallel binding pose to the nanosurface (as shown in Fig. 6j) because the tilt angle is located at around 90° (Fig. 6j, s4). Such a binding mode maximizes the intimate contacts between $CO₂$ and the substrates, representing the stronger attraction from the nano-surface than from IL.

It is notable that the three tilt angles are highly conserved in the substrate-IL interfacial region, regardless of the IL thickness. This is also true for the tilt angles of the ethyl chains, which are restrained between 30° and 60° (Fig. 6c and d, s5), indicating that the ethyl chains of $[EMIM]^+$ point to the gas phase. This is in line with the finding that more ethyl groups are located at the IL-gas interfaces than at the imidazolium rings (Fig. S3†). Such structure is also consistent with previous experimental reports that, at the IL-gas interface, a longer alkyl chain of cations tends to point to the gas phase.³⁴ As for the $CO₂$ molecules at the IL-gas interface, a parallel orientation along the surface is also favored although the specific angle spreads in a range of $60-120^\circ$ (Fig. 6j-l, s6). Such orientation also ensures intimate binding of $CO₂$ with IL.³⁵ In general, from these analyses, the assembled $CO₂$ in two interfacial regions are accompanied by the directionally-oriented $CO₂$ and cations, indicating structural compatibility among them.

From the above-mentioned analyses, the orientations of $CO₂$ in IL are highly regulated by the two interfacial regions. In addition, the dynamic characteristics of the $CO₂$ molecules in the two interfacial regions are also different, as revealed by the residence time of $CO₂$ (Fig. S4†). The $CO₂$ residence time at the IL-gas interface distributes mainly around 40–50 ps in both GRA systems and C_3N systems, which indicates the stable CO_2 adsorption at the IL-gas interfaces. In contrast, the residence time at the substrate-IL interface is more dispersed than that at the IL-gas interface (Fig. S4†). It is found that the binding at the substrate-IL interface can either last for more than 100 ps (mainly representing $CO₂$ molecules in the first adsorption layer

Fig. 6 Definition and distribution of the tilt angles of the ethyl side chain of cations (a–d), imidazolium ring of cations (e–h) and CO₂ (i–l) in three IL films attached to GRA with the IL thickness of 0.57 nm, 2.41 nm and 4.09 nm, respectively.

of the substrate-IL interface) or less than 10 ps, which represent $CO₂$ molecules that cannot form a significant residence in the IL-gas interface region.

4. Conclusion

Using all atomic molecular dynamics simulations, we studied the structural and energetic characteristics of SILM with $CO₂$ capture. By coating different number of $[EMIM][BF₄]$ pairs onto the nanosheet, the influences of the IL film thickness are systematically discussed. First, the $CO₂$ accumulation at the ILgas interfacial region dominates the $CO₂$ capture by SILM, suggesting the minor role of specific substrate materials. In general, the $CO₂$ capturing efficiency of supported IL increases with a decrease in the IL film thickness. A sub-nanometer IL film possesses the highest $CO₂$ capture efficiency because of the formation of a reduced bulk IL region. The nanomaterial surfaces can effectively accelerate the diffusion of $CO₂$, which is beneficial for the $CO₂$ mass transfer. The current study provides deep understanding of the molecular interactions at the interfaces of SILMs, and paves the way for the design and fabrication of optimized supporting nanomaterials of SILM for the CCS usage.

Conflicts of interest

There are no conflicts to declare.

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