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# New adsorbent materials based on PILs for Freon refrigerants

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The development of refrigerant adsorbent materials is not only essential for enhancing the efficiency of refrigeration systems but also plays a pivotal role in environmental conservation and addressing global warming challenges. However, traditional adsorbent materials are often limited in widespread applications in industrial scales due to various disadvantages, such as low adsorption efficiency, difficulties in desorption, and poor reusability. In this context, three distinct PILs, P[EVIM][PF<sub>6</sub>], P[BVIM] [PF<sub>6</sub>] and P[HVIM][PF<sub>6</sub>], were synthesized and characterized. In addition, their structure as well as adsorption capacities towards three different Freon refrigerants (R12, R22 and R134a) were explored. The results indicated that the synthesized PILs had high thermal stability and exceptional adsorption capabilities, with P[EVIM][PF<sub>6</sub>] demonstrating the best adsorption performance. These PILs consistently maintain a stable saturated adsorption capacity throughout nine consecutive adsorption—desorption cycles, and the desorption rate of the adsorbent tubes consistently exceeded 96%. Thus, the superior recyclability of these PILs was verified. These PILs provide a promising route for efficient adsorption of Freon refrigerants, highlighting their potential significance in pertinent industries and environmental conservation efforts.

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## 1 Introduction

Global warming is a pressing environmental issue, primarily attributable to greenhouse gas emissions. Various industries are seeking methods to reduce carbon emissions in response to this challenge. Particularly in the refrigeration industry, recycling and regenerating refrigerants for reuse not only aids in addressing the pressure of greenhouse gas emission reductions, but also effectively alleviates the demand pressure for refrigerants. The Montreal Protocol, a global accord on environmental matters, was designed to safeguard the ozone layer by gradually eliminating substances that reduce it.1 This protocol underscores the importance of refrigerant recycling and reuse to manage these compounds properly, reducing environmental pollution and resource wastage.2 In implementing the Montreal Protocol, many countries have adopted a series of innovative strategies and technologies to promote the recycling, reuse and regeneration of refrigerants. These strategies include establishing recycling systems, enhancing recovery efficiency, and developing alternative refrigerants that have less impact on the ozone layer and climate. Through these efforts, the international community is advancing towards a more sustainable and

environmentally friendly refrigeration industry for long-term environmental conservation and global climate stability.<sup>3,4</sup>

Against this broad backdrop, the recovery of chlorofluorocarbon refrigerants from discarded equipment has become an important research direction in environmental protection. In adsorption separation, ionic liquids (ILs) provide a new solution for green chemical processes. Lee et al.5 prepared an ionic liquid based on 1-hexyl-3-methylimidazolium (HMIM) and evaluated its performance in selectively absorbing refrigerants R134a and R1234yf. It was found that this material had superior refrigerant absorption performance for R134a compared to R1234yf, suggesting the possibility of selective separation using ionic liquids. Nevertheless, in the practical application of ILs, researchers have found that additional steps may be required to separate the target product from the ILs when ILs exist in a liquid state. Meanwhile, the difficulty in shaping ILs adversely affects the efficiency of gas adsorption. These issues limit the effectiveness of these materials in practical applications.

Fortunately, scholars discovered a significant increase in adsorption capacity when the ionic liquid was transformed into a PIL compared to the ILs.<sup>6</sup>

Polymeric ionic liquids (PILs) are new materials developed on the basis of ILs. Similar to ionic liquids (ILs), polymeric ionic liquids (PILs) not only exhibit superior surface properties but also demonstrate significant thermal stability, chemical stability, and radiolytic stability.<sup>7-11</sup> These characteristics endow

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PILs with tremendous potential in a variety of application fields, particularly in environments with high temperatures, harsh chemical reactions, and radiation exposure.<sup>12,13</sup> Compared to ILs (existing in liquid or molten states at low temperatures), PILs usually exist in solid states, <sup>14</sup> with a small amount in-gel states. Like ILs, PILs possess favorable surface properties.<sup>11,12</sup> Additionally, PILs exhibit specific characteristics of polymers (*e.g.*, excellent mechanical stability).<sup>15</sup> These properties may make some PILs competitive candidates for the best refrigerant adsorbents. PILs have be extensively applied to gas adsorption separation, especially in the capture and utilization of carbon dioxide and carbon monoxide.<sup>16-18</sup> Nevertheless, the researches

on the use of ionic liquids in adsorptive separation of refriger-

Therefore, this study aims to prepare a novel refrigerant adsorbent material based on PILs. In this study, three types of linear vinylimidazolium class polyionic liquids, including P [EVIM][PF<sub>6</sub>], P[BVIM][PF<sub>6</sub>], and P[HVIM][PF<sub>6</sub>], were synthesized, using the polymerization of 1-vinyl-3-alkylimidazolium hexafluoro-phosphate ([CnVIM][PF<sub>6</sub>], n = 2, 4, 6, 8) ILs double bonds, through the free radical polymerization method. The adsorption and desorption properties of the three types of PolyIonic Liquids (PILs) materials were evaluated under three Freon refrigerants R22, R134a, and R12. Concurrently, the adsorption heat and desorption activation energy of the three gases on the three PILs materials were determined in a scholarly and professional manner according to Inverse Gas Chromatography (IGC) and intrinsic kinetic methods. The adsorption and desorption effects, reusability of the three PILs materials adsorption tubes and activated carbon adsorption tubes for the three refrigerants were evaluated experimentally. The in-depth study and optimization of PIL provide a more effective solution for the recovery of refrigerants from waste equipment.

# 2 Experiment

ants are still relatively limited.

#### 2.1 Materials

monomers included 1-ethyl-3-vinylimidazolium hexafluorophosphate ([EVIM][PF<sub>6</sub>]), 1-butyl-3-vinylimidazolium hexafluorophosphate ([BVIM][PF6]) and 1-hexyl-3-vinylimidazolium hexafluorophosphate ([HVIM][PF<sub>6</sub>]). The IL monomers with a mass fraction of more than 0.99 were purchased from Chengjie Chemical Co. Ltd, China. In the free-radical polymerization, the initiator and solvent used were 2,2'-azobis (isobutyronitrile) (AIBN) and N,N-dimethyl-formamide (DMF) respectively. Ethanol was used as a solvent for precipitation. This study used Freon refrigerant gases purchased from DuPont Trading Co. Ltd, including dichloro-difluoromethane (R12), chlorodifluoromethane (R22), and 1,1,1,2-tetrafluoromethane (R134a). All of which had mass fractions exceeding 0.99. All the solvents and probes utilized for the test were of high purity and obtained from J&K Scientific without further purification.

#### 2.2 Preparation and characterization of PILs

The essential PILs were produced *via* free radical polymerization process of equivalent monomers, as shown in Fig. 1. The

"R" substituent represents an alkyl chain attached to the imidazole ring. The value of "n" indicates the number of carbon atoms in the alkyl chain, reflecting the length of the side chain. Specifically, when n equals 2, "R" denotes an ethyl group. "X\*" signifies the average degree of polymerization of the polymer chain. The asterisk (\*) indicates that this value represents an average.

For instance, the preparation of P[EVIM][PF<sub>6</sub>] involved sequential additions of the ionic liquid monomer, azobisisobutyronitrile (around 1% of the monomer weight) and N,N-dimethylformamide to a round-bottom flask, which was stirred at ambient temperature for thorough mixing. The mixture was heated to 343.15 K in an oil bath with  $N_2$  safeguarding and magnetically stirred continuously for a reaction duration of 24 hours. After the reaction, the compound was transferred to a beaker, and a sufficient volume of ethanol was introduced to induce polymer precipitation. Prior to precipitation, the residue was rigorously rinsed by ethanol to eliminate the instigator and unprocessed ILs monomer. The resulting dense solid was then dried in a vacuum oven and kept at 343.15 K for 24 hours. Once dried, the substance was crushed in a mortar and pestle. It was rinsed three times with ethanol before drying again.

In order to characterize the prepared PILs, several analytical techniques were employed: Fourier transform infrared spectrometer was used to analyze the infrared spectra of compounds to reveal their structures and functional groups. Considering the type of adsorbate, temperature, pressure, and other operational conditions, the specific surface area and pore size distribution of the samples were measured by using a BSD-PM1 high-performance surface area and micropore analyzer. Molecular weight analysis was performed with the help of a GPC-LS Gel Permeation Chromatograph under specific conditions such as solvent type, flow rate, and column type. MacroMR12-150H-I Nuclear Magnetic Resonance spectrometer was utilized to determine the structure of PILs by measuring the magnetic resonance frequencies of their atomic nuclei.

In this study, Thermogravimetric Analysis (TGA) was conducted on three different PILs to assess their thermal stability under a helium atmosphere. Equipped with platinum crucibles, the experiments were performed by using a TA Instruments Q50 thermogravimetric analyzer. Approximately 8 mg of the PILs sample was loaded into the crucible for each experiment. These experiments were carried out in a helium atmosphere, with a gas flow rate of 40 mL min<sup>-1</sup> and a heating rate set at 10 K min<sup>-1</sup>. The temperature range for the experiments extended from 303.15 K to 973.15 K.<sup>19,20</sup>

$$[C_{n}VIM][PF_{6}] (n=2, 4, 6, 8)$$

$$24h, 343.15 \text{ K}$$

$$DMF, AIBN$$

$$R$$

$$P[C_{n}VIM][PF_{6}] (n=2, 4, 6, 8)$$

$$P[C_{n}VIM][PF_{6}] (n=2, 4, 6, 8)$$

Fig. 1 The polymerization process for corresponding IL monomers.

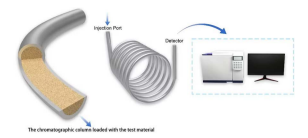


Fig. 2 Schematic diagram of the process for characterizing PILs based on IGC technology.

**2.2.1 Preparation of adsorption tube.** About 200 mg of PILs material (sized between 40 and 60 mesh), was packed into a stainless-steel adsorption tube (with an inner diameter of 1200 mm  $\times$  2 mm), and sealed at both ends with degreased cotton and a stainless-steel mesh (ID. 90 mm  $\times$  15 mm). The JX-1 type thermal analyzer was purged to 523.15 K at nitrogen flow rate of 20 mL min<sup>-1</sup> for two hours under a nitrogen atmosphere to remove any potentially volatile impurities and substances the material might have adsorbed. After processing, the ends were sealed with sealing caps for future use.

#### 2.3 IGC experimental method

Unlike conventional chromatographic techniques, Inverse Gas Chromatography (IGC) uses the material to be tested as a stationary phase. It employs a series of organic solvents with known properties as probe molecules. The retention time of the probe molecules can be measured at different temperatures and carrier gas flow rates to further calculate the thermodynamic parameters of the material to be tested. <sup>21,22</sup> IGC is capable of conducting quantitative analyses because of high sensitivity and rapid analytical capability. Hence, IGC offers significant advantages in detecting and investigating material adsorption properties. <sup>23,24</sup> Fig. 2 shows a schematic diagram of the process for characterizing PILs based on IGC technology in this work.

The IGC experiment utilized a QP-2010 Gas Chromatograph equipped with a Flame Ionization Detector (FID). It was operated with a high-purity nitrogen gas flow rate of 30 mL min<sup>-1</sup>. During the experiment, the injector and detector temperatures were maintained at 553.15 K, and the testing temperature was between 303.15 K and 343.15 K. The stationary phase was dried at 363.15 K until it achieved a constant mass and subsequently packed into a stainless-steel chromatographic column. The stationary phase consisted of 25–30% of the sample, with an analytical mass between 0.65 and 0.71 g. The packed column was aged at 363.15 K for 24 hours before testing. Each test was performed three times to ensure repeatability, and the calculation values were averaged.

This mass was dried at 363.15 K until its weight stabilized. Before undergoing testing, the now-filled column was aged at the same temperature at 363.15 K for 24 hours.

#### 2.4 Preparation of the chromatographic packed column

Before measurement, all samples were vacuum-treated for 48 hours to minimize moisture and volatile compounds. Afterwards,

they were dissolved in acetonitrile, loaded onto 102 monomer carriers, and dried at 363.15 K. Around 200 mg of PILs material was packed into a stainless-steel tube and purged at 523.15 K under nitrogen to remove impurities. After sealing the tube, the stationary phase was loaded into the chromatographic column using a vacuum pump. The mass of PILs was determined based on the change in the column's mass. This preparation is vital to the accuracy of subsequent chromatographic analyses.

#### 2.5 Adsorption heat and measurement

Adsorption heat, denoted as  $\Delta H$ , is crucial in understanding the interaction between gas molecules and solid materials. It quantifies the energy change during the adsorption process and provides insights into the nature of the interaction, whether it is physisorption or chemisorption. Accurate measurement of adsorption heat is pivotal for evaluating the efficiency and feasibility of adsorption processes in various applications, including separation and storage of gas.

The restriction of interaction between gas molecules and the packed solid material in a column was confined to surface interactions as the gas flows through. By measuring the retention time of gas molecules passing through the packed column at different temperatures,  $^{25}$  the heat of saturated adsorption ( $\Delta H$ ) between the gas molecules and the material under test could be determined using eqn (1).

$$\ln \frac{F_j(t_1 - t_2)}{m} = -\frac{\Delta H}{RT} + C \tag{1}$$

The mass of the packing material, represented as m(g) in the equation, works with the ideal gas constant R, column temperature T, and retention times  $t_1$  and  $t_2$  for the gas molecule and reference gas. C is constant, and  $F_j$  indicates the corrected flow rate.  $F_i$  can be calculated through eqn (2).

$$F_j = F \frac{3(P_i/P_o)^2 - 1}{2(P_i/P_o)^3 - 1}$$
 (2)

where  $P_i$  and  $P_o$  (MPa) respectively represent the inlet and outlet pressures of the packed column to calculate the carrier gas flow rate, F (mL min<sup>-1</sup>). Under the experimental conditions, variables included  $P_i$ ,  $P_o$ , and m remained constant. The net retention time  $t_{\rm N}$ , was given by  $t_1$ – $t_2$ , and eqn (1) was allowed to transformed.

$$\ln t_{\rm N} = -\frac{\Delta H}{RT} + C \tag{3}$$

By conducting experiments at different temperatures, we could establish the net retention time under different experimental conditions. When  $\ln t_{\rm N}$  was graphed in relation to 1/T, the slope of the resulting fitted line allows for the calculation of  $\Delta H$ .<sup>26</sup>

# 2.6 Desorption activation energy and measurement methods

The TPD technique was grounded on the intrinsic kinetic model, and the desorption rate of the adsorbate typically adhered to the following kinetic eqn (4) and (5).<sup>27,28</sup>

$$\frac{r_{\rm d}}{N_{\rm s}} = -\frac{{\rm d}\theta_{\rm A}}{{\rm d}t} = k_{\rm d}\theta_{\rm A} \tag{4}$$

$$k_{\rm d} = k_{\rm o} \exp\left(\frac{-E_{\rm d}}{RT}\right) \tag{5}$$

During the temperature ramping process, with a specific heating rate  $\beta_H$ , the relationship between temperature T and time is as follows:

$$T = T_{o} + \beta_{H}t \tag{6}$$

As the temperature increased, the desorption rate increases. When the temperature reached a specific point  $T_p$ , the desorption rate attained its maximum value. The relationship can be expressed as follows:

$$\begin{split} \frac{1}{N_{\rm s}} \frac{\mathrm{d}r_{\rm d}}{\mathrm{d}t} &= k_{\rm o} \frac{\mathrm{d}\theta}{\mathrm{d}t} \exp\left(-\frac{E_{\rm d}}{RT_{\rm p}}\right) + k_{\rm o}\theta \frac{E_{\rm d}}{R} \frac{1}{T_{\rm p}^2} \exp\left(-\frac{E_{\rm d}}{RT_{\rm p}}\right) \frac{\mathrm{d}T}{\mathrm{d}t} \\ &= 0 \end{split}$$

(7)

After rearrangement and taking the logarithm, we obtain:

$$\ln \frac{\beta_{\rm H}}{RT_{\rm p}^2} = -\frac{E_{\rm d}}{RT_{\rm p}} - \ln \frac{E_{\rm d}}{k_{\rm o}} \tag{8}$$

This equation identifies  $r_d$  (mol min<sup>-1</sup>) as the rate of desorption,  $N_s$  (mol cm<sup>-2</sup>) denotes the maximum capacity of adsorption for the adsorbate on a singular adsorption surface,  $\beta_{\rm H}$  (K min<sup>-1</sup>) is the heating rate, and  $E_d$  (kJ mol<sup>-1</sup>) is the desorption activation energy.  $k_0$  is the pre-exponential factor for the desorption rate,  $k_d$ indicates the desorption rate constant, and  $\theta$  represents the coverage of the component on the unit adsorbent surface. Different temperature-programmed desorption (TPD) curves and  $T_{\rm p}$  values can be obtained under different heating rate conditions. The value of  $E_{\rm d}$  could be assessed by graphing  $\ln(\beta_{\rm H}/RT_{\rm p}^{2})$  in relation to  $1/RT_p$  and analyzing the slope of the resulting line.

Specific experiments to measure the desorption activation energy were performed as follows:29 the gas to be tested was first injected into the chromatographic column containing the aforementioned PILs materials until the saturated adsorption was reached. Subsequently, nitrogen was used to purge any remaining gases at a flow rate of 30 mL min<sup>-1</sup>. The chromatography was then set with an initial temperature of 303.15 K and subjected to heating rates of 4, 5, 6, 8, and 10 K min<sup>-1</sup>, up to a terminal temperature of 453.15 K. The experimenter calibrated the rotameter. The rotameter was calibrated with nitrogen as the carrier gas at a rate of 20 mL min<sup>-1</sup>. Multiple iterations were performed to obtain desorption isotherms, and the average value derived from three separate trials was employed to calculate the desorption activation energy  $(E_d)$ .

#### Adsorption tube evaluation method

The performance of PILs material tubes in adsorbing R12, R22, and R134a was evaluated based on their adsorption capacity at different temperatures, linear range at room temperature, desorption rate, repeatability, and practicality.

- 2.7.1 Methods for measuring adsorption capacity. Penetration experiments were conducted to determine the adsorption capability of tubes for the three Freon gases at different temperatures. The initial temperatures of instruments were set at 278.15 K, 288.15 K, 298.15 K, 308.15 K, and 318.15 K at the beginning of the experiment and maintained for more than five minutes. The adsorption capacity for the three Freon refrigerants at different temperatures was determined by examining the introduction of different volumes of standard gas.
- 2.7.2 Standard curve analysis. The method for constructing the standard calibration curve involved collecting three adsorption tubes and testing their desorption by injecting different volumes of standard gas, and then averaging the desorption peak areas of the three tubes under the same volume of gas injected to determine the final peak area. The standard working curve was established based on the relationship between the peak area and the theoretical gas mass. Finally, the linearity of the response was analyzed by comparing the linear correlation coefficients of the curve.
- 2.7.3 **Desorption rate analysis method.** The desorption rate (Q) was calculated according to eqn (9). In this equation,  $S_1$  and S2 represent the target gas peak areas detected in the chromatographs from the first and second desorptions, respectively.

$$Q(\%) = \frac{S_1 - S_2}{S_1} \times 100\% \tag{9}$$

- 2.7.4 Repeatability analysis method. At 298.15 K, each adsorption tube was injected with supersaturated standard gas for nine consecutive adsorption and desorption cycles. Repeatability was discussed based on the adsorption capacity of each tube.
- 2.7.5 **Practicality analysis.** In order to assess the practicality of the three types of PILs adsorption tubes, we compared their adsorption capacity, standard curves, desorption rate, and repeatability for each refrigerant against those of activated carbon adsorption tubes under the same conditions at 298.15 K.

#### 3 Results and discussion

This section describes how to evaluate the performance of adsorption tubes filled with PILs material, focusing on methods to calculate desorption rates, analyze repeatability and assess overall practicality.

#### 3.1 Synthesis results and characterization analysis of PILs

Three types of PILs were synthesized in the experiment, i.e., P [EVIM][PF<sub>6</sub>], P[BVIM][PF<sub>6</sub>], and P[HVIM][PF<sub>6</sub>], with yields of approximately 73%, 64% and 83% respectively. P[OVIM][PF<sub>6</sub>] could not polymerize under similar conditions, possibly due to steric hindrance from the octyl group, making linking vinyl double bonds difficult.

Based on an analysis conducted in a wavelength range of 4000-400 cm<sup>-1</sup>, Fig. 3(a)–(c) presents

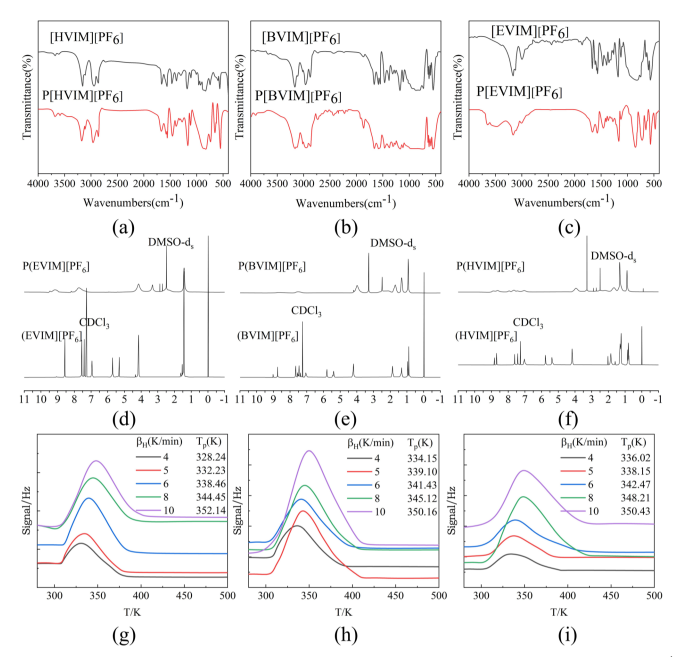


Fig. 3 (a-c) Infrared characterization (a) P[EVIM][PF<sub>6</sub>] and [EVIM][PF<sub>6</sub>], (b) P[BVIM][PF<sub>6</sub>] and [BVIM][PF<sub>6</sub>], (c) P[HVIM][PF<sub>6</sub>] and [HVIM][PF<sub>6</sub>]. (d-f)  $^1$ H NMR spectra (d) P[EVIM][PF<sub>6</sub>] and [EVIM][PF<sub>6</sub>], (e) P[BVIM][PF<sub>6</sub>] and [BVIM][PF<sub>6</sub>], (f) P[HVIM][PF<sub>6</sub>] and [HVIM][PF<sub>6</sub>]. (g-i) TPD curves of R12 (g) P [EVIM][PF<sub>6</sub>], (h) P[BVIM][PF<sub>6</sub>], (i) P[HVIM][PF<sub>6</sub>].

characterization results. It portrays the stretching vibration peak of the vinyl C=C bond at 1660 cm<sup>-1</sup>, and the bending vibration peak of the vinyl C-H bond at 961.88 cm<sup>-1</sup>, both of which almost disappear. This reveals the polymerization of double bonds in IL monomers.<sup>30</sup> The chemical shifts of different protons in PILs and ILs monomers are depicted in Fig. 3(d)-(f). For example, in <sup>1</sup>H NMR of P[EVIM][PF<sub>6</sub>] and [EVIM][PF<sub>6</sub>], the disappearance of corresponding peaks indicates that the double bonds in the IL monomers have polymerized. As illustrated in Fig. 3(d)-(f), taking the <sup>1</sup>H NMR of P [EVIM][PF<sub>6</sub>] and [EVIM][PF<sub>6</sub>] as examples, the protons at 9.42-8.71 (br, 1H), 7.96-6.98 (br, 2H), 4.34-3.95 (s, 2H), and 1.68-1.08

(s, 3H) represent the protons in the structural unit of P[EVIM] [PF $_6$ ]. The protons corresponding to the vinyl group in [EVIM] [PF $_6$ ] include 7.00–6.87 (dd, 1H), 5.75–5.62 (dd, 1H), and 5.35–5.26 (dd, 1H). The disappearance of these corresponding peaks demonstrates the polymerization of the double bond in the ILs monomer.

A series of characterization experiments were carried out to obtain a deeper understanding of the physical and chemical properties of the synthesized three types of PILs. Initially, through molecular weight characterization, it was concluded that all three PILs were low oligomers with a number-average degree of polymerization ranging between 6 and 7, as detailed

Table 1 Molecular weight characterization of PILs

PILs	Number-average molecular	Weight weight-average molecular	Weight Z-average molecular	Weight dispersity index	Number-average degree of polymerization
P[EVIM][PF <sub>6</sub> ]	1689	3533	8271	2.09	6.35
$P[BVIM][PF_6]$	1789	8669	50 906	4.85	6.08
P[HVIM][PF <sub>6</sub> ]	2233	11 326	58 973	5.07	6.93

Table 2 BET analysis of PILs

PILs	Specific surface area/ $(m^2 g^{-1})$	Pore size/(nm)	Pore volume/(cm <sup>3</sup> g <sup>-1</sup> )
P[EVIM][PF <sub>6</sub> ] P[BVIM][PF <sub>6</sub> ] P[HVIM][PF <sub>6</sub> ]	2.353	5.448 7.337 7.432	0.005 0.004 0.003

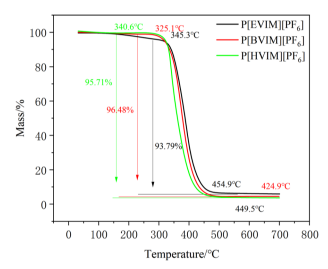


Fig. 4 Thermogravimetric curves  $P[EVIM][PF_6]$ ,  $P[BVIM][PF_6]$ ,  $P[HVIM][PF_6]$ .

in Table 1. Further BET analysis results, as referenced in Table 2, indicate that elongating alkyl side chains on the polymer backbone leads to a decrease in surface area, an increase in pore size, and a reduction in pore volume. This finding aligns with the research of Zulfiqar *et al.*, who proposed that variations in cations, anions, polymer backbones, alkyl substituents, porosity, cross-linking, molecular weight, and moisture content could account for the performance differences observed in PILs.<sup>31</sup>

As described in Fig. 4, Thermogravimetric analysis results indicate that these PILs exhibit no significant mass loss up to 573.15 K, demonstrating high thermal stability. However, they all underwent substantial mass loss, primarily attributed to the degradation of the polymer backbone within 573.15–773.15 K. Notably, the mass remained stable between 773.15 and 973.15 K without further loss.

#### 3.2 Adsorption heat

The detailed introduction and meticulous measurement of Adsorption heat are paramount in comprehending the interactions and assessing the efficacy of the PILs in adsorption processes. In the experimental temperature, the net retention time of three Freon refrigerants on three types of PILs filled columns showed different degrees of reduction as the observed temperature increased. This phenomenon indicates that the adsorption interaction between Freon and the surface of the synthesized PILs materials is an exothermic process. Fig. 5(a) displays the assessed values of the heat of saturated adsorption experimented under specific conditions.

The magnitude of the heat of adsorption serves as a critical indicator for distinguishing between physical and chemical adsorption processes.32 Physical adsorption typically occurs under the influence of weaker intermolecular forces, such as van der Waals interactions, resulting in relatively lower heats of adsorption, approximately around 40 kJ mol<sup>-1</sup>. This lower heat of adsorption signifies weaker interactions between the adsorbate and the adsorbent, characteristic of physical adsorption phenomena. Conversely, chemical adsorption involves the formation of stronger chemical bonds, thereby exhibiting significantly higher heats of adsorption, generally ranging from 84 to 417 kJ  $\text{mol}^{-1}$ . The elevated heat of adsorption in chemical adsorption indicates more robust chemical bonding between the adsorbate and the adsorbent, a hallmark of chemical adsorption processes. Thus, by assessing the heat of adsorption, one can effectively differentiate between the distinct mechanisms of physical and chemical adsorption.

Among the three types of PILs, P[EVIM][PF<sub>6</sub>] exhibited the highest  $\Delta H$  values for all three Freons (50.41–58.95 kJ mol<sup>-1</sup>), while P[HVIM][PF<sub>6</sub>] had the lowest (29.60–32.12 kJ mol<sup>-1</sup>). P [EVIM][PF<sub>6</sub>] and P[BVIM][PF<sub>6</sub>] both undergo both physical and chemical adsorption, except for P[HVIM][PF<sub>6</sub>], which undergoes only physical adsorption.

This may stems from the differences in the basicity of the three types of PILs. Since R12, R22, and R134a are acidic, PILs with stronger basicity interact more with these gases.

#### 3.3 Desorption activation energy

 $T_{\rm p}$  values can be calculated in accordance with TPD curves measured at different heating rates. Using R12 as the sample, TPD curves at different heating rates are depicted in Fig. 3(g)–(i). There is only one desorption peak on each TPD curve, indicating uniform adsorption that does not involve multiple adsorption centers or re-adsorption. According to eqn (8), the calculated results for  $E_{\rm d}$  are illustrated in Fig. 5(b). Experimental

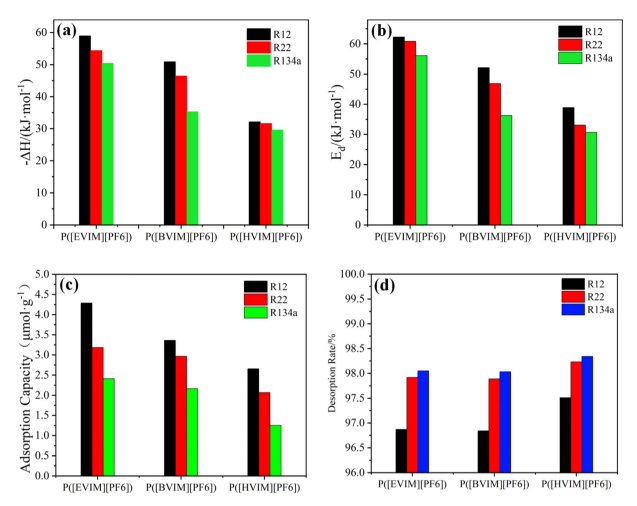


Fig. 5 (a) Adsorption heat of Freon gas on the surface of three PlLs. (b) Desorption activation energy of Freon gas on the surface of three PlLs. (c) Adsorption capacities of PlLs for refrigerants R12, R22, and R134a, measured at an adsorption time of 15 minutes under constant conditions of 298.15 K and a flow rate of 5 mL min<sup>-1</sup>. (d) Desorption rate analysis of Freon gas on PlLs.

**Table 3** Comparison of the  $E_d$  and  $\Delta H$  values

	R12		R22		R134a	
PILs	$-\Delta H/(kJ \text{ mol}^{-1})$	$E_{\rm d}/({\rm kJ~mol}^{-1})$	$-\Delta H/(kJ \text{ mol}^{-1})$	$E_{\rm d}/({\rm kJ~mol}^{-1})$	$-\Delta H/(kJ \text{ mol}^{-1})$	$E_{\rm d}/({\rm kJ~mol}^{-1})$
P([EVIM][PF <sub>6</sub> ])	58.95	62.27	54.37	60.85	50.41	56.05
$P([BVIM][PF_6])$	50.92	52.14	46.48	46.85	35.28	36.23
$P([HVIM][PF_6])$	32.12	38.89	31.62	33.05	29.60	30.64

results indicate that the  $E_{\rm d}$  values for the three refrigerants are highest on P[EVIM][PF<sub>6</sub>] (56.05–62.27 kJ mol<sup>-1</sup>), followed by P [BVIM][PF<sub>6</sub>] (36.23–52.14 kJ mol<sup>-1</sup>), and are lowest on P[HVIM] [PF<sub>6</sub>] (30.64–38.89 kJ mol<sup>-1</sup>). A greater  $E_{\rm d}$  value indicates that the adsorbate needs more energy to detach from the adsorbent, suggesting a stronger adsorption. For each type of PILs, the  $E_{\rm d}$  values for the three fluorocarbon refrigerants follow R134a < R22 < R12. Table 3 compares the measured  $E_{\rm d}$  and  $\Delta H$  values for R12, R22, and R134a on the three types of PILs. It is clear that the  $E_{\rm d}$  values are all higher than the  $\Delta H$  values. This suggests

that the adsorption rates of the three fluorocarbon gas molecules on the three types of PILs surfaces are more significant than their desorption rates. In our study, we observed significant differences in adsorption heats and desorption activation energies even among structurally similar PILs. This finding aligns with the research conducted by Yuan *et al.*, who emphasized the impact of intrinsic differences in the chemical structures and synthesis trends of PILs on their adsorption heats and desorption activation energies.<sup>17</sup>

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5 5 5.0 4.5 - si lomi 3.5 Adsorption Capacity 2.0 3mL/min 5mL/min 7mL/min 10mL/min 0.5 0.0 30 33 36 39 T/min

Fig. 6 Profiles of R22 adsorption capacity of P[EVIM][PF<sub>6</sub>] adsorption tube with different flow rates at 298.15 K.

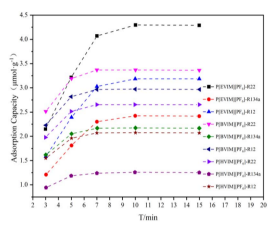
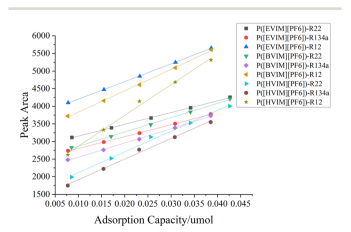


Fig. 7 Adsorption capacity of the PILs adsorption tube to the refrigerants with different adsorption time at 298.15 K.

#### Adsorption capacity

The adsorption of refrigerants on PILs tubes was investigated at 298.15 K. For instance, the process of R22 adsorption onto a P [EVIM][PF<sub>6</sub>] adsorption tube is depicted in Fig. 6. An decrease in

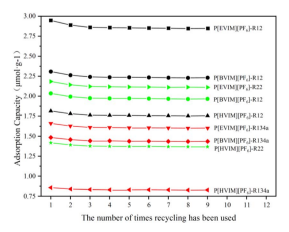


the equilibrium adsorption capacity of P[EVIM][PF6] was observed with the increase in R22 flow rate, which aligns with the findings of Tao Wang et al. This decrease was attributed to the reduced contact time between the adsorbate and adsorbent at higher flow rates.33

This study also examined the changes in adsorption capacity over time for three types of refrigerants on three kinds of PILs adsorption tubes at 5 mL min<sup>-1</sup> and 298.15 K. As can be seen in Fig. 7, the adsorption capacity of the three types of PILs adsorption tubes remains consistent for the refrigerants at an adsorption time duration of 10 minutes. The findings suggest that the adsorption tubes of the three types of PILs maintain a steady adsorption capacity for refrigerants during a 10 minute adsorption timeframe. At the same time, Fig. 5(c) displays the adsorption capacities of R12, R22, and R134a on these adsorption tubes within a temperature scope of 298.15 K and a gas flow velocity of 5 mL min<sup>-1</sup>. There is an inverse correlation between the adsorption capacity of the three Freon gases on the three PILs adsorption tubes and the elongation of the alkyl side chain on the polymer backbone. This observation aligned with the trend identified during the synthesis process, where the elongation of the alkyl side chain on the polymer backbone led to a reduction in specific surface area, an increase in pore diameter, and a decrease in pore volume. Additionally, the adsorption capacities of the three PILs adsorption tubes for the refrigerants follow the sequence R134a < R22 < R12, consistent

Table 4 Standard working curves and correlation coefficients of the Freon gas on PILs adsorption tube

-			
PILs	Freon gas	Fitting equation	$R^2$
P([EVIM][PF <sub>6</sub> ])	R12	Y = 50497X + 3706	0.9995
(E 3E 03)	R22	Y = 33431X + 2826	0.9996
	R134a	Y = 33739X + 2471	0.9995
$P([BVIM][PF_6])$	R12	Y = 60632X + 3243	0.9924
	R22	Y = 40406X + 2475	0.9986
	R134a	$Y = 405\ 10X + 2162$	0.9984
$P([HVIM][PF_6])$	R12	Y = 87944X + 1990	0.9946
	R22	$Y = 592\ 25X + 1523$	0.9927
	R134a	$Y = 587\ 58X + 1327$	0.9933



Standard working curves of Freon gas on PILs adsorption tube. Fig. 9 Repeatability analysis of Freon gas on PILs adsorption tubes.

Table 5 Comparison of activated carbon adsorption tube and PILs adsorption tube at 298.15 K

Adsorbent material	Name of the gas	Linear equation	$R^2$	Desorption rate (%)
Activated carbon	R12	Y = 5113X + 2033	0.9724	88.36
	R134a	Y = 5872X + 1779	0.9650	92.72
	R22	Y = 6469X + 2697	0.9530	89.60
P[EVIM][PF <sub>6</sub> ]	R12	Y = 50497X + 3706	0.9996	96.87
35 3	R134a	Y = 33739X + 2471	0.9995	98.05
	R22	Y = 33431X + 2826	0.9995	97.92
P[BVIM][PF <sub>6</sub> ]	R12	Y = 60632X + 3243	0.9986	96.84
E 3E ~3	R134a	Y = 40510X + 2162	0.9984	98.03
	R22	Y = 40406X + 2475	0.9924	97.89
P[HVIM][PF <sub>6</sub> ]	R12	Y = 87944X + 1990	0.9927	97.51
L 3L 33	R134a	Y = 58758X + 1327	0.9933	98.34
	R22	Y = 59225X + 1523	0.9946	98.23

Table 6 Saturated adsorption capacity of various materials adsorption tube

		Adsorption capacity (μmol g <sup>-1</sup> )			
Materials	$T/\mathrm{K}$	R22	R134a	R12	
Activated carbon	298.15	4.413	2.728	3.345	
$P[EVIM][PF_6]$	298.15	2.862	1.613	2.123	
P[BVIM][PF <sub>6</sub> ]	298.15	2.243	1.445	1.978	
P[HVIM][PF <sub>6</sub> ]	298.15	1.765	0.834	1.378	

with the adsorption heat analysis results for these gases on the PILs surfaces. The higher the adsorption heat value, the stronger the interaction between the gas and the PILs. In summary, the interaction strength between the three Freon gases and the three PILs materials predominantly determines the adsorption capacity.

#### 3.5 Standard curve analysis

Standard operation curves at a 298.15 K temperature was established and the relationship between peak area and hypothetical gas mass was examined (Fig. 8). Table 4 displays the equations reflecting fit and matching correlation coefficients of the standard curve. The table below reveals a strong linear correlation between R12, R22, and R134a across the three PILs adsorption tube types across the three PILs adsorption tube types each exceed 0.99. This evident linear relationship suggests that the adsorption effect can be predicted and controlled, thereby aiding in designing efficient refrigerant recovery or separation systems.

#### 3.6 Desorption rate analysis

At 298.15 K, the practical applicability of PILs adsorption tubes was comprehensively evaluated by comparing their performance with activated carbon adsorption tubes under the same conditions, specifically in terms of adsorption capacity, standard curves, and desorption rates for three types of Freon gas refrigerants. The experimental results are shown in Fig. 5(d). The table clearly shows that the linearity and desorption rates in

the three PILs adsorption tubes are superior to those found in the activated carbon adsorption tubes. With respect to saturated adsorption capacity, the activated carbon adsorption tubes hold a higher capacity than the three different PILs adsorption tubes due to their more complex pore structure and a larger specific surface area compared to PILs materials.

#### 3.7 Reproducibility analysis

Each PILs adsorption tube underwent nine successive adsorption and desorption cycles at 298.15 K, with the adsorption capacity serving as the standard for analyzing reproducibility. The findings from the experiment are depicted in Fig. 9. As repeated uses of the three types of PILs adsorption tubes increase, the saturated adsorption capacity of the refrigerants remains relatively stable.

#### 3.8 Practicality analysis

At 298.15 K, the practical applicability of PILs adsorption tubes was comprehensively evaluated by comparing their performance with activated carbon adsorption tubes under the same conditions, specifically in terms of adsorption capacity, standard curves, and desorption rates for three types of Freon gas refrigerants. The experimental results are listed in Table 5. It is obvious that the linearity and desorption rates of the three types of PILs adsorption tubes are superior to those of the activated carbon adsorption tubes. According to Table 6, the saturated adsorption capacity of activated carbon adsorption tubes is higher than that of PILs manufactured from the three different types of materials. This can be attributed to that activated carbon has a more developed pore structure and a larger specific surface area than PILs.

#### 4 Conclusion

In current study, three linear PILs materials were synthesized using four ILs as polymerization monomers through a free radical polymerization reaction. The elongation of the alkyl side chain impedes monomer polymerization, which is unfavorable for the synthesis of linear PILs. All three linear PILs are oligomers with a number-average degree of polymerization between

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6 and 7, without mass loss at 575.15 K. The adsorption properties were experimentally evaluated by preparing adsorbent tubes using the three PIL materials. It was found that apart from P[HVIM][PF<sub>6</sub>], which exhibits physical adsorption, both P[EVIM] [PF<sub>6</sub>] and P[BVIM][PF<sub>6</sub>] undergo simultaneous physical and chemical adsorption. For each PILs, the  $\Delta H$  and  $E_d$  values for the three refrigerants follow R134a < R22 < R12. Moreover, as the alkyl side chain on the polymer backbone elongates, the adsorption efficacy of the PILs materials for the three tracers decreases. After nine consecutive adsorption and desorption cycles, the saturation adsorption capacity on the three PILs adsorption tubes remains stable. The desorption rates of the three PILs adsorption tubes surpass those of activated carbon tubes with excellent linear response. As a result, these three PILs exhibit commendable performance in adsorption linearity, adsorption-desorption properties, and repeatability, suggesting that they have excellent prospects for practical applications.

# Data availability

Data will be made available on request.

### Author contributions

Mingyuan Xin: conceptualization; methodology; experiments; manuscript writing. Qiang Wang: conceptualization; experiments; data curation; manuscript reviewing. Qiang WangJ: experiments; data curation; manuscript reviewing. Haoyu Wang: methodology; manuscript reviewing. Furqan Muhammad: supervision; manuscript reviewing. Guanze Nie: conceptualization; supervision; manuscript reviewing.

## Conflicts of 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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