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# Pillar-layer Zn–triazolate–dicarboxylate frameworks with a customized pore structure for efficient ethylene purification from ethylene/ethane/acetylene ternary mixtures†

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The selective adsorption of C<sub>2</sub>H<sub>6</sub> and C<sub>2</sub>H<sub>2</sub> over C<sub>2</sub>H<sub>4</sub> from C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> ternary mixtures for one-step C<sub>2</sub>H<sub>4</sub> purification represents a crucial yet challenging task in industry. The pore structure of the adsorbents must be finely tailored to meet the demanding requirements for the separation considering the very similar physicochemical properties of the three gases. Herein, we report a Zn–triazolate–dicarboxylate framework, HIAM-210, featuring a novel topology which possesses one-dimensional channels decorated with adjacent uncoordinated carboxylate–O atoms. The suitable pore size and customized pore environment enable the compound to selectively capture C<sub>2</sub>H<sub>6</sub> and C<sub>2</sub>H<sub>2</sub> with high C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub> selectivities of both 2.0. Breakthrough experiments show that polymer-grade C<sub>2</sub>H<sub>4</sub> can be directly harvested from C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> (34/33/33 and 1/90/9) ternary mixtures. The underlying mechanism of the preferential adsorption was uncovered by grand canonical Monte Carlo simulations and DFT calculations.

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## Introduction

Ethylene (C<sub>2</sub>H<sub>4</sub>), the raw material for the production of various organic products, exceeded an annual global production of 200 million tons in 2022.<sup>1–3</sup> In the petrochemical industry, C<sub>2</sub>H<sub>4</sub> is mainly produced through thermal cracking of naphtha or ethane where the crude products include ethane (C<sub>2</sub>H<sub>6</sub>) and acetylene (C<sub>2</sub>H<sub>2</sub>) as impurities.<sup>4</sup> The separation of C<sub>2</sub>H<sub>4</sub> from C<sub>2</sub>H<sub>6</sub> or C<sub>2</sub>H<sub>2</sub> is currently a complex and energy-consuming process that involves solvent extraction or catalytic hydrogenation (C<sub>2</sub>H<sub>2</sub> removal) and subsequent low-temperature distillation (C<sub>2</sub>H<sub>6</sub> removal).<sup>5,6</sup> To this end, it is necessary and urgent to develop a simpler separation approach with lower energy consumption for C<sub>2</sub>H<sub>4</sub> purification.<sup>7,8</sup>

Adsorptive separation based on physical adsorbents has emerged as an alternative means for industrial hydrocarbon separation.<sup>9</sup> However, the simultaneous removal of C<sub>2</sub>H<sub>2</sub> and C<sub>2</sub>H<sub>6</sub> from C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> mixtures remains a great challenge as the quadrupole moment and kinetic diameter of C<sub>2</sub>H<sub>4</sub> (1.5 × 10<sup>–26</sup> esu cm<sup>2</sup> and 4.1 Å) lie between those of C<sub>2</sub>H<sub>2</sub> (7.2 × 10<sup>–26</sup> esu cm<sup>2</sup> and 3.3 Å) and C<sub>2</sub>H<sub>6</sub> (0.65 × 10<sup>–26</sup> esu cm<sup>2</sup> and 4.4 Å) (Table S1†). Traditional adsorbents such as zeolites and

activated carbons have not yet demonstrated the potential to meet the stringent requirements for the separation, mainly due to their limited structural tunability. In this context, metal–organic frameworks (MOFs), for which the pore size and chemical environment can be precisely adjusted, are considered promising candidates to address this challenging process.<sup>10,11</sup> In general, a non-polar surface favors C<sub>2</sub>H<sub>6</sub> while smaller pore size prefers C<sub>2</sub>H<sub>2</sub>. However, since C<sub>2</sub>H<sub>2</sub> has the lowest polarizability among the three hydrocarbons, a non-polar pore surface is also unfavorable for C<sub>2</sub>H<sub>2</sub> adsorption, and most previously reported MOFs with high C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub> selectivity also possess low C<sub>2</sub>H<sub>2</sub> uptake, insufficient for the efficient separation of C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> mixtures. Thus, the key to constructing MOFs for one-step C<sub>2</sub>H<sub>4</sub> purification is to find the “sweet spot” by creating an optimal chemical environment on the pore surface and suitable pore size so as to achieve simultaneously high selectivities for both C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub>.

In this work, we present a new pillar-layer MOF constructed *via* zinc ions, triazole and a dicarboxylate linker (denoted as HIAM-210, HIAM = Hoffman Institute of Advanced Material), which combines suitable pore dimensions and a pore chemical environment for one-step separation of C<sub>2</sub>H<sub>4</sub> from C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> ternary mixtures. The adjacent uncoordinated carboxylate–O atoms decorated on the pore surface act as strong binding sites for preferential adsorption toward C<sub>2</sub>H<sub>6</sub> and C<sub>2</sub>H<sub>2</sub> over C<sub>2</sub>H<sub>4</sub>, enabling HIAM-210 to have relatively high adsorption selectivities for C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub> (2.0) and C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub> (2.0), which exceed most of the MOFs previously tested for C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>2</sub> separation, such as TJT-100 (1.8 and 1.2), ZJNU-115 (2.05 and

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1.56), NUM-9 (1.50 and 1.62), NPU-2 (1.25 and 1.52), UPC-66 (1.05 and 1.65), *etc.*<sup>12–16</sup> Breakthrough experiments indicated that polymer-grade (99.95+%) C<sub>2</sub>H<sub>4</sub> can be directly produced from ternary C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> mixtures. Grand canonical Monte Carlo (GCMC) simulations and DFT calculations provide important insights into the preferential adsorption mechanism at the molecular level.

## Results and discussion

Yellow rod-shaped crystals of HIAM-210 were obtained by solvothermal reactions of Zn(NO<sub>3</sub>)<sub>2</sub>·6H<sub>2</sub>O, 1,2,4-triazole (TZ) and 1,4-naphthalene dicarboxylic acid (NDC) in *N,N*-dimethylformamide (DMF)/H<sub>2</sub>O solution at 423 K (Fig. 1a and S1, see the ESI† for detailed synthesis). Single crystal X-ray diffraction revealed that HIAM-210 crystallized in the monoclinic crystal system with a space group of *Pm* (Table S2†). The asymmetric unit of the structure contains ten independent Zn(II) centers coordinated in three fashions (Fig. S2†). Zn1, Zn3, Zn4, Zn7, Zn8, and Zn9 are tetrahedrally coordinated to two oxygen atoms from two NDC and two N atoms from two TZ ligands respectively; Zn2 and Zn6 are octahedrally bonded to two oxygen atoms from two NDC and four N atoms from four TZ ligands respectively; Zn5 and Zn10 are pentahedrally coordinated to three oxygen atoms from two NDC and two N atoms from two TZ ligands respectively. As depicted in Fig. 1b, each TZ ligand connects to three Zn(II) ions through μ<sub>1,2,4</sub>-bridging mode to form a 1D Zn–TZ chain along the *b*-axis, and the 1D chains are interconnected through NDC linkers to form a 2D Zn–TZ–NDC layer. The layers are further pillared by the linear ditopic

carboxylate NDC by linking to two independent Zn(II) ions (Fig. 1c), constructing a 3D structure with 1D channels (Fig. 1d). It is noteworthy that this structure is different from the widely reported Zn–triazolate–dicarboxylate frameworks where the 2D layer was formed only by metal–triazolate coordination.<sup>17–21</sup> The compound features an unprecedented topology with point symbols {4<sup>3</sup>·5<sup>2</sup>·7}·{4<sup>3</sup>·5<sup>3</sup>}<sub>2</sub>·{4<sup>3</sup>}·{4<sup>6</sup>·5<sup>2</sup>·6<sup>8</sup>·7·8<sup>4</sup>}{4<sup>6</sup>·6<sup>6</sup>·8<sup>3</sup>} (Fig. S3†). The 1D channels of HIAM-210 were partitioned by benzene rings of NDC into arrays of interconnected cages with approximate sizes of 4.5 × 14.5 Å (Fig. 1e and S4†). As shown in Fig. 1f, multiple uncoordinated carboxylate–O atoms are unevenly distributed in four different cages, which have been believed to act as strong adsorption sites for hydrocarbons.<sup>3,22,23</sup>

The powder X-ray diffraction (PXRD) patterns of the as-synthesized and activated sample of HIAM-210 matched well with the simulated pattern, confirming its phase purity and structural integrity upon activation (Fig. S5†). N<sub>2</sub> adsorption–desorption measurement at 77 K was carried out to evaluate the porosity of HIAM-210 (Fig. 2a). The Brunauer–Emmett–Teller (BET) surface area and pore volume were calculated to be 565.96 m<sup>2</sup> g<sup>−1</sup> and 0.24 cm<sup>3</sup> g<sup>−1</sup>, respectively and the pore size was centered at 4.02 Å established by the Horvath–Kawazoe method (inset, Fig. 2a), which is close to the kinetic diameter of C<sub>2</sub> hydrocarbons. Thermogravimetric (TG) analysis of the as-synthesized and activated HIAM-210 revealed that the latter has a longer plateau from 353 to 453 K. A thermal stability test by treating the compound under high temperature indicated that HIAM-210 maintained its structural integrity upon heating at 423 K for 3 days (Fig. S7†). To further verify its chemical stability, PXRD patterns were collected on samples immersed in

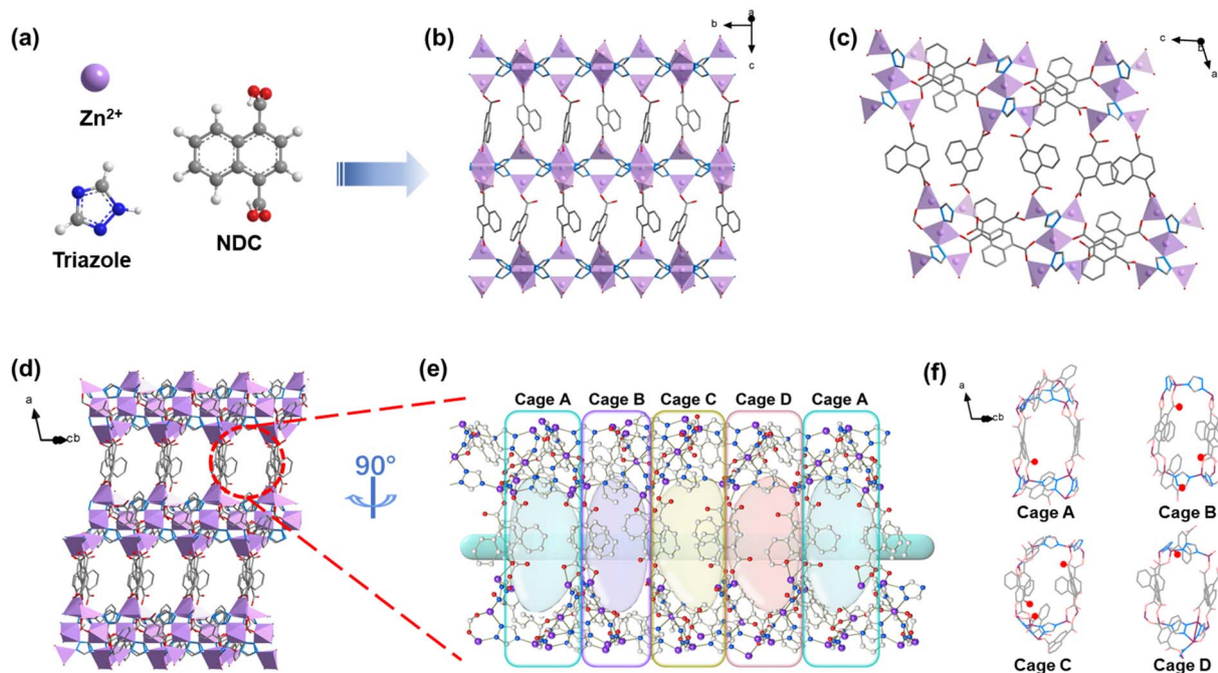


Fig. 1 Structure of HIAM-210. (a) The building blocks; (b) the 2D Zn–TZ–NDC layer structure; (c) the pillaring mode by NDC; (d) the structure viewed along the 1D channels; (e) a close-up view of the array of the four different cages in the channel; (f) the distribution of uncoordinated carboxylate–O atoms in the four cages.



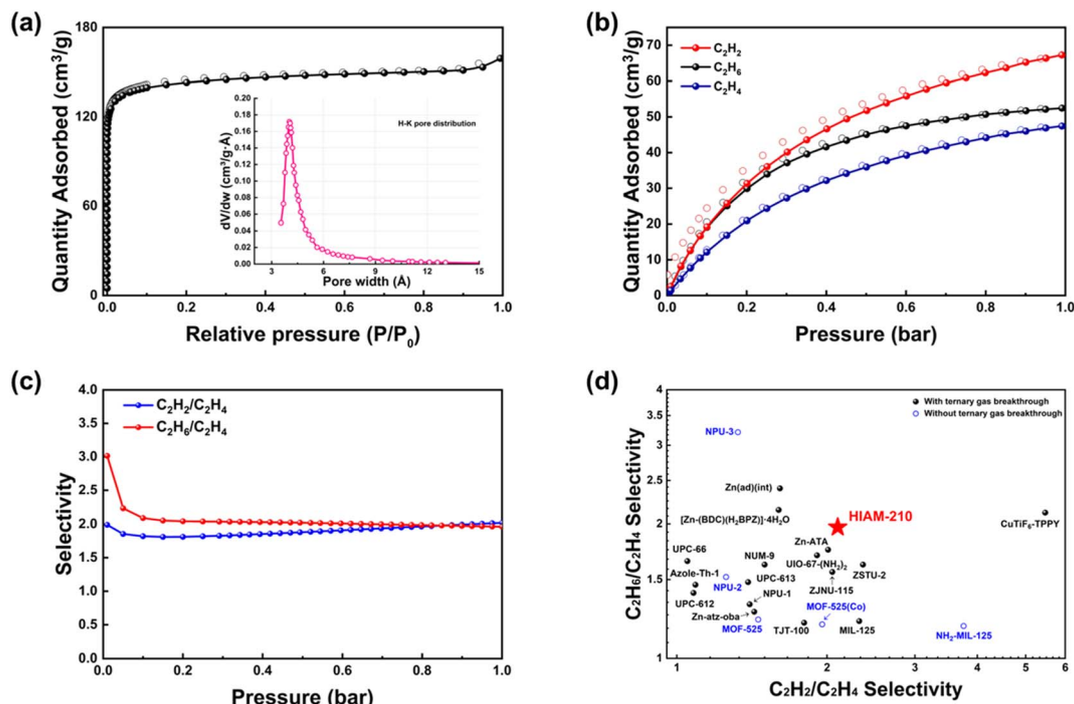


Fig. 2 Gas adsorption on HIAM-210. (a) N<sub>2</sub> adsorption–desorption isotherms at 77 K and pore size distribution (inset); (b) C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> adsorption–desorption isotherms at 298 K; (c) IAST selectivity of equimolar C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub> mixtures at 298 K; (d) comparison of the separation performance of representative MOFs (C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>: 50/50, v/v; and C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub>: 50/50, v/v).

water solutions with different pH values, and the results show that the structure of HIAM-210 remained intact under pH = 3 to 12 (Fig. S7†). The good stability, suitable pore size and delicate pore environment prompted us to further investigate the adsorption and separation ability of HIAM-210 for C<sub>2</sub> hydrocarbons.

Single-component adsorption isotherms of C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> for HIAM-210 were collected at 298 K and up to 1 bar, as shown in Fig. 2b. HIAM-210 adsorbs 52.44 cm<sup>3</sup> g<sup>-1</sup> (7.04 wt%) of C<sub>2</sub>H<sub>6</sub> and 67.26 cm<sup>3</sup> g<sup>-1</sup> (7.82 wt%) of C<sub>2</sub>H<sub>2</sub> at 1 bar and 298 K, higher than that of C<sub>2</sub>H<sub>4</sub> (47.43 cm<sup>3</sup> g<sup>-1</sup>, 5.94 wt%) under identical conditions, showing preferential adsorption of C<sub>2</sub>H<sub>2</sub> and C<sub>2</sub>H<sub>6</sub> over C<sub>2</sub>H<sub>4</sub>. To access the separation capability for C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> mixtures, ideal adsorbed solution theory (IAST)<sup>24</sup> selectivities for the two binary mixtures (C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub>) were calculated and the dual-site Langmuir–Freundlich fitting parameters are shown in Table S3.† The calculated selectivities of C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>6</sub>/C<sub>2</sub>H<sub>4</sub> at 298 K and 1 bar are both 2.0, which are higher than those of most of the MOFs previously tested for C<sub>2</sub>H<sub>2</sub>/C<sub>2</sub>H<sub>4</sub>/C<sub>2</sub>H<sub>6</sub> separation (Fig. 2d).<sup>3,12–16,25–32</sup> The isosteric heats of adsorption (*Q*<sub>st</sub>) of C<sub>2</sub>H<sub>2</sub>, C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>6</sub> on HIAM-210 were determined by using the Virial equation (Fig. S8 and S9†). The initial *Q*<sub>st</sub> of C<sub>2</sub>H<sub>6</sub> and C<sub>2</sub>H<sub>2</sub> was 31.24 and 34.31 kJ mol<sup>-1</sup>, respectively, higher than that of C<sub>2</sub>H<sub>4</sub> (22.72 kJ mol<sup>-1</sup>), suggesting that C<sub>2</sub>H<sub>6</sub> and C<sub>2</sub>H<sub>2</sub> have a relatively stronger interaction with the adsorbent than C<sub>2</sub>H<sub>4</sub>. It is worth noting that the *Q*<sub>st</sub> curve of C<sub>2</sub>H<sub>2</sub> shows a decrease followed by an increase with the gas loading. The decrease at low coverage is caused by the fact that adsorption sites with higher interaction

energy are first occupied before the sites with lower energy. In contrast, the increase at higher coverage should be attributed to the attractive interactions between the adsorbed molecules.<sup>33,34</sup>

To reveal the preferential adsorption behavior of C<sub>2</sub>H<sub>6</sub> and C<sub>2</sub>H<sub>2</sub> over C<sub>2</sub>H<sub>4</sub>, GCMC simulation was carried out to study the preferential adsorption sites (see the ESI† for simulation detail). The simulation shows that the primary adsorption site is located at Cage C with two adjacent uncoordinated carboxylate–O atoms which can capture all three hydrocarbon molecules (Fig. 3a–c). The C<sub>2</sub>H<sub>2</sub> molecule interacts strongly with the framework by forming multiple C–H⋯O hydrogen bonds with two uncoordinated O atoms (2.57 and 2.93 Å) and one coordinated O atom (2.70 Å). For C<sub>2</sub>H<sub>6</sub>, the adsorbed molecule forms multiple bonding interactions involving one C–H⋯π van der Waals (vdW) interaction (3.52 Å) with TZ, and three C–H⋯O hydrogen bonds with three O atoms (2.84–3.00 Å, including two uncoordinated and one coordinated one). In contrast, we found C<sub>2</sub>H<sub>4</sub> is weakly adsorbed by longer bonding distances including three C–H⋯O hydrogen bonds with three O atoms (3.06–3.54 Å, including two uncoordinated and one coordinated one) and one C–H⋯π vdW interaction (3.43 Å) with TZ. The different interaction patterns between the gas and framework induced different binding energies with the trend of C<sub>2</sub>H<sub>2</sub> (21.44 kJ mol<sup>-1</sup>) > C<sub>2</sub>H<sub>6</sub> (19.29 kJ mol<sup>-1</sup>) > C<sub>2</sub>H<sub>4</sub> (16.39 kJ mol<sup>-1</sup>), obtained from DFT calculations. This trend is consistent with the sequences of adsorption isotherms and heats of adsorption. Moreover, we noticed a secondary adsorption site located in Cage B, which only adsorbs the C<sub>2</sub>H<sub>2</sub> molecule (Fig. 3d). In Cage B, the C<sub>2</sub>H<sub>2</sub> molecule also interacts with the uncoordinated



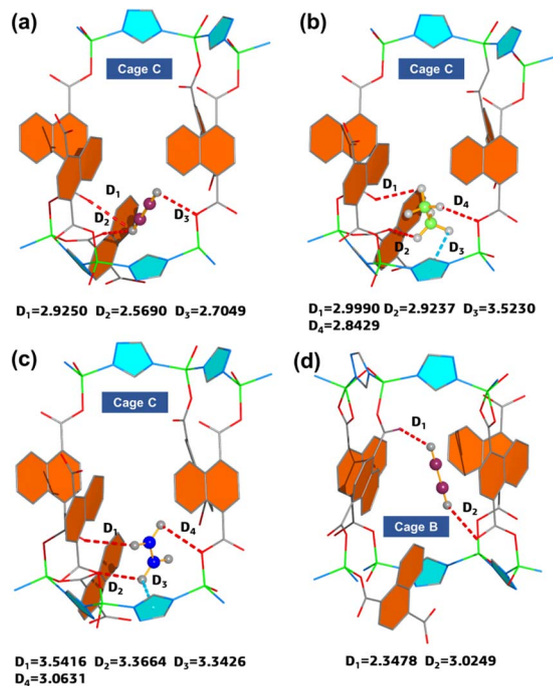


Fig. 3 The preferential adsorption sites for (a)  $C_2H_2$ , (b)  $C_2H_6$ , (c)  $C_2H_4$  in Cage C and (d) preferential adsorption site for  $C_2H_2$  in Cage B, and the  $C-H\cdots\pi$  and  $C-H\cdots O$  interactions are represented by blue and red dashed lines respectively.

carboxylate-O atoms to form short  $C-H\cdots O$  hydrogen bonding interactions (2.35 and 3.02 Å). The above phenomenon indicates that the adjacent uncoordinated carboxylate-O atoms along the channels of HIAM-210 are crucial for the preferential recognition of  $C_2H_2$  and  $C_2H_6$  over  $C_2H_4$ .

To further evaluate the competitive adsorption of  $C_2H_2$  and  $C_2H_6$  on HIAM-210, GCMC simulation was conducted by introducing an equimolar binary mixture into the framework. The results show that  $C_2H_2$  still occupies the primary binding sites within Cage C near the two adjacent uncoordinated carboxylate-O atoms. In contrast,  $C_2H_6$  molecules tend to bind with uncoordinated carboxylate-O atoms in Cage A and Cage D (Fig. S10†). The results indicated that the binding energy of  $C_2H_2$  in Cage C is higher than that of  $C_2H_6$  at similar adsorption sites, which is consistent with the previous binding energy calculations.

Dynamic breakthrough experiments were further carried out to evaluate the feasibility of using HIAM-210 for the separation of  $C_2$  binary and ternary mixtures. As depicted in Fig. 4a, b and S11,† HIAM-210 is capable of efficiently separating  $C_2H_4$  from  $C_2H_2/C_2H_4$  (50/50 and 1/99, v/v) and  $C_2H_6/C_2H_4$  (50/50 and 10/90, v/v) binary mixtures.  $C_2H_4$  always first eluted through the column to produce polymer-grade (>99.95%)  $C_2H_4$ . Breakthrough tests for ternary mixtures were subsequently performed. As shown in Fig. 4c, this compound can separate  $C_2H_4$  from the equimolar ternary mixture. And breakthrough measurement with a gas mixture of  $C_2H_2/C_2H_4/C_2H_6$  (1/90/9, v/v) that is close to the gas compositions in real applications was carried out. It was observed that  $C_2H_4$  eluted out first at the 30th min, followed by  $C_2H_6$  at the 38th min and  $C_2H_2$  at the 39th min, producing polymer-grade  $C_2H_4$  during the interval between the breakthrough of  $C_2H_4$  and  $C_2H_6$  with a productivity of 2.56 L  $kg^{-1}$  (Fig. 4d). Considering the potential interference and competitive adsorption of moisture in practical processes, we carried out a breakthrough experiment under a humid environment of 50% relative humidity (RH). As shown in Fig. 4e, the breakthrough curves and productivity of pure  $C_2H_4$  under humid and dry environments were similar, demonstrating the

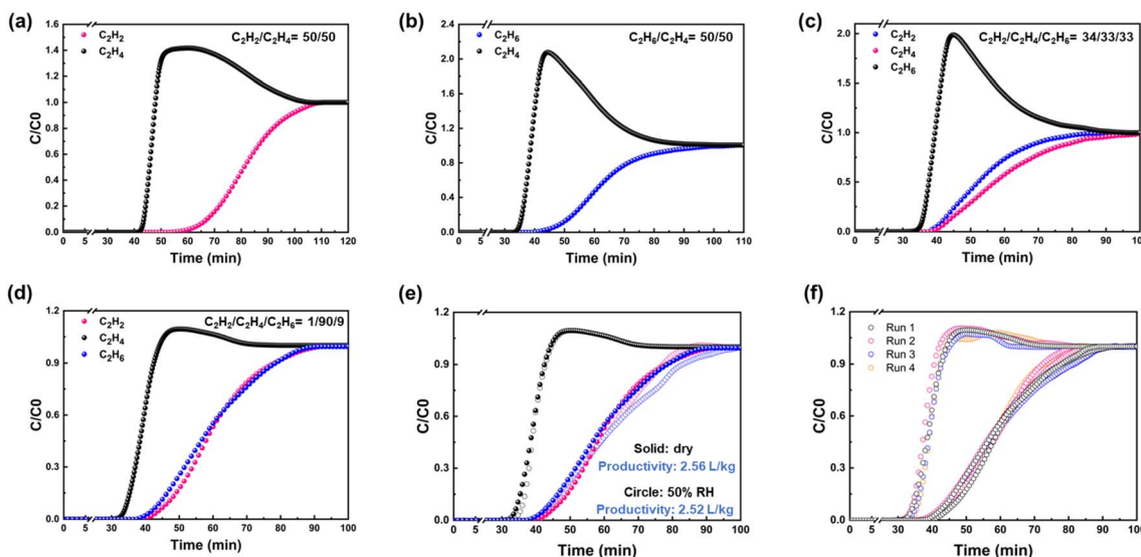


Fig. 4 Dynamic breakthrough tests on HIAM-210. The breakthrough curves for binary mixtures with the composition of (a)  $C_2H_2/C_2H_4$  (50/50, v/v) and (b)  $C_2H_6/C_2H_4$  (50/50, v/v), the breakthrough curves for ternary mixtures with the composition of (c)  $C_2H_2/C_2H_4/C_2H_6$  (34/33/33, v/v/v) and (d)  $C_2H_2/C_2H_4/C_2H_6$  (1/90/9, v/v/v), all collected at 298 K; (e) comparison of the breakthrough curves under humid and dry environments; (f) four consecutive breakthrough cycles at 298 K.



excellent water resistance of HIAM-210. To further evaluate the recyclability of the material, the adsorption column was regenerated under a helium flow ( $10 \text{ mL min}^{-1}$ ) at 373 K, and four consecutive breakthrough cycles were carried out. The results displayed no noticeable decrease of the separation capability of the adsorbent (Fig. 4f). The above tests suggest the durability of HIAM-210 for  $\text{C}_2\text{H}_4$  purification under real-world industrial conditions.

## Conclusions

In summary, we have developed a novel pillar-layer MOF (HIAM-210) combining a suitable pore size and optimal pore environment for simultaneous trapping of  $\text{C}_2\text{H}_2$  and  $\text{C}_2\text{H}_6$  from  $\text{C}_2\text{H}_2/\text{C}_2\text{H}_4/\text{C}_2\text{H}_6$  ternary mixtures. HIAM-210 shows high selectivities for both  $\text{C}_2\text{H}_6/\text{C}_2\text{H}_4$  and  $\text{C}_2\text{H}_2/\text{C}_2\text{H}_4$ . Dynamic breakthrough tests confirmed the outstanding separation performance of HIAM-210 for one-step purification of  $\text{C}_2\text{H}_4$  from  $\text{C}_2\text{H}_2/\text{C}_2\text{H}_4/\text{C}_2\text{H}_6$  ternary mixtures, with high recyclability and strong resistance to moisture. GCMC calculations and DFT calculations demonstrated that the uncoordinated carboxylate-O atoms were the origin of the preferential adsorption. This work not only provides a promising candidate for  $\text{C}_2\text{H}_4$  purification but also offers insights into tailoring the pore environments of MOFs for highly efficient gas separations.

## Data availability

The data that support the findings of this study are available in the ESI† of this article.

## Author contributions

H. Wang and J. Li designed the project. J. Liu synthesized the compounds, conducted and analyzed the PXRD, adsorption, and breakthrough tests. J. Liu, H. Wang, and J. Li wrote the first draft and all authors contributed to the discussion of the final draft.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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