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Linker Desymmetrisation Directs Low-polar Cages in an Anion-Pillared MOF for Acetylene and Ethylene Purification from Ternary Mixtures

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Anion-pillared metal-organic frameworks (APMOFs) constitute a promising class of porous adsorbents, yet precise organization of anionic groups to maximize their strong electronegative character while circumventing highly polar pore environments and high adsorption enthalpies remains a substantial challenge. Here, we report a novel APMOF, CTGU-45, featuring an atypical low-polar cage surface distinct from reported APMOFs and then explore its C₂H₂ and C₂H₄ purification from ternary C₂H₂/CO₂/CH₄ and C₂H₂/C₂H₄/C₂H₆ mixtures. Theoretical calculation results revealed the mechanisms of simultaneous recognition of C₂H₂ and C₂H₆ molecules from C₂ hydrocarbons: the SiF₆²⁻ pillars provide strong binding sites for C₂H₂ via C-H...F hydrogen bonds, while the TIB-derived low-polar surface offers an optimal environment for C₂H₆ adsorption through synergistic C-H...π and C-H...N interactions. Dynamic breakthrough experiments further confirm the efficient separation of C₂H₂/CO₂, C₂H₂/CO₂/CH₄, and notably one-step production of polymer-grade C₂H₄ (≥ 99.95%, 9.57 L kg⁻¹) from ternary C₂H₂/C₂H₄/C₂H₆ mixtures with low regeneration energy. Crucially, CTGU-45 can be synthesized on a 500 mL scale with retained performance, underscoring its practical feasibility and laying a solid foundation for future industrial applications.

Introduction

Acetylene (C₂H₂) and ethylene (C₂H₄) are pivotal feedstocks in the chemical industry.¹ C₂H₂ is typically generated by partial oxidation of methane (CH₄) or thermal cracking of hydrocarbons, which co-produces significant amounts of carbon dioxide (CO₂).²⁻⁵ Similarly, C₂H₄ is obtained via steam cracking or ethane (C₂H₆) dehydrogenation, where C₂H₆ and trace C₂H₂ will unavoidably be produced as by-products.⁶⁻⁷ These impurities severely compromise the purity and productivity of polymer-grade C₂H₄ or high-purity C₂H₂, necessitating highly efficient separation processes.⁸⁻⁹ However, the similar physicochemical properties of these gas molecules make separations (C₂H₂/CO₂/CH₄, C₂H₂/C₂H₄/C₂H₆) particularly difficult (Table S2). Conventional methods like catalytic hydrogenation and cryogenic distillation are complex and energy-intensive, thus driving the urgent need for efficient, alternative separation technologies.¹⁰⁻¹²

Adsorptive separation using porous materials presents a promising, energy-efficient alternative to traditional cryogenic distillation.¹³⁻¹⁶ Among various adsorbents, metal-organic frameworks (MOFs) have garnered immense attention due to their designable pore structures and tunable surface

functionalities. In particular, anion-pillared MOFs (APMOFs) have emerged as an important subclass of MOF adsorbents. These materials utilize inorganic anions (e.g., SiF₆²⁻, GeF₆²⁻, NbOF₅²⁻, MoO₄²⁻) as structural pillars to construct robust frameworks with strong electrostatic potential fields and abundant electronegative atoms that act as hydrogen-bond (H-bond) acceptors, enabling highly selective recognition of C₂H₂ with strong positive H atoms and C₂H₆ with more H atoms. Initially, APMOFs are constructed from linear bidentate ligands to form one-dimensional (1D) channels.¹⁷⁻²⁰ The subsequent introduction of tri- and tetra-dentate ligands created more open APMOFs with a "cage + window" structure, which provides high capacity from large cages and high selectivity from narrow windows. Certainly, these APMOFs have been considered as valuable candidates capable of addressing the "trade-off" between capacity and selectivity in gas separation.

To date, only three highly symmetrical tridentate modules (Tripp, 2,4,6-tris(4-pyridyl)pyridine; TPA, Tri(pyridin-4-yl)amine; and TPT, 2,4,6-tri(4-pyridyl)-1,3,5-triazine) have been successfully employed to construct APMOFs and used for gas separation, such as C₂H₂/CO₂, CO₂/N₂, C₃H₄/C₃H₆, Xe/Kr, and C₆ hydrocarbons. These ligands share critical structural features: their terminal coordinating units are all pyridine rings, which enforce a nearly ideal equilateral triangular arrangement of the three coordination N atoms. Although limited rotation of the pyridine rings is possible, the spatial positions of these key N donors remain essentially fixed, enabling a predictable assembly of frameworks. Interestingly, replacing the terminal pyridyl moiety with less symmetric imidazole rings would introduce significant conformational flexibility. The free rotation around the C-N single bonds linking the central

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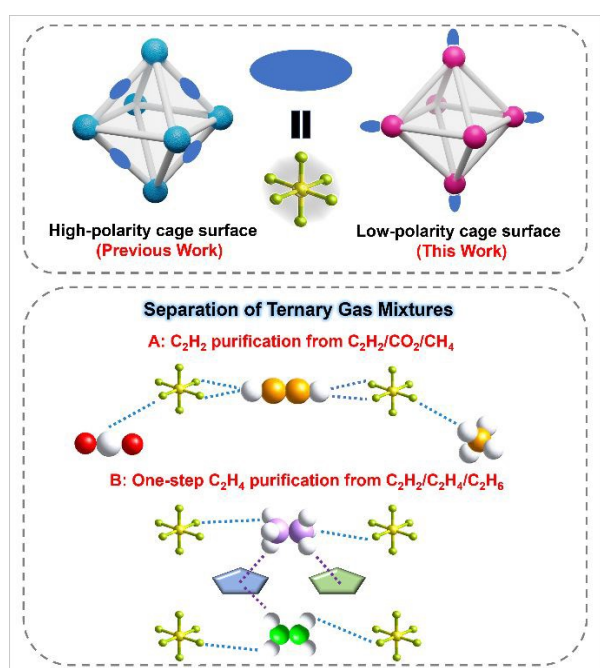
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benzene ring to the imidazole arms results in variable spatial orientations of the coordination N atoms, thereby breaking the inherent topological symmetry and creating opportunities for constructing APMOFs with unique pores. Moreover, certain challenging separations, e.g., one-step C_2H_4 purification from ternary C_2 mixtures, are seldom realized in APMOFs owing to intrinsic polarity mismatches, suggesting that a reconfiguration of functional sites within APMOFs is required. Since the coordination geometry and configuration of inorganic anions are relatively well-defined, the linker desymmetrisation offers a powerful means to systematically reorganize the functional sites within APMOFs. By changing the symmetry of the organic linkers, it becomes possible to create frameworks in which the chemical environment around each coordination site can be independently tuned. This controlled asymmetry allows for the introduction of complementary binding sites, modulation of local polarity, and the rational design of preferential adsorption domains, ultimately enhancing the selectivity and efficiency of challenging gas separations.



Scheme 1 Comparison of cage polarity and using CTGU-45 for the separation of ternary gas mixtures. Unlike the high-polarity cages in typical APMOFs (left), CTGU-45 (right) features a low-polarity cage surface constructed exclusively from Cu^{2+} nodes and TIB ligands. The SiF_6^{2-} pillars served as strong basic sites for C_2H_2 , and non-polar cage surfaces provided the interaction for C_2H_6 , which promotes CTGU-45 to realize the effective C_2H_2 and C_2H_4 purification from $C_2H_2/CO_2/CH_4$ and $C_2H_2/C_2H_4/C_2H_6$ mixtures, respectively.

Keep this in mind, herein, by utilizing a tridentate TIB linker (1,3,5-tri(1H-imidazol-1-yl)benzene) with C_3 symmetry, a novel SiF_6^{2-} anion hybrid Cu-MOF (SIFSIX-Cu-TIB, CTGU-45, CTGU =

China Three Gorges University) has been successfully constructed. Single-crystal X-ray diffraction (SCXRD) analysis reveals that the activated CTGU-45 features uniform cage-like pores with an internal cage diameter of approximately 7.6 Å and a pore window size of about 6.5 Å. In contrast to most reported cage-structured APMOFs, the main pore surface of CTGU-45 is defined solely by TIB ligands and Cu(II) nodes, resulting in a low-polar pore environment that provides optimized adsorption space for C_2H_6 molecules. This establishes a foundation for the one-step production of high-purity C_2H_4 from ternary C_2 hydrocarbon mixtures ($C_2H_2/C_2H_4/C_2H_6$). Meanwhile, the marginalized F atoms of the inorganic SiF_6^{2-} pillars serve as strong basic sites, engaging in pronounced interactions with the acidic H atoms of C_2H_2 , thereby facilitating efficient C_2H_2/CO_2 , $C_2H_2/CO_2/CH_4$, and C_2H_2/C_2H_4 separations (**Scheme 1**). The outstanding separation performances, as well as the underlying adsorbate-adsorbent interactions, have been comprehensively verified by a combination of experimental static adsorption/dynamic breakthrough tests and theoretical calculations. This work not only creates CTGU-45 as an efficient adsorbent for challenging C_2 hydrocarbons separation but also demonstrates the great potential of ligand engineering to break the inherent limitations of conventional APMOFs, opening up new avenues for developing advanced adsorbents.

Results and discussion

Blue octahedral single crystals of CTGU-45 were successfully synthesized by reacting $Cu(OAc)_2 \cdot H_2O$ and TIB ligand in a mixed solvent of *N,N*-dimethylacetamide (DMA)/ H_2O , with HBF_4 added in a sealed glass tube (Fig. S3). SCXRD analysis revealed that CTGU-45 crystallizes in a three-dimensional (3D) framework with the cubic system space group $Pm\bar{3}n$. Each Cu(II) cation is coordinated in a square-planar geometry by four N atoms from imidazole groups of four distinct tridentate TIB ligands (Fig. S2). The SiF_6^{2-} anions as 2-connected nodes occupy the two axial positions to complete the octahedral coordination. Notably, the SiF_6^{2-} anions originated from the in situ etching of the borosilicate glass vessel by HF (from HBF_4 hydrolysis). Due to the free rotation of the C-N bonds between the central benzene and imidazole moieties, the TIB ligands in CTGU-45 exhibit substantial disorder. Upon removal of the minor disordered components, the real structure shows a uniform cage-like architecture with a pore diameter of approximately 7.6 Å (Fig. 1f). Each incomplete octahedral cage displays four identical triangular windows consisting of three TIB linkers and three Cu^{2+} ions. The effective aperture of these windows, after considering the van der Waals radii of H atoms, is only about 6.5 Å, allowing the entry of C_1 - C_2 hydrocarbon molecules (Fig. S1). These cages are further linked by SiF_6^{2-} anion, whose F atoms act as potential Lewis basic sites, which may work synergistically with the confined space and aromatic surface to enhance the binding of gas molecules. In contrast to



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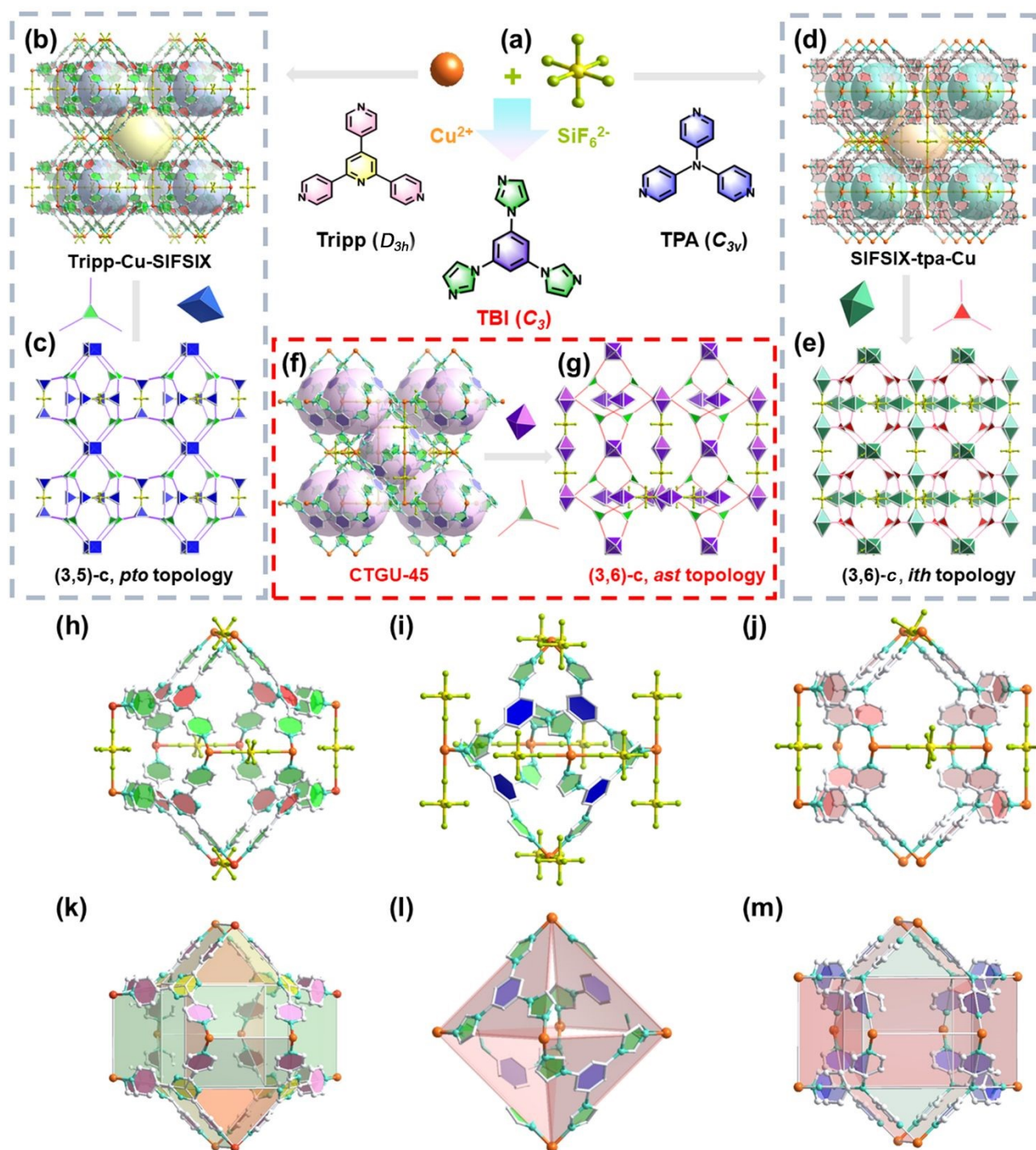
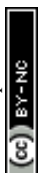


Fig. 1 (a) The frameworks of Tripp-Cu-SIFSIX, SIFSIX-Cu-TPA, and CTGU-45 are based on the assembly of SiF_6^{2-} , Cu^{2+} , and Tripp/TPA/TIB ligands, respectively. (b, c) The crystal structure and (3,5)-connected *pto* topology of Tripp-Cu-SIFSIX. (d, e) Crystal structure and (3,6)-connected *ith* topology of SIFSIX-Cu-TPA. (f, g) The crystal structure and (3,6)-connected topology of CTGU-45. Cage structures within (h)/(k) Tripp-Cu-SIFSIX, (i)/(l) CTGU-45, and (j)/(m) SIFSIX-Cu-TPA. H atoms are omitted for clarity.



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most reported APMOFs, which typically consist of bidentate ligands and anion pillars forming 1D linear channels (e.g., SIFSIX-1-Cu,²¹ SIFSIX-3-Ni²²), the structure presented here features a well-defined windowed cage architecture. Furthermore, APMOFs constructed from tridentate ligands are exceptionally rare. To our knowledge, only three tridentate ligands (Tripp, TPA, and TPT) have been reported for constructing APMOFs, such as Tripp-Cu-SIFSIX²³ (Fig. 1b) and SIFSIX-Cu-TPA (Fig. 1d),²⁴ both of which exhibit two distinct types of cages. Tripp-Cu-SIFSIX displays octahedra cages (~7.2 Å) and icositetrahedron cages (~8.9 Å), while SIFSIX-Cu-TPA possesses icosahedral cages (~6.0 Å) and tetrahedral cages (~4.7 Å). Though the pyridine rings of Tripp (D_{3h} , excluding the non-coordinating pyridine N atom) and TPA (C_{3v}) ligands could rotate around C(pyridine)-C(pyridine) or N-C(benzene) single bonds, their three pyridyl N donors were locked into a fixed, equilateral triangular geometry. In contrast, the rotation of C(benzene)-N(imidazole) bonds in the TIB (C_3) linkers results in dynamic positions of coordination imidazole N atoms, breaking the perfect symmetry, as presented in Figs. 1a and Fig. S4. Such ligand asymmetry easily leads to a distorted coordination geometry and complicates the crystallization process, hindering the predictable assembly of conventional networks. Concretely, in the Tripp-Cu-SIFSIX, SIFSIX-Cu-TPA, and CTGU-45, all the organic linkers act as 3-connected topological nodes with D_{3h} (Tripp/TPA-node) and C_3 (TIB-node) symmetry, respectively. Meanwhile, the coordination numbers of Cu^{2+} are either 5 or 6, exhibiting square pyramidal (C_{4v}) or octahedral (D_{4h}) coordination geometries. The combination of organic nodes and metal nodes ultimately gives rise to (3,5)-c or (3,6)-c topological frameworks in three APMOFs (Fig. 1c/1g/1e). It is noteworthy that although Cu^{2+} acts as a 6-c node in both SIFSIX-Cu-TPA and CTGU-45, the four pyridine rings coordinated to the Cu^{2+} center in the former exhibit relative dihedral angles close to 90°, whereas the four imidazole rings coordinated to Cu^{2+} in the latter are severely twisted. This causes the 6-c nodes in CTGU-45 to link adjacent nodes in a more distorted fashion, breaking the D_{4h} extension symmetry of the 6-c nodes observed in SIFSIX-Cu-TPA. This, together with the 3-c nodes with C_3 symmetry, facilitates the formation of a geometrically self-consistent fundamental cage-like network in the absence of SiF_6^{2-} anions (Fig. S4). The foregoing analysis has validated the critical influence of ligand symmetry on the formation of anionic hybrid frameworks. Overall, compared to Tripp-Cu-SIFSIX and SIFSIX-Cu-TPA, CTGU-45 shares the common feature of incorporating SiF_6^{2-} anions, whose electronegative F atoms serve as Lewis-basic sites for interacting with acidic H atoms of C_2H_2 . However, a fundamental distinction lies in their pore architecture: in conventional APMOFs, the SiF_6^{2-} anions are integral components that participate directly in cage formation,

resulting in highly polar pore surfaces dominated by F atoms (Figs. 1h/1k and 1j/1m). In contrast, in CTGU-45, the pore cages are constructed exclusively by the TIB ligands and $Cu(II)$ nodes, while the SiF_6^{2-} anions merely serve as inter-cage pillars (determined by the topology, Fig. 1i/1l). This unique arrangement creates a low-polar pore surface rich in aromatic rings and N atoms, which not only provides optimal van der Waals interactions with highly polarizable C_2H_6 molecules but also enables synergistic binding through multiple C-H... π and C-H...N interactions. This distinctive pore environment, combined with the selective C-H...F binding sites at the SiF_6^{2-} pillars, allows CTGU-45 to achieve cooperative adsorption of both C_2H_2 and C_2H_6 , thereby providing a robust foundation for one-step C_2H_4 purification from ternary mixtures. The phase purity of the as-synthesized CTGU-45 was confirmed by powder X-ray diffraction (PXRD), as the experimental pattern matches well with simulated ones (Fig. S5). The thermogravimetric analysis (TGA) revealed the high thermal stability of activated CTGU-45, which remains stable up to 350 °C (Fig. S6). The initial weight loss below 100 °C is ascribed to solvent removal, while the subsequent major loss above 350 °C corresponds to the decomposition of the MOF framework. The N_2 adsorption isotherm of CTGU-45 at 77 K displayed a typical Type I profile, and the corresponding HK pore size distribution suggested ultra-micropores of 6.1 Å, collectively confirming its microporous characteristic. The N_2 adsorption uptake of 125.5 $cm^3 g^{-1}$ at 1 kPa and the calculated BET surface area of 544.8 $m^2 g^{-1}$ further corroborated its porous structure (Fig. 2a). Moreover, the experimental pore volume of CTGU-45 was calculated to be 0.21 $cm^3 g^{-1}$, which is consistent with the theoretical value from crystal structure (0.24 $cm^3 g^{-1}$). Combined with its excellent thermal stability, these integrated properties collectively endorse CTGU-45 as a highly promising adsorbent for gas separation applications.

Single-component adsorption isotherms of CH_4 , CO_2 and C_2-C_3 hydrocarbons on CTGU-45 were measured at 273, 283 and 298 K (Figs. 2b, 2c and S7-S18). CTGU-45 exhibited an obviously preferential adsorption of C_2H_2 over CO_2 and CH_4 , with an uptake of 2.83 $mmol g^{-1}$ at 298 K and 101 kPa, significantly exceeding those of CO_2 (1.64 $mmol g^{-1}$) and CH_4 (0.74 $mmol g^{-1}$). Notably, the uptakes of C_2H_2 (2.00 $mmol g^{-1}$) and C_2H_6 (1.88 $mmol g^{-1}$) are significantly higher than that of C_2H_4 (1.75 $mmol g^{-1}$) at 30 kPa, thus showing great potential for one-step C_2H_4 purification. The C_2H_2 uptake capacity of CTGU-45 is higher than that of many reported APMOFs, such as NBOFFIVE-dps-Cu (1.65 $mmol g^{-1}$),²⁵ sql-SIFSIX-bpe-Zn (1.79 $mmol g^{-1}$),²⁶ DICRO-4-Ni-I (1.92 $mmol g^{-1}$),²⁷ ZNU-15 (1.61 $mmol g^{-1}$),²⁸ BSF-2 (1.85 $mmol g^{-1}$),²⁹ and BSF-4 (2.38 $mmol g^{-1}$).³⁰



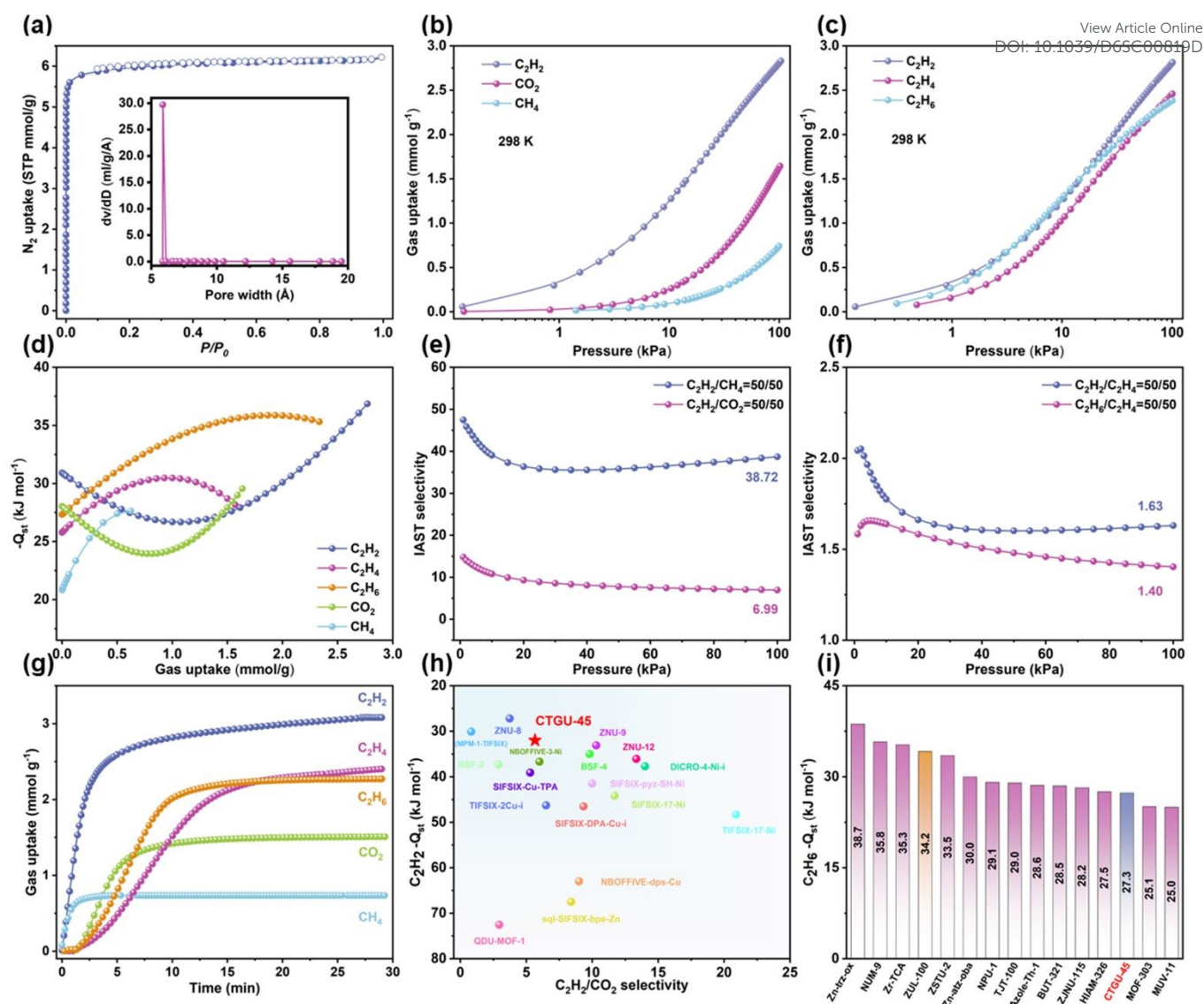


Fig. 2 (a) N_2 adsorption-desorption isotherms recorded at 77 K. The inset represents pore size distribution analyzed based on the H-K method. The single-component adsorption isotherms of (b) C_2H_2 , CO_2 , CH_4 and (c) C_2H_2 , C_2H_4 , C_2H_6 on CTGU-45 at 298 K. (d) Adsorption heats of CH_4 , CO_2 , and C_2 hydrocarbons. IAST selectivity of (e) C_2H_2/CO_2 (v/v, 50/50), C_2H_2/CH_4 (v/v, 50/50) mixtures and (f) C_2H_2/C_2H_4 (v/v, 50/50), C_2H_6/C_2H_4 (v/v, 50/50) mixtures at 298 K. (g) Adsorption kinetics profiles of CTGU-45 for C_2H_2 , C_2H_4 , C_2H_6 , CO_2 , and CH_4 . (h) Comprehensive comparison of C_2H_2/CO_2 selectivity and $C_2H_2-Q_{st}$ value of CTGU-45 with the reported APMOFs. (i) A comparison of near-zero coverage isosteric heats for C_2H_6 among representative porous materials employed for one-step C_2H_4 purification from C_2 hydrocarbon mixtures.

Meanwhile, the CO_2 adsorption amount is markedly lower than that of top-performing APMOFs, including SIFSIX-Cu-TPA (4.79 mmol g^{-1}),²⁴ QDU-MOF-1 (5.70 mmol g^{-1}),³¹ TIFSIX-2-Cu-I (4.3 mmol g^{-1}),³² ZNU-9 (4.32 mmol g^{-1}),³³ and ZNU-12 (4.33 mmol g^{-1}).³⁴ According to the virial equation, the isosteric heat of adsorption (Q_{st}) values at near-zero loading for C_2H_2 , C_2H_4 , C_2H_6 , CO_2 , and CH_4 on CTGU-45 are 30.9, 25.8, 27.3, 28.0, and 20.9 kJ mol^{-1} , respectively (Figs. 2d and S24-S28). The higher Q_{st} value for C_2H_2 and C_2H_6 , consistent with the steeper slope of its single-component adsorption isotherm, indicates a stronger binding affinity between the framework and C_2H_2 and C_2H_6 molecules. Overall, except for ZNU-8 (27.2 kJ mol^{-1})³³ and MPM-

1-TIFSIX (30.1 kJ mol^{-1}),³⁵ CTGU-45 exhibits the lowest Q_{st} value for C_2H_2 among reported APMOFs, which is significantly lower than that of QDU-MOF-1 ($72.52 \text{ kJ mol}^{-1}$),³¹ TIFSIX-17-Ni (48.3 kJ mol^{-1}),³⁶ NBOFFIVE-dps-Cu (53.6 kJ mol^{-1}),³⁷ SIFSIX-Cu-TPA (39.1 kJ mol^{-1})²⁴ and so on (Fig. 2h). Given that only one APMOF (ZUL-100)³³ has been reported for C_2H_4 purification from C_2 mixtures, the C_2H_6 Q_{st} value of CTGU-45 (27.3 kJ mol^{-1}) is not only lower than that of ZUL-100 (34.2 kJ mol^{-1}),³³ but also lower than those of most previously reported C_2H_2/C_2H_6 -selective MOFs for one-step C_2H_4 purification, such as Zn-trz-ox (38.7 kJ mol^{-1}),³⁸ NUM-9 (35.8 kJ mol^{-1}),³⁹ Zn-atz-oba (30.0 kJ mol^{-1}),⁴ TJT-100 (29.0 kJ mol^{-1}),⁴⁰ and BUT-321 (28.5 kJ mol^{-1})⁴¹ (Fig. 2i). This low Q_{st} value ensures easy desorption and low-energy regeneration, which



might significantly reduce the operating cost in practical separation cycles.

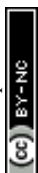
In addition, since practical gas separation is a non-equilibrium process, the kinetic adsorption behavior should be evaluated synchronously. Therefore, we studied the kinetic diffusion rate of C_2H_2 , C_2H_4 , C_2H_6 , CO_2 , and CH_4 in CTGU-45. As shown in Fig. 2g, the time required to reach adsorption equilibrium follows the order: $T_{ads}(CH_4) < T_{ads}(C_2H_2) < T_{ads}(CO_2) < T_{ads}(C_2H_6) < T_{ads}(C_2H_4)$. In detail, the calculated diffusion time constants (D_c/r_c^2) of these gases on CTGU-45 are 0.0380 (CH_4), 0.2628 (C_2H_2), 0.0284 (CO_2), 0.0141 (C_2H_6), and 0.0104 s^{-1} (C_2H_4), respectively (Fig. S29). Interestingly, in the desorption kinetics tests (Fig. 30), while CH_4 and CO_2 still exhibited the fastest desorption, the desorption rate sequence of the C_2 gases ($T_{des}(C_2H_4) < T_{des}(C_2H_6) < T_{des}(C_2H_2)$) was exactly opposite to their respective adsorption rate sequence. This observation directly reflects the relative strength of their interactions with the framework, consistent with the assessments discussed earlier.

To further assess the separation potential of CTGU-45, the adsorption selectivity for equimolar C_2H_2/CO_2 , C_2H_2/CH_4 , C_2H_2/C_2H_4 , and C_2H_6/C_2H_4 mixtures at 298 K was calculated based on the Ideal Adsorbed Solution Theory (IAST) after fitting isotherms to the single-site/dual-site Langmuir-Freundlich equation with excellent accuracy (Fig. S19-S23, $R^2 > 0.99999$). As illustrated in Fig. 2e-2f, the IAST selectivity of CTGU-45 at 298 K and 101 kPa reached 6.99 for C_2H_2/CO_2 and 38.72 for C_2H_2/CH_4 ; notably, the selectivity for C_2H_2/C_2H_4 (50/50, v/v) (1.63) and C_2H_6/C_2H_4 (50/50, v/v) (1.40) further underscore its potential for one-step C_2H_4 purification. The C_2H_2/CO_2 selectivity is moderate among benchmark APMOFs, being comparable to those of NBOFFIVE-3-Ni (6.0) and TIFSIX-2-Cu-I (6.5), and higher than SIFSIX-Cu-TPA (5.3), QDU-MOF-1 (2.95), and ZNU-8 (3.74) (Fig. 2h). Meanwhile, the moderate C_2H_2/CO_2 selectivity, combined with the high C_2H_2/CH_4 selectivity, suggests that CTGU-45 is a viable candidate for separating C_2H_2/CO_2 and C_2H_2/CH_4 mixtures. Furthermore, the calculated C_2H_2 and CO_2 uptake (2.20 and 0.32 $mmol\ g^{-1}$) in the equimolar mixture closely matches the corresponding single-component adsorption capacities at 50 kPa (2.37 and 1.02 $mmol\ g^{-1}$), confirming the consistency between mixture prediction and gas adsorption measurement.

Grand Canonical Monte Carlo (GCMC) simulations and density functional theory with dispersion corrections (DFT-D) calculations were employed to elucidate the gas adsorption mechanisms in CTGU-45, revealing two primary binding sites within the framework: one near the SiF_6^{2-} anion pillars and another inside the pore cavity (Fig. 3). In general, the region adjacent to the SiF_6^{2-} pillars serves as a strong adsorption site for CO_2 , CH_4 , C_2H_2 , and C_2H_4 , whereas the pore cavity functions as the preferred strong binding region for C_2H_6 . Taking C_2H_2 as a representative case, the first corresponds to a strong adsorption site, where the C_2H_2 molecule was confined by two SiF_6^{2-} with a binding energy of $-53.42\ kJ\ mol^{-1}$. Specifically, each H atom of C_2H_2 forms two C-H...F hydrogen bonds (H-bonds) with F atoms from two adjacent SiF_6^{2-} pillars, resulting in a total of four H-bonds with distances ranging from 2.34 to 3.69 Å.

These are complemented by $C\equiv C\cdots H$ interactions (distances: 2.58-3.45 Å) between the π -electrons of C_2H_2 and H atoms of imidazole rings. The second site corresponds to a weaker interaction environment within the pore cavity, involving $C\equiv C\cdots H$ (distances: 3.85-3.94 Å), C-H (C_2H_2)...N(imidazole) (2.91-3.82 Å), and C-H (C_2H_2)... π (imidazole/benzene) interactions with the distances of 3.10-3.92 Å. The binding energy for this weak-interaction site is $-19.92\ kJ\ mol^{-1}$, substantially lower than that of the above-mentioned site. A similar distribution of adsorption sites was observed for CO_2 , CH_4 , and C_2H_4 with strong binding near the SiF_6^{2-} pillars and weaker interactions in the pore cavity. For CO_2 , the electronegative O atoms participated in $C=O\cdots H$ and $C=O\cdots N$ contacts with H and N atoms of the imidazole rings, as well as $C=O\cdots\pi$ interactions with the aromatic systems. Simultaneously, its electropositive C atom interacts with F atoms of the SiF_6^{2-} pillars via $O=C\cdots F$ interactions. In the case of CH_4 , C_2H_4 , and C_2H_6 , adsorption was governed by C-H...F and C-H...N interactions with the pillar F and imidazole N atoms, respectively, along with C-H... π contacts involving the aromatic rings of the TIB ligands. Notably, C_2H_6 exhibits its strongest adsorption within the pore cavity rather than near the anion pillars. Consequently, the binding energies of CO_2 and CH_4 at both types of adsorption sites are substantially lower than those of C_2H_2 , while those of C_2H_2 and C_2H_6 are higher than those of C_2H_4 . This energetic trend ($C_2H_2 > CO_2 > CH_4$; $C_2H_2 > C_2H_6 > C_2H_4$) further rationalizes the preferential adsorption of C_2H_2 over CO_2 and CH_4 , and of C_2H_2 and C_2H_6 over C_2H_4 by CTGU-45. The stronger intermolecular interactions between the framework and C_2H_2/C_2H_6 directly correlate with its higher adsorption affinity, which is consistent with the observation on Q_{st} and single-component adsorption data.

Dynamic breakthrough experiments were conducted to further evaluate the practical separation performance of CTGU-45 for C_2H_2/CO_2 , $C_2H_2/CO_2/CH_4$, and C_2 hydrocarbon mixtures. As depicted in Fig. 4a, CTGU-45 can achieve efficient separation of a C_2H_2/CO_2 (50/50, v/v) mixture at a total gas flow rate of 4.0 $mL\ min^{-1}$, wherein CO_2 gases were detected in $\sim 9\ min\ g^{-1}$ while C_2H_2 breakthrough occurred at $\sim 22\ min\ g^{-1}$. Based on the integration of the curve, the dynamic CO_2 and C_2H_2 capture amounts of CTGU-45 were calculated to be 6.10 and 47.68 $cm^3\ g^{-1}$, corresponding to approximately 26 and 90% of their respective static uptake values at 50 kPa (23.04 and 53.12 $cm^3\ g^{-1}$), respectively. After adsorption saturation, the column was regenerated by purging with helium at 10 $mL\ min^{-1}$ under ambient temperature. The desorption profiles indicated rapid release of both gases, with CO_2 completely desorbed within 23 $min\ g^{-1}$. Beyond this point, high-purity C_2H_2 ($> 95.2\%$) was obtained at a productivity of 22.1 $L\ kg^{-1}$ (Fig. 4a). Subsequently, we further investigated the separation capacity of CTGU-45 for actual ternary $C_2H_2/CO_2/CH_4$ mixtures (33/33/33, v/v/v) with flow rates of 6 and 3 $mL\ min^{-1}$. Owing to its higher C_2H_2 uptakes and selectivity over CO_2 and CH_4 , efficient separation of C_2H_2 from $C_2H_2/CO_2/CH_4$ mixtures can be fulfilled by CTGU-45 under different temperatures (298 and 308 K, Fig. S31).



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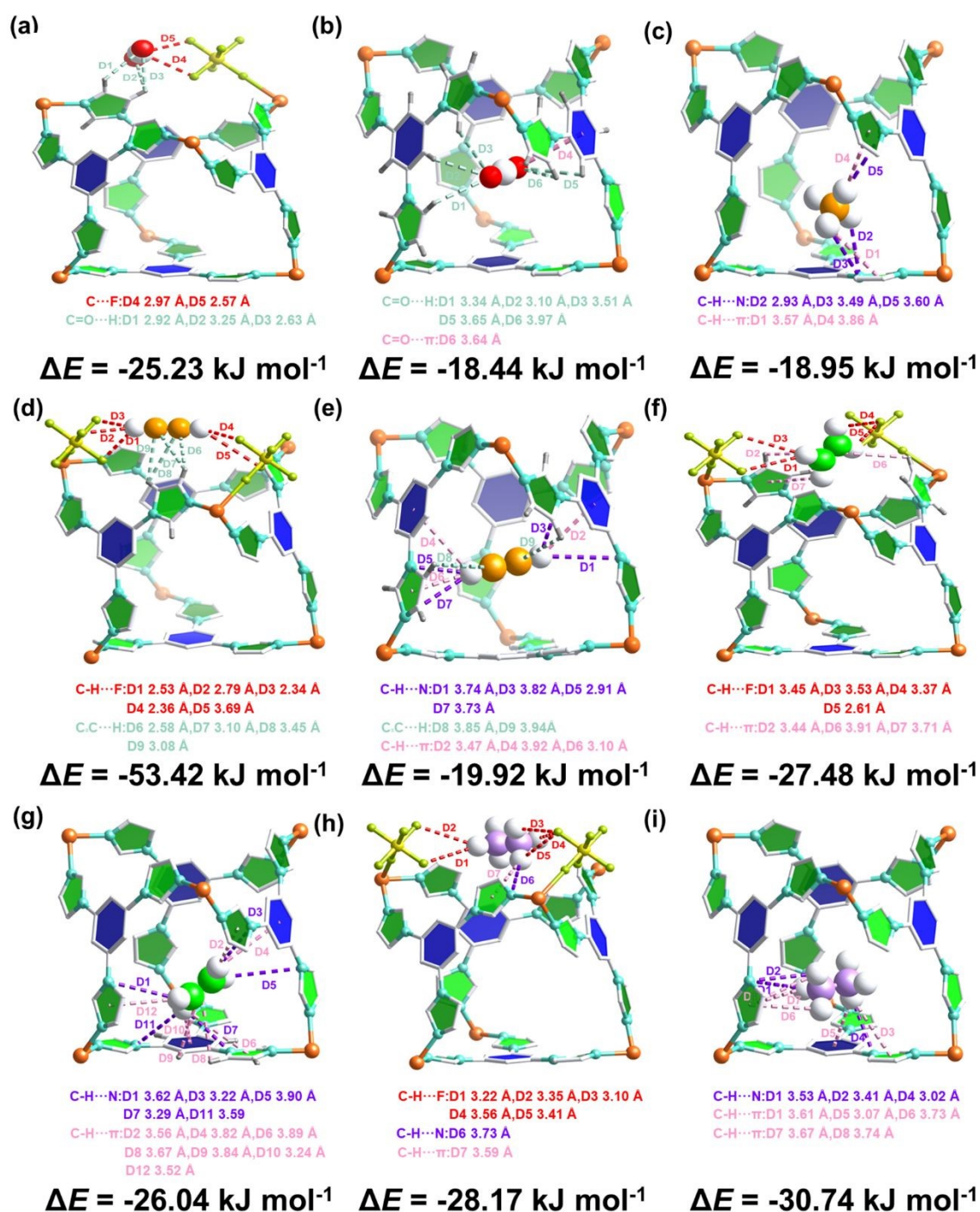


Fig. 3 Optimal binding sites and adsorption configurations of (a-b) CO_2 , (c) CH_4 , (d-e) C_2H_2 , (f-g) C_2H_4 , and (h-i) C_2H_6 molecules within CTGU-45 defined by DFT-D calculations.

As illustrated in Fig. 4b, CH_4 first eluted through the fixed bed at 3 min g^{-1} , followed by CO_2 at 4 min g^{-1} , whereas the C_2H_2 was retained in the packed column over 18 min g^{-1} . Despite CO_2

and CH_4 exhibiting a faster adsorption rate than C_2H_2 in CTGU-45, the stronger host-guest interactions between the framework and C_2H_2 ($\text{C}-\text{H}\cdots\text{F}$, $\text{C}-\text{H}\cdots\text{N}$, and $\text{C}-\text{H}\cdots\pi$ interactions)



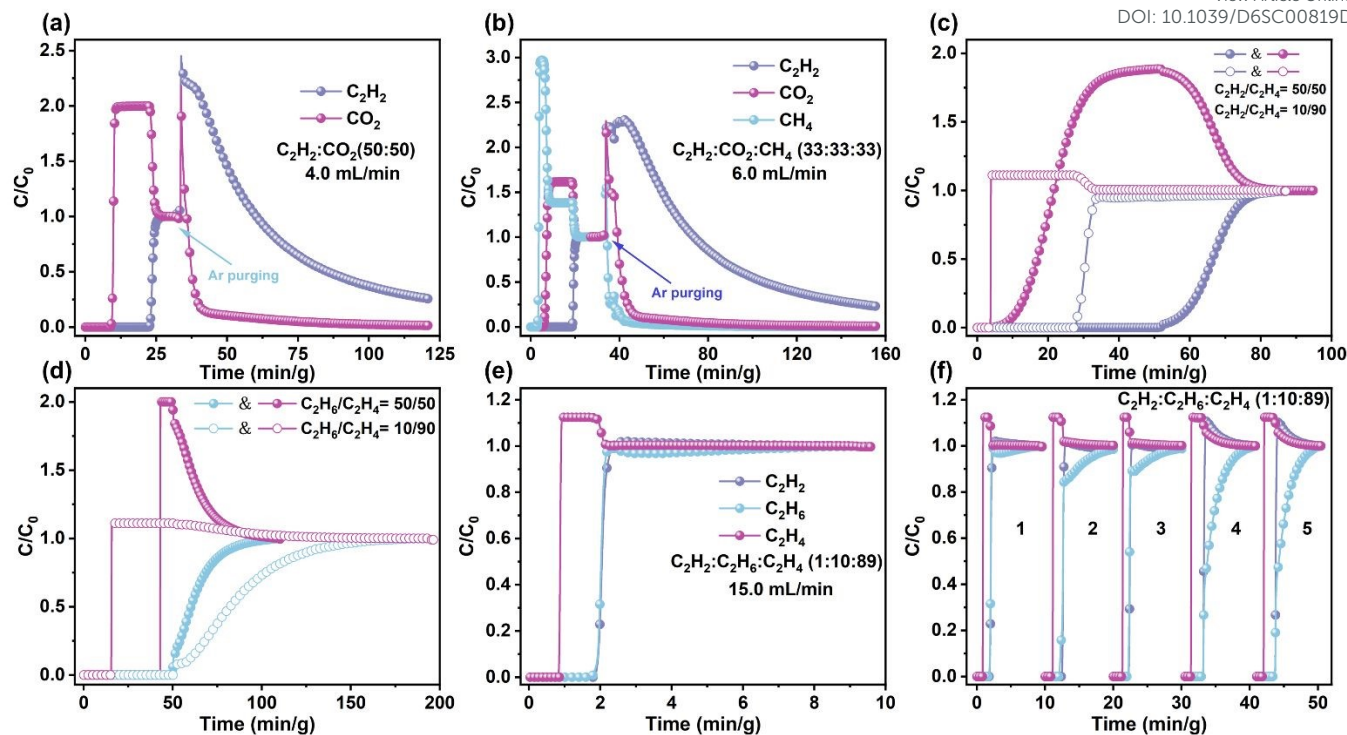


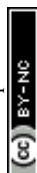
Fig. 4 Experimental column breakthrough curves and the desorption curves for (a) C_2H_2/CO_2 (50/50, v/v) and (b) $C_2H_2/CO_2/CH_4$ (33/33/33, v/v/v) on CTGU-45 at 298 K with a total flow of (a) 4 and (b) 6 $mL\ min^{-1}$. The dynamic breakthrough curves for (c) C_2H_2/C_2H_4 (50/50 and 10/90, v/v), (d) C_2H_6/C_2H_4 (50/50 and 10/90, v/v), and (e) $C_2H_2/C_2H_6/C_2H_4$ (1/10/89, v/v/v) on CTGU-45 at 298 K with a total flow of (c) 1, (d) 1, and (e) 15 $mL\ min^{-1}$. (f) Breakthrough cycling test of CTGU-45 for $C_2H_2/C_2H_6/C_2H_4$ (1/10/89, v/v/v, 298 K, 15 $mL\ min^{-1}$).

confer thermodynamic dominance in the actual separation process, leading to preferential C_2H_2 capture and the longest breakthrough time. The desorption curve further confirmed that polymer-grade C_2H_2 can be recovered at a productivity (> 95.2%) of 19.7 $L\ kg^{-1}$.

For C_2H_2/C_2H_4 and C_2H_6/C_2H_4 mixtures with different compositions (50/50 or 10/90, v/v), C_2H_4 first eluted from the packed column while C_2H_2 or C_2H_6 remained strongly retained within the CTGU-45, demonstrating its ability to directly produce polymer-grade C_2H_4 products. In detail, C_2H_4 molecules broke through the fixed bed and were detected at 4.2 (3.7) and 42.9 (15.5) $min\ g^{-1}$ for the 50/50 (10/90) C_2H_2/C_2H_4 and C_2H_6/C_2H_4 mixtures, respectively, whereas C_2H_2 and C_2H_6 were adsorbed in the column over 51.7 (27.6) and 49.9 (51.4) $min\ g^{-1}$. As depicted in Figs. 4c-d, the productivity of 99.9% pure C_2H_4 from the 50/50 (10/90) C_2H_2/C_2H_4 and C_2H_6/C_2H_4 mixtures reached 28.6 (47.7) and 6.0 (33.3) $L\ kg^{-1}$, respectively. Building on the excellent separation performance observed for these binary mixtures, breakthrough experiments were further conducted using a ternary $C_2H_2/C_2H_6/C_2H_4$ (1/10/89, v/v/v) mixture. Notably, C_2H_4 eluted (0.9 $min\ g^{-1}$) prior to both C_2H_2 (1.8 $min\ g^{-1}$) and C_2H_6 (1.7 $min\ g^{-1}$), enabling the direct production of high-purity C_2H_4 (Fig. 4e). After calculation, 9.57 $L\ kg^{-1}$ of C_2H_4 with $\geq 99.95\%$ purity can be directly obtained in a single cycle (298 K, 15 $mL\ min^{-1}$). For both ternary mixtures, CTGU-45 maintains reliable efficiency in C_2H_2 and C_2H_4

purification with steady gas productivities when the operating temperature is raised from 298 to 308 K, or when the flow rate is decreased from 6 to 3 $mL\ min^{-1}$ (15 to 14 $mL\ min^{-1}$, Fig. S31). Moreover, repeated breakthrough experiments over five consecutive cycles showed no discernible decay in either C_2H_4 purity or productivity (9.24~11.0 $L\ kg^{-1}$), confirming the stable one-step C_2H_4 purification capability of CTGU-45 (Fig. 4f, Fig. S32). Finally, PXRD measurements reveal that CTGU-45 displays moderate chemical stability across a wide pH range (1–13) and in aqueous environments (Fig. S33). The combination of high separation efficiency, excellent adaptability, good thermal/chemical stability, and mild regeneration requirements (room-temperature He purging) underscores the potential of CTGU-45 for practical C_2H_2 and C_2H_4 purification.

In addition to the impressive separation properties, CTGU-45 crystals can be successfully synthesized at scale-up levels using glass containers with volumes ranging from 50 to 500 mL (Fig. 5a). PXRD measurements confirm the basic phase purity of the CTGU-45 sample obtained via scaled-up synthesis (Fig. 5b). Although the N_2 adsorption capacity of samples synthesized in 50-, 100-, and 200-mL vessels were only slightly lower than that of the 20 mL reference (Fig. 5c), and the sample prepared in a 500 mL bottle exhibits a 25.2% reduction in N_2 uptake. To further assess the impact on functional performance, C_2H_2 adsorption isotherms were measured at 298 K (Fig. 5d). The results indicate that the C_2H_2 uptake of samples from 20- and 50-mL vessels is nearly identical, while those from 100- and 200-



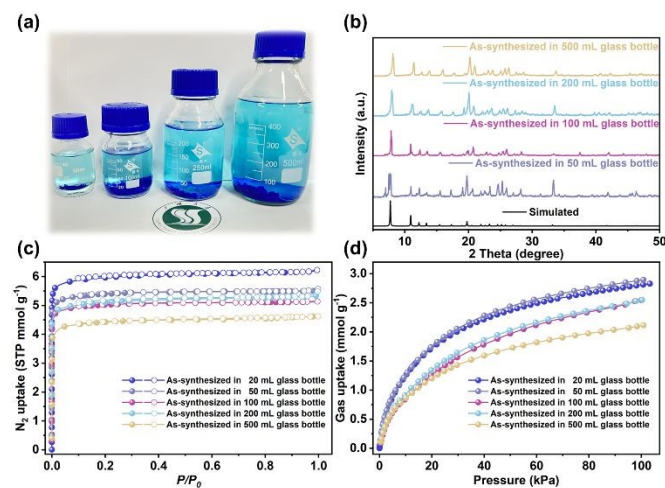


Fig. 5 (a) Scale-up synthesis of CTGU-45 at scales of 50, 100, 200, and 500 mL. (b) PXRD patterns, (c) N₂ adsorption-desorption isotherms, and (d) C₂H₂ adsorption isotherms of CTGU-45 synthesized at different scales via a solvothermal method.

mL vessels show an approximate 8.8% decrease. In contrast, the sample synthesized in the 500 mL bottle exhibits a 25.4% reduction in C₂H₂ adsorption capacity, consistent with the trend observed in N₂ adsorption. Through additional analysis of the N₂ ads/des-orption isotherms, pore size distributions (Fig. S34), and PXRD patterns of samples synthesized at different scales, we found that the decrease in adsorption capacity may arise not only from undesired structural defects but also from the presence of by-products (amorphous, non-porous, and non-adsorbing). Since MOF synthesis is a complex self-assembly process governed by both thermodynamic and kinetic factors, changes in the scale of the reaction system—that is, the microscopic reaction conditions—can perturb the reaction pathway, leading to the formation of unknown by-products. To mitigate the formation of amorphous by-products and structural defects during industrial production for CTGU-45, future efforts might focus on two key strategies: (i) The implementation of continuous-flow synthesis or microwave-assisted heating can ensure uniform nucleation and growth conditions, thereby suppressing kinetic traps that lead to amorphous phases; (ii) dynamic adjustment of reaction parameters (e.g., reactant concentration, temperature gradients, local mass transfer) to steer the self-assembly toward the desired crystalline product.

Conclusions

In summary, we reported a SiF₆²⁻ embedded MOF, CTGU-45, featuring abundant electronegative F atoms and a low-polar cage environment, which enables one-step C₂H₄ purification via the simultaneous removal of C₂H₂ and C₂H₆. CTGU-45 exhibits high C₂H₂ and C₂H₆ uptake, markedly exceeding those for CO₂, CH₄, and C₂H₄. The low-polar pore environment effectively addresses the critical challenge of achieving C₂H₆-selective adsorption in C₂H₆/C₂H₄ separation, a behavior that is

exceedingly rare among reported APMOFs. As a result, ~9.57 kg⁻¹ of polymer grade C₂H₄ (≥ 99.95%) can be directly produced from ternary C₂H₂/C₂H₆/C₂H₄ (1/10/89, v/v/v) mixtures in a single step. DFT calculations reveal that strong C-H...F interactions between the inorganic SiF₆²⁻ pillars and C₂H₂ enable CTGU-45 to not only efficiently purify C₂H₂ from C₂H₂/CO₂/CH₄ mixtures but also, in synergy with its C₂H₆-trapping capability, to achieve one-step production of high-purity C₂H₄ from ternary C₂ hydrocarbon mixtures. In addition, the low C₂H₆ Q_{st} (27.3 kJ mol⁻¹) ensures facile regeneration with low energy input. Notably, CTGU-45 can be synthesized on a scalable level (up to 500 mL) with well-retained adsorption capacity. Overall, this work demonstrates that breaking ligand symmetry in APMOFs can fundamentally alter the spatial role of anion pillars, thereby reshaping pore surface chemistry and enabling rare one-step C₂H₄ purification from C₂ hydrocarbon mixtures.

Author contributions

J.-J. Wu., P.-D. Zhang., X.-Q. Wu., and D.-S. Li. conceived this project and designed the experiments. J.-J. Wu., S.-H. Huang., W.-W. Dong. and Y.-P. Wu conducted the experiments. J.-J. Wu., P.-D. Zhang. and X.-Q. Wu. drafted the work. D.-S. Li. revised and finalised the manuscript. All authors approved the submitted version.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article are available in the SI file, which includes experimental procedures, TGA curves, gas adsorption/separation isotherms, crystallographic data, and theoretical computational details. See DOI:***. CCDC 2522768 contains the supplementary crystallographic data for this paper.

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The data that support the findings of this study are openly available in the Supporting Information. [View Article Online](#)
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