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Confinement-modulated diffusion of alkenes in NU-1000 framework material

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Metal—organic frameworks (MOFs) and similar molecular frameworks are increasingly investigated in catalysis for energy applications due to the tunability of these materials for spatially controlled positioning of catalytic active sites, combined with tailor-made porosity. However, because of the complex interplay of catalytic reactions and diffusion processes, the influence of the framework is not well understood. In this study, we used molecular dynamics (MD) and grand canonical Monte Carlo (GCMC) simulations to explore the confinement effects of the NU-1000 MOF on the diffusivity and adsorption of α -olefins. We provide a direct comparison of diffusion coefficients under non-confined conditions and a reduction in diffusion constants primarily governed by MOF—olefin interactions rather than olefin—olefin interactions, with higher mobility along the channel axis than perpendicular to it.

MOFs are promising carrier materials for single atom catalysts (SAC)¹⁻³ due to their tunable composition, structural stability, SAC stabilization at the atomistic level,⁵ and ease of product separation.6 They are beneficial especially for reactions demonstrating the crucial role of selectivity, e.g. ethylene oligomerization. For example, Ni-grafted MOFs demonstrated activity levels, raising ethylene conversion, that are one to two orders of magnitude higher than classical porous materials like zeolites and oxides. The elevated activity is influenced by confinement effects and the coordination of metals.8 Similar observations were reported for NU-1000 MOF9 (schematically shown in Fig. 1A and B) after its functionalization with SAC. 10,11 This MOF features a hierarchical pore structure with mesochannels and microchannels, enabling selective reagent adsorption and diffusion, making it well-suited for catalysis and ethylene oligomerization, 10-12 previously studied in NU-1000 at ~300 K and 1-15 bar. 10,12 Note that common conditions reported for porous materials span wider ranges, for example 300-600 K and 1-50 bar. 13

While metal catalysts exploit local electronic properties, MOF confinement additionally governs selectivity, kinetics, and separation *via* diffusion and adsorption. Haag¹⁴ showed cracking chain-length effects stem from stronger adsorption of longer hydrocarbons in zeolites rather than variations in zeolite acidity. Bickel and Gounder¹⁵ reported the change in catalytic performances by the dynamic composition of in-pore species and additional diffusional constraints on reactants and products during chain-growth reactions in porous catalysts. However, detailed investigations of these effects remain scarce: recent reviews note that studies beyond binary mixtures are rare and overwhelmingly centered on alkanes.^{16,17}

Here, we employ GCMC/MD simulations to investigate the co-adsorption and diffusion of a full homologous series of α -olefins (C_2 - C_{10}) within the hierarchical pores of NU-1000, considering typical experimental conditions used for ethylene oligomerization. Due to its proven thermophysical and vapor-liquid equilibrium accuracy, 18,19 we perform simulations by combining the Extension of the Universal Force Field for Metal-Organic Frameworks (UFF4MOF) for unmodified NU-1000 MOF and Transferable Potentials for Phase Equilibria (TraPPE) force field for guest molecules (see SI for more details).

For NU-1000, like other mesoporous materials, type IV or V isotherms are typical according to the International Union of Pure and Applied Chemistry classification. PNU-1000 shows type IV isotherms for N_2 , indicating weak adsorbate—adsorbent interactions. For the adsorption of C_4H_8 and C_6H_{12} in NU-1000, we observe a type I isotherm (Fig. 1C), which is typically associated with microporous materials and strong adsorbate—adsorbent interactions. This behavior arises from significantly stronger van der Waals interactions of olefins with the MOF scaffold and the contribution of microchannels of NU-1000, as we discuss further. A similar type I-like isotherm profile was reported for n-butane in NU-1000 at pressure below saturation, and for 1-butene in MOF-74. Heavier olefins in the liquid phase (Fig. 1E), as well as C_4H_8 at $P > P_0$, exhibit negligible pressure dependence, indicating pore saturation.

In olefin mixtures, the mutual influence of other molecules is evident, for example, when comparing the adsorption of

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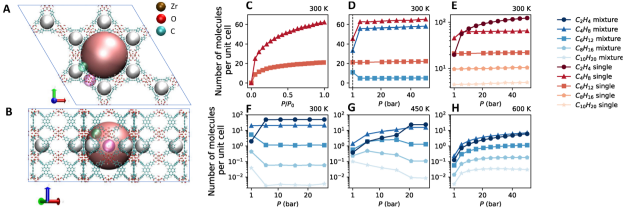


Fig. 1 Visualization of $2 \times 2 \times 2$ unit cell of NU-1000: (A) viewed along the z-direction (blue arrow) and (B) viewed along the in x-y-plane (red and green arrows). Hexagonal mesochannels of 32 Å, (pink spheres) triangle microchannels of 12 Å (white spheres), windows between mesochannels of 8 Å (purple transparent sphere), and a microchannel and mesochannel (green transparent sphere) with the size of 7 Å are marked for clarity. GCMC-simulated adsorption isotherms of olefins: (C) for single-component C_4H_8 (saturation pressure $P_0 = 3.14$ bar) and C_6H_{12} ($P_0 = 0.27$ bar) at 300 K. (D) Comparison of single-component and equimolar systems of two components C_4H_8 and C_6H_{12} at 300 K. (E) Single component systems $C_2H_4-C_{10}H_{20}$ at 300 K. (F)–(H) Equimolar systems of five components olefins ranging from C_2H_4 to $C_{10}H_{20}$ at 300 K, 450 K, and 600 K, respectively.

C₄H₈ and C₆H₁₂ independently (i.e. as single components) versus their equimolar mixture (Fig. 1D). While C₆H₁₂ alone shows negligible pressure dependence, in the presence of C₄H₈, its adsorption atypically decreases with increasing pressure from 5 to 10 bars. This behavior stems from competitive adsorption, where increased C₄H₈ uptake at high pressures displaces C₆H₁₂. Such mutual influence is also observed in equimolar mixtures of more than two olefins, as shown in Fig. 1F. Lighter components (C₂H₄ to C₄H₈) replace heavier olefins in NU-1000 until the plateau corresponding to saturation is reached. While lighter olefins exhibit a monotonic increase in adsorption, heavier ones, such as C₁₀H₂₀, show a monotonic decrease. Olefins with saturation pressures within this range (see Section S5 in SI), such as C₆H₁₂, display nonmonotonic behavior, increasing until P = 15.7 bar before decreasing as the phase transitions occur (Fig. 1G). As the temperature elevates, more olefins exhibit a monotonic behavior with their loading increasing consistently with the pressure

increase (see Fig. 1H). In addition, typically, the adsorption capacity diminishes with the chain length increase, *i.e.* as observed for olefins ranging from C_4H_8 to $C_{10}H_{20}$. However, in the case of C_2H_4 , the adsorption capacity is lower than that of C_4H_8 . In the provided examples (Fig. 1F and G), within the pressure range of 25 to 50 bars, there is no substantial variation in the adsorption behavior (see Fig. S5).

To elucidate the adsorption sites of olefins in NU-1000, we analyzed the spatial distribution of molecules from GCMC simulations at 300 K using $\rm C_4H_8$ as an example, an olefin with the highest adsorption loading (see Fig. 2A). It can be seen that the molecules are more evenly distributed throughout the triangle microchannels, while adsorption in the hexagonal mesochannel occurs primarily near the MOF linkers rather than in the channel center or near the metal node in mesochannel. A similar tendency was reported in the binding energies and adsorption enthalpies from the density functional theory calculations of $\rm CH_4$ and $\rm C_2H_6$ in NU-1000, showing a

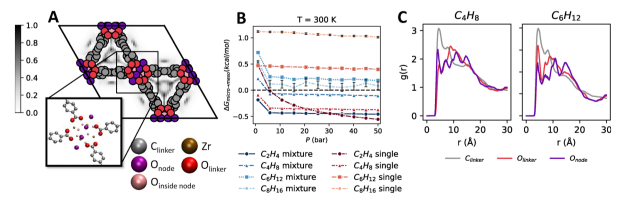


Fig. 2 (A) Heat map displaying the normalized probability distribution of C_4H_8 adsorption in NU-1000 MOF at 300 K and 50 bars in z-direction; darker regions indicate higher probability of finding a molecule at specific position. (B) The free energy change associated with transferring an olefin's carbon atoms from a triangle microchannel to a hexagonal mesochannel as a function of loading corresponding to specific pressure. The statistical uncertainty of ΔG is given in Fig. S4. (C) The radial distribution function (RDF) of distances between the olefin and the framework considering different atomic sites.

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preference for microchannel corners over metal nodes, 22 and for pore edges over the pore center.26 Moreover, olefins may interact with the pyrene-based linkers in NU-1000 via π -based interactions,²⁷ explaining adsorbate distribution near the aromatic rings. There is a probability for C₄H₈ adsorption directly on the metal node within the microchannel; however, a different perspective (Fig. S6A) reveals that olefins are actually concentrated in the windows between the mesochannels. This behavior can be linked to the common placement of catalytic sites in NU-1000 nodes in windows between mesochannels (see Fig. S7).

To quantify the probability of the fractional occupancy of olefins within different channels, the free energy difference for moving of olefin's carbon atoms from a micro to mesochannel was calculated using the equation: $\Delta G = -k_{\rm B}T \ln \frac{\rho_{\rm meso}}{\rho_{\rm micro}}$, where

 $ho_{
m micro}$ and $ho_{
m meso}$ denote the probabilities of an olefin's carbon atoms being in a microchannel or mesochannel. These probabilities were calculated from GCMC snapshots (see Section S3 in SI). In Fig. 2B, values below zero (the dashed line) indicate a preference for adsorption in mesochannel, whereas values above the dashed line indicate a preference for adsorption in microchannel. The results show that single-component C₂H₄ at 1 bar prefers to occupy the microchannel. As pressure increases, enhanced ethylene adsorption saturates the microchannel, shifting olefin uptake to the mesochannel regions. It is consistent with the observation of microchannels saturation reported by Vergas L.24 For C4H8, mesochannel adsorption is already favored at 1 bar, with this preference becoming even stronger as pressure rises. For heavier olefins (C₆H₁₂-C₈H₁₆) in single component systems, the distribution of their fractional occupancy remains consistent across the pressure range as adsorption loading remains constant, indicating a strong preference for microchannel.

In mixed-component systems, the distribution of fractional occupancies of reagents across different channel types is mutually influenced. Compared to single-component systems, atoms of C₄H₈ tend to shift toward the microchannels, thereby displacing other species into the mesochannels. This effect is evident in Fig. 2B, where the single-component curves generally lie above their corresponding blue curves, except for C₄H₈. This trend is even clearer when contrasted with coadsorption scenarios lacking C₄H₈ (see Fig. S9). Across all pressures, fractional occupancies of olefins exhibit a greater $\Delta G_{\text{micro}->\,\text{meso}}$ compared to those olefins that include C₄H₈. A similar, though less pronounced, behavior is observed for C2H4 at pressures above 30 bar and for C₆H₁₂ at 1 bar. Furthermore, microchannel saturation is observed for all olefins at pressures between 1 and 5 bar.

To further investigate the microscopic differences in olefin interactions near the metal nodes, we analyzed the RDF of the selected atomic distances between adsorbates single system component C₄H₈ and C₆H₁₂ and a MOF using MD simulations that incorporate GCMC loading data at T = 300 K under various pressures. Since the differences of RDF across loadings were negligible, the results at P = 50 bar are demonstrated in Fig. 2C

(see more details in SI). The RDF analysis identified three primary adsorption sites in NU-1000: the carbon and oxygen atoms of the linkers, and the oxygen atom of the metal node (see Fig. 2A and C). For olefin adsorption site, the terminal sp² carbon of the α -olefin double bond was selected for the RDF analysis due to its role in the olefin oligomerization reactions.

From the analysis of the distances between atoms of the adsorbate and the MOF, the most favorable adsorption site for C₄H₈ and C₆H₁₂ was found near carbon atoms of the linker (in gray in Fig. 2C) at a distance of 4.4 Å. It complements the result of the normalized probability distribution of olefin adsorption obtained in GCMC (Fig. 2A and Fig. S6 in SI). For C₄H₈, the RDF reveals a second peak corresponding to the metal node oxygen at 4.1 Å with a peak height of 2.3, indicating a close interaction between the olefin and the node oxygen. In contrast, the peak associated with the linker oxygen atom is observed at 4.6 Å with a lower height of 1.8. However, for C_6H_{12} , the peak, indicating the close contact between adsorbates and the oxygen atom from the linker, dominates over the oxygen of the metal node with a closer distance at 4.3 Å and an RDF peak height of 3.2, while the peak at the node oxygen has a lower height of 2.4. In general, shorter distances between C₆H₁₂ and the oxygen atoms are more populated than with C₄H₈. We attribute the closer peak positions for node oxygen to the large hexagonal channel topology, which features a wide 120° opening that provides ample space for olefins to interact with the metal nodes. Similarly, olefins located in the windows between mesochannels benefit from this wide opening, facilitating their interaction with the metal nodes. Moreover, the higher peaks observed for C₆H₁₂ near the oxygen atoms of the linkers are attributed to a stronger preference of the microchannels for C₆H₁₂ in single-component systems compared to C₄H₈ (see Fig. 2B). Lighter olefins exhibit lower peak intensities across all atomic sites (see Fig. S8 in SI), indicating higher mobility. This enhanced mobility is advantageous, as ethylene plays a critical role in the oligomerization process.

Olefin mobility in the MOF can be estimated by the calculation of self-diffusion coefficients. To this end, we have computed the mean squared displacement (MSD) using 100 ns MD simulations at various loadings according to GCMC data (see SI for details). To assess the effect of MOF confinement on selfdiffusion, we also calculated diffusion coefficients for nonconfined systems (a box containing pure olefin). Due to limitations in the MSD approach, which applies only to molecules with sufficiently fast diffusion, we report diffusion coefficients only for olefins up to C_8H_{16} in single component systems. They are depicted in Fig. 3. Due to spatial restrictions in NU-1000, olefin diffusion coefficients drop by two orders of magnitude compared to non-confined conditions for C2H4-C4H8 and one order for C₆H₁₂-C₈H₁₆. Unlike in non-confined systems, diffusion in NU-1000 decreases more gradually with chain length, highlighting their distinct confinement effects.

Due to NU-1000's anisotropic structure with onedimensional channels aligned along the crystallographic axes anisotropic diffusion is expected (Fig. 3B). Our findings confirm this expectation: the diffusion coefficient along the Communication ChemComm

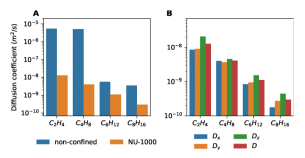


Fig. 3 (A) Diffusion coefficients of olefins in non-confined and confined environments at 1 bar and 300 K. (B) Anisotropic diffusion of olefins in D_{x} , D_{v} , D_{z} directions and overall diffusion D.

channel axis, D_z , exceeds those in the perpendicular directions, D_x and D_y . For instance, C_2H_4 exhibits diffusion along D_z that is 2.4 times higher than in the perpendicular directions, while for C_4H_8 it is 1.2, for C_6H_{12} it is 1.7, and for C_8H_{16} it is 2. These results, obtained within loading boundaries, are in good agreement with Vargas et al.24

By explicitly accounting for multicomponent interactions, our findings demonstrate that the confinement effect of NU-1000 leads to a selective segregation of α-olefins, driven by their complex co-adsorption behavior, where lighter olefins displace heavier ones. Moreover, the unique topology of NU-1000 induces an internal separation: heavier olefins preferentially occupy the microchannels, while lighter ones are more abundant in the mesochannels. Our investigation suggests that positioning the catalytic sites on the metal nodes oriented toward the mesochannels is advantageous, as these sites benefit from the high accessibility afforded by the mesochannel topology. The confinement effect results in a significant decrease in diffusion coefficients and induces anisotropic diffusion compared to non-confined environments, thereby facilitating product separation. These insights are valuable for developing kinetic models that incorporate mass transfer limitations and product-specific concentration profiles.

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Conflicts of interest

There are no conflicts to declare.

Data availability

Data generated is available within the NOMAD Repository. https://doi.org/10.17172/NOMAD/2025.07.08-1.

Computational details, free energy changes in fourcomponent systems, and statistical measures of uncertainty and variability is available. See DOI: https://doi.org/10.1039/ d5cc01329a

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