Splitting Mono- and Di-Branched Alkane Isomers by a Robust Aluminum-Based Metal-Organic Framework Material with Optimal Pore Dimensions

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Supporting Information Placeholder

ABSTRACT: The separation of alkanes with different degrees of branching, particularly mono- and di-branched isomers, represents a challenging yet important industrial process for the production of premium gasoline blending components with high octane number. We report here the separation of linear/mono-branched and di-branched alkanes through complete molecular sieving by a robust aluminum-based MOF material, Al-bttotb (H₂bttotb= 4,4',4'''-(benzene-1,3,5-triyltris(oxy))tribenzoicacid). Single- and multi-component adsorption experiments reveal that the material adsorbs linear and mono-branched alkanes, but fully excludes their di-branched isomers. Adsorption sites of alkanes within the MOF channels have been identified by single-crystal X-ray diffraction studies, and adsorption mechanism explored through computational calculations and modeling. The highly selective adsorption of the MOF should be attributed to its optimal pore dimensions.

Naphtha streams represent the distillate collected in the range of 35 - 200 °C during oil refinement in petrochemical industry. Further distillation of Naphtha produces light naphtha streams consisting mostly C5 and C6 alkanes, which are subject to catalytic isomerization reactions generating isomers with different degrees of branching.1 The resultant C5 and C6 alkane isomers can be used as valuable gasoline blending components, however, they must undergo further separation so as to guarantee sufficiently high octane rating.2,3 Typically, more branched alkanes possess higher research octane number (RON) than their less branched isomers. For example, the RONs for n-hexane (nHEX), 2-methylpentane (2MP), 3-methylpentane (3MP), 2,2-dimethylbutane (22DMB), and 2,3-dimethylbutane (23DMB) are 24.8, 73.4, 74.5, 91.8, and 101.7, respectively. Thus, linear alkanes must be removed, or in a more ideal scenario, linear and monobranced alkanes are separated from their dibranched isomers which have the highest octane number and are premium gasoline blending components.

The separation of alkane isomers represents a challenging process because of their similar chemical and physical properties. Conventional heat-driven distillations could accomplish the separation but with huge capital investment for the equipment and high energy consumption. Alternatively, adsorptive separation technology using porous solids have been proven to be cost efficient and environmentally beneficial, as it would reduce the energy input and suppress carbon dioxide emission. Having the ability to completely separate linear alkanes from their branched isomers, zeolite 5Å remains the benchmark material for this separation process.2,4 However, since zeolite 5Å does not adsorb any branched alkanes, it cannot be used to achieve further improved octane number by differentiating mono- and di-branched isomers, which limits its wider application. To this end, seeking new types of adsorbents with enhanced separation performance is imperative.

Metal-organic frameworks (MOFs) are particularly desirable as adsorbents for industrially challenging separations because of their diverse structures and highly tunable pore systems.5 MOFs have demonstrated exceptional performance for the separation of hydrocarbons, such as paraffins/olefins and alkane isomers.6-14 Several MOF materials, including Y-fum,6 Fe₃(BDP)₃,2 Zr-bptc, Zr-abtc,7 Ca-tcpb,8 have shown similar or even better performance for the separation of alkane isomers compared to zeolite 5Å, with respect to adsorption capacity or selectivity. However, to date, none of the reported materials have reached optimum performance. For example, Fe₃(BDP)₃ and Zr-abtc are capable of discriminating mono- and di-branched alkane isomers thermodynamically, but with relatively low selectivity. Ca-tcpb has the ability to separate
linear, mono-branched, and di-branched alkanes through temperature programming by making use of its structure flexibility. While an ideal size-exclusion based separation was achieved, it was limited to specific temperature and pressure which may not represent the ideal conditions under the current industrial setting. In this work, we show that Al-bttotb, an ultramicroporous metal-organic framework with one dimensional (1D) channels, adsorbs linear and mono-branched alkanes only and excludes their di-branched isomers. Adsorption sites of linear and mono-branched alkanes have been crystallographically identified and DFT calculations reveal that the selective adsorption can be attributed to the optimal channel dimension of the MOF.

Al-bttotb was prepared solvothermally with Al(NO$_3$)$_3$·9H$_2$O, H$_3$btotb, formic acid in DMF at 150 °C. Single-crystal X-ray diffraction reveals the compound crystallizes in the orthorhombic crystal system with a space group of Ccca. In its crystal structure, each Al(III) center is octahedrally coordinated to O atoms from four carboxylates and two bridging hydroxyl anions, forming 1D array of corner-sharing AlO$_6$ polyhedra (Figure 1). The 1D chains are further interconnected through bttotb$^3$ forming 3D network with two types of 3D channels with a square-shaped cross section. Thermogravimetric analysis (TGA) of the as-synthesized material displays a continuous weight loss of 15 wt% up to 180 °C corresponding to the solvent molecules in the pore (Figure S1). Methanol exchange allows full activation of the material under milder conditions (Figure S1-S2). Nitrogen adsorption at 77 K of Al-bttotb reveals a typical Type I profile, giving a BET surface area of 572 m$^2$/g and a pore volume of 0.22 cm$^3$/g (Figure S3). Pore size distribution shows that the material has a uniform pore size of 5.6 Å (Figure S4). It is well recognized that aluminum-based MOFs are generally robust, attributed to the strong Al-O bonds. Our experiments confirm that the structure of Al-bttotb is well-retained after being heated at 180 °C in open air or being exposed to 90% humidity or boiling water for 1 week (Figure S5).

The measured pore size of Al-bttotb falls between the kinetic diameters of mono- and di-branched C$_6$ alkanes (3MP: 5.5 Å, 22DMB: 6.2 Å). This encourages us to explore its adsorption properties toward these alkane isomers. Single-component vapor adsorption analysis reveals that the material adsorbs linear and mono-branched alkanes (nHEX and 3MP) with uptakes of 151 and 94 mg/g, respectively, at 30 °C (Figure 2a). However, it shows essentially no adsorption toward their di-branched isomer (22DMB). To the best of our knowledge, this represents the first porous material with rigid framework that shows complete separation between mono- and di-branched alkanes, which cannot be accomplished by the benchmark material zeolite 5A. The adsorbed alkanes could be desorbed completely at 120 °C and no loss of adsorption capacity was observed after six adsorption-desorption cycles. (Figure S11-S12). To confirm it is a case of selective size exclusion rather than kinetic separation due to insufficient equilibration, we looked into the adsorption kinetics. As shown in Figure 2b, nHEX and 3MP reaches equilibrium in 20 s and 2 mins, respectively, while 22DMB shows no changes in uptake as equilibrium time increases, suggesting the molecule is fully excluded by the MOF.

Figure 1. Crystal structure of Al-bttotb. Left: 1D array of corner-sharing AlO$_6$ polyhedra. Right: 3D structure of Al-bttotb showing two types of channels.

Figure 2. (a) Adsorption isotherms and (b) adsorption kinetics of nHEX, 3MP, and 22DMB at 30 °C.

To evaluate its capability of separating alkane isomers, we performed multicomponent column breakthrough measurements on Al-bttotb with a ternary mixture of equimolar nHEX, 3MP, and 22DMB. The results show that the MOF can well separate the mixture into individual components (Figure 3). 22DMB breaks at the beginning without any retention, confirming the fact that it is not adsorbed by Al-bttotb. In contrast, 3MP and nHEX were retained in the column for 23 and 100 minutes, respectively, corresponding to dynamic adsorbed amounts of 28 and 129 mg/g. The dynamic adsorption capacity
is notably higher than that of zeolite 5A under identical conditions. Real-time RON detection indicates that the initial eluent has a RON of more than 90, well meeting the requirement for gasoline blending components. The notably longer retention of nHEX compared to 3MP could be a result of combined effect of thermodynamics and kinetics, agreeing with the results from single-component adsorption isotherms. This is also consistent with previously reported studies that linear alkanes have better contact with 1D channels compared to their branched isomers.

Figure 3. Multicomponent breakthrough curves of a ternary mixture of nHEX, 3MP, and 22DMB. Green curve depicts the RON of the eluent.

We further expanded the adsorbate mixture from three-component (nHEX, 3MP, 22DMB) to five-component (nHEX, 2MP, 3MP, 23DMB, and 22DMB) to evaluate if the separation ability of the material will be retained under a more complicated scenario. Single-component adsorption measurement indicates that the adsorbed amount and adsorption rate of 2MP fall between nHEX and 3MP, and that material shows moderate adsorption toward 23DMB with relative low uptake of 42 mg/g and slow adsorption kinetics (Figure S6-S7). This is not surprising as the kinetic diameter of 23DMB (5.8 Å) is comparable to the effective pore size of the MOF. Considering the inherent structure flexibility of MOF materials, it is reasonable that it can accommodate 23DMB molecules, but with certain diffusion restrictions. Five-component column breakthrough experiments reveal that the material can well separate the di-branched alkanes from other isomers (Figure S8). This is crucial as both di-branched isomers (23DMB and 22DMB) have high octane numbers. In addition, the structure of material was well-maintained after adsorption/separation measurements (Figure S9 and S10).

To further explore the adsorption behavior of Al-bttotb toward alkanes, we studied the shape and dimension of its channels through adsorption simulations. Helium simulation at 1 K and 1 bar was performed to sketch the profile of the channels. The packing of helium atoms reveals that both channels (A and B) are segmented (Figure S3). Adsorption simulation of nHEX and 3MP at 30 °C and 1 bar indicates that the adsorbed alkanes preferentially reside in channel B, leaving channel A unoccupied (Figure 4c and 4e). This could be attributed to the fact that the smaller chambers in channel B give rise to stronger interaction with nHEX or 3MP molecules.

To experimentally verify the aforementioned adsorption phenomenon, we attempted to resolve the crystal structure of adsorbate loaded MOF through single-crystal X-ray diffraction study. This is the most straightforward technique to study the adsorption sites, adsorbate-adsorbent interaction, and adsorption mechanisms. Fortunately Al-bttotb is robust and maintains its single-crystallinity upon guest removal and inclusion. By exposing activated MOF sample to nHEX, 3MP, and 22DMB and subsequent single-crystal X-ray diffraction analysis, we were able to determine the crystal structure of nHEX and 3MP loaded MOF. In contrast, no sign of 22DMB was detected in the MOF channels, confirming that it is fully excluded by the MOF. Looking into the structure of Al-bttotb with nHEX or 3MP adsorbed, we indeed observe that the alkane molecules are packed in channel B, consistent with the simulation results (Figure 4d and 4f). The adsorbed amount and packing of molecules for nHEX or 3MP agree well with the simulated structures. nHEX (and 3MP) are adsorbed in channel B in pairs, with two molecules residing laterally within each segment. Close contacts of nHEX molecules with the bridging hydroxyl from the MOF has been found, with a H-H distance of 2.19 Å. While the closest H-H distance from 3MP to the MOF is 2.27 Å (Figure S4).

To understand the underlying mechanisms for the experimentally observed preferential adsorption in channel B and its accommodation of nHEX and 3MP but not 22DMB, we perform a computational study at the density functional theory (DFT) level. To simulate the experimental situation, we model two guest molecules (nHEX, 3MP, 22DMB) in both channel A and B of the Al-MOF and explore the effect on the binding energy. Results for the binding energy per molecule of nHEX, 3MP and 22DMB are presented in Table S4. Indeed, we observe a preferred channel occupation for both nHEX and 3MP in channel B over channel A, which is in good agreement with our experimental observations. Due to the smaller chamber size of channel B, there is not only an interaction of the guest molecules with the ligands, but the guest molecules are also forced into a significant lateral interaction in-between them due to the tighter confinement. Our results for the binding energy of the 22DMB show that even this molecule theoretically binds in channel A and B with significant binding energy. However, one significant difference for 22DMB as compared to nHEX and 3MP has been observed: We do find that in the case of nHEX and 3MP, both molecules also bind to the necks with slightly reduced binding energy (as compared to the values given in Table S4, which show binding in the chambers). However, for 22DMB we find positive binding energies in the necks, i.e. a repulsion of the molecule from the necks. This results in a kinetic barrier for 22DMB—which is not present for nHEX or 3MP—that effectively excludes 22DMB from adsorbing in the MOF, explaining our experimental observation.
Adsorptive separation of similar molecules is challenging as it requires precise control of the pore structure and/or functionality of the adsorbent. The diversity of MOF pore structures offers unprecedented opportunity for highly efficient separation of similar molecules, such as alkane isomers. In this work, we show that the separation of monobranched and dibranched alkanes can be accomplished through selective molecular sieving by a robust microporous MOF. The underlying mechanism has been explored through single-crystal X-ray diffraction studies and computational calculation and modeling. The selective adsorption behavior of the material toward alkane isomers should be attributed to its optimal channel dimensions.

ASSOCIATED CONTENT
Supporting Information
Experimental details, synthesis of the MOF, PXRD data, TGA data, and adsorption data. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
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