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## PAPER

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# Selective complexation and efficient separation of cis/trans-1,2-dichloroethene isomers by a pillar[5] arene†

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The complexation and separation of industrially important cis- and trans-1,2-dichloroethene (cis- and trans-DCE) isomers using perethylated pillar[5]arene (EtP5) are described. EtP5 exhibits considerable binding capability for the trans-DCE isomer over the cis-DCE in organic solution. Furthermore, nonporous adaptive crystals (NACs) of EtP5 can efficiently separate trans-DCE from a 50 : 50 (v/v) cis/ trans-isomer mixture.

1,2-Dichloroethenes (DCEs), including cis-DCE and trans-DCE isomers, are high-value chemicals with a variety of uses in synthetic chemistry and chemical industry.<sup>1</sup> They are widely applied as low-temperature extraction solvents for heat sensitive substances and feedstocks for synthesizing copolymer materials with other unsaturated monomers.<sup>2</sup> trans-DCE and cis-DCE can be also used as an environment-friendly refrigerant and foaming additive agent, respectively. In the industrial production process, they are produced as a mixture of cis and trans-DCE isomers by direct chlorination of acetylene or by the pyrolytic dehydrochlorination of 1,1,2-trichloroethane.<sup>3</sup> In most cases, the two isomers must be used separately. However, the separation of cis- and trans-DCE isomers is difficult due to their similar molecular sizes and close boiling points. Industrially, although both isomers can be separated by fractional distillation with very high columns, this process is high energyconsumption and environmentally unfriendly. **PAPER**<br> **CALCRISE SECTIVE COMPLEXATION and efficient separation**<br>
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Crystalline porous materials such as zeolites,<sup>4,5</sup> metalorganic frameworks (MOFs)<sup>6-9</sup> and covalent organic frameworks (COFs)<sup>10</sup> have been described as promising adsorbents for the economical and energy-efficient separation of hydrocarbons (e.g., alkanes, alkenes and benzene derivatives). However, the examples of the physisorptive separation of cis- and transisomers are relatively scarce.<sup>11-14</sup> Recently, a novel class of macrocycle-based crystalline materials, termed as nonporous adaptive crystals (NACs) of pillar[n]arenes, has shown

interesting adsorption/separation properties.<sup>15-21</sup> Subsequently, some new NACs based on other important macrocycles<sup>22</sup> such as biphen[n]arene,<sup>23</sup> leaning towerarene,<sup>24,25</sup> tiararene,<sup>26</sup> geminiarene,<sup>27</sup> hybrid[3]arene,<sup>28</sup> naphthotubes<sup>29</sup> and cucurbit[6] uril<sup>30</sup> have been developed. In comparison with traditional porous materials, which usually possess large surface area, NACs are nonporous and structurally adaptive in the initial crystalline state. The intrinsic or extrinsic porosity inside macrocycle crystals could be opened by capturing preferable vaporized molecules, forming corresponding host–guest crystal structures along with a solid-state structural transformation. This unique feature enables NACs to work as adsorptive materials in adsorption and separation at the solid-vapor phase.

Very recently, our group used NACs of  $2,2',4,4'$ -biphen $[3]$ arene (MeBP3a) to separate cis-DCE from trans-DCE with a purity of 96.4% in the solid-vapor state.<sup>23</sup> Crystal structure of cis-DCE@MeBP3 reveals that cis-DCE molecules are not encapsulated into macrocycle cavities, but located in self-assembled extrinsic porosity. Here, we present that perethylated pillar[5] arenes (EtP5) display a totally opposite selectivity to preferentially bind trans-DCE over its cis-isomer not only in solution but also in the solid state. The selectivity comes from the suitability of size/shape between EtP5's cavity and trans-DCE. NACs of EtP5 (EtP5 $\alpha$ ) efficiently separate trans-DCE from the 50:50 (v/v) mixture of cis/trans-DCE isomers with 94.94% purity (Fig. 1), accompanied by adsorption induced crystalline structure transformation. More intriguingly, when EtP5a and MeBP3a are exposed to cis- and trans-DCE mixed vapor, they exhibit selfsorting adsorption behavior for the two isomers, i.e. producing trans-DCE@EtP5 and cis-DCE@MeBP3.

As we know, pillar $[n]$ arenes, as important family of supramolecular macrocycles with prism-like geometries and  $\pi$ -rich cavities, have shown interesting cavity host–guest properties towards a variety of guest molecules.<sup>31-41</sup> The most peculiar complexation behavior of pillar $[n]$ arenes is that pillar $[5]$ arenes

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Fig. 1 Chemical structures of (a) EtP5 and (b) cis-DCE and trans-DCE. (c) Schematic representation of the cis/trans-DCE isomers separation and the structural transformation.

can strongly bind suitable neutral molecules in organic solution compared with other popular macrocycles.<sup>42-46</sup>

First, we tested the host–guest complexation between cis- or *trans-*DCE and EtP5 in solution by  $^{1}$ H NMR spectroscopy (Fig. S1 and S2, ESI†). As shown in Fig. 2, when equimolar EtP5 was added into an  $o$ -xylene- $d_{10}$  solution containing cis- or trans-DCE (5.0 mM), the signal related to the protons on trans-DCE exhibits a very significant upfield shift of  $-0.52$  ppm as well as an extensive broadening effect compared with the free trans-DCE. The reason is that trans-DCE molecule was encapsulated in the cavity of EtP5 forming a threaded structure. Simultaneously, the aromatic protons on EtP5 moved downfield with slight chemical shift as a result of the interactions with trans-DCE. However, no obvious signal changes were observed when cis-DCE and EtP5 were mixed in o-xylene- $d_{10}$ , suggesting that cisDCE can not be contained in the cavity of EtP5. trans-DCE molecule with stretched structure is more suitable for the cavity of EtP5.<sup>45</sup> It was observed that the inclusion-induced upfield shift of *trans*-DCE ( $\Delta \delta$  = -0.25 ppm for H<sub>a</sub>) in CDCl<sub>3</sub> was smaller than those observed in  $o$ -xylene- $d_{10}$  (Fig. S3 and S4, ESI†), suggesting that stronger host–guest interactions occurred in the non-polar  $o$ -xylene- $d_{10}$  solution. For comparison purpose, the interaction between perethylated pillar[6]arene (EtP6) and cis- or trans-DCE was also investigated. No obvious complexation was detected (Fig. S5 and S6, ESI†), which is reasonable that the guests are too small in comparison with EtP6's cavity. To determine the binding affinity of EtP5 to trans-DCE, <sup>1</sup>H NMR titration methods were employed with the concentration of EtP5 kept constant at 0.50 mM and that of trans-DCE varied from 0 to 33.8 mM in  $o$ -xylene- $d_{10}$  (Fig. S7, ESI†). The association constant  $(K_{\rm a})$  was determined to be  $(1.03\pm0.12)\times 10^2$  M $^{-1}$ by a nonlinear curve-fitting method, which is larger than that in CDCl<sub>3</sub> (31.5  $\pm$  4.1 M<sup>-1</sup>, Fig. S8, ESI†).

X-ray crystallography further confirmed the formation of inclusion complex. Crystals of trans-DCE@EtP5 were successfully obtained by slow evaporation of a solution of EtP5 in trans-DCE (Table S1, ESI†). The crystal structure of trans-DCE@EtP5 is shown in Fig. 3a and S9.† One trans-DCE molecule threads through the cavity of EtP5 to form an inclusion complex in the solid state, which is stabilized by triple C-H $\cdots$  $\pi$  forces and triple weak C-H $\cdots$ Cl interactions (Fig. S10, ESI†). In addition, in the stacking mode of trans-DCE@EtP5, we find extra trans-DCE molecules lie in the channel formed by two adjacent EtP5 molecules (Fig. S9, ESI†). Although no host–guest interactions were found between EtP5 and cis-DCE in solution, we attempted to grow the crystals of cis-DCE@EtP5. Interestingly, when EtP5 was crystallized from cis-DCE solution, the obtained crystal structure of EtP5 did not contain cis-DCE molecules and formed a new structure (Fig. 3b and Table S2†). The experimental and simulated powder X-ray diffraction (PXRD) pattern of EtP5



Fig. 2 Partial proton  ${}^{1}H$  NMR spectra (500 MHz) of (a) EtP5, (b) EtP5 + trans-DCE, (c) trans-DCE, (d) EtP5 + cis-DCE, and (e) cis-DCE at 5.0 mM in  $o$ -xylene- $d_{10}$ . "\*" indicates the solvent peaks.



Fig. 3 Molecular structures of (a) trans-DCE@EtP5 and (b) cis-DCE– free EtP5 obtained from cis-DCE solution in the solid-state.

crystals crystallized in cis-DCE was a perfect match for EtP5a (Fig. S11, ESI†), suggesting that EtP5a has the same structure as EtP5 crystallized in cis-DCE.

Based on the above highly selective host–guest complexation both in solution and in the solid state, we considered the selective adsorption of *cis*- and *trans*-DCE isomers vapor by EtP5a. Single-component time-dependent solid-vapor sorption experiments were examined to test the adsorption capacity using crystalline EtP5 $\alpha$  by <sup>1</sup>H NMR spectroscopy. As shown in Fig. S12,† EtP5a can rapidly capture trans-DCE vapor with an adsorption amount of about one trans-DCE/EtP5 (Fig. S14, ESI†). By contrast, EtP5a did not take up cis-DCE vapor (Fig. S13 and S15, ESI†). Thermogravimetric analysis (TGA) measurement was further confirmed the uptake amount of trans-DCE (Fig. S16, ESI†). However, the single-crystal structure of trans-DCE@EtP5 shows two host molecules and four trans-DCE, which is different from the results of vapor adsorption. This could be reasonable that during the removal of physical surface adsorption before NMR and TGA measurements (for details see ESI†), the guest molecules located at the outside of EtP5 were easily desorbed, while those stabilized in the cavities remained.<sup>47</sup> RSC Advances **Consulting the consulting the consulting the consultant and the particle in the consulting the signifi** 

In order to investigate the mechanism for the uptake of trans-DCE vapor by EtP5a, PXRD experiments were carried out. The PXRD pattern of EtP5 $\alpha$  after adsorption of trans-DCE was different from that of the EtP5a but almost identical to that of the simulated pattern determined from trans-DCE@EtP5 (Fig. 4a), suggesting that the uptake of trans-DCE induces the structural transformation of EtP5a into trans-DCE@EtP5. As expected, exposing EtP5a to cis-DCE vapor did not result in any structure changes (Fig. 4a).



Fig. 4 (a) PXRD patterns of EtP5: (I) original EtP5a; (II) after uptake of cis-DCE vapor; (III) after uptake of cis/trans-DCE mixed vapor; (IV) after uptake of trans-DCE vapor; (V) simulated from single-crystal structure of trans-DCE@EtP5. (b) Time-dependent vapor-solid sorption plot of EtP5a for cis- and trans-DCE (50 : 50 v/v) mixed vapor and the relative uptake ratios determined by GC. (c) Time-dependent selfsorting sorption plot of EtP5a and MeBP3a for cis- and trans-DCE (50 : 50 v/v) mixed vapor. (d) The relative uptake ratios determined by GC.

Considering the remarkable adsorption selectivity in the single-component sorption experiments, we wondered whether crystalline EtP5 $\alpha$  materials would be able to separate trans-DCE from cis-DCE. Therefore, a time-dependent EtP5a solid-vapor sorption experiment for the cis-/trans-DCE (50 : 50 v/v) mixture was performed. As shown in Fig. 4b, the adsorption of trans-DCE in EtP5a was very fast. Meanwhile, the uptake amount of cis-DCE by EtP5a was tiny. According to gas chromatography (GC), the adsorption percentages of trans-DCE is up to 94.94% at the saturated adsorption point (Fig. S17, ESI†). These results indicate that EtP5a favors to capture trans-DCE over cis-DCE vapor from their isomers mixture. Then, we measured the recycling adsorption capacity of EtP5 $\alpha$ . After five adsorption– desorption cycles, no distinct loss of performance in the selective of the trans-DCE uptake was observed (Fig. S18 and S19, ESI†), showing that EtP5a crystals have excellent separation abilities.

Owing to the opposite adsorption selectivity for cis-/trans-DCE isomers of EtP5 $\alpha$  and MeBP3 $\alpha$ <sup>23</sup>, we considered the use of co-adsorbents to achieve the self-sorting discrimination of the two isomers. We then conducted the self-sorting uptake experiments (details see ESI†). As expected, when exposing EtP5 $\alpha$  and MeBP3 $\alpha$  in the saturated vapor of *cis*- and *trans*-DCE (50 : 50 v/v) mixture (Fig. 4c), EtP5a selectively took up trans-DCE, and MeBP3a tended to adsorb cis-DCE (Fig. S20 and S22†), giving the uptake ratios of 95.27% and 93.80%, respectively (Fig. 4d, S21 and S23†).

In summary, we have demonstrated the cis-/trans-selective complexation of industrially important DCE isomers by EtP5 macrocycle both in solution and in the solid state. Its adaptive crystals, EtP5a, are able to efficiently separate trans-DCE from a cis- and trans-DCE isomers mixture. The selectivity derives from the size/shape-fit host–guest complexation and the vaporinduced crystalline structure transformation. Interestingly, a self-sorting adsorption method to simultaneously separate cisand trans-isomers by crystalline EtP5 $\alpha$  and MeBP3 $\alpha$  materials has been presented. Future work will attempt to achieve more separation of configurational isomers using extended biphen $[n]$ arene<sup>48</sup> and functional macrocycles.<sup>49</sup>

#### Conflicts of interest

There are no conflicts to declare.

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#### Notes and references

- 1 K. A. Marshall, Chlorocarbons and Chlorohydrocarbons, Survey, Kirk-Othmer Encyclopedia of Chemical Technology, Wiley-Interscience, New York, 2003.
- 2 F. Richter, Beilsteins Handbuch der organischen Chemie, EIII, 1958, vol. 1, p. 652.
- 3 J. W. Scroggins, US20070191653, 2007.
- 4 M. Yu, R. D. Noble and J. L. Falconer, Acc. Chem. Res., 2011, 44, 1196–1206.
- 5 Y. Yang, P. Bai and X. Guo, Ind. Eng. Chem. Res., 2017, 56, 14725–14753.
- 6 H. Furukawa, K. E. Cordova, M. O'Keeffe and O. M. Yaghi, Science, 2013, 341, 1230444.
- 7 H. Wang and J. Li, Acc. Chem. Res., 2019, 52, 1968–1978.
- 8 E. D. Bloch, W. L. Queen, R. Krishna, J. M. Zadrozny, C. M. Brown and J. R. Long, Science, 2012, 335, 1606–1610.
- 9 Z. R. Herm, E. D. Bloch and J. R. Long, Chem. Mater., 2014, 26, 323–338.
- 10 Y. Zhao, Chem. Mater., 2016, 28, 8079–8081.
- 11 L. Alaerts, M. Maes, M. A. van der Veen, P. A. Jacobs and D. E. De Vos, Phys. Chem. Chem. Phys., 2009, 11, 2903–2911.
- 12 H. Liu, Y. He, J. Jiao, D. Bai, D. Chen, R. Krishna and B. Chen, Chem.–Eur. J., 2016, 22, 14988–14997.
- 13 A. Luna-Triguero, J. M. Vicent-Luna, A. Poursaeidesfahani, T. J. H. Vlugt, R. Sánchez-de-Armas, P. Gómez-Álvarez and S. Calero, ACS Appl. Mater. Interfaces, 2018, 10, 16911–16917.
- 14 M. Maes, L. Alaerts, F. Vermoortele, R. Ameloot, S. Couck, V. Finsy, J. F. M. Denayer and D. E. De Vos, J. Am. Chem. Soc., 2010, 132, 2284–2292.
- 15 K. Jie, Y. Zhou, E. Li and F. Huang, Acc. Chem. Res., 2018, 51, 2064–2072.
- 16 T. Ogoshi, R. Sueto, K. Yoshikoshi, Y. Sakata, S. Akine and T.-a. Yamagishi, Angew. Chem., Int. Ed., 2015, 54, 9849–9852.
- 17 M. Wang, J. Zhou, E. Li, Y. Zhou, Q. Li and F. Huang, J. Am. Chem. Soc., 2019, 141, 17102–17106.
- 18 T. Ogoshi, K. Saito, R. Sueto, R. Kojima, Y. Hamada, S. Akine, A. M. P. Moeljadi, H. Hirao, T. Kakuta and T. Yamagishi, Angew. Chem., Int. Ed., 2018, 57, 1592–1595.
- 19 K. Jie, M. Liu, Y. Zhou, M. A. Little, S. Bonakala, S. Y. Chong, A. Stephenson, L. Chen, F. Huang and A. I. Cooper, J. Am. Chem. Soc., 2017, 139, 2908–2911.
- 20 Y. Zhou, K. Jie, R. Zhao, E. Li and F. Huang, J. Am. Chem. Soc., 2020, 142, 6957–6961.
- 21 X. Sheng, E. Li, Y. Zhou, R. Zhao, W. Zhu and F. Huang, J. Am. Chem. Soc., 2020, 142, 6360–6364.
- 22 J.-R. Wu and Y.-W. Yang, Angew. Chem., Int. Ed., 2020, 59, DOI: 10.1002/anie.202006999.
- 23 Y. Wang, K. Xu, B. Li, L. Cui, J. Li, X. Jia, H. Zhao, J. Fang and C. Li, Angew. Chem., Int. Ed., 2019, 58, 10281–10284.
- 24 J.-R. Wu, B. Li and Y.-W. Yang, Angew. Chem., Int. Ed., 2020, 59, 2251–2255.
- 25 J.-R. Wu and Y.-W. Yang, CCS Chem., 2020, 2, 836–843.
- 26 W. Yang, K. Samanta, X. Wan, T. U. Thikekar, Y. Chao, S. Li, K. Du, J. Xu, Y. Gao, H. Zuilhof and A. C.-H. Sue, Angew. Chem., Int. Ed., 2020, 59, 2–8.
- 27 J.-R. Wu and Y.-W. Yang, J. Am. Chem. Soc., 2019, 141, 12280– 12287.
- 28 J. Zhou, G. Yu, Q. Li, M. Wang and F. Huang, J. Am. Chem. Soc., 2020, 142, 2228–2232.
- 29 H. Yao, Y.-M. Wang, M. Quan, M. U. Farooq, L. P. Yang and W. Jiang, Angew. Chem., Int. Ed., 2020, 59, 19945–19950.
- 30 Q. Li, K. Jie and F. Huang, Angew. Chem., Int. Ed., 2020, 59, 5355–5358.
- 31 T. Ogoshi, S. Kanai, S. Fujinami, T.-a. Yamagishi and Y. Nakamoto, J. Am. Chem. Soc., 2008, 130, 5022–5023.
- 32 D. Cao, Y. Kou, J. Liang, Z. Chen, L. Wang and H. Meier, Angew. Chem., Int. Ed., 2009, 48, 9721–9723.
- 33 T. Ogoshi, T.-a. Yamagishi and Y. Nakamoto, Chem. Rev., 2016, 116, 7937–8002.
- 34 N. Song, T. Kakuta, T.-a. Yamagishi, Y.-W. Yang and T. Ogoshi, Chem, 2018, 4, 2029–2053.
- 35 K. Wang, J. H. Jordan, K. Velmurugan, X. Tian, M. Zuo, X.-Y. Hu and L. Wang, Angew. Chem., Int. Ed., 2020, DOI: 10.1002/anie.202010150.
- 36 G. Yu, J. Zhou, J. Shen, G. Tang and F. Huang, Chem. Sci., 2016, 7, 4073–4078.
- 37 B. Li, Z. Meng, Q. Li, X. Huang, Z. Kang, H. Dong, J. Chen, J. Sun, Y. Dong, J. Li, X. Jia, J. L. Sessler, Q. Meng and C. Li, Chem. Sci., 2017, 8, 4458–4464. Public and Licence Access Article is article in 21 December 2020. Downloaded on 21 December 2020. Downloaded on 21 December 2020. The Creative Commons Article is licensed under a creative Commons Article is licensed under
	- 38 J. Chen, H. Ni, Z. Meng, J. Wang, X. Huang, Y. Dong, C. Sun, Y. Zhang, L. Cui, J. Li, X. Jia, Q. Meng and C. Li, Nat. Commun., 2019, 10, 3546.
	- 39 X.-B. Hu, Z. Chen, G. Tang, J. Hou and Z.-T. Li, J. Am. Chem. Soc., 2012, 134, 8384–8387.
	- 40 Q. Hao, Y. Chen, Z. Huang, J.-F. Xu, Z. Sun and X. Zhang, ACS Appl. Mater. Interfaces, 2018, 10, 5365–5372.
	- 41 S.-H. Li, H.-Y. Zhang, X. Xu and Y. Liu, Nat. Commun., 2015, 6, 7590.
	- 42 X. Shu, S. Chen, J. Li, Z. Chen, L. Weng, X. Jia and C. Li, Chem. Commun., 2012, 48, 2967–2969.
	- 43 C. Li, Chem. Commun., 2014, 50, 12420–12433.
	- 44 Y. Wang, G. Ping and C. Li, Chem. Commun., 2016, 52, 9858– 9872.
	- 45 X. Zhang, X. Wang, B. Wang, Z.-J. Ding and C. Li, Chin. Chem. Lett., 2020, DOI: 10.1016/j.cclet.2020.02.037.
	- 46 B. Li, S. Li, B. Wang, Z. Meng, Y. Wang, Q. Meng and C. Li, iScience, 2020, 23, 101443.
	- 47 W. Zhu, E. Li, J. Zhou, Y. Zhou, X. Sheng and F. Huang, Mater. Chem. Front., 2020, 4, 2325–2329.
	- 48 B. Li, B. Wang, X. Huang, L. Dai, L. Cui, J. Li, X. Jia and C. Li, Angew. Chem., Int. Ed., 2019, 58, 3885–3889.
	- 49 K. Xu, Z.-Y. Zhang, C. Yu, B. Wang, M. Dong, X. Zeng, R. Gou, L. Cui and C. Li, Angew. Chem., Int. Ed., 2020, 59, 7214–7218.