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## CO<sub>2</sub>-enhanced TADF of an ultra-stable Cu(I) cluster via guest–host $\pi$ – $\pi$ interaction†

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Efficient and reversible luminescence detection for CO<sub>2</sub> without solvent assistance is of great significance but remains challenging to achieve, due to the lack of efficient interaction between CO<sub>2</sub> molecules and the host emitting center. Benefiting from the abundant host–guest interactions, metal clusters provide a platform for detecting small molecules. However, the insufficient chemical stability of most metal clusters limits their practical applications. Here, we report a hydrophobic Cu(I) cluster (denoted as CuIDPO) with one-dimensional channels. Notably, it displays exceptional chemical stability in both acidic and alkaline aqueous solutions (pH = 1–14). More importantly, CuIDPO shows remarkable CO<sub>2</sub>-induced luminescence enhancement (up to 385% under 1 bar CO<sub>2</sub>), which can be applied to analyze CO<sub>2</sub> content (LOD = 7.7 mbar). Crystallographic analysis and theoretical calculations suggest the mechanism of CO<sub>2</sub>-locking rotation of the phenyl groups in the Cu(I) cluster through guest–host  $\pi$ – $\pi$  interaction, which is quite unique when compared to the known acid–base neutralization and framework flexibility adjustment mechanisms. Such luminescence CO<sub>2</sub> sensing shows advantages like ultrafast response and good reversibility. Additionally, CuIDPO-loaded membranes were fabricated for spatially resolved 2D visual detection.

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### Introduction

The detection of CO<sub>2</sub> is crucial in various fields like agriculture,<sup>1</sup> biology,<sup>2</sup> carbon emissions<sup>3–5</sup> and so on. Traditional methods primarily rely on electrochemical technology,<sup>6,7</sup> Fourier-transform infrared (FT-IR) spectroscopy,<sup>8</sup> mass spectrometry techniques,<sup>9,10</sup> and luminescence analysis methods. Notably, luminescence detection offers the advantages of non-electrical connection, convenient operation, high sensitivity and two-dimensional visual detection.<sup>11,12</sup> Reported luminescence CO<sub>2</sub> detecting cases usually proceed in solutions, based on acid–base neutralization reactions, as CO<sub>2</sub> causes weak acidity.<sup>13–16</sup> However, such a process usually requires the assistance of solvents and is hard to reverse, which is not favourable for the regeneration of optical probes after absorbing CO<sub>2</sub>. In the meantime, there are also several reports that explore CO<sub>2</sub>

sensing performance in porous solid-state materials, for example, flexible metal–organic frameworks (MOFs), via employing the flexible changes of the framework caused by the adsorption/desorption of CO<sub>2</sub> to achieve luminescence changes. These materials no longer require the assistance of solvents, and the activation of fluorescence probes is further facilitated. However, their sensitivity and device fabrication still need enhancement, because of the lack of proper interaction between the CO<sub>2</sub> molecule and the host emitting center, as well as a simple synthesis method.<sup>17–19</sup> Therefore, the exploration of efficient luminescent probes for CO<sub>2</sub> with new sensing mechanisms is an essential and attractive topic.

Clusters of d<sup>10</sup> metal ions like Cu(I) and Ag(I) possess excellent luminescence stimulus-responsive properties due to the rich transition modes,<sup>20</sup> making them highly promising for sensing temperature,<sup>21</sup> pressure,<sup>22</sup> gas,<sup>23</sup> and solvent molecules.<sup>24</sup> The capping effect of ligands can effectively enhance the water stability of d<sup>10</sup> metal clusters.<sup>25–27</sup> However, due to the relatively weak coordination bonding between the ligands and d<sup>10</sup> metal ions, these clusters are more susceptible to most acids and bases compared with other coordination compounds.<sup>28</sup> Although the increasing hydrophobicity of metal clusters can enhance their resistance to acids and bases, there are still very few related reports.<sup>29–31</sup>

Here, we report a new discrete Cu(I) cluster with one-dimensional pores. This solid compound possesses excellent stability in both acidic and alkaline aqueous solutions (pH = 1–

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14), due to its strong hydrophobicity. Additionally, it displays blue thermally activated delayed fluorescence (TADF), which can be efficiently enhanced by CO<sub>2</sub> (by 385% under 1 bar CO<sub>2</sub>). Such a characteristic is then further used for efficient luminescence CO<sub>2</sub> detection with a fast response and good reversibility, as well as high sensitivity and a low limit of detection (LOD = 7.7 mbar). Single-crystal X-ray diffraction (SCXRD) analysis and theoretical calculations clearly suggest that a guest–host  $\pi$ – $\pi$  interaction is formed between CO<sub>2</sub> and the phenyl groups in the Cu(I) cluster, which restricts the host molecular rotation and reduces non-radiative transitions, thus generating unique CO<sub>2</sub>-enhanced luminescence. This unique mechanism provides a new direction for the design of sensitive optical probes for detecting CO<sub>2</sub>. Finally, this metal cluster is also successfully loaded into membranes to achieve spatially resolved two-dimensional visual detection.

## Results and discussion

The Cu(I) cluster compound, [Cu<sub>2</sub>I<sub>2</sub>(DPO)<sub>2</sub>]·4CH<sub>2</sub>Cl<sub>2</sub> (DPO = bis(2-diphenylphosphinophenyl)ether), denoted as CuIDPO·CH<sub>2</sub>Cl<sub>2</sub>, was obtained by the reaction of the DPO ligand and CuI in CH<sub>2</sub>Cl<sub>2</sub> solvent (Fig. S1†). SCXRD data show that CuIDPO·CH<sub>2</sub>Cl<sub>2</sub> crystallizes in the monoclinic *P*2<sub>1</sub>/*c* space group with the asymmetric unit consisting of one CuI, one DPO ligand and two CH<sub>2</sub>Cl<sub>2</sub> guest molecules (Table S1 and Fig. S2†). The discrete [Cu<sub>2</sub>I<sub>2</sub>] cluster is formed by two tetrahedrally coordinated Cu(I) ions, which are bridged by two I<sup>−</sup> ions and further chelated by DPO ligands. These clusters stack with each other through C–H··· $\pi$  interactions in the lattice (Fig. S3†), while CH<sub>2</sub>Cl<sub>2</sub> molecules occupy hydrophobic channels through van der Waals interactions (Fig. S4†). Remarkably, CH<sub>2</sub>Cl<sub>2</sub> guests can be completely removed from CuIDPO·CH<sub>2</sub>Cl<sub>2</sub> through a single-crystal to single-crystal (SC–SC) transformation under gentle operation like vacuuming at room temperature, yielding a guest-free structure (denoted as CuIDPO, Fig. S5†). After removing the CH<sub>2</sub>Cl<sub>2</sub> molecules, although the cell volume decreases from 3711.92(10) to 3235.0(8) Å<sup>3</sup> (Table S1†), CuIDPO still retains the one-dimensional (1D) wavy pores with 4.7% porosity (Fig. S6†), generating two distinct micropores (cavity A: 3.8 Å; cavity B: 2.7 Å; Fig. 1a). Additionally, no significant change in the configuration of the DPO ligand is found after guest desorption, whereas a decrease in the intramolecular Cu–Cu distance from 3.55 Å (CuIDPO·CH<sub>2</sub>Cl<sub>2</sub>) to 3.31 Å (CuIDPO) is observed (Fig. S7†).

Commonly, d<sup>10</sup> metal clusters are highly susceptible to acids and bases, which usually lead to the collapse of cluster structures.<sup>32</sup> Exceptionally, after immersion in aqueous solutions with a wide pH range (1.03–14.05) for 1 month, CuIDPO can still retain the original crystalline phase (Fig. 1b, S8a and b†), due to its high hydrophobicity with a contact angle of 124.75° (Fig. S8c†). In addition, CuIDPO can retain the original crystalline phase even after long-term exposure (for 8 months) to ambient air conditions (Fig. S9†), further proving its chemical stability. This feature corresponds to facing-out petal-like aromatic rings of DPO that encapsulate the [Cu<sub>2</sub>I<sub>2</sub>] cluster (Fig. S10†). Moreover, as shown in Fig. S11,† the crystallographic planes (100),

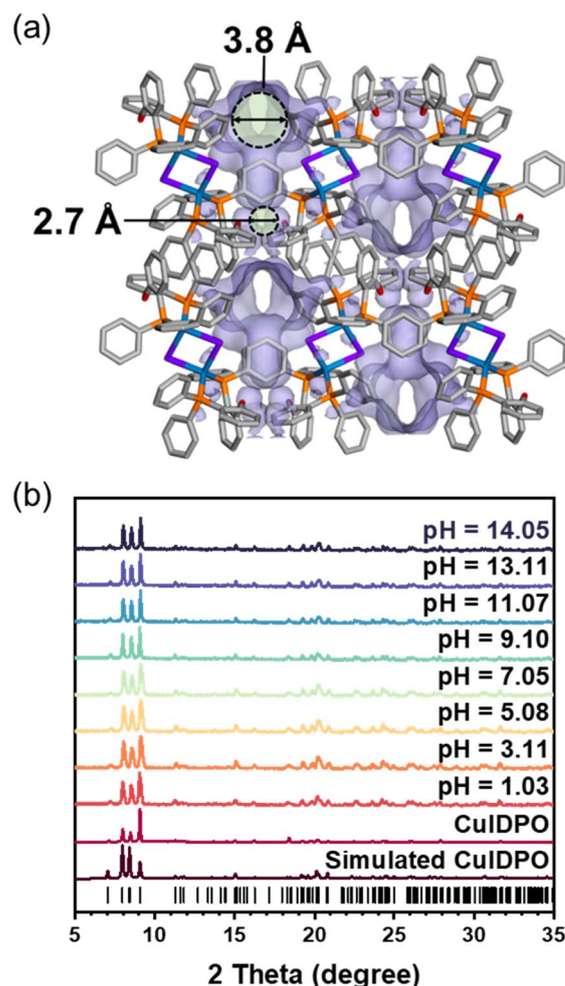


Fig. 1 (a) Crystal structure and void surface of CuIDPO viewed along the *b*-axis. Hydrogen atoms are omitted for clarity. Color codes: Cu, blue; I, purple; P, orange; C, grey; O, red. (b) PXRD patterns of CuIDPO after immersion in aqueous solutions with different pH values.

(010) and (001) are all hydrophobic, so [Cu<sub>2</sub>I<sub>2</sub>] clusters are well shielded and effectively stabilized in the framework when immersed in acidic and basic aqueous solutions. In addition, the relatively hydrophobic pore surface in CuIDPO is also beneficial for high stability (Fig. S12†). Thermogravimetric analysis (TGA) shows that after exposure to humid N<sub>2</sub> (Fig. S13†), the weight change of CuIDPO is minimal (less than 0.25%), which further demonstrates the hydrophobic nature of the CuIDPO channels. Moreover, TGA of CuIDPO shows that the structure does not collapse till 340 °C (Fig. S14†), suggesting its high thermostability.

CuIDPO shows negligible N<sub>2</sub> adsorption at 77 K (0.125 mmol g<sup>−1</sup> at *P*/*P*<sub>0</sub> = 0.90, Fig. S15†), attributed to the extremely narrow channel apertures and quasi-discrete pores. In contrast, CO<sub>2</sub> adsorption follows a type-I isotherm at 195 K (Fig. 2a). The experimental saturated CO<sub>2</sub> uptake is 2.92 mmol g<sup>−1</sup> at *P*/*P*<sub>0</sub> = 0.94, which is slightly lower than the empirically calculated value (3.25 mmol g<sup>−1</sup>) with a Langmuir surface area of 374.95 m<sup>2</sup> g<sup>−1</sup> at 195 K (Fig. 2a and S16†). Additionally, at 273 K and



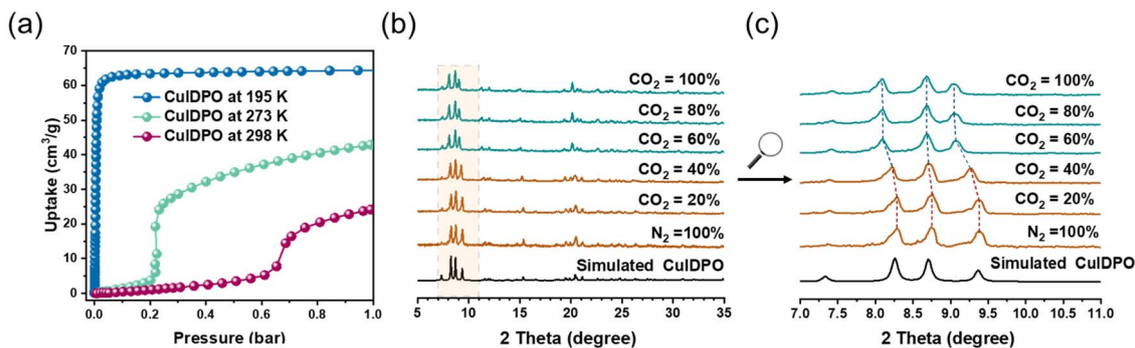


Fig. 2 (a) CO<sub>2</sub> sorption isotherms of CuIDPO at 195, 273 and 298 K. (b) PXRD patterns of CuIDPO in CO<sub>2</sub>/N<sub>2</sub> mixtures of different ratios (v/v). (c) Enlarged patterns of (a) for  $2\theta = 7\text{--}11^\circ$ .

298 K, the CO<sub>2</sub> sorption isotherms exhibit an S-shaped profile, revealing significant breathing or gate-opening behaviour with the gate-opening pressures ( $P_{go}$ ) of 0.2 and 0.6 bar at 273 and 298 K, respectively. Specifically, at 298 K, the saturated CO<sub>2</sub> uptake is 1.06 mmol g<sup>-1</sup>, corresponding to 1.55 CO<sub>2</sub> molecules per unit cell. For 273 K, these values increase to 1.92 mmol g<sup>-1</sup> and 2.80 CO<sub>2</sub> molecules per unit cell, respectively. To further elucidate the gate-opening process of CuIDPO, PXRD patterns in 1 bar gas mixtures with different ratios (v/v) of CO<sub>2</sub> and N<sub>2</sub> were recorded (Fig. 2b). The PXRD patterns remain unchanged after exposure to 0–40% CO<sub>2</sub> and are consistent with the simulated pattern of CuIDPO, illustrating that no gate-opening effect occurs at low CO<sub>2</sub> content. However, upon further increasing the CO<sub>2</sub> content ( $\geq 60\%$ ), a variation can be observed in the PXRD pattern. As shown in Fig. 2c, the peak at 9.4° shifts to 9.0°, indicating that a new phase has emerged.

The photophysical properties of CuIDPO were subsequently investigated. An absorption band at 200–400 nm is found in the UV-vis adsorption spectrum of CuIDPO (Fig. S17†), corresponding to the white appearance of the powder sample under ambient light. Density functional theory (DFT) and time-dependent density functional theory (TDDFT) calculations show that the highest energy absorption ( $S_0 \rightarrow S_2$ ) can be assigned to the mixed metal/halogen to ligand charge transfer (M/XLCT, Fig. S18†). Moreover, as seen in the crystal structure of CuIDPO, the Cu···Cu distance is 3.38 Å (much larger than 3 Å), indicating no significant Cu···Cu interaction (Fig. S7 and Table S3†).<sup>33</sup> Under excitation of 365 nm LED light, the maximum emission wavelength ( $\lambda_{em}$ ) of CuIDPO is located at 432 nm with a lifetime of 0.47  $\mu$ s in air via  $T_1 \rightarrow S_0$  transitions from M/XLCT states (Fig. S19†). Temperature-dependent (80 to 360 K) emission spectra show that, with the increase of temperature, the emission of CuIDPO decreases in intensity and exhibits a blue shift (Fig. S20†). Such thermochromic behaviour resembles those of highly emissive solid-state Cu(I) complexes, which is ascribed to the TADF mechanism.<sup>33,34</sup> In the analysis of the decay curve for luminescence lifetime in a vacuum, by fitting the long-lived component to eqn S1,† the results of  $\Delta E(S_1 - T_1) = 0.1076$  eV ( $< 0.2$  eV), indicating a low energy gap for reverse intersystem crossing (RISC),  $\tau(S_1) = 52$  ns and  $\tau(T_1) = 224$   $\mu$ s can be obtained (Fig. S21†). Such narrow  $\Delta E(S_1 - T_1)$  is also confirmed by TDDFT calculations (Fig. S22†).

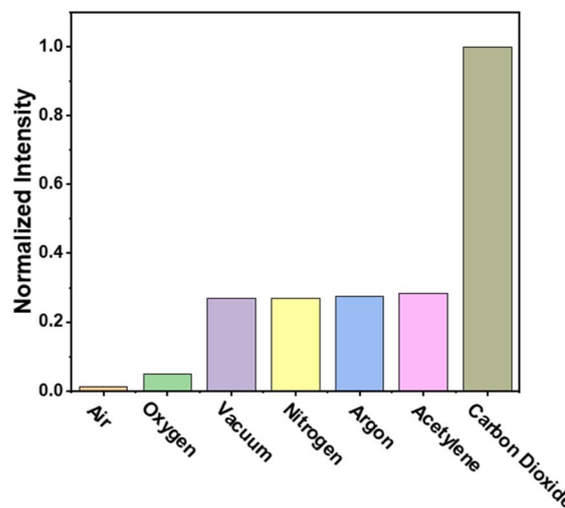


Fig. 3 Emission intensities of CuIDPO in different gases and a vacuum, excited with a 365 nm LED and detected at 432 nm.

Interestingly, under 1 bar CO<sub>2</sub>, an obvious enhancement of luminescence intensity is observed for CuIDPO, which is about 5 times that collected in a vacuum (Fig. 3 and S23†). Additionally, other common gases, such as O<sub>2</sub>, air, N<sub>2</sub> and Ar, cannot enhance the emission of CuIDPO, demonstrating its high selectivity for CO<sub>2</sub> sensing (Fig. 3 and S23†). Even C<sub>2</sub>H<sub>2</sub>, which is a linear molecule with a high quadrupole moment, cannot affect its fluorescence (Fig. 3 and S23†). As this CO<sub>2</sub> light-up emission is quite unique in reported solid complexes, the corresponding study is performed in detail next.

Upon alternating exposure to CO<sub>2</sub> and vacuum, CuIDPO exhibits reversible switching between bright blue emission (“on” state) and near-quenched emission (“off” state) under 365 nm illumination (Fig. 4a). No obvious decay of the intensity appears even after 10 alternating cycles, indicating its high optical stability (Fig. S24†). Moreover, the CO<sub>2</sub> sensing shows very rapid responding/recovering speeds. Specifically, the response time ( $t_{res}$ ) and the recovery time ( $t_{rec}$ ) are defined as the time when the luminescence intensity of the detecting intensity changes more than 90%. The “measured”  $t_{res}$  and  $t_{rec}$  are 1.5 and 1.7 s, by switching the atmosphere between CO<sub>2</sub> and



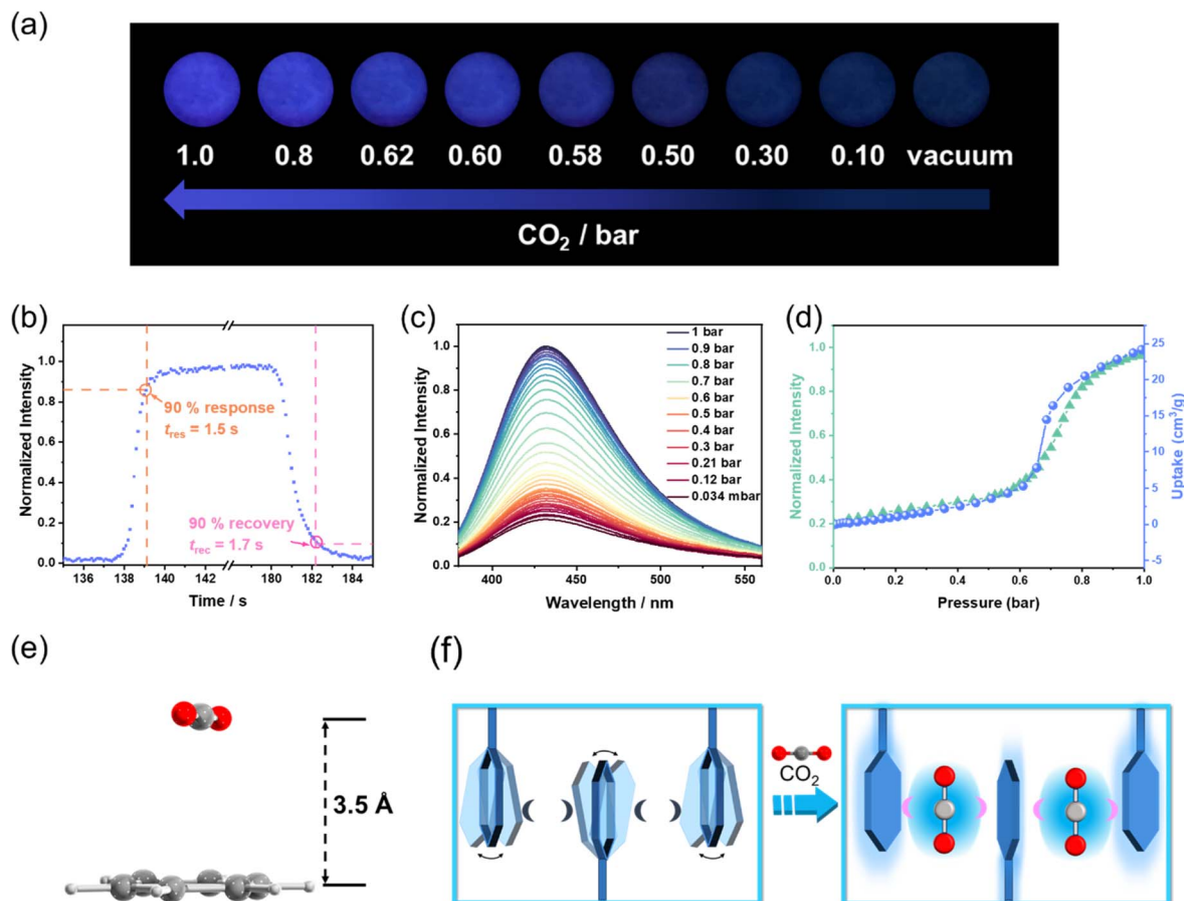


Fig. 4 (a) Photographs of CuIDPO at different CO<sub>2</sub> pressures and room temperature, excited with a 365 nm UV lamp. (b) Enlarged view of one cycle of the kinetic scan of CuIDPO under alternating vacuum and CO<sub>2</sub> conditions, excited at 365 nm and detected at 432 nm. (c) Emission spectra of CuIDPO at different CO<sub>2</sub> pressures, excited at 365 nm. (d) Comparison between the luminescence intensity (excited at 365 nm and detected at 432 nm) and CO<sub>2</sub> uptakes (298 K) of CuIDPO at different CO<sub>2</sub> pressures. (e) The  $\pi$ - $\pi$  interaction between CO<sub>2</sub> and the phenyl ring of CuIDPO·CO<sub>2</sub>. (f) Schematic diagram of the CO<sub>2</sub> sensing mechanism. The pink and dark crescents represent the presence of and the lack of interaction between the phenyl groups and the guest molecules, respectively.

vacuum, respectively (Fig. 4b). Actually, the visual color transition appears instantaneous. When gradually increasing the CO<sub>2</sub> pressure from 0 to 1 bar at room temperature, a monotonic luminescence intensity enhancement can be observed across the entire process (excited at 365 nm, Fig. 4c). In particular, the pressure-intensity curve is almost coincided with the sorption isotherms of CuIDPO for CO<sub>2</sub> at 298 K (Fig. 4d), confirming a direct correlation between CO<sub>2</sub> uptake and emission enhancement. Moreover, the luminescence intensities of CuIDPO exhibit a good linear relationship with the concentration of CO<sub>2</sub> in the range of 0–0.2 bar. Hence, the LOD concentration for CO<sub>2</sub> can be calculated to be 7.7 mbar based on eqn (S2) (Fig. S25).<sup>†</sup> Additionally, CuIDPO can retain the original crystalline (CuIDPO) phase after CO<sub>2</sub> absorption and CO<sub>2</sub> response experiments (Fig. S26<sup>†</sup>), further confirming its high stability. Except for the luminescence intensity enhancement, CO<sub>2</sub> also helps to increase the lifetime of CuIDPO, which can be increased to 13.59  $\mu$ s when under 1 bar CO<sub>2</sub> (Fig. S27<sup>†</sup>). Moreover, even after soaking CuIDPO in aqueous solutions with pH = 1 (HCl solution) and pH = 14 (NaOH solution), respectively,

for as long as one week, CuIDPO still retains its fluorescence sensing ability for CO<sub>2</sub> (Fig. S28 and S29<sup>†</sup>), further demonstrating the chemical stability of CuIDPO.

To further confirm the mechanism of CO<sub>2</sub>-enhanced emission, the interaction between CO<sub>2</sub> and CuIDPO was investigated. SCXRD data of CuIDPO were collected under 1 bar CO<sub>2</sub>. Crystallographic analysis reveals that, at 291 K, CO<sub>2</sub> molecules are absorbed at cavity A, which is highly disordered (Fig. S30<sup>†</sup>). After cooling to 150 K, the CO<sub>2</sub> molecules became ordered (Fig. S31<sup>†</sup>), and this CO<sub>2</sub>-absorbed structure is denoted as CuIDPO·CO<sub>2</sub>. The PXRD pattern of CuIDPO under 1 bar CO<sub>2</sub> is consistent with the simulated PXRD pattern of CuIDPO·CO<sub>2</sub> (Fig. S32<sup>†</sup>). CuIDPO·CO<sub>2</sub> crystallizes in the monoclinic  $P2_1/n$  space group, and the cell volume (3381.77(6) Å<sup>3</sup>) expands, compared with that of CuIDPO (150 K, 3235.0(8) Å<sup>3</sup>, Table S2<sup>†</sup>). Though the Cu···Cu distance extends from 3.31 Å (CuIDPO at 150 K) to 3.51 Å (CuIDPO·CO<sub>2</sub> at 150 K,  $\gg 3$  Å), it was still too long to generate effective Cu···Cu interaction (Fig. S33 and Table S3<sup>†</sup>). Interestingly, CO<sub>2</sub> is very close (3.5 Å) to one of the phenyl rings in the framework, which is close enough to form



a  $\pi$ - $\pi$  interaction (Fig. 4e). This  $\pi$ - $\pi$  interaction locks the rotation of the free phenyl ring, significantly reducing non-radiative transitions and enhancing luminescence (Fig. 4f). DFT calculations show that the binding energy between  $\text{CO}_2$  and this phenyl ring is about  $-13.92 \text{ kJ mol}^{-1}$  (Table S4†), which is close to the common energy level of reported  $\text{CO}_2$ -phenyl  $\pi$ - $\pi$  interaction.<sup>35</sup> Moreover, when CuIDPO absorbs  $\text{CO}_2$ , the  $\lambda_{\text{em}}$  of CuIDPO remains unchanged, proving that the configuration of its luminescent center has remained essentially unchanged (Fig. 4c). This fact excludes the formation of exciplexes and the process of intramolecular charge transfer, which require the influence to the transition of the molecular orbitals in CuIDPO, attributed to the nonpolar nature of  $\text{CO}_2$ . Therefore, the luminescence enhancement solely originates from the  $\text{CO}_2$ -caused restriction of the molecular rotation of phenyl groups in CuIDPO (Fig. 4f), similar to the restriction of intramolecular

motion (RIM) effect in the aggregation-induced emission luminogens (AIEgens). Similar to  $\text{CO}_2$ , the incorporation of  $\text{CH}_3\text{CN}$  into the framework induces a comparable luminescence enhancement, suggesting a similar host-guest interaction mechanism (Fig. S34 and S35†). The crystal structure of the  $\text{CH}_3\text{CN}$ -absorbed compound ( $\text{CuIDPO}\cdot\text{CH}_3\text{CN}$ ) shows that C-H $\cdots\pi$  interactions between  $\text{CH}_3\text{CN}$  and the metal cluster are formed with a binding energy of  $-16.02 \text{ kJ mol}^{-1}$  (Fig. S36 and Table S4†), further illustrating the importance of RIM.

To further increase the portability and visualizability of CuIDPO in optical  $\text{CO}_2$  sensing, membrane sensors based on glass fiber (GF) paper were fabricated. First, a  $\text{CH}_3\text{CN}$  solution of CuI was sprayed onto GF paper, allowing the GF paper to fully absorb CuI. After drying, it was soaked in a  $\text{CH}_2\text{Cl}_2$  solution of DPO to conduct a reaction between CuI and DPO within the GF paper, resulting in the formation of CuIDPO@GF (Fig. 5a and

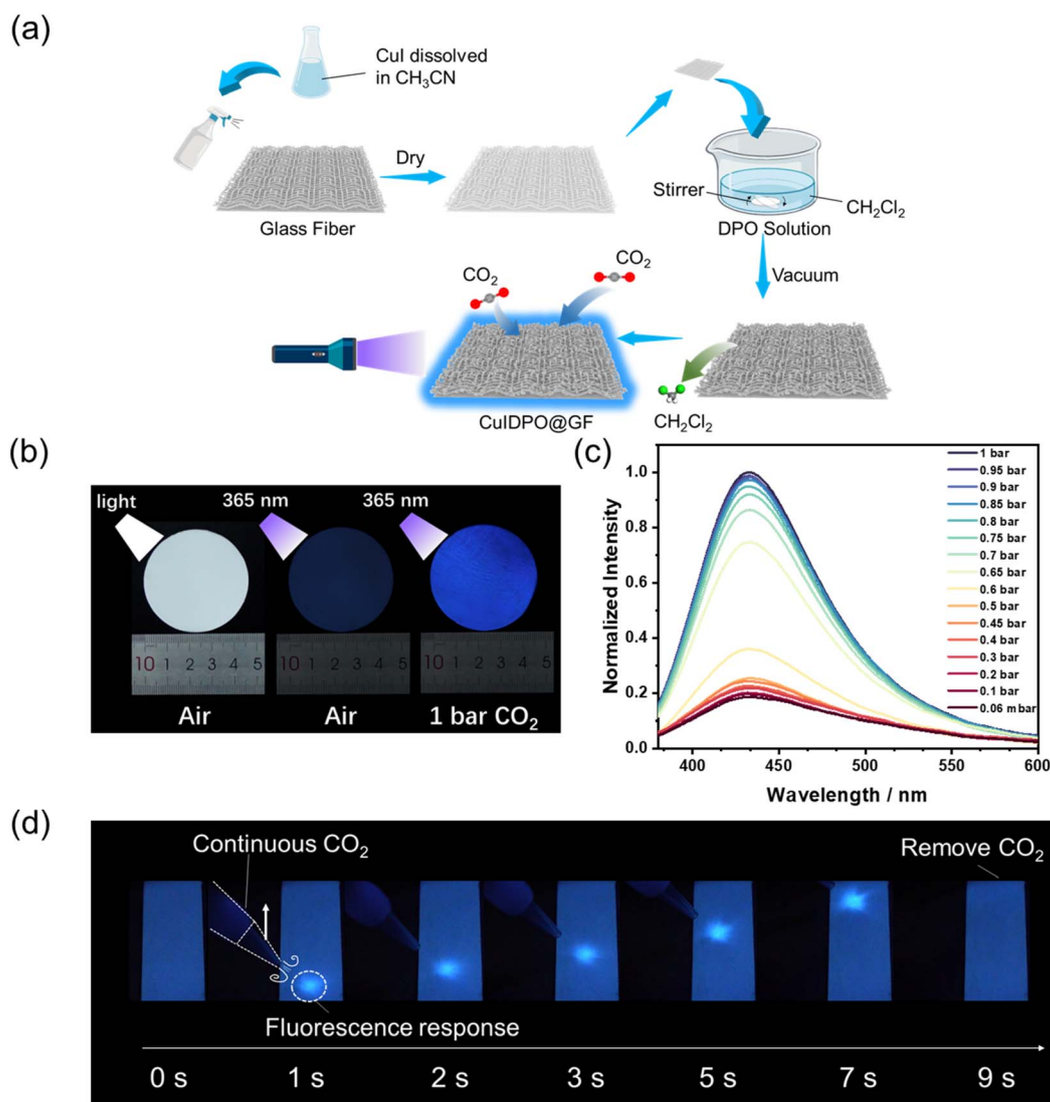


Fig. 5 (a) Schematic illustration of *in situ* synthesis of CuIDPO@GF. Color codes for  $\text{CO}_2$  and  $\text{CH}_2\text{Cl}_2$ : C, grey; O, red; Cl, green; H, white. (b) Photographs of CuIDPO@GF, excited by daylight and 365 nm UV light, respectively, in air and under 1 bar  $\text{CO}_2$ . (c) Emission spectra of CuIDPO@GF at different  $\text{CO}_2$  pressures, excited with a 365 nm UV LED. (d) Luminescence photographs of CuIDPO@GF under  $\text{CO}_2$  purged using a glass needle, excited with a 365 nm UV LED.



b). The PXRD pattern shows that CuIDPO on the GF paper maintains its crystalline phase (Fig. S37<sup>†</sup>). The scanning electron microscope (SEM) photographs show that the CuIDPO particles adhere to the GF paper well (Fig. S38<sup>†</sup>). CuIDPO@GF exhibit not only similar photophysical properties to CuIDPO, but also high sensitivity and fast response to CO<sub>2</sub> (Fig. 5c, d and S39<sup>†</sup>). This membrane achieves two-dimensional visual detection of CO<sub>2</sub>. As shown in Fig. 5d and Video S1,<sup>†</sup> when exposed to a CO<sub>2</sub> flow, a bright spot is immediately observed in the corresponding site of CuIDPO@GF. However, the spot turns dark as soon as the CO<sub>2</sub> flow is removed, indicating a great possibility for displaying the pressure of CO<sub>2</sub> in two dimensions. To test the responsiveness of CuIDPO@GF in high humidity (82% RH) environments, we bubbled CO<sub>2</sub> through water before it came into contact with CuIDPO@GF (Fig. S40<sup>†</sup>). After 10 cycles, CuIDPO@GF retains its responsiveness without any decay (Fig. S41<sup>†</sup>). Delightfully, CuIDPO@GF retains a rapid response time even under high humidity (82% RH) conditions (Fig. S42<sup>†</sup>). These facts indicate that CuIDPO@GF has great potential to operate in more complex environments.

## Conclusions

A Cu(I) cluster with blue TADF was synthesized, which exhibits excellent stability in acidic and basic environments. Its emission can be selectively enhanced by CO<sub>2</sub>, based on which a new CO<sub>2</sub> sensor with rapid response, high selectivity and good reversibility is developed. Experimental and theoretic studies indicate that the luminescence enhancement phenomenon caused by CO<sub>2</sub> can be attributed to a new response mechanism, that is, strong  $\pi$ - $\pi$  interaction between CO<sub>2</sub> and the phenyl groups of CuIDPO. Such interaction restricts the molecular rotation of CuIDPO, resulting in the reduction of non-radiative transitions, thus enhancing the luminescence intensity. Furthermore, to achieve spatially resolved two-dimensional visual detection, CuIDPO was successfully loaded onto GF paper to form membrane sensors.

## Data availability

All data have been included in the main text and ESI.<sup>†</sup>

## Author contributions

Jia-Wen Ye designed the research. Hong-Jin Zhang performed syntheses and most of the measurements. Zong-Ren Chen and Ji-Tong Xu assisted with crystallographic data. Jia-Wen Ye, Ling Chen and Xiao-Ming Chen analysed data and wrote the manuscript.

## Conflicts of interest

The authors declare that they have no conflict of interest.

## Acknowledgements

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## Notes and references

- 1 S. Rojas, A. Rodríguez-Diéguez and P. Horcajada, *ACS Appl. Mater. Interfaces*, 2022, **14**, 16983.
- 2 M. Zhang, C. Zhu, Y. Y. Duan, T. B. Liu, H. P. Liu, C. Su and Y. Lu, *Nat. Cell Biol.*, 2022, **24**, 1029.
- 3 Z. Liu, Z. Deng, S. J. Davis and P. Ciais, *Nat. Rev. Earth Environ.*, 2024, **5**, 253.
- 4 F. M. Schwandner, M. R. Gunson, C. E. Miller, S. A. Carn, A. Eldering, T. Krings, K. R. Verhulst, D. S. Schimel, H. M. Nguyen, D. Crisp, C. W. O'Dell, G. B. Osterman, L. T. Iraci and J. R. Podolske, *Science*, 2017, **358**, 7.
- 5 F. S. Rossi, J. L. Della-Silva, L. P. R. Teodoro, P. E. Teodoro, D. C. Santana, F. H. R. Baio, W. B. Morinigo, L. G. T. Crusiol, N. La Scala Jr and C. A. da Silva Jr, *Sci. Rep.*, 2024, **14**, 11.
- 6 M. Struzik, I. Garbayo, R. Pfenninger and J. L. M. Rupp, *Adv. Mater.*, 2018, **30**, 10.
- 7 A. Sharma, S. B. Eadi, H. Noothalapati, M. Otyepka, H. D. Lee and K. Jayaramulu, *Chem. Soc. Rev.*, 2024, **53**, 2530.
- 8 H. J. Zhang, T. Wu, Q. Wu, W. D. Chen, C. W. Ye, M. Y. Wang and X. D. He, *Anal. Chem.*, 2023, **95**, 18479.
- 9 L. Chen, J. M. Chen, W. W. Fu, J. Y. Chen, D. Wang, Y. K. Xiao, S. B. Xi, Y. F. Ji and L. Wang, *Nat. Commun.*, 2024, **15**, 14.
- 10 M. Zoccali, P. Q. Tranchida and L. Mondello, *TrAC, Trends Anal. Chem.*, 2019, **118**, 444.
- 11 W. J. Zhang, W. T. Chen, C. H. Li, W. Z. Sun, J. W. Ye, L. Chen, H. P. Wang and X. M. Chen, *Inorg. Chem. Front.*, 2023, **10**, 6909.
- 12 W. Q. Zhou, J. W. Ye, W. J. Zhang, M. J. Huang, L. Chen and X. M. Chen, *Anal. Chem.*, 2023, **95**, 8239.
- 13 H. Wang, S. I. Vagin, B. Rieger and A. Meldrum, *ACS Appl. Mater. Interfaces*, 2020, **12**, 20507.
- 14 X. Zhou, S. Lee, Z. C. Xu and J. Yoon, *Chem. Rev.*, 2015, **115**, 7944.
- 15 J. L. Zhu, P. P. Jia, N. Li, S. Y. Tan, J. H. Huang and L. Xu, *Chin. Chem. Lett.*, 2018, **29**, 1445.
- 16 N. Nakamura and Y. Amao, *Sens. Actuators, B*, 2003, **92**, 98.
- 17 N. Yanai, K. Kitayama, Y. Hijikata, H. Sato, R. Matsuda, Y. Kubota, M. Takata, M. Mizuno, T. Uemura and S. Kitagawa, *Nat. Mater.*, 2011, **10**, 787.
- 18 Z. Wang, H. Ma, T. L. Zhai, G. Cheng, Q. Xu, J. M. Liu, J. K. Yang, Q. M. Zhang, Q. P. Zhang, Y. S. Zheng, B. Tan and C. Zhang, *Adv. Sci.*, 2018, **5**, 7.
- 19 X. L. Qi, R. B. Lin, Q. Chen, J. B. Lin, J. P. Zhang and X. M. Chen, *Chem. Sci.*, 2011, **2**, 2214.



- 20 J. Troyano, F. Zamora and S. Delgado, *Chem. Soc. Rev.*, 2021, **50**, 4606.
- 21 L. Chen, X. B. Dong, Z. W. Mo, H. P. Wang, J. W. Ye, K. Zhang and X. M. Chen, *Adv. Opt. Mater.*, 2021, **9**, 10.
- 22 W. T. Chen, L. Chen, Z. Y. Liang, Z. W. Mo, J. W. Ye and X. M. Chen, *Adv. Opt. Mater.*, 2023, **11**, 9.
- 23 R. W. Huang, Y. S. Wei, X. Y. Dong, X. H. Wu, C. X. Du, S. Q. Zang and T. C. W. Mak, *Nat. Chem.*, 2017, **9**, 689.
- 24 C. Y. Liu, X. R. Chen, H. X. Chen, Z. Niu, H. Hirao, P. Braunstein and J. P. Lang, *J. Am. Chem. Soc.*, 2020, **142**, 6690–6697.
- 25 P. P. Sun, M. Xie, L. M. Zhang, J. X. Liu, J. Wu, D. S. Li, S. F. Yuan, T. Wu and D. Li, *Angew. Chem., Int. Ed.*, 2022, **61**, 8.
- 26 J. Peng, B. L. Chen, Z. C. Wang, J. Guo, B. H. Wu, S. Q. Hao, Q. H. Zhang, L. Gu, Q. Zhou, Z. Liu, S. Q. Hong, S. F. You, A. Fu, Z. F. Shi, H. Xie, D. Y. Cao, C. J. Lin, G. Fu, L. S. Zheng, Y. Jiang and N. F. Zheng, *Nature*, 2020, **586**, 390.
- 27 B. B. Su, J. Jin, Y. H. Peng, M. S. Molocheev, X. B. Yang and Z. G. Xia, *Adv. Opt. Mater.*, 2022, **10**, 10.
- 28 Q. C. Peng, Y. B. Si, J. W. Yuan, Q. Yang, Z. Y. Gao, Y. Y. Liu, Z. Y. Wang, K. Li, S. Q. Zang and B. Z. Tang, *Angew. Chem., Int. Ed.*, 2023, **62**, 6.
- 29 Y. Z. Wang, W. J. Zhao, Y. Y. Guo, W. B. Hu, C. X. Peng, L. Li, Y. Wei, Z. B. Wu, W. D. Xu, X. Y. Li, Y. D. Suh, X. W. Liu and W. Huang, *Light:Sci. Appl.*, 2023, **12**, 9.
- 30 C. Wang, J. Huang, R. K. Huang, Z. M. Ye, Z. W. Mo, S. Y. Liu, J. W. Ye, D. D. Zhou, W. X. Zhang, X. M. Chen and J. P. Zhang, *Inorg. Chem.*, 2019, **58**, 3944.
- 31 M. L. Ding, X. C. Cai and H. L. Jiang, *Chem. Sci.*, 2019, **10**, 10209.
- 32 S. Li, N. N. Li, X. Y. Dong, S. Q. Zang and T. C. W. Mak, *Chem. Rev.*, 2024, **124**, 7262.
- 33 M. E. Moussa, S. Evariste, H. L. Wong, L. Le Bras, C. Roiland, L. Le Polles, B. Le Guennic, K. Costuas, V. W. W. Yam and C. Lescop, *Chem. Commun.*, 2016, **52**, 11370.
- 34 T. Hofbeck, U. Monkowius and H. Yersin, *J. Am. Chem. Soc.*, 2015, **137**, 399.
- 35 M. L. Foo, R. Matsuda, Y. Hijikata, R. Krishna, H. Sato, S. Horike, A. Hori, J. G. Duan, Y. Sato, Y. Kubota, M. Takata and S. Kitagawa, *J. Am. Chem. Soc.*, 2016, **138**, 3022.

