Stable radical anions generated from a porous perylenediimide metal-organic framework for boosting near-infrared photothermal conversion

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Radical anions of electron-deficient systems are widely used, but are easily reoxidized upon exposure to air. Therefore, the stabilization of radical anions under ambient conditions is of great significance, but still remains a scientific challenge. Herein, perylenediimide is employed to prepare a crystalline metal-organic framework for stabilizing radical anions without extensive chemical modification. The porous, three-dimensional framework of perylenediimide can trap electron donors such as amine vapors and produce radical anions in-situ through photo-induced electron transfer. The radical anions are protected against quenching by shielding effect in air and remain unobstructed in air for at least a month. Because of the high yield and stability of the radical anions, which are the basis for near-infrared photothermal conversion, the framework shows high near-infrared photothermal conversion efficiency ($\eta = 52.3\%$). The work provides an efficient and simple method towards ambient stable radical anions and affords a promising material for photothermal therapy.
Near-infrared (NIR) photothermal materials convert absorbed NIR light into thermal energy by inhibiting the radiative relaxation of materials. Because of the deep penetration ability and dark field viewing properties of NIR light, the NIR photothermal effect has been of interest in applications such as NIR photothermal ablation, NIR laser-assisted photothermal therapy, and night vision sensors. Conventional organic NIR-absorbing molecules may not only require complex synthesis procedures but also easily suffer from photobleaching under light irradiation. These drawbacks lead to high costs and a risk of performance decay in photothermal processes. Thus, it is essential to devise an organic photothermal agent with stable photothermal capacity.

Perylenediimides (PDIs) are an outstanding class of organic dye molecules that have been utilized for nearly a century. They are chosen because the reported Zr-based MOFs exhibit remarkable stability and high porosity. Zr-PDI forms a fascinating 3D MOF, with a yield of 58%. Single-crystal analysis revealed that Zr-PDI crystallized in the tetragonal space group I41/a. In the Zr6(μ3-O)4(μ3-OH)4 octahedral cluster formed, the six vertices of the octahedron are occupied by Zr(IV) centers and the eight triangular faces are alternatively capped by four μ3-OH and four μ3-OH, as shown in Fig. 1a. The Zr6(μ3-O)4(μ3-OH)4 cluster is fully coordinated by 12 carboxylate units (i.e. 8 P-2COOH ligands and 4 acetic acid molecules).

Framework formation and structural characterization. The solvothermal reaction of P-2COOH (Supplementary Figs 1 and 2) and ZrCl4 in dimethyl formamide (DMF) in the presence of acetic acid as a modulator (a coordination modulator controls coordination kinetics and the resultant crystal size) produced Zr-PDI in a yield of 58%. A similar stability originates from a highly symmetric Zr6(μ3-O)4(μ3-OH)4(CO2)12 secondary building unit and organic ligands. Such useful NIR absorbers play an important role as photothermal agents for photothermal therapy. Photothermal therapy has significant advantages over chemotherapy and radiotherapy due to its superior local efficacy and fewer systemic side effects. PDI- has already been used to regulate the balance of bacterial flora. However, these applications, especially NIR photothermal conversion, are adversely affected by the instability of the RAs under ambient conditions. The formation and use of a stable PDA thus remain of prime importance. Many efforts have been devoted to improving the stability of PDIs, such as covalently modifying PDI cores and incorporating PDIs into films or supramolecular assemblies. These efforts are often demanding from both a synthesis and a processing point of view. Except for an imidazolium-modified zwitterionic PDI, the RAs are rapidly reoxidized upon exposure to air. Functionalized PDI materials without extensive chemical modification, but yielding stable RAs under ambient conditions, would be highly desirable.

Metal-organic frameworks (MOFs) are a class of robust, porous, crystalline hybrid materials that have received much attention due to their myriad of properties and simple synthesis. It is indeed possible to obtain rationally designed functional MOFs by selecting various metal ions and organic ligands or by adjusting the synthesis conditions. Among other functions, three-dimensional (3D) porous MOFs can serve as stable cages. However, the known MOFs do not on their own produce PDIs because of the lack of strongly electron-deficient aromatic ligands. We reasoned that cages composed of PDI ligands could produce PDIs in situ with improved stability due to a pronounced shielding effect. PDIs composed of PDI ligands have not been reported because 1D PDIs with coordinating carboxyl or pyridyl functions exhibit poor solubility in common solvents and 2D PDIs tend to form one-dimensional or two-dimensional assemblies with close π-π stacking, which inhibits porosity.

To elucidate this concept, we report a PDI-based 3D MOF, Zr-PDI, composed of a N,N'-di(4-benzoic acid)-1,2,6,7-tetrachloroperylene-3,4,9,10-tetracarboxylic acid diimide (P-2COOH ligand and Zr6(μ3-O)4(μ3-OH)4 clusters. The non-planar structure of P-2COOH enhances its solubility in polar solvents, thus allowing the successful solvothermal synthesis of Zr-PDI. Due to the reported Zr-based MOFs exhibit remarkable stability and high porosity, Zr-PDI forms a fascinating 3D porous network and can serve as cage to trap electron donors such as organic amines to furnish RAs in situ by photo-induced electron transfer (PET). Being confined in a framework, Zr-PDI has superior stability in air and remains intact for at least a month. To the best of our knowledge, Zr-PDI- is the first example of isolated RAs that are stable at ambient conditions without modification. A similar stability cannot be obtained by using the ligand P-2COOH under the same conditions. Zr-PDI, with NIR absorbance, in addition to high yield and stability, shows an exceptionally high NIR photothermal conversion efficiency (η = 52.3%) and good recyclability. In comparison, P-2COOH shows poor NIR photothermal conversion efficiency and recyclability.

Optical properties and formation of RAs. The arrangement of P-2COOH molecules within the framework makes it a remarkable model to study the optical properties of Zr-PDI (Supplementary Figs 8–12). In the UV–Vis–NIR absorption spectrum shown in Fig. 2a, Zr-PDI exhibits three peak absorbances in the 450–560 nm range, which are similar to the absorptions of alkyl-substituted PDI thin films. Zr-PDI also gives a red-shift of the broad absorption in the visible region from that of the
corresponding monomer in DMF, and this shift is due to a combination of excitonic coupling and charge transfer interactions between the π-stacked PDI chromophores (Supplementary Fig. 13)\(^2\). Upon irradiation with blue light (455 nm) in the presence of TEA vapor as an electron donor, the red Zr-PDI powder turned black in 10 min. The absorption spectrum of Zr-PDI\(^{-}\) displays three new peaks at 778, 917, and 1008 nm, which are characteristic peaks of PDI\(^{-}\) produced by PET\(^9,20\). The new peaks could be assigned to the D\(_{0}\) → Dn transition with a complex electronic vibration. In comparison, when treated similarly, the P-2COOH crystalline powder (Supplementary Fig. 14) produced only a low yield of P-2COOH\(^{-}\), as presented in its absorption spectrum (Supplementary Fig. 15)\(^13\).

Fluorescence quenching of Zr-PDI\(^{-}\) confirmed the formation of RAs (Fig. 2b). There are two fluorescence peaks of Zr-PDI: one is a monomer emission peak at around 578 nm and the other is a broad peak at around 657 nm (Supplementary Figs 16–18). The latter can be regarded as an excimer emission from the perylene groups\(^42\). All experiments and calculations support the proposed PET process demonstrated in Fig. 2d: the excited state Zr-PDI\(^{+}\) is reductively quenched by TEA to afford Zr-PDI\(^{-}\) and the radical cation TEA\(^{+}\).

The facile production of Zr-PDI\(^{-}\) provides evidence that Zr-PDI has an outstanding ability to adsorb TEA. To calculate the exact sorption capacity of TEA, we measured the TEA sorption test of activated Zr-PDI (MOF without solvent in the pore) sample at 298 K (Fig. 3a). Zr-PDI gave type I isotherms, with steep uptakes at low absolute pressures. The total TEA uptake capacity at 60 millibars was 4.44 mmol g\(^{-1}\) (or 0.45 g g\(^{-1}\)) for Zr-PDI (Supplementary Fig. 19). Considering the size and molecular weight of TEA, Zr-PDI exhibits high amine capacities with an equilibrium uptake of 9.06 molecules of TEA per P-2COOH molecule\(^43\). The excellent sorption capacity ensures the efficient PET reaction between TEA and Zr-PDI. The TPA sorption was also measured (Supplementary Fig. 20), the sorption isotherm of TPA being similar to that of TEA. The desorption and adsorption curves nearly coincide, indicating that there is no specific interaction between MOF and amines. The incorporation of TEA inside the framework was then probed by \(^1\)H nuclear magnetic resonance (NMR) spectroscopy. After adsorbing TEA vapor, Zr-PDI was dissolved in NaOH/D2O solution, and two additional chemical shifts corresponding to the proton signals of -CH\(_2\) and -CH\(_3\) of TEA are observed in Fig. 3b. After the TEA-loaded Zr-PDI was irradiated with 455 nm light, two new peaks belonging to TEA\(^{+}\) emerged. Besides, only broadened resonances belonging to PDI after irradiation were detected due to the paramagnetic nature of Zr-PDI\(^{-}\).

Electron paramagnetic resonance (EPR) spectroscopy was carried out for Zr-PDI samples before and after PET to support the formation of PDI\(^{-}\). As presented in Fig. 3c, Zr-PDI, after treatment with TEA and irradiation, displays a typical EPR signal, confirming the existence of free radicals. The intense resonance has no resolved hyperfine coupling, which is in accordance with literature data\(^20\). Zr-PDI\(^{-}\) and P-2COOH\(^{-}\) have the same g value of 2.0046 (Supplementary Figs 21 and 22), pointing toward the formation of PDI\(^{-}\)\(^{14}\). It has been reported that the integration of the EPR signal is proportional to the yield of RAs\(^13\). The integration of the EPR signal was calculated to be 21.7 for P-2COOH\(^{-}\) and 3639.1 for Zr-PDI\(^{-}\). Therefore, the yield of Zr-PDI\(^{-}\) was about 168 times that of P-2COOH\(^{-}\), consistent with the above UV–Vis–NIR data.

**RAs stable under ambient conditions.** Most anionic PDI species are readily reoxidized upon exposure to air\(^{16–26}\). To evaluate the stability of Zr-PDI\(^{-}\), we kept TEA-loaded Zr-PDI\(^{-}\) under ambient conditions (open air and laboratory light) or at different temperatures. Encouragingly, under ambient conditions, Zr-PDI\(^{-}\) remained almost unaffected for over a month (Fig. 2a), while the RAs produced by P-2COOH or P-2COOH/ZrCl\(_4\).
mixtures disappeared completely in 12 h (Supplementary Fig. 23). Furthermore, Zr-PDI\(^{-}\) was rather stable (detected at 778 nm in UV–Vis–NIR spectra) at high temperatures (retention rates of 89% at 120 °C and 76% even at 180 °C) after 1 h (Supplementary Figs 24 and 25), while under the same conditions, P-2COOH\(^{-}\) was completely destroyed by oxidation and high temperatures (Supplementary Fig. 26). The porous framework of Zr-PDI appears to stabilize RAs under ambient conditions and at high temperatures. The PET process leading to Zr-PDI\(^{-}\) crystalline powder was performed under ambient conditions and required neither protective measures nor low temperatures often necessary for the crystallization of radical ions. Direct formation of Zr-PDI\(^{-}\) by P-2COOH\(^{-}\) was also attempted, but failed to provide Zr-PDI\(^{-}\). Obviously, the solvothermal reaction requires high temperature and modulators, so that the RAs will be quenched.

There are three state-of-the-art ambient stable aromatic diimide RAs obtained by covalent modification of aromatic cores\(^{17,20,44}\). The displayed stability of Zr-PDI\(^{-}\) under ambient conditions is not inferior to that reported in previous studies\(^{17,20}\), but without any extra modifications in the tetrachlorinated aromatic core. The present result calls for a closer look on the improved stability of Zr-PDI\(^{-}\). The PET processes between Zr-PDI and different amines were investigated (Fig. 3d). Zr-PDI\(^{-}\) is not supposed to form when large amines (~15 Å) are used (Supplementary Fig. 27), simply because they cannot enter the pores of Zr-PDI. Some amine-loaded Zr-PDIs\(^{-}\) are presented in Fig. 3f. DEA, EDA, and TEA have comparable sizes of 6.8, 5.4, and 6.6 Å (Supplementary Fig. 28), respectively, which are much smaller than the pore sizes of Zr-PDI (~12 and ~15 Å). Nevertheless, the three amine-loaded Zr-PDIs\(^{-}\) display large differences in stability (Supplementary Fig. 29), so the size of the amine used is irrelevant to the stability of RAs. After TEA-loaded Zr-PDI\(^{-}\) were heated for 120 °C, some TEA leaked from Zr-PDI\(^{-}\). But, the amount of TEA\(^+\) was constant, and the stability of the residual Zr-PDI\(^{-}\) remained almost unchanged for another month under ambient conditions (Supplementary Figs 30 and 31). Thus, the loading of the amine used is irrelevant to the stability of RAs. By comparing amines with different boiling points, we found that the higher the boiling point of the amine used the higher the stability of the Zr-PDI\(^{-}\), as further confirmed by \(^1\)H NMR (Supplementary Table 2 and Supplementary Figs 32 and 34). Amines with low boiling points have the tendency to leak from Zr-PDI\(^{-}\) under ambient conditions, resulting in the instability of RAs. However, amines with high boiling points enable to stay in Zr-PDI under ambient conditions and even at high temperatures. The interpenetrating nature of Zr-PDI leads to a highly porous network with small cages\(^{43}\), so it is most likely that high boiling point amines together with amine radical cations are trapped in these cages to form stable adducts. Also, the reduced π–π stacking of PDIs in the framework prevents RAs to dimerize and quench\(^{13}\). Therefore, one can not only improve the yield of RAs but achieve a pronounced stabilization by confining them in the MOF cages.

**Photothermal conversion of Zr-PDI\(^{-}\).** Zr-PDI\(^{-}\) with NIR absorbance can be produced facilely and has extraordinary stability. To further verify and use this stability, we utilized
Fig. 3 Characterizations of Zr-PDI\(^{-}\). a TEA adsorption and desorption of activated Zr-PDI; inset shows TEA vapor in Zr-PDI channel. b \(^{1}\)H NMR spectra of TEA-loaded Zr-PDI before (black line) and after (red line) irradiation; inset shows the peaks of PDI. c EPR spectra of solid Zr-PDI\(^{-}\) and P-2COOH\(^{+}\); inset shows the magnified spectrum of P-2COOH\(^{+}\). d Stability of Zr-PDI\(^{-}\) loaded with different amines under ambient conditions after a week. The ordinate represents the retention rates of the absorption peak at 1008 nm. TEA triethylamine, EDA ethylenediamine, TPA tripropylamine, DEA diethylamine, IPA isopropylamine, TMA trimethylamine.

Fig. 4 Photothermal conversion performance of Zr-PDI\(^{-}\). a Illustration of the colored PDI\(^{-}\) formation and photothermal conversion of Zr-PDI\(^{-}\). b Photothermal conversion curves of Zr-PDI\(^{-}\) film on quartz glass under laser irradiation (808 nm, 0.7 W cm\(^{-2}\)). c Temperature rises of Zr-PDI\(^{-}\) at different NIR laser intensities. Inset shows average temperature rise (\(\Delta T\)) as a function of NIR laser power. d Photothermal cycling curve of Zr-PDI\(^{-}\) film (808 nm, 0.7 W cm\(^{-2}\)). e IR camera images of Zr-PDI-patterned letters BUCT (808 nm, 0.7 W cm\(^{-2}\)).
Zr-PDI as a NIR photothermal material (Fig. 4a). Under irradiation with NIR laser (808 nm, 0.7 W cm$^{-2}$), the temperature of Zr-PDI powder reached 160 °C in 10 s (Supplementary Movie 1). Then to accurately measure the photothermal conversion, Zr-PDI powder was fixed on a piece of quartz glass to form a uniform film. Under irradiation with 808 nm laser light (0.7 W cm$^{-2}$), the temperature of the quartz glass was sharply increased to as high as 114 °C, a temperature rise of over 89 °C within 200 s (Supplementary Movie 2). In comparison, with NIR laser irradiation, the temperature of a piece of blank quartz glass rose only 2.4 °C, and the temperature of a piece of quartz glass coated with a film of Zr-PDI rose only 9 °C due to the lack of NIR absorbance (Fig. 4b). These results imply that Zr-PDI has fascinating photothermal conversion properties, which is comparable to recently reported photothermal conversions of MOFs using UV–Vis light. Lower temperature increases were recorded for P-2COOH because of the low yield of RAs, indicating that it cannot serve as a photothermal conversion material. The cooling curve of Zr-PDI film, from which the conversion efficiency can be calculated (details are shown in Supplementary Note 1), is presented in Supplementary Fig. 35. An exceptionally high conversion efficiency (52.3%) was obtained, much higher than most reported photothermal materials such as Au nanorods (21.0%), organic cocrysals (18.8%), and selenophene-derived polymer films (40%) (Supplementary Table 3). Fig. 4c demonstrates that the photothermal effect is linearly dependent on the NIR laser power from 0.25 to 1 W cm$^{-2}$, indicating an indication of a thermal control performance. Correspondingly, there is no significant light emission in Zr-PDI (Supplementary Fig. 36), demonstrating the dominant non-radiative transition, in agreement with the high photothermal conversion. Notably, the PXRD pattern of Zr-PDI after laser irradiation is in agreement with the powder data (Fig. 2e), indicating that the crystal structure of Zr-PDI remains intact upon laser irradiation. Zr-PDI after photothermal conversion has a slight decrease in the $N_2$ adsorption, because of the residual TEA in the framework (Supplementary Fig. 37). The cycling test results (Fig. 4c) also demonstrate the high stability of Zr-PDI at high temperatures, which was attributed to the effective NIR absorbance of TEA, TPA, EDA-loaded Zr-PDI (Supplementary Figs 24 and 25). However, P-2COOH was unstable at high temperatures and showed inferior cycle test results (Supplementary Fig. 38). The high photothermal conversion of Zr-PDI convincingly displays the potential of the RAs in photothermal imaging, an illustration of which is shown in Fig. 4e. Under irradiation with NIR light, photothermal materials reveal a significant temperature increase, which can be captured with an IR thermal camera. We then fabricated the letter pattern BUCT with Zr-PDI powder. Under irradiation with NIR light (808 nm, 0.7 W cm$^{-2}$), the powder pattern became imageable with a high resolution. The temperature increase of Zr-PDI is large enough in bio-imaging and biomedical applications such as photothermal therapy. Also, the PET process under formation of Zr-PDI can be initiated by medically active dopamine (Supplementary Figs 39 and 40). To the best of our knowledge, this is the first direct NIR photothermal conversion of a MOF without extra chemical modification by using RAs.

In summary, a PDI-based 3D MOF (Zr-PDI) with ultrastable structure and outstanding thermal stability have great potential in biomedical applications such as bio-imaging and photothermal therapy, and the excited RAs may be useful as photothermochemical catalysts. This work not only provides an efficient method towards ambient stable RAs, but also gives insights into the development of novel PDI materials.

**Methods**

**General procedure for Zr-PDI preparation.** A mixture of P-2COOH (13.8 mg, 0.018 mmol), ZnCl$_2$ (4.3 mg, 0.018 mmol) was dissolved in 10 mL DMF and 0.45 mL acetic acid. The resulting mixture was heated in a 20 mL bottle with cap at 90 °C for 3 days, and then allowed to cool slowly to room temperature. Red crystals were collected and washed with chloroform for three times and dried at vacuum. Yield: 58% (based on P-2COOH). Anal. Calcd (%) for Zr-PDI (C$_{80}$H$_{36}$Cl$_8$N$_4$O$_{27}$Zr$_3$): C, 48.53; H, 2.28; N, 2.69.

**Gas adsorption isotherms of Zr-PDI.** Nitrogen adsorption isotherms were measured by a volumetric method using a Micromeritics ASAP 2460 gas sorption analyzer. A sample of ca. 111.6 mg Zr-PDI pre-activated at 100 °C to remove all residual solvent. Nitrogen isotherms were measured using UHP grade nitrogen. All nitrogen analyses were performed using a liquid nitrogen bath at −195.8 °C. TEA adsorption isotherms were measured by a gravimetric method using an IA002 vapor sorption analyzer. A sample of ca. 33.8 mg Zr-PDI, pre-activated at 100 °C to remove all residual solvent. TEA adsorption analyses were performed using a water bath at 25 °C and under vapor pressure of TEA.

**Photothermal conversion properties measurement.** One milligram Zr-PDI powders were dispersed in 0.1 mL acetone; the mixture was drop-cast carefully on a quartz glass (0.8 × 0.8 cm, 320 mg) to form a uniform film. The film was then treated with TEA vapor and blue light (455 nm) for 10 min to produce Zr-PDI. The 808 nm laser beam was irradiated at a power density from 0.25 to 1 W cm$^{-2}$ and 1 W cm$^{-2}$. The temperature was monitored every 10 s by a Fluke (TI400) thermal imaging camera. Photothermal conversion efficiency of the Zr-PDI was calculated by recording the change in the temperature of the quartz glass coated with Zr-PDI powders as a function of time under continuous irradiation of a 808 nm laser (0.7 W cm$^{-2}$) until the glass reached a steady-state temperature.

**Data availability**

The X-ray crystallographic coordinates for structures reported in this study have been deposited at the Cambridge Crystallographic Data Centre (CCDC), under deposition numbers 1859441. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

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