DHPA-Containing Cobalt-Based Redox Metal-Organic Cyclohelicates as Enzymatic Molecular Flasks for Light-Driven H₂ Production

Liang Zhao, Jian Wang, Pengyan Wu, Cheng He, Xiangyang Guo & Chunying Duan

The supramolecular assembly of predesigned organic and inorganic building blocks is an excellent tool for constructing well-defined nanosized molecular cavities that catalyse specific chemical transformations. By incorporating a reduced nicotinamide adenine dinucleotide (NADH) mimic within the ligand backbone, a redox-active cobalt-based macrocycle was developed as a redox vehicle for the construction of an artificial photosynthesis (AP) system. The cyclohelicate can encapsulate fluorescein within its cavity for light-driven H₂ evolution, with the turnover number (TON) and turnover frequency (TOF) reaching 400 and 100 moles H₂ per mole redox catalyst per hour, respectively. Control experiments demonstrated that the reactions were potentially occurred within the cavity of the cyclohelicates which were inhibited in the presence of adenosine triphosphate (ATP), and the redox-active NADH mimic dihydropyridine amido moieties within the ligands played an important role in photocatalytic proton reduction process.

Metal-organic macrocycles represent a unique class of functional molecular containers that display interesting recognition properties and fascinating reactivity similar to natural enzymes. The architectures generating well-defined cavities provided specific inner environments for the selective bonding of guest molecules and catalysing their reactions. Inspired by the pocket feature of natural enzymes, functional coordination cages with various structures and catalytic activities have been developed to achieve the excellent catalytic ability of natural enzymes. However, few Werner-type capsules have been used as mimics of highly evolved and finely tuned molecular photosynthetic systems, despite their importance in living systems and sustainable solar energy conversion. As artificial photosynthesis (AP) systems always involve a photosensitizer for light absorption, a catalyst for H₂ generation, and an electron donor, the intrinsic difficulties in mimicking photosynthetic systems include the feasibility of processing in an aqueous solution and a well-defined cavity to bring the proton reduction centre and photosensitizer closer together. Most importantly, metal-organic nanocages should comprise at least one of the two basic functional units that exhibit redox activity and/or light-harvesting ability for the possible construction of highly efficient and easily operated supramolecular AP systems.

Reduced nicotinamide adenine dinucleotide (NADH) plays an important role in the reduction-oxidation metabolism of some most important coenzymes found in living cells. The dihydropyridine amido (DHPA) group is described as the key structure in NADH models and an important part in electron transfer. Therefore, the incorporation of a DHPA group into the ligand backbone as the active site should be a powerful approach to mimic the activity of these enzymes, i.e., the cofactor in [FeFe]-hydrogenase that controls the redox levels by sharing the effect of electron gain, loss and distribution.

Through modulation of tridentate N₂O units containing amide groups on a central dihydropyridine ring at the meta sites (Fig. 1), we developed new cobalt-based redox-active helical triangles to encapsulate a photosensitizer for light-driven H₂ production. We reasoned that the amide groups in the positively charged cages matched the functional requirements and could offer hydrogen bonding interactions for the recognition of fluorescein (Fl).
The confinement of the cavity possibly enforced the proximity between the redox-active cobalt(II) centres and Fl, enhancing the PET efficiency to avoid unwanted energy transfer or reverse-ET reactions. The mild redox couple of DHPA close to the H₂/H⁺ couple and the geometric position of DHPA close to the redox catalyst centre made this supramolecular system a more complete working model of AP systems. Control experiments based on the reference compound that has the similar structural feature and almost same coordination geometries, as well as redox potential with that of the original, but without the DHPA fragments were also carried out for a comparison.

**Results and Discussion**

The ligand H₂ZPB containing two tridentate coordinated units was obtained from the reaction of 2-pyridyl aldehyde with malono-hydrazide in an ethanol solution. Evaporating a solution containing equivalent molar ratios of H₂ZPB and Co(NO₃)₂·6H₂O in the presence of NaClO₄ for several days led to the formation of the compound Co–ZPB. ESI-MS spectrum of the formed Co–ZPB solid exhibited intense peaks at m/z = 947.19 and m/z = 996.66, with the isotopic distribution patterns separated by 0.50 ± 0.01 Da, and a comparison with the simulation results based on natural isotopic abundances suggested that these peaks are assigned to [Co₃(H₂ZPB)₃·ClO₄]²⁺ and [Co₃(H₂ZPB)₂(H₂ZPB)·2ClO₄]²⁺, respectively, indicating the successful assembly of a Co-based M₃L₃ molecular macrocycle (Fig. 2a). Tridentate (N₃O) coordinated units sharing two five-membered chelating rings are one kind of efficient building blocks that have been widely used to construct stable and functional discrete architectures with regular structure and high symmetry. According to our previous work, each...
of three cobalt centres typically coordinated with two planar tridentate N2O chelators to form a mer configuration molecular macrocycle with considerable stabilities. In the presence of Fl, the ESI-MS spectrum of Co–ZPB exhibited a new peak at \( m/z = 1112.22 \) that was assigned to \[ \text{[Co}_3(\text{H}_2\text{ZPB})_3\text{ClO}_4]^-\text{Fl}^{2+} \] through comparison with the simulation results obtained based on natural isotopic abundances (Fig. 2b), which indicates the ability of Co–ZPB to encapsulate Fl within its cavity.

\(^1\)H NMR spectrum of Co–ZPB recorded after the addition of a 1.0 molar ratio of Fl exhibited significant upfield shifts of protons H\(_3\),6 (\( \delta = 0.15 \) ppm) and other protons, suggesting that Fl was encapsulated within the electron-rich cavity of Co–ZPB (Figure S5 in supporting information). UV-Vis titration of Co–ZPB upon addition of Fl caused a significant absorption enhancement at 510 nm. The titration curve of this band reflected the formation of 1:1 stoichiometric ratio of the host-guest complexation, with a calculated association constant of \( 2.19 \times 10^5 \) M\(^{-1} \) (Figure S9 in supporting information)\(^{32} \). It is postulated that the amide groups located within the positively charged macrocycle introduced geometric and functional properties that are beneficial to the recognition of the organic dye\(^{33,34} \).

The cyclic voltammogram of Co–ZPB in CH\(_2\)CN exhibited broad peak at \(-0.88 \) V (vs. Ag/AgCl). Because redox potentials of the cobalt centres with same coordination environment in compound Co–QDB (\textit{vide infra}) and of the DHPA moiety are very close to this value (Figures S20 and S21 in supporting information), the peak was assigned to the overlap of Co\(^{II}\)/Co\(^{I}\) reduction reaction with the reduction reaction of the DHPA moiety. Clearly, Co–ZPB is well suited to explore the redox-induced reactions that occur near the H\(_2\)/H\(^+\) couple (Fig. 3a).

When the addition of increasing amounts of Et\(_3\)NH\(^+\) triggered the appearance of a new irreversible cathodic wave near the Co\(^{II}\)/Co\(^{I}\) response. Increasing the acid concentration raised the height of the new wave with a linear relationship and shifted it to more negative potentials. The new wave was assigned to proton reduction, suggesting that Co–ZPB can reduce protons in a catalytic reaction\(^{35,36} \). Moreover, as the oxidation potential of Fl in its photoexcited state (\( \text{FI}^* \rightarrow \text{FI}^+ + e^- \)) and ground state (\( \text{FI} \rightarrow \text{FI}^+ + e^- \)) are \(-1.55 \) V and \(0.87 \) V (vs SCE)\(^{37} \), respectively, the photoexcited state of Fl (\( \text{FI}^* \)) has sufficient capability to reduce Co(II) to Co(I) directly. In the meantime, Co–ZPB was also an efficient quencher of the photosensitizer Fl (Fig. 4a and Figure S12 in supporting information). The addition of Co–ZPB to the solution of Fl (10 \( \mu \)M) in 1:1 CH\(_2\)CN/H\(_2\)O caused significant emission quenching. The quenching behaviour is considered a photoinduced electron transfer process from the excited state of Fl (\( \text{FI}^* \)) to Co–ZPB, enabling the activation of Co–ZPB by Fl for H\(_2\) production in solution\(^{38,39} \).
The photocatalytic activities of Co–ZPB (0.1 mM) assembled with Flu (0.1 mM) towards evolution of molecular hydrogen were evaluated in an acetonitrile/water solution at room temperature in the presence of 5% triethylamine (TEA) as the sacrificial electron donor. The volume of H₂ was quantified at the end of the photolysis by GC of the headspace gases. Our system could work at pH range from 10.5 to 12.5, with maximal H₂ evolution at pH 11.0 (Figure S15 in supporting information). The initial calculated turnover frequency (TOF) was approximately 100 moles H₂ per mole catalyst per hour, with a turnover number (TON) of approximately 400 moles H₂ per mole of catalyst (Fig. 3b). Notably, the TON for Flu and the redox catalyst was obtained in a stoichiometric catalyst/photosensitizer ratio. Compared to the intermolecular systems in which the TON value of one component is optimized with the other component in greater excess, the TON in the stoichiometric system reflects the true activity of the AP system. Meantime, ESI-MS spectrum of the Co–ZPB after reaction exhibited intense peaks at m/z = 947.21, 1062.71 and 1112.20, with the isotopic distribution patterns separated by 0.50 ± 0.01 Da. The peaks were assigned to host and host-guest complex, respectively, indicating the Co–ZPB/Fl system has sufficient structural stabilities during the reaction (Figure S2 in supporting information).

At a fixed Flu concentration (0.1 mM), the initial rates of H₂ generation increased with the [Co–ZPB] at lower concentrations (<0.1 mM) (Figure S19 in supporting information). When [Co–ZPB] was fixed 0.1 mM and the Flu concentration was varied, the TOF plateaued at 0.1 mM; further addition of Flu did not increase the lifetime or TON of Co–ZPB. In all cases, the optimal conditions consisted of a constant molar ratio of Co–ZPB/Fl. An increase in the Co–ZPB/Fl ratio decreased the TON, and a decrease in the Co–ZPB/Fl ratio hardly increased the TONs of Flu or Co–ZPB. A 1:1 stoichiometric ratio of Co–ZPB/Fl complexation species apparently dominated the photosynthetic system. Control experiments demonstrated that Flu, Co–ZPB and light are essential for H₂ generation.

To confirm whether the photoinduced H₂ production occurred within the cavity of Co–ZPB or through a normal homogeneous system, the photocatalytic reaction was inhibited by the addition of a non-reactive species, adenosine triphosphate (ATP), to the reaction mixture because previous work showed that a cobalt-based cyclohelicate recognized ATP. As expected, the presence of the molecular host Co–ZPB led to obvious upfield shifts of the aromatic protons on the adenosine ring, suggesting that ATP was encapsulated within the cavity of the macrocyclic complex (Figure S6 in supporting information). The ESI-MS spectrum of Co–ZPB in the presence of ATP exhibited an intense peak at m/z = 1149.69, with the isotopic distribution patterns separated by 0.50 ± 0.02 Da. This peak was assigned to [Co₃(HZPB)₃(H₂ZPB) ⊃ ATP]⁺, indicating the stable existence of Co–ZPB in solution and the successful encapsulation of ATP within the cavity of Co–ZPB (Fig. 2c).
Importantly, the addition of ATP to replace the photosensitizer or redox catalyst Co–ZPB did not result in any H₂ production, but the presence of 0.3 mM ATP effectively stopped the photocatalytic H₂ production of the Co–ZPB (0.1 mM)/FI (0.1 mM) system (Fig. 4b). This competitive inhibition behaviour was described as enzymatic-like and suggested that the H₂ production possibly occurred within the cavity of Co–ZPB. The UV-Vis titration of Co–ZPB after the addition of ATP caused a significant decrease in absorption at 510 nm. The titration curve confirmed the 1:1 stoichiometric host-guest behaviour with an association constant of 3.64 × 10⁶ M⁻¹ (Figure S17 in supporting information). This value was thirty fold larger than that of the encapsulation of FI, demonstrating the possibility of ATP to substitute for FI to encapsulate the cavity of the metallohelicate. The H₂ production likely occurred within the cavity of Co–ZPB, rather than in a normal homogeneous system.⁴³

The incorporation of a DHPA group into the ligand backbone as the active site seemed being a powerful approach to adjust the overpotential of the metal sites for proton reduction by sharing the effect of electron gain, loss and distribution. To further investigate the important role of the NADH model in the proton reduction process, a new metallohelicate Co–QDB that has the similar molecular structural features and coordination geometries of cobalt centers with that of Co–ZPB, but without fragment of the DHPA group was synthesized and structurally characterized for comparison. The ligand H₂QDB was synthesized through the reaction of 2-quinoinecarboxaldehyde with 5-(dibenzylamino)soprophaloyldrazide according to the literature method (Fig. 5)⁴⁴. Co–QDB was prepared in a yield of 75% by layering a methanol solution of H₂QDB in the presence of NH₄PF₆. The ESI-MS spectrum of Co–QDB exhibited intense peaks at m/z = 1088.05, 1119.55 and 1161.03, with the isotopic distribution patterns separated by 0.5 ± 0.01 Da, and a comparison with the simulation results based on natural isotopic abundances suggested that the peaks are assigned to [Co₃(QDB)(H₂QDB)₃(NO₃)]³⁺, [Co₃(H₂QDB)(NO₃)₂]²⁺ and [Co₃(H₂QDB)(PF₆)]²⁺, respectively, revealing the same structure and stability of the Co–QDB in solution (Figure S3 in supporting information). Single-crystal X-ray analysis confirmed the formation of a pseudo-C₄ symmetric macrocyclic helicate with three cobalt ions and three deprotonated H₂QDB ligands connected in an alternating pattern (Figure S1 in supporting information). Each cobalt centre was coordinated by two tridentate N₂O chelating groups in a mer geometry, with pairs of O atoms and amide N atoms trans to each other, further indicating the mer configuration of Co–ZPB. The measured C–O, C–N and N–N bond distances were all within the normal range of single and double bonds, pointing to the extensive electron delocalization over the entire molecular skeleton (Table S1 in supporting information)⁴⁵,⁴⁶. The separations between cobalt ions were 9.58 Å on average, and the average separation between the tertiary amine N atoms was 11.36 Å. The presence of four counter anions revealed that only two of the amide groups lost their protons during the coordination. These amid groups provided geometric and functional properties beneficial to the recognition of organic dyes, as observed in our previous works.⁴³

The cyclic voltammogram of Co–QDB recorded in DMF exhibited one reversible reduction of Co³⁺/Co²⁺ at −1.08 V (vs. Ag/AgCl). This potential falls well within the redox range of reducing a proton in aqueous media⁴⁷, enabling the host to be a redox catalyst for proton reduction (Figure S21 in supporting information). Co–QDB was also demonstrated to be an efficient quencher of the excited state of FI through photoinduced electron transfer (Fig. 6a and Figure S14 in supporting information). Photolysis of a solution of 0.04 mM FI and 0.08 mM Co–QDB in a solvent mixture containing TEA (5% v:v) in DMF/CH₂CN/H₂O resulted in H₂ generation, with optimal photocatalysis at pH 10.0 (Figure S16 in supporting information). As shown in Fig. 6b, the initial TOF was approximately 40 moles H₂ per mole catalyst per hour, with a TON of approximately 250 moles H₂ per mole of catalyst. The TON and TOF of the Co–QDB/FI system is obviously lower than those of the Co–ZPB/FI system.

Interestingly, the Co–QDB/FI ratio is crucial: the TON plateau at a 2:1 stoichiometric ratio of Co–QDB/FI under the optimal conditions. At a fixed Co–QDB concentration (0.08 mM), the decrease in the Co–QDB/FI ratio decreased the TON, and the increase in the Co–QDB/FI ratio hardly increased the TON of FI or Co–QDB, suggesting that a potential 2:1 stoichiometric ratio of the Co–QDB/FI complexation species dominated the photosynthetic system (Fig. 6b). Additionally, glutathione (GSH), an important compound in natural systems that is inactive toward hydrogenation, was chosen as an inhibitor because our previous work showed that an

![Figure 5](image-url)
isostructural cyclohelicate could recognize GSH well. When the addition of 0.2 mM GSH to the 2:1 Co–QDB (0.08 mM)/Fl (0.04 mM) system directly stopped the photocatalytic H₂ production. Since GSH does not exhibit any suitable redox potential for H₂ production, this competitive inhibition suggested that the H₂ production occurred within the cavity of Co–QDB.

The ESI-MS spectrum of Co–QDB in the presence of Fl exhibited intense peaks at \(m/z \approx 2360.27\) assigned to \([K[Co(HQDB)(QDB)]_2 \cdot Fl]^2^+\), providing additional proof for the 2:1 stoichiometric complexation behaviour (Figure S3 in supporting information). After irradiating the system for 6 hours, ESI-MS spectrum of the Co–QDB also exhibited intense peaks at \(m/z = 1088.05, 1161.03, \) and 2341.27, with the isotopic distribution patterns separated by \(0.50 \pm 0.01\) Da. The peaks are assigned to host and host-guest complex, respectively, indicating the Co–QDB system also has sufficient structural stabilities in the reaction process (Figure S4 in supporting information). UV-Vis titration of Co–QDB after addition of Fl supported the 2:1 stoichiometry of the host-guest complexation, with an association constant of \(8.32 \times 10^9\) M⁻² (Figure S10 in supporting information). The ¹H NMR spectrum of Co–QDB after the addition of a 0.5 molar ratio of Fl exhibited significant upfield shifts of protons \(H_3,6 (\delta = 0.13\) ppm\) and other protons, reflecting the encapsulation of Fl within the cavity of the macrocycle Co–QDB (Figure S7 in supporting information). Of course Co–QDB was able to recognize GSH in similar aqueous media (Figure S8 in supporting information). UV-Vis absorption titration of Co–QDB after the addition of GSH also induced quenching and suggested the formation of a 1:1 stoichiometry of the host-guest complexation with an association constant of \(5.05 \times 10^5\) M⁻¹ (Figure S18 in supporting information). At a fixed Co–QDB concentration of 0.08 mM, the presence of GSH could substitute for Fl to occupy the cavity of Co–QDB. It is hypothesized that Co–ZPB and Co–QDB are true molecular flasks, within which AP systems are assembled through encapsulation of an organic dye as a photosensitizer.

**Conclusion**

In summary, we have reported the preparation of a redox-active cobalt-based macrocycle through the incorporation of an NADH mimic within the ligand backbone and a new strategy for the construction of AP systems. The metal–organic cyclohelicate is an enzymatic molecular flask and encapsulated Fl within its cavity for light-driven H₂ evolution with a TON and TOF that reached 400 and 100 moles H₂ per mole redox catalyst per hour, respectively. The reaction was inhibited by the presence of ATP and occurred within the cavity of the cyclohelicate. The control experiments indicated that the redox-active dihydropyridine amido group of the NADH mimic was
helpful for the photocatalytic proton reduction process. By incorporating other redox-active or photocatalytic functional groups, this strategy can be extended to highly active AP systems.

Materials and Methods

Materials. All chemicals were reagent grade, obtained from commercial sources and used without further purification. The elemental analyses of C, H and N were performed on a Vario EL III elemental analyser. 1H NMR spectra were measured on a Varian INOVA 400 MHz spectrometer. ESI mass spectra were obtained on an HPLC-Q-TOFMS instrument using methanol as the mobile phase. UV-Vis spectra were measured on an HP 8453 spectrometer. The solution fluorescence spectra were obtained using an FLS920 spectrometer (Edinburgh Instruments). Both the excitation and emission slit widths were 2 nm. The solutions of Co–ZPB (1.0 × 10−3 M) and Co–QDB (4.0 × 10−3 M) were prepared in CH3CN and DMF, respectively. Stock solutions of Fl (1.0 × 10−4 M) were prepared directly in CH3CN and were excited at 460 nm.

All electrochemical measurements were carried under nitrogen at room temperature on a CHI 1130 (CH Instrument Co., Shanghai) electrochemical analyser with a conventional three-electrode system consisting of a homemade Ag/AgCl electrode as the reference electrode, a platinum wire electrode with a 0.5 mm diameter as the counter electrode, and a glassy carbon electrode as the working electrode. Cyclic voltammograms were recorded at solution concentrations of 0.1 mM and 1.0 mM for Co–ZPB and Co–QDB, respectively, and 0.1 M for the supporting electrolyte, (n-Bu4N)ClO4. The electrodes were polished on an MD-Nap polishing pad. A 0.2 M solution of Et3NHCl was added via a syringe.

General Procedure for Hydrogen Production. For photoinduced hydrogen evolution, varying amounts of the catalyst and the 1:1 CH3CN/H2O solution containing Fl and TEA were added in a total volume of 5.0 mL for Co–ZPB, and varying amounts of catalyst, Fl and TEA in DMF/CH3CN/H2O (1/4/4) were added to a total volume of 5.0 mL for Co–QDB. The pH of this solution was adjusted by adding HCl or NaOH and measured with a pH metre56,57. Typically, the Co–ZPB sample contained Co–ZPB (1 × 10−4 M), Fl (1 × 10−4 M) and 5% TEA as the sacrificial electron donor at pH 11.0, and the Co–QDB sample contained Co–QDB (8 × 10−5 M), Fl (4 × 10−5 M) and 5% TEA as the sacrificial electron donor at pH 10.0. The flask was sealed with a septum, protected from light, and degassed by bubbling nitrogen for 15 min under atmospheric pressure at room temperature. Next, the samples were irradiated by a 500 W xenon lamp; the reaction temperature was maintained at 293 K using a water filter to absorb heat. The generated photoproduction of H2 was characterized on a 7890 TGC instrument with a 5 Å molecular sieve column (0.6 m × 3 mm), a thermal conductivity detector, and nitrogen as the carrier gas. The amount of hydrogen generated was determined by the external standard method. The hydrogen in the resulting solution was not measured, and the slight effect of the hydrogen gas generated on the pressure of the Schlenk bottle was neglected in the calculation of the volume of hydrogen gas.

Synthesis of Co–ZPB. Dimethyl 1-benzyl-4-phenyl-1,4-dihydropyridine-3,5-dicarboxylate. Methyl propiolate (1.68 g, 20 mmol), benzaldehyde (1.06 g, 10 mmol), and benzylationine (1.07 g, 10 mmol) in glacial acetic acid (2.0 mL) were heated at 80 °C for 30 min52. After cooling, the mixture was poured into water (20 mL) and CH3OH/CH2Cl2 (1/1, v/v) to give a red solution. After addition of NaClO4, red precipitates formed were isolated from the Schlenk bottle was neglected in the calculation of the volume of hydrogen gas.

1-benzyl-4-phenyl-1,4-dihydropyridine-3,5-dicarbonyldrazide. A mixture solution of 80% hydrazine hydrate (50 mL) and dimethyl 1-benzyl-4-phenyl-1,4-dihydropyridine-3,5-dicarboxylate (3.63 g, 10 mmol) was stirred at 85 °C over 12 h. The white precipitate was formed, which was collected by filtration, washed with methanol and dried under vacuum. Yield: 68%. 1H NMR (400 MHz, DMSO-d6, δ, ppm): δ 7.51 (s, 2H), 7.45–7.41 (m, 2H), 7.37–7.34 (m, 3H), 7.20–7.16 (m, 2H), 7.13–7.08 (m, 3H), 4.82 (s, 2H), 4.70 (s, 1H), 3.53 (s, 6H).

Synthesis of Co–QDB. 1-Benzyl-4-phenyl-1,4-dihydropyridine-3,5-dicarbonyldrazide (3.63 g, 10 mmol) was added to a ethanol solution (50 mL) containing 2-pyrdylaldehyde (2.35 g, 22 mmol). After 5 drops of acetic acid was added, the mixture was heated at 85 °C under magnetic stirring for 12 h according to the ref.53. The yellow solid product was filtered and washed with Et2O (3×30 mL) to give pure dimethyl 1-benzyl-4-phenyl-1,4-dihydropyridine-3,5-dicarboxylate. Yield: 1.91 g, 52.3%. 1H NMR (400 MHz, DMSO-d6, δ, ppm): δ 7.74 (m, 3H), 7.20–7.16 (m, 2H), 7.13–7.08 (m, 3H), 4.82 (s, 2H), 4.70 (s, 1H), 3.53 (s, 6H).

Synthesis of H2ZPB. 1-Benzyl-4-phenyl-1,4-dihydropyridine-3,5-dicarbonyldrazide (3.63 g, 10 mmol) was added to an ethanol solution (50 mL) containing 2-pyridylaldehyde (2.35 g, 22 mmol). After 5 drops of acetic acid was added, the mixture was heated at 85 °C under magnetic stirring for 12 h according to the ref.53. The yellow solid product was filtered, washed with methanol and dried in vacuum. Yield: 1.72 g, 47.3%. 1H NMR (400 MHz, DMSO-d6, δ, ppm): δ 8.67 (s, 2H), 7.42–7.39 (m, 2H), 7.35–7.32 (m, 3H), 7.20 (s, 2H), 7.17–7.12 (m, 4H), 7.09–7.05 (m, 1H), 5.00 (s, 1H), 4.60 (s, 2H), 4.12 (s, 4H).

Synthesis of Co–ZPB. Co(NO3)2·6H2O (29.1 mg, 0.10 mmol) and H2ZPB (54.2 mg, 0.10 mmol) were dissolved in CH3OH/CH2Cl2 (1/1, v/v) to give a red solution. After addition of NaClO4, red precipitates formed were isolated and dried under vacuum. Yield: 68%. 1H NMR (400 MHz, DMSO-d6, δ, ppm): δ 11.36 (s, 2H), 8.57 (d, 2H), 8.21 (s, 2H), 7.83–7.78 (m, 4H), 7.51–7.47 (m, 2H), 7.46–7.41 (m, 4H), 7.39–7.35 (m, 3H), 7.27–7.20 (m, 4H), 7.14–7.09 (m, 1H), 5.35 (s, 1H), 4.77 (s, 2H). Anal calc. for Co3H17N3O8·2H2O: C 48.90, H 3.46, N 13.75%, Found: C 48.73, H 3.48, N 13.72%. ESI-MS calcd for Co3H17N3O8·2H2O, 454.36, found 454.23 [M+H]+, 564.21 [M+Na]+.

Synthesis of Co–QDB. Co(NO3)2·6H2O (30.1 mg, 0.10 mmol) and H2QDB (70 mg, 0.10 mmol) were dissolved in DMF and then stirred for 2h. The solution was left for several days at room temperature to give X-ray quality black block crystals. Yield: 65%. 1H NMR (400 MHz, DMSO-d6, δ, ppm): δ 12.24 (s, 2H), 8.61 (s, 2H), 8.46 (m, 2H), 8.12 (m, 2H), 8.05 (m, 4H), 7.79 (m, 2H), 7.76 (s, 1H), 7.66 (m, 2H), 7.46 (s, 2H), 7.40–7.27 (m, 10H), 4.85 (s, 4H).
References

1. Brown, C. J. et al. Supramolecular catalysis in metal–ligand cluster hosts. *Chem. Rev.* **115**, 3012–3035 (2015).
2. Hooley, R. J. Taking on the turnover challenge. *Nat. Chem.* **8**, 202–204 (2016).
3. Wang, Q. Q. et al. Self-assembled nanospheres with multiple endohedral binding sites pre-organize catalysts and substrates for highly efficient reactions. *Nat. Chem.* **8**, 225–230 (2016).
4. Wiester, M. J., Ullmann, P. A. & Mirkin, C. A. Carotenoid and pheophytin on semiconductor surface: self-assembly and photoinduced electron transfer. *Angew. Chem., Int. Ed.* **50**, 114–137 (2011).
5. Neel, A. J., Hilton, M. J., Sigman, M. S. & Toste, F. D. Exploiting non-covalent π interactions for catalyst design. *Nature* **543**, 637–646 (2017).
6. Deraedt, C. & Astruc, D. Supramolecular nanoreactors for catalysis. *Coord. Chem. Rev.* **324**, 102–122 (2016).
7. Wang, Z. J. et al. A supramolecular approach to combining enzymatic and transition metal catalysis. *Nature Chem.* **5**, 100–103 (2013).
8. Mahata, K., Frischmann, P. D. & Würthner, F. Giant electroactive M4L4 tetrahedral host self-assembled with Fe(II) vertices and perylene bisimide dye edges. *J. Am. Chem. Soc.* **135**, 15565–15561 (2013).
9. Meng, W. J. et al. A self-assembled M4L4 cubic cage that selectively encapsulates large aromatic guests. *Angew. Chem. Int. Ed.* **50**, 3479–3483 (2011).
10. Gray, H. B. & Maverick, A. W. Solar chemistry of metal complexes. *Science* **214**, 1201–1205 (1981).
11. Lubitz, W. & Tumas, W. Hydrogen: an overview. *Chem. Rev.* **107**, 3990–3993 (2007).
12. Han, Z. & Eisenberg, R. Fuel from Water: The photochemical generation of hydrogen from water. *Acc. Chem. Res.* **47**, 2537–2544 (2014).
13. Moonschiram, D. et al. Tracking the structural and electronic configurations of a cobalt proton reduction catalyst in water. *J. Am. Chem. Soc.* **138**, 10586–10596 (2016).
14. Stoll, T. et al. Photo-induced redox catalysis for proton reduction to hydrogen with homogeneous molecular systems using rhodium-based catalysts. *Coord. Chem. Rev.* **304-305**, 20–27 (2015).
15. Jing, X. et al. A metal—organic tetrahedron as a redox vehicle to encapsulate organic dyes for photocatalytic proton reduction. *J. Am. Chem. Soc.* **137**, 3967–3974 (2015).
16. Jagadesan, P. et al. Photochemical reaction containers as energy and electron-transfer agents. *Org. Lett.* **15**, 1326–1329 (2013).
17. Frischmann, P. D. M. & Würthner, K. F. Powering the future of molecular artificial photosynthesis with light-harvesting metallosupramolecular dye assemblies. *Chem. Soc. Rev.* **42**, 1847–1870 (2013).
18. Klüwer, A. M. et al. Self-assembled biomimetic [2Fe2S]-hydrogenase-based photocatalyst for molecular hydrogen evolution. *Proc. Natl. Acad. Sci. USA* **106**, 10460–10465 (2009).
19. Saraste, M. Oxidative Phosphorylation at the fin de siècle. *Science* **283**, 1488–1493 (1999).
20. Ying, W. H. NAD+/NADH and NADP+/NADPH in cellular functions and cell death: regulation and biological consequences. *Antioxid. Redox Signal.* **10**, 179–206 (2008).
21. Houtkooper, R. H. et al. The Secret Life of NAD+: An old metabolite controlling new metabolic signaling pathways. *Endocr. Rev.* **31**, 194–223 (2010).
22. Chen, Q. et al. Dihydrophenanthridine: a new and easily regenerable NAD(P)H model for biomimetic asymmetric hydrogenation. *J. Am. Chem. Soc.* **134**, 2442–2448 (2012).
23. Darenbourg, M. Y. & Bethel, R. D. Biomimetic chemistry: Merging the old with the new. *Science* **319**, 1269–1272 (2008).
24. Carroll, M. E. et al. Synthetic models for the active site of the [FeFe]-Hydrogenase: catalytic proton reduction and the structure of the doubly protonated. *J. Am. Chem. Soc.* **134**, 18843–18852 (2012).
25. Liu, Y. C. et al. Electron delocalization from the fullerene attachment to the dirnion core within the active-site mimics of [FeFe] Hydrogenase. *Inorg. Chem.* **51**, 5997–5999 (2012).
26. Rau, S., Walther, D. & Vos, J. G. Inspired by nature: light driven organometallic catalysis by heterooligomeric Ru(II) complexes. *Dalton Trans.* **915–919** (2007).
27. Schulz, M. et al. The role of the bridging ligand in photocalytic supramolecular assemblies for the reduction of protons and carbon dioxide. *Coord. Chem. Rev.* **256**, 1682–1705 (2012).
28. Camara, J. M. & Rauchfuss, T. B. Combining acid-base, redox and substrate binding functionalities to give a complete model for the [FeFe]-hydrogenase. *Nat. Chem.* **4**, 26–30 (2012).
29. Liu, Z. et al. Reduction of quinones by NADH catalyzed by organizerdinium complexes. *Angew. Chem. Int. Ed.* **52**, 4194–4197 (2013).
30. He, C. et al. Metal-Tunable Nanocages as Artificial Chemosensors. *Angew. Chem. Int. Ed.* **47**, 877–881 (2008).
31. Wu, H. et al. Metallohelical Triangles for Selective Detection of Adenosine Triphosphate in Aqueous Media. *Inorg. Chem.* **48**, 408–410 (2009).
32. Connors, K. A. Binding Contants, (John Wiley, New York, 1987).
33. Bent, H. A. Structural chemistry of donor-acceptor interactions. *Chem. Rev.* **68**, 587–648 (1968).
34. Huysekins, P. L. Factors governing the influence of a first hydrogen bond on the formation of a second one by the same molecule or ion. *J. Am. Chem. Soc.* **99**, 2578–2582 (1977).
35. Kasumadha, H. I., Chang, C. J. & Long, J. R. A molecular molybdenum-oxo catalyst for generating hydrogen from water. *Nature* **464**, 1329–1333 (2010).
36. McNamara, W. R. et al. A cobalt–dithioline complex for the photocatalytic and electrocatalytic reduction of protons. *J. Am. Chem. Soc.* **133**, 15368–15371 (2011).
37. Romero, N. A. & Nicewicz, D. A. Organic Photoredox Catalysts. *Chem. Rev.* **116**, 10075–10166 (2016).
38. Aldana, J. et al. Size-dependent dissociation pH of thiolate ligands from cadmium chalcogenide nanocrystals. *J. Am. Chem. Soc.* **127**, 2496–2504 (2005).
39. Wang, F. et al. A highly efficient photocatalytic system for hydrogen production by a robust hydrogenase mimic in an aqueous solution. *Angew. Chem. Int. Ed.* **50**, 3193–3197 (2011).
40. Han, Z. et al. Nickel pyridinecarboxylate complexes as catalysts for the light-driven production of hydrogen from aqueous solutions in noble-metal-free systems. *J. Am. Chem. Soc.* **135**, 14659–14669 (2013).
41. Hartley, C. L. et al. Iron polypyridyl complexes for photocatalytic hydrogen generation. *Inorg. Chem.* **55**, 8865–8870 (2016).
42. Wu, H. M. et al. metallohelical triangles for selective detection of adenosine triphosphate in aqueous media. *Inorg. Chem.* **48**, 854–860 (2009).
43. McKe, T. & McKe, J. R. Biochemistry: The Molecular Basis of Life, McGraw-Hill, New York, 3rd edn, 2003.
44. Wang, J. et al. Metal-organic cyclohexilates as optical receptors for glutathione: syntheses, structures, and host-guest behaviors. *Chem. Asian J.* **6**, 1225–1233 (2011).
45. Balakrishnarajan, M. M. & Hoffmann, R. Electron-deficient bonding in rhomoidal rings. *J. Am. Chem. Soc.* **126**, 13119–13131 (2004).
46. Zhao, Y. G. et al. A mixed-valence (Fe^{II})_2(Fe^{III})_2 square for molecular expression of quantum cellular automata. *Chem. Commun.* 5725–5727 (2008).
47. Artero, V., Chavarot-Kerlidou, M. & Fontecave, M. Splitting water with cobalt. *Angew. Chem. Int. Ed.* 50, 7238–7266 (2011).
48. Ueda, Y., Ito, H., Fujita, D. & Fujita, M. Permeable self-assembled molecular containers for catalyst isolation enabling two-step cascade reactions. *J. Am. Chem. Soc.* 139, 6090–6093 (2017).
49. Ward, M. D. & Raithby, P. R. Functional behaviour from controlled self-assembly: challenges and prospects. *Chem. Soc. Rev.* 42, 1619–1636 (2013).
50. Zhang, P. et al. Homogeneous photocatalytic production of hydrogen from water by a bioinspired [Fe_{5}S_{5}] catalyst with high turnover numbers. *Dalton Trans.* 39, 1204–1206 (2010).
51. Dong, J. F. et al. Promoting effect of electrostatic interaction between a cobalt catalyst and a xanthene dye on visible-light-driven electron transfer and hydrogen production. *J. Phys. Chem. C* 115, 15089–15096 (2011).
52. Mai, A. et al. Study of 1,4-dihydropyridine structural scaffold: discovery of novel sirtuin activators and inhibitors. *J. Med. Chem.* 52, 5496–5504 (2009).
53. Ranford, J. D., Vittal, J. J. & Wang, Y. M. Dicopper(II) complexes of the antitumor analogues acylbis (salicylaldehyde hydrazones) and crystal structures of monomeric [Cu_{2}(1,3-propanedioyl bis(salicylaldehyde hydrazone))(H_{2}O)]_{2}·(ClO_{4})_{2}·3H_{2}O and polymeric \([Cu_{2}(1,6-hexanediyl bis(salicylaldehydehydrazone))(C_{2}H_{5}OH)]_{2}·(ClO_{4})_{2}·m(C_{2}H_{5}OH)\). *Inorg. Chem.* 37, 1226–1231 (1998).

**Acknowledgements**
We gratefully acknowledge the financial support from the Natural Science Foundation of China (21501019, 21531001, and 21421005).

**Author Contributions**
C.D. and C.H. designed the project; L.Z. and J.W. synthesized the compounds and carried out the experiments; P.W. and X.G. synthesized part of the compounds. L.Z. and C.D. wrote the manuscript. All authors discussed the results and commented on the manuscript.

**Additional Information**
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-017-14728-8.

**Competing Interests:** The authors declare that they have no competing interests.

**Publisher’s note:** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© The Author(s) 2017