ORGANIC CHEMISTRY







FRONTIERS

RESEARCH ARTICLE

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Cite this: *Org. Chem. Front.*, 2022, **9**, 5485

Photoinduced host-to-guest electron transfer in a self-assembled coordination cage†

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A $[Pd_2L_4]$ coordination cage, assembled from electron-rich phenothiazine-based ligands and encapsulating an electron-deficient anthraquinone-based disulfonate guest, is reported. Upon excitation at 400 nm, transient absorption spectroscopy unveils photoinduced electron transfer from the host's chromophores to the guest, as indicated by characteristic spectral features assigned to the oxidized donor and reduced acceptor. The structure of the host-guest complex was characterized by NMR spectroscopy, mass spectrometry and single-crystal X-ray analysis. Spectroelectrochemical experiments and DFT calculations both agree with the proposed light-induced charge separation. A kinetic analysis of the involved charge transfer channels reveals, besides a guest-independent LMCT path, 44% efficiency for the host-guest charge transfer (HGCT).

Received 22nd August 2022, Accepted 24th August 2022 DOI: 10.1039/d2qo01339h

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Introduction

In nature, photosynthesis is the fundamental reaction cascade that transforms light into chemical energy, involving lightinduced charge separation within precisely arranged chromophore assemblies. Taking this as an inspiration, the quest to build efficient artificial molecular photosystems that harvest solar energy and transform it into an electrical potential difference or power the formation of high-energy fuel compounds is a key scientific challenge. 1-4 In this respect, organic and inorganic donor-acceptor (D-A) systems have gained considerable attention due to their ability to promote charge separation upon irradiation.⁵⁻⁸ A wide variety of potent approaches to design efficient D-A systems has been introduced over the years.6,9,10 In most reported discrete systems, donor and acceptor moieties are connected through some kind of linker by conventional covalent chemistry and examined for light-triggered charge transfer between excited donor and acceptor. However, connecting multiple donors/acceptors via covalent bonds under precise control over their optimal distance and spatial orientation can be synthetically challenging and tedious. Recent progress in supramolecular chemistry provides

Self-assembled coordination cages,^{20–24} featuring a confined yet accessible inner space, have been of interest as a nanoscopic container for guest encapsulation,^{25–27} as a molecular transport vehicle,^{28,29} as a stabilizer of reactive compounds³⁰ and as a supramolecular catalytic center.³¹ Furthermore, the interaction between cages and guest molecules in terms of photophysical processes was also explored before.^{32–40} Self-assembled supramolecular systems represent an ideal platform for realizing photoinduced electron transfer (PET) processes due to the possibilities to arrange donor and acceptor functionalities in short through-space distance, achieving maximum local concentration, and defined orientation. ^{18,35,36,38,41–44}

The choice of the D-A pair and the spatial arrangement of the D and A moieties play crucial roles to effectively achieve a charge separated state with long lifetime, a prerequisite for application of such systems as a basis for active materials in photovoltaic devices. As donor components, electron-rich aromatics such as phenothiazines (PTZ), arylamines as well as thiophene and carbazole derivatives are frequently chosen. Anthraquinones (AQ), benzothiadiazoles, squaraines and various cyano- and keto-functionalized heterocycles have been

alternative approaches to conventional multistep synthesis by connecting donor- and acceptor-functionalized building blocks through noncovalent or coordinative interactions, allowing the simple and high yielding assembly of D–A combinations in a modular fashion with determined stoichiometry and spatial arrangement. ^{11–13} Such supramolecular D–A systems have been explored to act as artificial photosynthetic systems that harvest light energy, as electron-transfer systems, hydrogen evolution catalysts, and further photo-redox-functional devices. ^{9,13–19}

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[†] Electronic supplementary information (ESI) available: Experimental procedures, characterization, NMR, mass, UV-Vis spectra, X-ray crystallography, and computational details. CCDC 2114030 and 2114031. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d2q001339h

employed as complementary partners to construct a D-A system. As a popular combination, PTZ and AQ derivatives are often implemented as D-A pairs owing to their suitable redox potentials and HOMO-LUMO energy gap characteristics. Act of the complementary partners to construct a D-A system.

Previously, we introduced a self-assembly approach to amalgamate PTZ- and AQ-based building blocks in a D-A system based on interpenetrated [Pd₄L₈] cage assemblies. 48,49 We first demonstrated that the self-assembly of banana-shaped PTZ- or AQ-based bismonodentate pyridyl ligands (named D or A, respectively) with Pd(II) cations yielded homoleptic catenated double-cages $[3BF_4@Pd_4L_8]^{5+}$ (L = D or A). Further, we showed that the equimolar mixture of these ligands (D:A = 1:1) with Pd(II) ions resulted in a statistical combination of mixed-ligand interpenetrated cages $[Pd_4D_mA_{8-m}]^{8+}$ (m = 8...0), containing all possible stoichiometries and stereoisomeric combinations. By far most of the components of the resulting mixture were mixed-ligand double-cages, containing at least one donor or one acceptor component, that contain densely packed PTZ and AQ functionalities in the immediate vicinity within the highly entangled supramolecular assemblies. Upon irradiation with light of 385 nm wavelength, electron transfer from the excited state phenothiazine donor onto the anthraquinone acceptor to obtain a charge-separated state could be unambiguously corroborated by a combination of transient absorption experiments in the UV-Vis and IR spectral regions. However, the fact that the examined samples in this study consisted of a statistical mixture of mixed-ligand cages hampered interpretation of the photophysical kinetics of the decisive species and restricts the understanding of structure-function relationships and the establishment of design principles for next-generation D-A systems.

Here, we report an alternative approach towards the rational design of discrete D-A systems composed of a metallosupramolecular host, assembled from electron-rich donor ligands, and an electron-deficient acceptor compound, serving as a guest. We elucidate the formation and structure of the donor-functionalized coordination cage and its host-guest complex by NMR spectroscopy, mass spectrometry, and single crystal X-ray analysis. The photophysical and electrochemical properties of the host-guest complex are described. Transient absorption experiments show that the donor-functionalized cage is able to transfer electrons to the incarcerated guest molecule within the host-guest assembly upon irradiation with light of proper wavelength.

Results and discussion

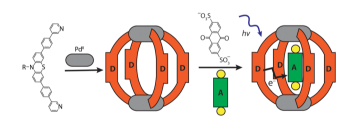
Design of the system

Ligand L was synthesized by Suzuki–Miyaura coupling of the dibromo derivative of a hexyl-substituted phenothiazine (PTZ) backbone with a boronic ester of phenyl pyridine in good yield (Scheme S1 and Fig. S1–S5†). The ligand is comparable to a PTZ ligand, previously reported by us.⁵⁰ However, the ethynyl linker was replaced with a 1,4-phenylene bridge which resulted

in the increase of the ligand's length (in terms of distance between coordinating nitrogens) by about 3 Å. Self-assembly of the ligand with palladium precursor [Pd(CH₃CN)₄](BF₄)₂ in 2:1 ratio resulted in quantitative formation of desired donor cage [Pd₂L₄](BF₄)₄ 1 in DMSO solution (Scheme 1). The formation of the cage was unambiguously confirmed by ¹H NMR spectroscopy (Fig. 1, and Fig. S6-S9†) and high-resolution mass spectrometry analysis (Fig. 2a, and Fig. S10†). In the ¹H NMR spectrum of cage 1, signals of the pyridine rings (H_i, H_h, H_f, and H_g) were downfield shifted compared to the free ligand, characteristic of the metal-mediated self-assembly. Signals corresponding to di-, tri-, and tetra-cationic complexes of the cage 1 with a differing number of BF₄ counter anions were identified in the ESI mass spectrum (Fig. 2). ¹H DOSY NMR also supported the formation of a single species with a hydrodynamic radius of 13.6 Å (Fig. S26†). All these studies confirm the successful assembly of four electron-rich ligands into a 3-dimensional cage 1. Next, cage 1 was explored in encapsulating electron-poor guest molecules.

Host-guest chemistry

Anthraquinones (AQs) are found in important compound classes such as natural pigments and redox carriers, anticancer drugs, and pesticides. They are also extensively studied in the context of redox flow batteries and photocatalysis, owing to their advantageous acceptor properties, reversible redox behaviour, and fast electrochemical kinetics. In this study,



Scheme 1 Self-assembly of donor cage 1 and host-guest complex $[G1@1]^{2+}$.

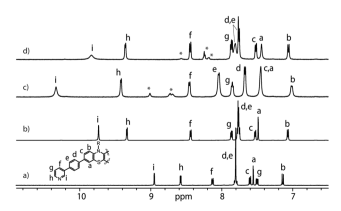


Fig. 1 Partial 1 H NMR spectra (500 MHz, DMSO- d_{6} , 298 K) of L (a), donor-functionalized cage 1 (b), host–guest assemblies [G1@1] $^{2+}$ (c) and [G3@1] $^{3+}$ (d). Signals of G1 and G3 are indicated by *.

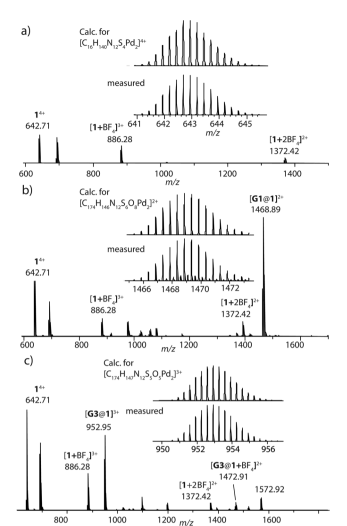


Fig. 2 HR-ESI mass spectra of donor cage 1 (a), host-guest assemblies $[G1@1]^{2+}$ (b) and $[G3@1]^{3+}$ (c).

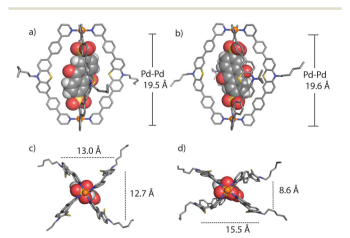
sodium salts of anthraquinone-2,6-disulfonate (G1), anthraquinone-2,7-disulfonate (G2), and anthraquinone-2-sulfonate (G3) were examined as suitable guest molecules for the donor-functionalized host 1 in DMSO. From ¹H NMR studies it was observed that only G1 and G3 were encapsulated inside cage 1 (Fig. 1), but not G2 (Fig. S13†). The size and shape of G2 are probably not compatible with the cavity of cage 1.51

The addition of one equivalent of G1 or G3 to a solution of cage 1 resulted in the formation of host-guest assemblies $[\mathbf{G1}@\mathbf{1}]^{2^+}$ and $[\mathbf{G3}@\mathbf{1}]^{3^+}$, respectively, within 5 min at room temperature, as confirmed by analysing the corresponding ¹H NMR spectra (Fig. 1). Compared to the parental species, a significant change was observed in the ¹H NMR spectrum of $[\mathbf{G1} \otimes \mathbf{1}]^{2+}$. Particularly, protons pointing towards the cavity (i.e. H_i and H_e) of the host are most affected after the addition of the guest molecule G1. Proton H_i, which points inside the cavity, is downfield shifted by about 0.64 ppm, suggesting strong interactions with the guest's sulfonate groups. In contrast, phenothiazine backbone signals (Ha, Hb and Hc) along with signal H_d were found somewhat upfield shifted. In the case of [G3@1]³⁺, only proton H_i is downfield shifted by about 0.1 ppm and broadened compared to free cage 1. A competitive ¹H NMR experiment was performed by combining **G1**, **G2**, and G3 in equimolar ratio (1:1:1) with cage 1 (1 equiv.) to check guest preference and selective uptake of G1 was observed (Fig. S24†).

Careful stoichiometric titration of guests G1 or G3 to host 1 revealed fast exchange kinetics on the NMR timescale, resulting in a single set of signals for both host and guest molecules upon guest addition (Fig. S19, and S20†). An estimation of the binding constant for G1 and G3 was derived from the titration experiments using the Bindfit online tool. 52,53 As expected, the binding constant of dianionic G1 ($\approx 1.16 \times 10^5 \text{ M}^{-1}$) is superior to monoanionic G3 (\approx 140 M⁻¹; Fig. S21, S22 and compare S50†). DOSY analysis suggested that the diffusion coefficient and size of the host are not much affected by encapsulation of the guest molecule inside the cavity (Fig. S26-S28†). Association between host and guest was also observed in the ESI-MS results. The ESI mass spectra of the host-guest assemblies show strong signals at m/z = 1469 and 953, assigned to cations $[G1@1]^{2+}$ and $[G3@1]^{3+}$, respectively (Fig. 2). Hence, all spectroscopic results indicate that the host-guest complexes of the anthraguinone acceptors inside the phenothiazine donor cages are the predominant species in DMSO solution at millimolar concentrations.

Crystal structure analysis

We succeeded in obtaining single-crystal X-ray diffraction data for ligand L (Fig. S29†) and host-guest complex [G1@1]²⁺ (Fig. 3). Slow diffusion of 1,4-dioxane into a DMSO solution of [G1@1]²⁺(with an excess of G1) over a period of four weeks resulted in tiny needle-shaped single crystals that proofed suitable for synchrotron X-ray diffraction. The host-guest assembly crystallized in triclinic form, showing space group $P\bar{1}$ (2). The



Two conformational isomers in the crystal structure of the hostguest assembly [G1@1]²⁺: conformer A (HG-A, left) and conformer B (HG-B, right), in side view along the Pd-Pd axis (a and b), and downward the Pd-Pd axis (c and d) (hydrogen substituents, solvent molecules and other G1 anions are omitted for clarity).

asymmetric unit contains two host-guest assemblies [G1@1]²⁺ besides two unbound G1 dianions and fourteen DMSO molecules (Fig. S30-S36†). The Pd···Pd distance within the cage is about 19.6 Å. The central cavity of the cage is occupied by one guest molecule. The inward pointing α -protons of the pyridine units are in close contact to the oxygen atoms of sulfonate groups (the shortest distance between S-O···H-C is 2.27 Å). The ligands' phenylene linkers also seem to help stabilizing the guest inside the cavity with similar interactions (the shortest distance between S-O···H-C is 2.37 Å). The AQ moiety is surrounded by four PTZ-based ligands of the cage assemblies, however, the guest's aromatic system does not show any significant direct noncovalent interactions with the host structure in both conformers found in the X-ray structure result. The minimum and maximum distances between the centroids of the PTZ and AQ aromatic rings are about 7.6 and 7.9 Å, respectively. One of the two cage conformers in the asymmetric unit is more extended, almost obeying a fourfold symmetry when inspected downwards the Pd-Pd axis (Fig. 3a and c), while the other conformer shows a more folded shape (Fig. 3b and d). In the latter case, the AQ major plane is positioned between the concave faces of two of the bent PTZ units while the other two PTZ units are facing towards the AQ unit with their edges in a slightly angled way. Overall, the acceptor AQ moieties of both host-guest conformers are surrounded by the four donor PTZ units in direct spatial proximity while not directly engaging in any close π - π -interaction. Owing to the fact that the NMR spectra of the host-guest assemblies do not show any guest-induced signal splitting of the host protons, free and rapid rotation of the guest inside the host can be anticipated in solution, most probably in a way that the guest's sulfonate-sulfonate axis coincides with the host's Pd-Pd axis.

Theoretical studies

Molecular models of the cage with only two counter anions [1 + $2BF_4$ ²⁺ and the host-guest assemblies [G1@1]²⁺ and [G3@1 + BF₄²⁺ were first optimized on B3LYP/def2-SVP level and the electronic properties were then calculated with B3LYP/def2-TZVP and implicit solvation (see ESI† for details).⁵⁴ As can be seen in Fig. 4, the fourfold degenerate HOMOs are spread over the PTZ moieties of the ligands L for every system. But while for $[1 + 2BF_4]^{2+}$ the twofold degenerate LUMOs are found at the palladium(II) centers, the LUMO for each host-guest

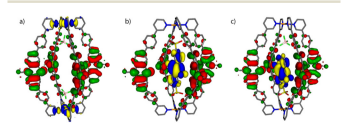


Fig. 4 Frontier molecular orbital visualization of (a) donor-functionalized cage 1 (containing two $\mathrm{BF_4}^-$ counter anions), and host-guest assemblies (b) $[G1@1]^{2+}$ and (c) $[G3@1 + BF_4]^{2+}$. The HOMOs are represented in red-green, LUMOs in yellow-blue.

assembly is spread over the acceptor guests G1 and G3, respectively. The calculated energy gaps between HOMO and LUMO are 1.95 eV for $[G1@1]^{2+}$ and 1.97 eV for [G3@1 + $BF_4|^{2+}$, while it is 2.92 eV for $[1 + 2BF_4]^{2+}$ (Table S2†).

UV-Visible spectroscopy studies

Absorption spectra of ligand L, cage 1, guests G1 and G3, and the host-guest assemblies were recorded in DMSO solution at ambient temperatures (Fig. S39†). The absorption spectra revealed maxima at 295 nm and 372 nm for L; 292 nm and 370 nm for cage 1; 284 nm and 333 for G1, 276 nm and 330 nm for G3. The absorption spectra of the host-guest assemblies are merely a sum of their components, indicating the absence of strong electronic communication between the donor and acceptor moieties at the ground state.

Cyclic voltammetry

The electrochemical properties of the D-A assemblies and their individual components were studied by cyclic voltammetry in DMSO solution at 298 K, with 0.1 M of TBAPF₆ serving as the electrolyte at a sweep rate of 100 mV s⁻¹ against an Ag/ AgNO₃ reference electrode, in a glovebox under Ar atmosphere. The resulting voltammograms are depicted in Fig. 5 and Fig. S41.† Ligand L shows a quasi-reversible wave for the PTZ moiety with an oxidation signal at 0.30 V and a reduction at about 0.22 V potential. Donor-functionalized cage 1 displays similar oxidation and reduction potentials for the PTZ moieties (0.30 and 0.22 V, respectively). The oxidation and reduction signals for the PTZ moieties in the host-guest assembly [G1@1]²⁺ are shifted to slightly lower positive potentials (0.29 and 0.21 V, respectively; $\Delta E_{\rm ox}$ = 0.01 V, and $\Delta E_{\rm red}$ = 0.01 V). Free guest G1 has its reduction and oxidation potentials at about -1.23 and -1.17 V, respectively. In the hostguest assembly $[G1@1]^{2+}$, the reduction potential (about -1.14

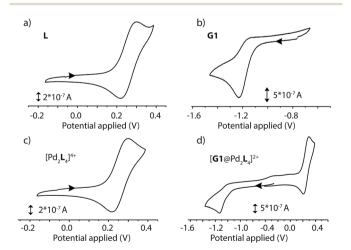


Fig. 5 Cyclic voltammograms of (a) ligand L (1.4 mM in DMSO), (b) guest G1 (0.35 mM in 0.1 M [NBu₄][PF₆] DMSO), (c) cage 1 (0.7 mM in 0.1 M [NBu₄][PF₆] DMSO) and (d) host-guest complex [G1@1] $^{2+}$, (0.7 mM in 0.1 M [NBu₄][PF₆] DMSO), recorded at a glassy carbon working electrode against a $Ag/AgNO_3$ reference electrode at 298 K. Scan rate 100 mV s⁻¹.

V) for the acceptor moiety is shifted towards a more positive potential in comparison to free **G1** ($\Delta E_{\text{red}} = 0.09 \text{ V}$).

Spectroelectrochemistry

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Spectroelectrochemical measurements of L, 1, G1, and [G1@1] were performed in a thin cell (1 mm optical path length) containing a Pt gauze and a reference electrode (Ag/AgNO₃) in combination with a diode array spectrophotometer. The resulting difference spectra are depicted in Fig. 6 and Fig. S42-S45.† The spectrum of oxidized $\mathbf{L}^{(\bullet+)}$ and $[\mathrm{Pd}_2\mathbf{L}_4^{(\bullet+)}]$ are similar, and a sharp absorption peak around 518 nm and a very broad peak around 691 nm is apparent in both cases. The reduced guest G1 revealed a broad signal around 574 nm. In the case of host-guest assembly [G1@1], the resulting spectra shows both signals for the oxidized donor and the reduced acceptor at respective voltages, similar to the free cage and guest at comparable potentials. Thus, donors and acceptor were found to maintain their characteristic electrochemical behaviours in the self-assembled host-guest complex and did not show any significant redox interplay in the ground state.

Transient absorption spectroscopy

Femtosecond pump–probe spectroscopy was applied to get insight into the dynamics following photoexcitation of the bare donor ligand **L** as well as the host cage $[Pd_2L_4]^{4+}$, with and without the presence of equimolar amounts of guest molecules **G1** or **G3**. All compounds were investigated in DMSO solution. They were excited at $\lambda_{\rm exc}$ = 400 nm (where **G1** and **G3** do not absorb) and probed in the UV-Vis at 390–730 nm using a white-light continuum.

Pump pulse-induced difference spectra for the free donor ligand L (see Fig. S46†) indicate population of a long-lived S_1 state with maxima at 450 and 613 nm and a minimum at around 500 nm, produced by superposition of excited state absorption and stimulated emission. The stimulated emission

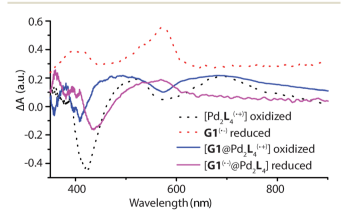


Fig. 6 Electrochemically generated difference absorption spectra for the oxidized $[Pd_2L^{(\cdot+)}_4]$, $[G1@Pd_2L^{(\cdot+)}_4]$ and reduced $G1^{(\cdot-)}$, $[G1^{(\cdot-)}@Pd_2L_4]$ species (number of oxidized L units and overall charges not specified in formulas). UV-Vis spectra of samples recorded during cyclic voltammetry with a scan rate of $0.1~V~s^{-1}$, Pt gauze as working electrode and Ag/AgNO $_3$ as reference electrode.

component exhibits a dynamic Stokes shift of 1000 cm⁻¹ within several ps. The relaxed excited state lifetime of >2 ns is consistent with a bright stationary fluorescence and similar to other phenothiazine compounds.⁵⁵

In the $[Pd_2L_4]^{4+}$ cage, the steady-state fluorescence of the phenothiazine ligand is completely quenched. This observation is consistent with the temporal evolution of the [Pd₂L₄]⁴⁺ transients presented in Fig. 7a. Directly after 400 nm excitation, a strong positive band at 613 nm characteristic for the S₁ state of the donor ligand (hereafter denoted as L^{S1}) appears, however, its lifetime is only ~ 0.7 ps. The minimum at 505 nm, indicating stimulated emission from L^{S1}, is less pronounced and gets even weaker and more red-shifted with time compared to the free ligand transients (Fig. S46†). In addition, above 700 nm the absorption slightly increases during the donor's S₁ decay and only at times >0.4 ps starts to drop with a characteristic time of ~1 ps. This is consistent with the formation of a short-lived radical cation $L^{(*)}$, exhibiting absorption bands at 500 and 700 nm (cf. spectra in Fig. 6). In our previous study on a [3BF₄@Pd₄L₈]⁵⁺ double cage, we have seen similar efficient fluorescence quenching of a related PTZ ligand derivative L due to a ligand-to-metal charge transfer (LMCT) to the bound Pd(II), followed by immediate back electron transfer (BET). 48,49 Fig. 7a suggests a similar mechanism operating in $[Pd_2L_4]^{4+}$, for which photo-induced LMCT leads to the intermediate $[Pd^{II}Pd^{I}L_3L^{(\bullet^+)}]^{4+}$. In Fig. 7a, the $L^{(\bullet^+)}$ characteristic absorption at 500 nm appears rather weak because (i) it is superimposed by the stimulated emission of the donor's S_1 state and (ii) BET is so fast that the $[Pd^{II}L_3Pd^{I}L^{(•+)}]^{4+}$ population is always low.

The spectral evolution of the photo-excited $[Pd_2L_4]^{4+}$ cage was found to be not affected by addition of the electron acceptor G3 to the cage at the same concentration used as for the host alone (0.35 mM in cage = 0.09 mM per PTZ; Fig. S47†). Given the low binding constant of G3, we suspected that the host-guest equilibrium is too far on the left side at the used concentrations which could be confirmed by an NMR experiment (Fig. S50†). However, the presence of equimolar amounts of stronger binding guest G1 leads to subtle but significant changes in the excited state dynamics (see Fig. 7b) compared to the bare host cage. Whereas for the latter, the 613 nm band assigned to LS1 stays constant at this wavelength while decaying (cf. vertical gray line in Fig. 7a), Fig. 7b indicates a blueshift of 13 nm starting at pump probe delays ≳1 ps (dashed grey line). We attribute the new band emerging at 600 nm to the reduced guest $G1^{(-)}$ being formed by electron transfer from one of the phenothiazine ligands in the coordination cage. The difference in peak position compared to the electrochemically generated spectrum of reduced G1 (λ_{max} = 575 nm, see Fig. 6) may be caused by the electric field inside the cage, associated with the proximity of two positively charge Pd2+ and one $\mathbf{L}^{(\bullet+)}$ ions. Formation of $\mathbf{G1}^{(\bullet-)}$ by host-to-guest charge transfer (HGCT) is also supported by a closer inspection of the decay kinetics of the 613 nm band. The grey data points in Fig. 7c show that with G1 the decay of this band is first faster and then turns over to a slower decay than without the guest (black

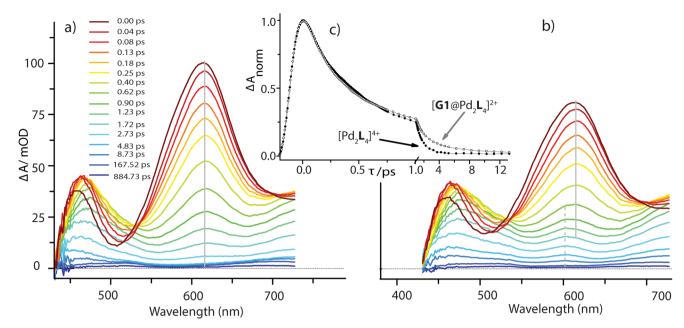


Fig. 7 Transient absorption spectra of the bare host cage [Pd₂L_a]⁴⁺ (a) and in the presence of equimolar amounts of G1 (b), following excitation at 400 nm. For both measurements the kinetic traces at a probe wavelength of 613 nm are compared in panel (c) (note that the scaling of time axis changes at 1.0 ps).

points). Also for other probe wavelengths, time traces were found to be altered upon encapsulation of G1 (Fig. S49†). The initial faster decay suggests that the presence of G1 opens up an additional reaction channel towards HGCT, accelerating depopulation of the donor's S₁ state. The following slower decay indicates that BET between the $[Pd_2L_4^{(*+)}]^{4+}$ - $G1^{(*-)}$ couple is significantly slower than for the $[Pd^{II}L_3Pd^{I}L^{(\bullet+)}]^{4+}$ intermediate.

Based on the reaction scheme depicted in Fig. 8, which involves the two identified charge transfer channels, we calculated time dependent absorption profiles of those species emerging in the transient spectra (L^{S1} , $L^{(*+)}$, and $G1^{(*-)}$). For four selected wavelengths these were fitted to measured time traces using rate constants and relative absorption cross sections $\sigma_i(\lambda)$ of the three species i as adjustable parameters (for details see the ESI†). In a first step, the kinetics of the bare

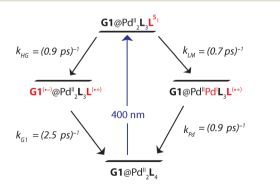


Fig. 8 Charge transfer channels of the photoexcited $[Pd_2L_4]^{4+}$ cage. The right branch represents LMCT with subsequent BET and is independent on whether G1 is present or not. The left branch corresponds to HGCT followed by BET in the presence of G1.

host cage was fitted (i.e. omitting the left branch of the reaction scheme in Fig. 8) yielding the rate constants $k_{\rm LM}$ = (0.7 ± $0.1 \text{ ps})^{-1}$ and $k_{Pd} = (0.9 \pm 0.2 \text{ ps})^{-1}$ for LMCT and corresponding BET, respectively, as well as the relative cross sections $\sigma(\mathbf{L}^{S1}, \lambda)$ and $\sigma(\mathbf{L}^{(*+)}, \lambda)$ (Fig. S48a†). Then, in a second step, the kinetics of [G1@Pd2L4] was fitted allowing for HGCT competing with LMCT, which yields $k_{\rm HG} = (0.9 \pm 0.2 \text{ ps})^{-1}$ and $k_{\rm G1} =$ $(2.5 \pm 0.3 \text{ ps})^{-1}$ for HGCT and corresponding BET, respectively, as well as the relative cross section $\sigma(G1^{(-)}, \lambda)$ (Fig. S48b†). The relative yield of $\Phi_{HG} = k_{HG}/(k_{HG} + k_{LM}) = 44\%$ shows that a significant fraction of excited state population undergoes host-toguest electron transfer. This analysis assumes that the LMCT dynamics do not change when G1 is taken up by the cage. This is supported by the absorption spectrum of [G1@Pd₂L₄] which is just the sum of the spectra of 1 and G1, indicating that there is no special interaction between host and guest that alters the electronic structure of the cage. In particular, there is no close π - π -interaction between host and guest aromatic parts. The main driving force for establishing decent host-guest affinity is the perfect arrangement of the guest's sulfonate groups close to both of the cage's inner Pd(pyridine)₄ faces, thereby replacing loosely bound BF₄ counter anions. Also, the crystal structure of the host-guest complex and DFT-based geometryoptimized models of the cage without guest are almost superimposable, i.e. the Pd-Pd distance is similar in both cases.

Conclusions

Herein, a self-assembled donor-functionalized cage based on electron-rich PTZ moieties is reported. The cage is capable of

encapsulating electron-deficient acceptor molecules such as anthraquinone-2,6-disulfonate and anthraquinone-2-sulfonate. The crystal structure of one of the host-guest assemblies [G1@1]²⁺ confirmed the presence of the guest in the central cavity of the cage. Theoretical studies of the host-guest assemblies suggested that HOMOs are spread over the donor cage components while the LUMOs are localized on the guest molecules with an energy gap of about 188 kJ mol⁻¹. Spectroelectrochemistry results revealed the characteristics of the radical ions of the donor and acceptor components to be maintained in the host-guest assembly and the absence of electronic communication between donor and acceptor in the ground state. However, the femtosecond transient UV-Vis absorption spectroscopy unambiguously shows the formation of a cage-based PTZ radical cation and a guest (G1) radical anion upon excitation of the host-guest assembly. Thus, we have demonstrated that the donor cage built from electronrich PTZ moieties is proficient in transferring electrons to the incarcerated acceptor G1 guest. The rational design of supramolecular donor-acceptor assemblies that promote photoinduced electron transfer will be of interest in the field of organic photovoltaics, given that modern self-assembly strategies allow to create complex and anisotropic molecular materials with potential to control the morphology of photoactive layers. Furthermore, photo-redox catalytic applications can benefit from host-guest chemistry via tuning substrate affinity and exchange kinetics with respect to the excited state life-time of a catalytic nanoconfinement.

Author contributions

S. Ganta and G. H. Clever conceived and designed the study. S. Ganta performed the synthesis and characterization of the materials. J. J. Holstein performed the X-ray structures analysis. C. Drechsler conducted the computational studies. J.-H. Borter and D. Schwarzer performed and analyzed femtosecond pump–probe spectroscopy experiments. S. Ganta wrote the initial draft and all authors reviewed and edited the paper.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank the Deutsche Forschungsgemeinschaft (DFG) for support through GRK2376 ("Confinement-controlled Chemistry", project number 331085229) and grant CL 489/2-2. We thank the European Research Council for ERC Consolidator Grant 683083 (RAMSES). We thank Dr. A. Baksi for the ESI-MS studies. Diffraction data of [G1@1]²⁺was collected at PETRA III, DESY (Hamburg, Germany), a member of the Helmholtz Association (HGF). We thank Eva Crosas for assistance at synchrotron beamline P11 ⁵⁶ (I-20200144). The

authors gratefully acknowledge the computing time provided on the Linux HPC cluster at TU Dortmund (LiDO3), partially funded in the course of the Large-Scale Equipment Initiative by the German Research Foundation (DFG) as project 271512359. We thank Dr. Christian Schürmann for help with the synthesis of ligand L.

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