Visible-light-responsive self-assembled complexes: improved photoswitching properties by metal ion coordination

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Abstract: A photoswitchable ligand based on azobenzene is self-assembled with palladium(II) ions to form a [Pd((E-L))3]4+ cage. Irradiation with 470 nm light results in the near-quantitative switching to a monomeric species [Pd(Z-L)]2+. The assembled structures improve selectivity of photoswitching towards the metastable isomer and extends the thermal half-life of the metastable isomer from 40 days to 850 days.

Introduction

Molecular assemblies that are addressable by non-destructive stimuli allow their properties to be adjusted at will. Visible light is an ideal stimulus that can be applied with high-resolution in time and space and, in principle, can be used without any build-up of chemical waste. One of the most appealing ways to introduce reversible light-responsive behavior to synthetic systems is by including molecular photoswitches.[1] Azobenzenes[2] and related compounds have been widely used to develop responsive receptors,[3] light responsive containers,[4] metal complexes with switchable magnetic properties,[5] photopharmacology,[6] polymers,[7] metal-organic frameworks,[8] and other types of assembled materials[9][10] including gels.[11] The binding of the different azobenzene isomers in macrocycles[12] has also allowed the formation of networks including an especially impressive example where such interactions control the assembly of macroscale gel pieces.[13]

While there are many examples of photoswitchable receptors,[14] there are relatively few examples of discrete assemblies that can be switched with visible light. Photoswitchable groups have been appended on the exterior[15] or interior[16] of assemblies that are self-assembled with metal ions[16,17] rather than the photoswitch acting as a structural element. This is due to, in part, that metal ion coordination[19] can limit photoswitching, especially for photoswitches that undergo significant geometry changes.[20] For example, when an azobenzene-appended ligand was self-assembled into a molecular sphere with palladium(II) ions, the azobenzene units could only be isomerized to form 17% of the metastable isomer, compared to 65% in the free ligand.[16] Macrocycles functionalized with azobenzenes self-assemble by hydrogen bonds to form dimeric[21] or hexameric capsules[22] where around 60-70% of the azobenzenes can be isomerized when irradiated.

The Clever group have developed a series of molecular cages with dithienylethene (DTE) photoswitches[23] that can be isomerized near quantitatively between open and closed forms with light. This switching causes a shape change of the cage and impressive changes to guest binding properties. The cages are mixtures of diastereoisomers as the closed isomer of the DTE photoswitch is chiral, which was later exploited using chiral guests to amplify the chirality of the cage.[23a] The first metal-template self-assembled cage with azobenzene switches as structural elements was reported by Hardie,[24] where the photoswitchable groups could be isomerized up to 40% in the cage, while the topology of the structure was maintained during switching. Cages formed by bridged azobenzenes (diazoacines) and palladium(II) ions can be isomerized to around 65% of the metastable isomer using visible light, essentially identical to the behavior of the free ligand.[25]

We recently reported the first example of self-assembled cages that were switchable with visible light.[26] ortho-Tetrafluoroazobenzenes[27][28] based ligands with appended pyridyl groups were assembled with palladium(II) ions to form a dynamic mixture of two species (trimer and tetramer). In that case, the most thermodynamically stable species could be selectively disassembled using light by isomerizing just 15% to the metastable isomer.[26] Herein, we report a new photoswitchable ligand that can be assembled into a single well-defined cage structure and reversibly switched to another discrete structure using visible light (Figure 1).

Results and Discussion

The photoswitchable ligand 1 was synthesized from readily available 4-bromo-2,6-difluoroniline following a modified literature procedure,[26][27] see Supporting Information S2 for details. The ligand was characterized using 1H, 19F and 13C NMR spectroscopy (Supporting Information S2), UV-vis absorption spectroscopy (Figure 2, Supporting Information S3), and mass spectrometry (see Supporting Information S6). A single crystal
suitable for X-ray diffraction was grown by the slow evaporation of a solution of E-1 in CH3CN/CHCl3/toluene. The solid state structure of E-1 (Figure 3a), is flat, with the phenyl rings perfectly co-planar, unlike in the structure of the parent \textit{ortho-tetrafluoroazobenzene} \cite{Supporting Information S3}.

We studied the photoswitching behavior of 1 in DMSO using NMR and UV-vis spectroscopy techniques (see Supporting Information S3 for details). We started by determining the photostationary state (PSS) distributions of the thermodynamically stable E-isomer and metastable Z-isomer using different irradiation wavelengths (Figure 2). Violet light centered at 405 nm provided the best selectivity towards E-1 (80% E), while red light centered at 617 nm provided the best selectivity towards Z-1 (90% Z). A 1:1 mixture of E-1 and Z-1 is generated by irradiating with blue light centered at 470 nm. When violet or blue light are used a PSS is rapidly reached, whereas a red-light source took ~2 hours to reach a PSS. Green light centered at 530 nm offers reasonable selectivity towards Z-1 (80% Z), while also generating a PSS within 5 min. Photoswitch 1 has poorer photo-selectivity compared to the parent \textit{ortho-tetrafluoroazobenzene} \cite{Supporting Information S3} resulting from smaller separation of the n-\pi* absorption bands of the two isomers, an observation that is in line with previous reports.\cite{Supporting Information S3, Supporting Information S4}

The stable E-isomer can be fully recovered using heat, with Z-1 having a 298 K thermal isomerization barrier of 111 kJ·mol⁻¹, corresponding to half-life of ~40 days (measured by variable temperature NMR, see Supporting Information S3.3). To prepare photoswitchable self-assembled structures, two equivalents of photoswitchable ligand 1 were reacted with one equivalent of [Pd(CH3CN)4][BF4]2 in DMSO-d6 (Supporting Information S4). The UV-vis absorption spectrum in DMSO is very similar to that of E-1 (Supporting Information S7). The \textsuperscript{1}H and \textsuperscript{19}F NMR spectra indicates the self-assembly of a symmetric single major species over 2 hours at room temperature (Figure 4Aii). Electrospray ionization mass spectrometry (ESI-MS) identified a [Pd(1)₂]\textsuperscript{4+} composition (Figure 4B, Supporting Information S6) and diffusion experiments show the structure has a 23 Å diameter (Supporting Information S5).

A single crystal suitable for X-ray diffraction was grown by slowly diffusing of a solution of [Pd(1)µ]\textsuperscript{4+} in CH3CN/CHCl3 into toluene. The helical structure of [Pd(1)µ]\textsuperscript{4+} is shown in Figure 3b. Both palladium(II) centers are square planar as expected. Three of the four ligands have whole-ligand disorder with each adopting two major conformations. It is clear some pedal motion \cite{Supporting Information S7} is occurring in the solid without significant change to the volume or shape of the overall assembly. Observing this motion demonstrates the flexible nature of the self-assembled cage. In solution the \textsuperscript{1}H, \textsuperscript{19}F and \textsuperscript{13}C NMR spectra are consistent with fast rotation of the phenyl rings, and/or of fast twisting of the helical structure from one helicity to the other. Such helicity inversion can occur without breaking any bonds and would change the cavity size and shape significantly, similar to the ability of flexible cages to accommodate guests by adjusting their cavity size.\cite{Supporting Information S8} In the solid state the central cavity accommodates acetonitrile solvent molecules, demonstrating capacity for guest binding.
When \([\text{Pd}(E-1)]_{2}(\text{BF}_4)_{2}\) is irradiated with green light centered at 530 nm a new major species is formed (Figure 4Aii). ESI-MS (Figure 4B, Supporting Information S6) and diffusion NMR experiments indicate that this new species comprises \([\text{Pd}(1)]^{2+}\) and has hydrodynamic diameter of 16 Å. This demonstrated that visible light can be used to cleanly convert \([\text{Pd}(E-1)]^{2+}\) to \([\text{Pd}(Z-1)]^{2+}\). Single crystals of \([\text{Pd}(Z-1)]^{2+}\) were grown by slowly diffusing of a solution of \([\text{Pd}(Z-1)]^{2+}\) in CH$_3$CN/CH$_2$Cl$_2$ into toluene. The crystals were used to determine the structure of \([\text{Pd}(Z-1)]^{2+}\), as shown in Figure 3c. The complex has one ordered ligand and one whole-ligand disorder, and confirms the trans-configuration of the ligand coordination to the palladium(II) center. The structure shows how the ligands encapsulate the palladium(II) ion, limiting access above and below the square-planar geometry which would be required for ligand exchange to occur. Heating a sample of \([\text{Pd}(Z-1)]^{2+}\) in DMSO (363 K, 16 h) regenerated \([\text{Pd}(E-1)]^{2+}\) (Figure 4Aiv).

The clean conversion of \([\text{Pd}(E-1)]^{2+}\) to \([\text{Pd}(Z-1)]^{2+}\) led us to investigate how the PSS distributions of photoswitch 1 are affected after the addition of palladium(II). We generated a 1:1 (48:52) mixture of E-1 and Z-1 (two equiv.) using blue light centered at 470 nm (Figure 5i) and added one equivalent of \([\text{Pd}(\text{CH}_3\text{CN})_2(\text{BF}_4)]_2\) to DMSO-d$_6$. The $^1$H and $^{19}$F NMR spectra showed the rapid formation of complex \([\text{Pd}(Z-1)]^{2+}\) within 5 minutes, followed by the slower formation of cage \([\text{Pd}(E-1)]^{2+}\) over 2 hours (Figure 5ii). Irradiating this sample with either light centered at 470 nm (Figure 5iii) or 530 nm (Figure 5v) causes the near-quantitative conversion to \([\text{Pd}(Z-1)]^{2+}\). To ensure the photoswitching is still reversible in the presence of palladium(II), we cycled through irradiating a sample with light centered at 405 nm and 530 nm (Figure 5v, v) to switch between mixtures of \([\text{Pd}(E-1)]^{2+}\) and \([\text{Pd}(Z-1)]^{2+}\), see Supporting Information S8 for details. As another control experiment, three separate samples of cage \([\text{Pd}(E-1)]^{2+}\) were irradiated with light centered at either 405 nm, 470 nm, or 530 nm.

Excess 4-dimethylaminopyridine (DMAP) in DMSO-d$_6$ was added to each sample to form \([\text{Pd}(\text{DMAP})_2]^{2+}\) and release ligand 1, allowing the relative abundance of E-1 and Z-1 to be determined using $^1$H NMR spectroscopy (Table 1). The sample irradiated with 470 nm light generated 95% \([\text{Pd}(Z-1)]^{2+}\), and after adding DMAP the free ligand was 89% Z-1.$^{[31]}$ In contrast, irradiating photoswitch 1 with the same light produces a PSS comprising just 52% Z-1, demonstrating unambiguously that metal ion coordination significantly improves the PSS selectivity towards the metastable isomer.

We considered the possibility of the ligands only isomerizing when dissociated from the palladium. If this were the case then an identical PSS should be reached whether palladium is present or not, provided the complexes do not significantly alter the spectrum of light available to the free ligands. To test this idea, a 1:1 mixture (46:54) sample of E-1:Z-1 (four equiv.) was prepared by irradiating 1 with 470 nm light, then one equivalent of \([\text{Pd}(\text{CH}_3\text{CN})_2(\text{BF}_4)]_2\) was added. This results in the rapid formation of \([\text{Pd}(Z-1)]^{2+}\) and the slower formation of \([\text{Pd}(E-1)]^{2+}\) over 12h to give a 1:1 (51:49) ratio of the two complexes (in terms of the ligand concentration), and leaving free E-1:Z-1 in 39:61 ratio (Supporting Information S8.4). This demonstrates the minor preference for the formation of palladium(II) complexes with the E-1 over Z-1. The sample was then irradiated with 470 nm light until a PSS was reached (20 min). The distribution of the free ligand E-1:Z-1 (45:55) at this PSS is the same as that for the free ligand in solution without added palladium (46:54). No \([\text{Pd}(E-1)]^{2+}\) was observed at the PSS, with the only complex being \([\text{Pd}(Z-1)]^{2+}\). This indicates that the photoswitching behavior of the free ligand is unchanged by the presence of the complexes and that the change in switching properties occurs either because switching can occur upon the coordinated ligands, or that the kinetics of ligand exchange of the two complexes is significantly different.

![Figure 4. A) Partial $^1$H (DMSO-d$_6$, 600 MHz, 298 K) and $^{19}$F (DMSO-d$_6$, 565 MHz, 298 K) NMR spectra of i) photoswitch E-1 ([I] = 10 mM, 2 equivalents); ii) the same sample after adding \([\text{Pd}(\text{CH}_3\text{CN})_2(\text{BF}_4)]_2\) (5.2 mM, 1 equivalent), iii) the same sample immediately after irradiating with 530 nm light for 15 min, and iv) the same sample after heating in the dark for 16 h at 90 °C and equilibrating at room temperature for 30 min. B) Zoom scans of select high resolution ESI-MS peaks for a sample of i) \([\text{Pd}(E-1)]_{2}(\text{BF}_4)_{2}\) ([I] = 16 mM), corresponding to \([\text{Pd}(1)]^{2+}\), and ii) the same sample after irradiating with 530 nm light for 20 min, corresponding to $[\text{Pd}(1)]^{2+}$.
](https://example.com/figure4.png)
Table 1. Photostationary state distributions for photoswitchable ligand 1 with and without added palladium.$^{[56]}$

| Entry | PSS 405 nm | PSS 470 nm | PSS 530 nm |
|-------|------------|------------|------------|
|       | % Z-1     | % Z-1     | % Z-1     |
| 1     | 20         | 52         | 80         |
| 1 + Pd(II)$^{[5]}$ | 54         | >95        | >98        |
| 1 + Pd(II) + DMAB$^{[5]}$ | 50         | 89         | 96         |

[a] Measured by $^1$H (600 MHz, DMSO-d$_6$, 298 K) and $^{19}$F NMR (565 MHz, DMSO-d$_6$, 298 K) data. [b] NMR spectra collected immediately after being irradiated for 30 min. [c] Immediately after the addition of ~50 equiv. of DMAB. See Supporting Information S8.3 for details.

The next significant difference in switching behavior was the slower thermal isomerization from $Z\rightarrow E$ in the presence of palladium(II). A sample of [Pd(Z-1)]$^{2+}$ was heated at 363 K and monitored by $^1$H NMR spectroscopy, see Figure 6 for details. At 363 K, [Pd(Z-1)]$^{2+}$ had a thermal half-life of 192 min and Z-1 alone had a half-life of 33 min, corresponding to room temperature (298 K) thermal half-lives of 850 days for [Pd(Z-1)]$^{2+}$ compared to only 40 days for free Z-1. When a sample of [Pd(Z-1)]$^{2+}$ was diluted, the thermal half-life was shorter, consistent with the proportion of free ligand increasing which can isomerize much faster than when it is coordinated (see Supporting Information S9 for details). The thermal relaxation from [Pd(Z-1)]$^{2+}$ to [Pd(E-1)]$^{4+}$ does not follow first-order kinetics, which is unsurprising given the mechanistic complexity of the process (see Supporting Information S9).

Most commonly, photoswitchable receptors are used to switch binding properties, rather than the binding properties being used to control photoswitching properties.$^{[14a, 14b, 32]}$ However, there are well known examples where intermolecular interactions do change photoswitching properties such as Shinkai’s classic azobenzene-linked crown ethers that can drive the PSS to 98% Z when Rb(I) is bound.$^{[33]}$ and Rebek’s azobenzene-containing receptor where the PSS modestly depends on the guest bound (from E 62% to 40–72% with guests).$^{[34]}$ An anion receptor also increased the rate of thermal Z→E isomerization when guests were bound.$^{[35]}$ The protonation of azobenzenes can significantly change their absorption and switching properties, such as the tetra-ortho-methoxy substituted azobenzenes$^{[36]}$ or azobenzazoles.$^{[37]}$ Protonation can also accelerate the Z→E thermal isomerization.$^{[38]}$ Hydrogen-bonded complexes have also been shown to accelerate Z→E isomerization,$^{[39]}$ while in other cases aggregation has slowed the Z→E isomerization time by two orders of magnitude.$^{[40]}$ Despite these examples, to the best of our knowledge the work here is the first where the rate of Z→E isomerization is decreased by coordination to transition metal ions, and offers a new method for controlling the photoswitching properties of azobenzenes.

Conclusion

In conclusion, we have shown how coordination of a molecular photoswitch to palladium(II) ions both increase the kinetic stability of the metastable Z isomer, and allow the photostationary state to be significantly improved. The resulting complexes have different properties–size, shape and reactivity—allowing visible light and heat to reversibly switch between two discrete self-assembled structures.

Figure 5. Partial $^1$H NMR (DMSO-d$_6$, 600 MHz, 298 K) spectra of i) A 1:1 mixture of E-1 and Z-1 generated using 470 nm light ([I] = 1.6 mM, 2 equivalents), ii) the same sample 2 hours after adding [Pd(CH$_3$CN)$_2$]$_2$(BF$_4$)$_2$ (0.82 mM, 1 equivalent), and iii) the same sample after irradiating with 470 nm light for 20 min, see Supporting Information S8 for details. iv) a mixture of [Pd(Z-1)]$^{2+}$ and [Pd(E-1)]$^{4+}$ ([I] = 7.1 mM) after irradiating with 405 nm light for 20 min, and v) the sample after irradiating with 530 nm light for 20 min, for more details see Supporting Information S8.

Figure 6. Comparison of thermal isomerization of Z-1 ([I] = 8.7 mM) and [Pd(Z-1)]$^{2+}$([BF$_4$])$^2$ ([I] = 7.9 mM), using $^1$H NMR (500 MHz, DMSO-d$_6$, 363 K) integrals of photoswitch Z-1 (9.14 ppm) and [Pd(Z-1)]$^{2+}$([BF$_4$])$^2$ (10.07 ppm) normalized to initial integrals. The barrier for the thermal isomerization of Z-1 at 363 K was measured at variable temperature (see Supporting Information S3.3 for details). The apparent barrier for the relaxation of [Pd(Z-1)]$^{2+}$([BF$_4$])$^2$ was determined by monitoring its depletion at 363 K.
Acknowledgements

This work was supported by the Australian Research Council (FT170100069). We acknowledge the Mark Wainwright Analytical Centre at UNSW Sydney for access to the NMR, X-ray diffraction and mass spectrometry facilities.

Keywords: photoswitch · self-assembly · metal-template · non-equilibrium · macrocyclic ligand
[35] K. Daźbrowa, P. Niedbala, J. Jurczak, Chem. Commun. 2014, 50, 15748-15751.

[36] a) A. A. Beharry, O. Sadovski, G. A. Woolley, J. Am. Chem. Soc. 2011, 133, 19684-19687; b) M. Dong, A. Babalhavaeji, S. Samanta, A. A. Beharry, G. A. Woolley, Acc. Chem. Res. 2015, 48, 2662-2670.

[37] A. D. W. Kennedy, I. Sandler, J. Andréasson, J. Ho, J. E. Beves, Chem.— Eur. J. 2020, 26, 1103-1110.

[38] a) R. S. L. Gibson, J. Calbo, M. J. Fuchter, ChemPhotoChem 2019, 3, 372-377; b) J. L. Greenfield, M. A. Gerkm, R. S. L. Gibson, G. G. D. Han, M. J. Fuchter, J. Am. Chem. Soc. 2021, 143, 15250-15257.

[39] R. Vulcano, P. Pengo, S. Velari, J. Wouters, A. De Vita, P. Tecilla, D. Bonifazi, J. Am. Chem. Soc. 2017, 139, 18271-18280.

[40] J. García-Amorós, M. C. R. Castro, S. Nonell, S. Vilchez, J. Esquena, M. M. M. Raposo, D. Velasco, J. Phys. Chem. C 2019, 123, 23140-