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Supramolecular-induced regiocontrol over the photochemical [4 + 4] cyclodimerization of NHC- or azole-substituted anthracenes†

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Thanks to the impressive control that microenvironments within enzymes can have over substrates, many biological reactions occur with high regio- and stereoselectivity. However, comparable regio- and stereoselectivity is extremely difficult to achieve for many types of reactions, particularly photochemical cycloaddition reactions in homogeneous solutions. Here, we describe a supramolecular templating strategy that enables photochemical [4 + 4] cycloaddition of 2,6-difunctionalized anthracenes with unique regio- and stereoselectivity and reactivity using a concept known as the supramolecular approach. The reaction of 2,6-azolum substituted anthracenes $H_4-L(PF_6)_2$ ($L = 1a-1c$) with Ag_2O yielded complexes *anti*- $[Ag_2L_2](PF_6)_4$ featuring an antiparallel orientation of the anthracene groups. Irradiation of complexes *anti*- $[Ag_2L_2](PF_6)_4$ proceeded under [4 + 4] cycloaddition linking the two anthracene moieties to give cyclodimers *anti*- $[Ag_2(2)](PF_6)_2$. Reaction of 2,6-azole substituted anthracenes with a dinuclear complex $[Cl-Au-NHC-NHC-Au-Cl]$ yields tetranuclear assemblies with the anthracene moieties oriented in *syn*-fashion. Irradiation and demetallation gives a [4 + 4] *syn*-photodimer of two anthracenes. The stereoselectivity of the [4 + 4] cycloaddition between two anthracene moieties is determined by their orientation in the metallosupramolecular assemblies.

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Introduction

Supramolecular approaches offer solutions to the problem of preorganization of the orientation and conformation of photo-reactive reactants,^{1,2} such that efficient photochemical reactions can also be extended to homogeneous systems.^{3,4} However, controlling the regio- and stereoselectivity of a photoinduced transformation while maintaining the high reactivity of the photoactive components remains an inherent challenge.⁵ For example, in order to improve the selectivity of photochemical [4 + 4] reactions of monosubstituted anthracenes (such as 2-anthracene carboxylic acid), much effort has been directed toward the application of various supramolecular environments.^{6–11} Although the photochemical [4 + 4] cycloaddition of anthracene derivatives is one of the well-studied subjects in

photochemistry,^{12–19} relatively few studies deal with the photodimerization of 2,6-difunctionalized anthracenes.²⁰ Generally, irradiation of 2,6-difunctionalized anthracene derivatives (AD) in solution leads to the formation of two [4 + 4] photodimerization products, the *anti*-dAD (mixture of enantiomers) and *syn*-dAD (achiral) photodimers (Fig. 1a). To the best of our knowledge, the controlled, selective formation of *anti*-dAD or *syn*-dAD in solution *via* photochemical [4 + 4] cycloaddition remains relatively unexplored, mainly due to microenvironments for a selective reaction.

Recently, we have demonstrated that NHC-functionalized (NHC = N-heterocyclic carbene) metallocycles are good candidates for efficient supramolecular-controlled photochemical [2 + 2] cycloaddition reactions in solution.²¹ We found that the antiparallel arrangement of photoreactive substrates with twisted structures can be formed by using suitable type-I units,^{21d} while a parallel arrangement of the reactants can be favored when type-II organometallic clips were employed (Fig. 1b).^{21b} In nature, many enzymes accelerate or facilitate reactions with high regio- and stereoselectivity by adjusting their microenvironments in the presence of different substrates. We thus envisaged that the flexibility of supramolecular templates, when appropriately introduced, would facilitate the preorganization of the spatial arrangement of anthracene moieties within their structures. Given the reduced translational, rotational, and conformational freedom of the

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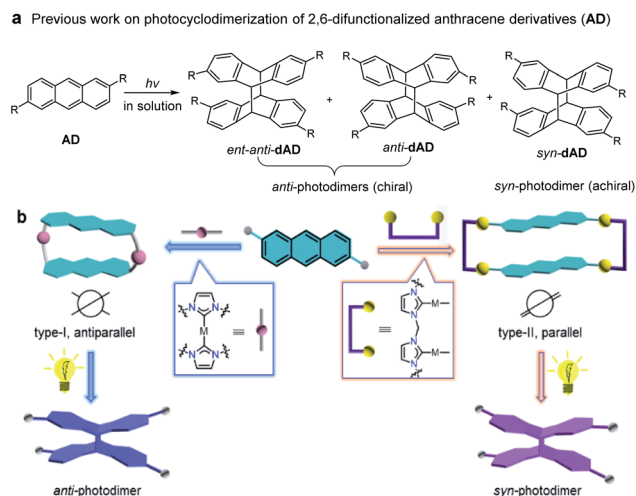
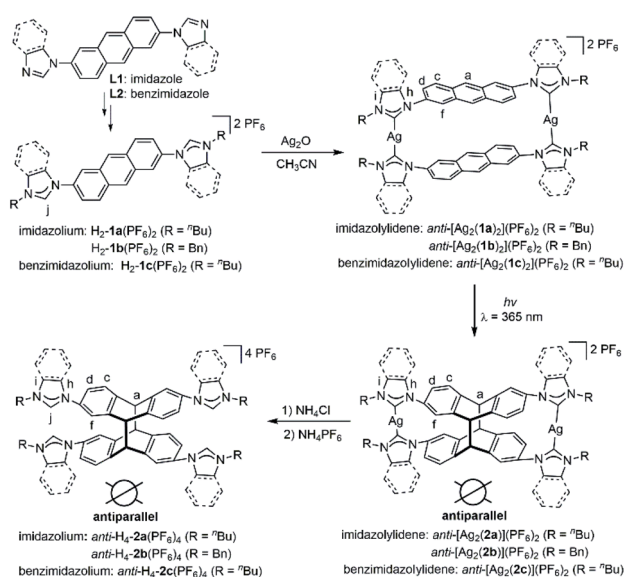


Fig. 1 (a) The mixture of isomeric products obtained by [4 + 4] cycloaddition of 2,6-substituted anthracenes. (b) The supramolecular-template-controlled regioselective [4 + 4] photochemical cycloaddition of 2,6-substituted anthracenes, in which the *anti*-photodimer is an enantiomer.

anthracene moieties and their close proximity within the metallosupramolecular structure, the selective synthesis of *anti*- and *syn*-photodimerization products from assemblies of types-I and -II appears possible. The two different supramolecular architectures should steer the [4 + 4] photochemical reaction of the anthracene groups towards different products with excellent regio- and stereoselectivity.

Herein, we demonstrate that a metallosupramolecular approach depicted in Scheme 1 allows to perform the photochemical [4 + 4] cycloaddition of 2,6-difunctionalized anthracene derivatives in solution with excellent regio- and



Scheme 1 Metallosupramolecular-controlled synthesis of tetrakisazolum salts *anti*-H₄-2(PF₆)₄ from complexes *anti*-[Ag₂(L)₂](PF₆)₂ by [4 + 4] cycloaddition.

stereoselectivity. In this supramolecular approach, the reaction outcome is determined by the preorganization of the reacting units rather than by their intrinsic reactivity. Irradiation of the different metallacycles led to the exclusive formation of a specific isomer in each case. The chiral *anti*-photodimers (*anti*-dAD and *ent-anti*-dAD in 1 : 1 molar ratio) were obtained in high yields after removal of the metal ions from the photo-products. Notably, *in situ* photolysis of metallarectangles of type-II led to the isolation of the desired achiral *syn*-photodimers (*syn*-dAD) without the need for a separate metal-removal step.

Results and discussion

Synthesis and characterization of *anti*-photodimers from type-I assemblies

The reaction of linear bisimidazolium salts with Ag_2O is known to produce dinuclear tetracarbene complexes $[\text{Ag}_2\text{L}_2]$.²¹ The 2,6-anthracene-bridged bisimidazolium salts $\text{H}_2\text{-L}(\text{PF}_6)_2$ ($\text{L} = \text{1a,b}$) and bisbenzimidazolium salt $\text{H}_2\text{-L}(\text{PF}_6)_2$ ($\text{L} = \text{1c}$) were synthesized from 2,6-di(1*H*-imidazol-1-yl)anthracene (**L1**) and 2,6-bis(1*H*-benzo[*d*]imidazol-1-yl)anthracene (**L2**) by *N*-alkylation and anion exchange (Scheme 1). The bis(benz)imidazole anthracenes **L1** and **L2** were synthesized in good yields from 2,6-dibromoanthracene and imidazole or benzimidazole by an Ullmann coupling reaction. The molecular structures of **L1** and **L2** were determined by X-ray diffraction studies (Fig. S1†). The ligands $\text{H}_2\text{-L}(\text{PF}_6)_2$ ($\text{L} = \text{1a-c}$) were characterized by NMR spectroscopy and electrospray ionization mass spectrometry (ESI-MS) (Fig. S11–S19†). Surprisingly, a solution of $\text{H}_2\text{-1a}(\text{PF}_6)_2$ was found to be photostable upon exposure to UV light ($\lambda = 365 \text{ nm}$) for 6 h. The anthracene motifs were found to be photoactive in solution only after metallation of the azolium groups with Ag^+ ions in a metallosupramolecular assembly.

The reaction of the 2,6-difunctionalized anthracenes $\text{H}_2\text{-L}(\text{PF}_6)_2$ ($\text{L} = \text{1a-c}$) with Ag_2O under exclusion of light in acetonitrile afforded the disilver(i) tetracarbene complexes *anti*-[$\text{Ag}_2(\text{L})_2$](PF₆)₂ ($\text{L} = \text{1a-c}$) in good yields (Scheme 1). The ¹H NMR spectrum of *anti*-[$\text{Ag}_2(\text{1a})_2$](PF₆)₂ showed a single set of signals with upfield shifts observed in the aromatic region relative to salt $\text{H}_2\text{-1a}(\text{PF}_6)_2$ (Fig. 2a and b). In addition, the disappearance of the imidazolium C2–H resonance (labeled H_j) was noted in the ¹H NMR spectrum upon metallation of $\text{H}_2\text{-1a}(\text{PF}_6)_2$ together with the appearance of the typical resonance at $\delta = 178.4 \text{ ppm}$ for the Ag^+ -bound carbene carbon atoms in the ¹³C{¹H} NMR spectrum (Fig. S21†). Similarly, the formation of carbene complexes *anti*-[$\text{Ag}_2(\text{L})_2$](PF₆)₂ ($\text{L} = \text{1b,c}$) were monitored by NMR spectroscopy (Fig. S26–S30† for *anti*-[$\text{Ag}_2(\text{1b})_2$](PF₆)₂ and ESI Fig. S32 and S33† for *anti*-[$\text{Ag}_2(\text{1c})_2$](PF₆)₂). The formation of *anti*-[$\text{Ag}_2(\text{L})_2$](PF₆)₂ ($\text{L} = \text{1a-c}$) was also confirmed by ESI-MS data. The HR-ESI mass spectra (positive ion mode) of the complexes showed the highest intensity peaks at $m/z = 530.1664$ (calcd for *anti*-[$\text{Ag}_2(\text{1a})_2$]²⁺ 530.1517), at $m/z = 598.1185$ (calcd for *anti*-[$\text{Ag}_2(\text{1b})_2$]²⁺ 598.1206) and at $m/z = 630.1703$ (calcd for *anti*-[$\text{Ag}_2(\text{1c})_2$]²⁺ 630.1832) with the correct isotopic patterns.



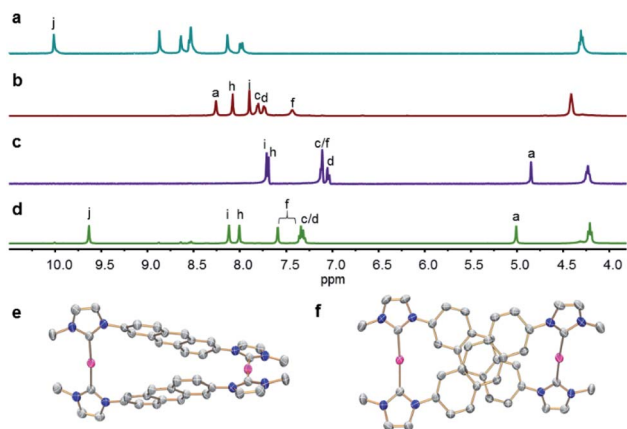


Fig. 2 Sections of the ^1H NMR spectra in $[\text{D}_6]\text{DMSO}$ of (a) bisimidazolium salt $\text{H}_2\text{-1a}(\text{PF}_6)_2$; (b) complex $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ before irradiation; (c) complex $\text{anti-}[\text{Ag}_2(\text{2a})](\text{PF}_6)_2$ obtained after irradiation; (d) tetrakisimidazolium salt $\text{anti-H}_4\text{-2a}(\text{PF}_6)_4$. (e) Side and (f) top views of complex cation $\text{anti-}[\text{Ag}_2(\text{1a})_2]^{2+}$ as determined by X-ray diffraction (N, blue; C, grey; Ag, purple) with hydrogen atoms omitted for clarity and only the first atom of each N -substituent is depicted.

Single crystals of $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{BPh}_4)_2$ were obtained by adding of an excess of NaBPh_4 to a CH_3CN solution of $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ and allowing the solution to stand at ambient temperature for several days. The X-ray diffraction analysis confirmed the formation of the disilver metallacycle (Fig. 2e and f). The coordination geometry around the silver(I) atoms is almost linear and the Ag-C_{NHC} distances (2.075(2)–2.079(2) Å) fall in the range previously reported for related silver polycarbene assemblies.²¹ The nonbonding $\text{Ag}\cdots\text{Ag}$ distance measures 10.925(3) Å.

As shown in Fig. 2f, the two anthracene skeletons are arranged in the expected antiparallel conformation, forming an X-shaped arrangement as seen from the top view. The distance between two central aromatic rings measures about 3.57 Å, which is within Schmidt's range proposed for a photoinduced cycloaddition reaction.²² It is worth mentioning that the intermolecular interactions of adjacent $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ assemblies were not observed in the unit cell, thus preventing intermolecular $[4 + 4]$ photochemical reactions.

A computational study was performed to gain insight into the preferred formation of complex $\text{anti-}[\text{Ag}_2(\text{1a})_2]^{2+}$ with the antiparallel orientation of the anthracene groups in solution. Density-functional theory (DFT) calculations at the BP86-D3(BJ)/def2-TZVPP level (see ESI† for details) revealed that the $\text{anti-}[\text{Ag}_2(\text{1a})_2]^{2+}$ species is thermodynamically more stable in CH_3CN solution than the syn -isomer (+2.4 kcal mol $^{-1}$, Fig. 3a). The calculated geometry of the anti -isomer matches the geometry of cation $\text{anti-}[\text{Ag}_2(\text{1a})_2]^{2+}$ determined by X-ray diffraction. Further calculations indicated that the rotation of the anthracene unit in $\text{anti-}[\text{Ag}_2(\text{1a})_2]^{2+}$ to give $\text{syn-}[\text{Ag}_2(\text{1a})_2]^{2+}$ features a high energy barrier, suggesting that this isomerization is not feasible.

Next, the photochemical $[4 + 4]$ cycloaddition of the anthracene units within metallacycles $\text{anti-}[\text{Ag}_2(\text{L})_2](\text{PF}_6)_2$ ($\text{L} = \text{1a-c}$) was investigated. A $[\text{D}_6]\text{DMSO}$ solution of metallacycle $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ ($c = 0.04 \text{ M}$) was irradiated with UV light (λ

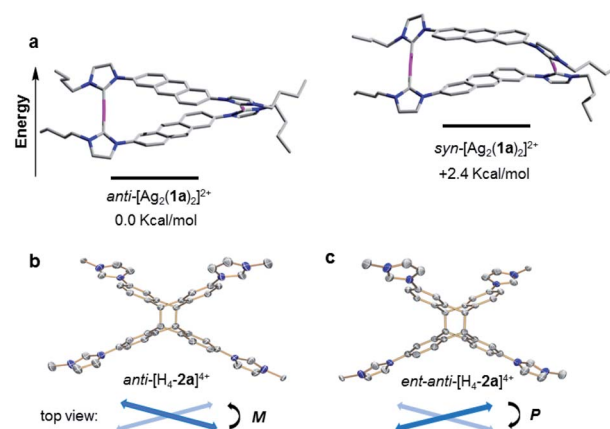


Fig. 3 (a) DFT calculated structures (in CH_3CN) and relative energies of $\text{anti-}[\text{Ag}_2(\text{1a})_2]^{2+}$ (left) and $\text{syn-}[\text{Ag}_2(\text{1a})_2]^{2+}$ (right). (b) Side view of the M -isomer of enantiomer of $\text{anti-}[\text{H}_4\text{-2a}]^{4+}$ and (c) side view of the P -isomer of enantiomer of $\text{anti-}[\text{H}_4\text{-2a}]^{4+}$ as determined by single-crystal X-ray diffraction (N, blue; C, grey; Ag, purple). Hydrogen atoms have been omitted for clarity, and only the first atom of each N -substituent is depicted.

$= 365 \text{ nm}$) at ambient temperature. ^1H NMR monitoring showed the quantitative conversion of $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ into a new single product after 30 min. Upon irradiation, the singlet resonance assigned to the H_a protons of $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ at $\delta = 8.26 \text{ ppm}$ disappeared (Fig. 2b) and a new singlet appeared at $\delta = 4.86 \text{ ppm}$, which can be assigned to H_a of the newly formed photodimer (Fig. 2c). No mass change was observed upon irradiation by mass spectrometry, confirming the intramolecular $[4 + 4]$ photodimerization within the assembly as the proceeding reaction (Fig. S25 and S43†). It should be noted that the photodimerization proceeded under mild conditions without exclusion of oxygen. The photochemical transformation can also be monitored by UV/Vis and fluorescence spectroscopy. The UV/Vis spectrum of $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ was measured in acetonitrile solution and exhibited a series of vibrationally spaced bands at wavelengths of $\lambda = 350\text{--}410 \text{ nm}$, which are assigned to the characteristic $\pi\text{-}\pi^*$ absorption of anthracene systems (Fig. S81†). Irradiation into these bands would lead to the $[4 + 4]$ photodimerization of the anthracene units, accompanied by the disappearance of these characteristic bands. Comparison of the fluorescence spectra of $\text{anti-}[\text{Ag}_2(\text{1a})_2](\text{PF}_6)_2$ before and after photodimerization exhibited a significantly decrease in the intensity for the photodimer $\text{anti-}[\text{Ag}_2(\text{2a})](\text{PF}_6)_2$, attributed to the reduction of $\pi\text{-}\pi$ stacking interactions of adjacent anthracene units (Fig. S82†).

Under the conditions described above, complex $\text{anti-}[\text{Ag}_2(\text{1b})_2](\text{PF}_6)_2$ featuring N -benzyl groups at the NHC pendants reacted upon irradiation over 25 min to give the quantitatively the photodimer $\text{anti-}[\text{Ag}_2(\text{2b})](\text{PF}_6)_2$ (Fig. S2†). The benzimidazol-2-ylidene disilver complex $\text{anti-}[\text{Ag}_2(\text{1c})_2](\text{PF}_6)_2$ also yielded photodimer $\text{anti-}[\text{Ag}_2(\text{2c})](\text{PF}_6)_2$ after irradiation for 45 min (Fig. S3†). Combined multinuclear and two-dimensional NMR and ESI-MS experiments confirmed the formation of the dinuclear photoproducts $\text{anti-}[\text{Ag}_2(\text{L})](\text{PF}_6)_2$ ($\text{L} = \text{2a-c}$).

Finally, the free tetrakisimidazolium salt *anti*-H₄-2a(PF₆)₄ has been readily obtained through demetallation and anion exchange by treating *anti*-[Ag₂(2a)](PF₆)₂ with NH₄Cl and NH₄PF₆ in methanol. The metal-free tetrakisimidazolium salt *anti*-H₄-2a(PF₆)₄ was isolated as colorless crystals in 84% yield by slow diffusion of ethyl ether into a solution of *anti*-H₄-2a(PF₆)₄ in methanol/acetonitrile at ambient temperature. The ¹H NMR spectrum suggested the presence of a single highly symmetrical species (Fig. 2d). The molecular structure of *anti*-H₄-2a(PF₆)₄ was unambiguously determined by X-ray crystallography (Fig. 3b and c). As expected, the *anti*-photodimer was obtained as a pair of *P* and *M* enantiomers (the "*P/M*" chirality is defined as the right- or left-handed screw arrangement of the longer axes of the anthracenes). Due to the generated 1,4-cyclohexadiene rings, both anthracene planes are slightly bent and the newly formed σ-bonds are rather long (up to 1.612(5) Å), which is consistent with previously reported similar compounds.²³

Synthesis and characterization of *syn*-photodimers from type-II assemblies

Encouraged by the previously described results, the photochemical conversion of the type-II assemblies (Fig. 1) was studied next. The combination of two organometallic digold clips 3 and two linear 2,6-imidazole-substituted anthracenes L1 in the presence of AgOTf resulted in the formation of the tetranuclear metallacycle *syn*-[4a](OTf)₄ in good yield (Scheme 2). NMR spectroscopy revealed downfield shifts of the imidazole proton resonances upon formation of metallacycle *syn*-[4a](OTf)₄ caused by the loss of electron density upon the coordination of the imidazole nitrogen atoms to the Au^I centers (Fig. 4a and b). The ESI-MS spectrum (positive ion mode) provided further support for the formation of *syn*-[4a](OTf)₄ by showing high-intensity peaks at *m/z* = 1113.1998 (calcd for [*syn*-[4a](OTf)₂]²⁺ 1113.2065), 692.4965 (calcd for [*syn*-[4a](OTf)]³⁺ 692.4868) and 482.1455 (calcd for [*syn*-[4a]⁴⁺ 482.1270). The peaks were isotopically resolved and the results are in perfect agreement with the theoretical distribution (Fig. S40†). Since attempts to obtain X-ray quality crystals of the complex *syn*-

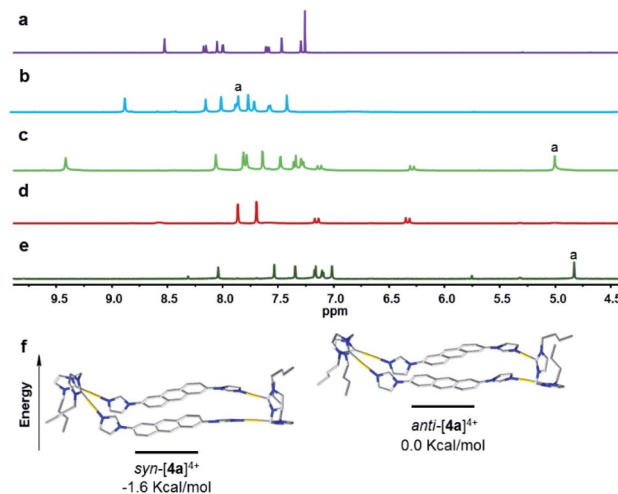
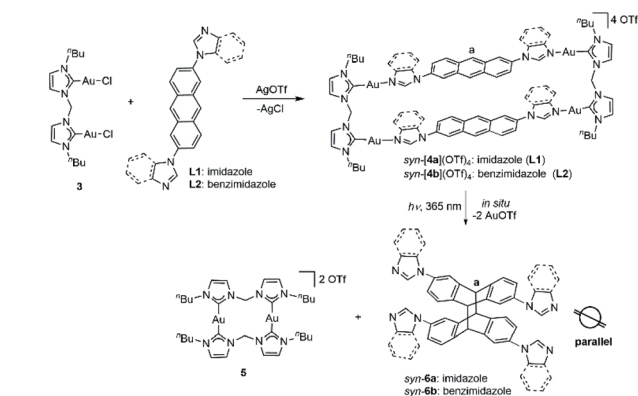


Fig. 4 Sections of the ¹H NMR spectra of (a) L1 in CDCl₃; (b and c) complex *syn*-[4a](OTf)₄ before and after irradiation in [D₆]DMSO; (d) complex 5 in [D₆]DMSO; (e) photodimer *syn*-6a in [D₆]DMSO. (f) Energy diagram with DFT calculated structures of *syn*-[4a]⁴⁺ (left), *anti*-[4a]⁴⁺ (right). The energies of the respective metallarectangles were calculated according to equations described in the ESI† (N, blue; C, grey; Au, yellow).

[4a](OTf)₄ failed, DFT calculations were performed in order to determine the relative energies of *syn*-[4a]⁴⁺ and *anti*-[4a]⁴⁺ regioisomers. These calculations suggested that the *syn*-[4a]⁴⁺ rectangle is the thermodynamically favored species by −1.6 kcal/mol^{−1} (Fig. 4f). A rotation of the anthracenes and thus the isomerization *syn*-[4a]⁴⁺ to *anti*-[4a]⁴⁺ is energetically unfavorable.

Next metallarectangle *syn*-[4a](OTf)₄ dissolved in [D₆]DMSO (*c* = 5.6 × 10^{−3} M) and irradiated (λ = 365 nm) in order to investigate the [4 + 4] cycloaddition of this derivative. Monitoring by ¹H NMR spectroscopy revealed the complete disappearance of the signal assigned to proton H_a of *syn*-[4a](OTf)₄ at δ = 7.81 ppm (Fig. 4b) after 3 hours of irradiation. Over this period a new singlet signal at δ = 5.01 ppm appeared, which can be assigned to the H_a resonance of the newly formed photodimer product *syn*-6a (Fig. 4c). In line with our previously reported results,^{21b} the purely organic compound *syn*-6a was formed under extrusion of the known dinuclear gold(i) species 5 (Fig. 4d and e). The organic photodimer *syn*-6a was identified by NMR spectroscopy and ESI-MS analysis (Fig. S61 and S62†).

Single crystals of *syn*-6a were obtained by slow diffusion of ethyl ether into a solution of the compound in a solvent mixture of dichloromethane and methanol at ambient temperature in 85% yield. The molecular structure and the *syn*-conformation of the photodimer was unambiguously determined by X-ray crystallography (Fig. 5a and b). Importantly, the conformation of *syn*-6a is consistent with the conformation that was calculated to be the thermodynamically favored tetranuclear complex cation *syn*-[4a]⁴⁺ (Fig. 4f). In addition, a tetranuclear metallarectangle was constructed from 2 equiv. each of L2 and 3 (Scheme 2). This self-assembly yielded complex *syn*-[4b](OTf)₄ which upon irradiation gave the photodimer *syn*-6b (Fig. 5c and d).



Scheme 2 Synthesis of metallarectangles *syn*-[4a](OTf)₄ and *syn*-[4b](OTf)₄ and their [4 + 4] photocycloaddition to afford photodimers *syn*-6a and *syn*-6b.

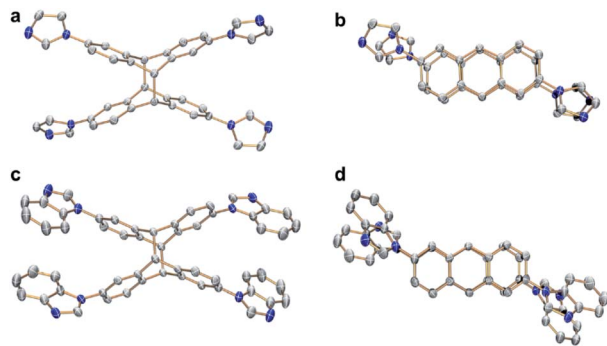


Fig. 5 (a) Side and (b) top views of *syn*-6a as determined by single-crystal X-ray diffraction. (c) Side and (d) top views of *syn*-6b (N, blue; C, grey). Hydrogen atoms have been omitted for clarity.

The tetrakisimidazole compounds *syn*-6a and *syn*-6b can be further functionalized at the imidazole moiety. Tetrakis-*N*-alkylation, for example, would lead to new tetrakisimidazolium salts which could be used for the preparation of unprecedented tetra-NHC complexes. We therefore investigated the preparation of a disilver tetracarbenic complex from *syn*-6a, a complex which would be attainable only with immense difficulties by organic chemistry procedures.

Tetrakis-*N*-alkylation of the metal-free compound *syn*-6a readily afforded the tetrakisimidazolium NHC-precursor *syn*-H₄-7a(PF₆)₄ with retention of the *syn*-conformation as confirmed by X-ray diffraction (Fig. 6a) and ¹H NMR

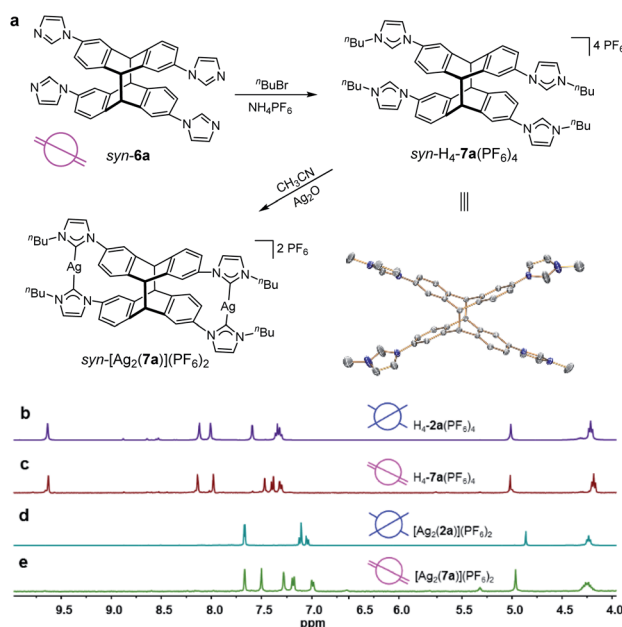


Fig. 6 (a) Synthesis of *syn*-H₄-7a(PF₆)₄ and of tetra-NHC complex *syn*-[Ag₂(7a)](PF₆)₂ and molecular structure of the tetracation *syn*-[H₄-7a]⁴⁺ (N, blue; C, grey). Hydrogen atoms have been omitted for clarity and only the first atom of each *N*-substituent is depicted. Sections of the ¹H NMR spectra of (b) tetrakisimidazolium salt *anti*-H₄-2a(PF₆)₄; (c) tetrakisimidazolium salt *syn*-H₄-7a(PF₆)₄; (d) complex *anti*-[Ag₂(2a)](PF₆)₂; (e) complex *syn*-[Ag₂(7a)](PF₆)₂ (all spectra were recorded in [D₆]DMSO).

spectroscopy (Fig. 6c). The photodimer *syn*-6a exhibited high thermal stability during the *N*-alkylation which proceeded at 110 °C for 36 h. The subsequent reaction of *syn*-H₄-7a(PF₆)₄ with Ag₂O led straightforward to the formation of the dinuclear tetracarbenic silver complex *syn*-[Ag₂(7a)](PF₆)₂ in high yield. This complex was fully characterized by ¹H NMR spectroscopy showing the disappearance of the resonance for the imidazolium C2-H protons (Fig. 6e) and by ¹³C and ESI-MS analysis (Fig. S75–S80†).

In addition, Fig. 6b depicts the ¹H NMR spectra of tetrakisimidazolium salts *anti*-H₄-2a(PF₆)₄ which was obtained by demetallation of complex *anti*-[Ag₂(2a)](PF₆)₂ (Fig. 6d) both featuring the *anti*-arrangement of the anthracene groups. Comparison of the ¹H NMR spectra of the *syn*-derivatives *syn*-H₄-7a(PF₆)₄ and *syn*-[Ag₂(7a)](PF₆)₂ to those of the *anti*-derivatives *anti*-H₄-2a(PF₆)₄ and *anti*-[Ag₂(2a)](PF₆)₂ reveals significant differences both for the tetrakisimidazolium salts as well as for the dinuclear silver complexes. These differences between the *syn*- and *anti*-photodimers may constitute a useful feature for the identification of stereoisomers in photochemical [4 + 4] dimerization of anthracene derivatives.

Conclusions

In summary, we have developed a supramolecular approach that enables the photochemical [4 + 4] cycloaddition of 2,6-difunctionalized anthracene derivatives with excellent regio- and stereoselectivity based on the preorganization of the reactants in metallosupramolecular assemblies. By using different metal templates for the construction of the assemblies, the spatial arrangement and orientation of anthracene units (*anti*-parallel or *parallel*) can be controlled, allowing subsequently the exclusive generation of either the *anti*- or *syn*-photodimer isomers. In addition, DFT calculations indicated that the geometric constraints imposed by the metal template and ligand components act synergistically to afford the specific isomer. Thus, our results provide promising approaches for accessing functional organic molecules with interesting and useful characteristics through the modification of supramolecular assemblies. Given the simplicity and high efficiency of the described strategy, we envisage this paradigm system could be considered as a key feature for new photo-induced material designs with different applications.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

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