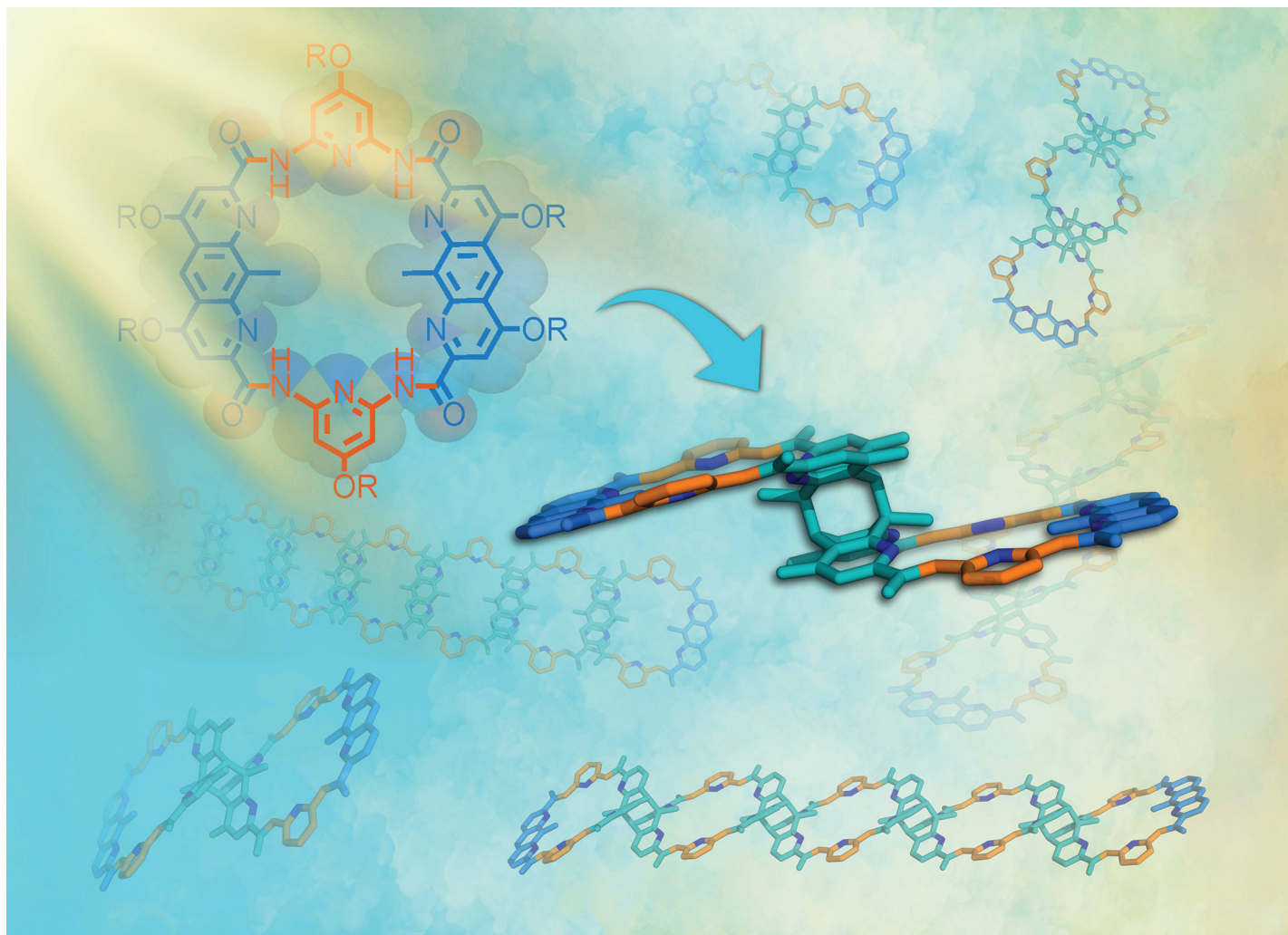




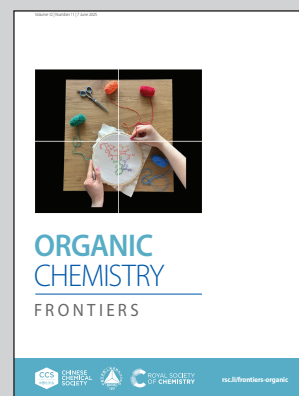
Showcasing research from Professor Bappaditya Gole's laboratory, Department of Chemistry, Shiv Nadar Institution of Eminence, Delhi NCR, India.

Unveiling stereoselective ladders *via* photo-oligomerization of a diazaanthracene macrocycle

Stereoselective oligomers are synthesized by leveraging steric interactions at remote sites from a photoresponsive macrocycle *via* [4 + 4] cycloaddition. Oligomers up to octamer were isolated and characterized. They retain reactivity to undergo further oligomerization, either independently or with fresh monomers.

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Unveiling stereoselective ladders *via* photo-oligomerization of a diazaanthracene macrocycle†Jinti Moni Kumar, ^a Ivan Huc, ^b Yann Ferrand ^c and Bappaditya Gole *^a

We present an efficient light-driven oligomerization of a flat aromatic tetra-amide macrocycle containing photoresponsive 1,8-diazaanthracenes. Oligomers with molecular weights exceeding 10 kDa were straightforwardly obtained, isolated using gel permeation chromatography (GPC), and characterized by NMR and mass spectrometry up to an octamer. We finely tuned the side chain size to exclusively favor the *exo*-[4 + 4] photoadduct over the *endo*-[4 + 4] photoadduct, thus controlling the stereoselectivity of the photoreaction to produce a single isomer, which yielded ladder-like architectures. The octamer is a ~10 nm long ladder with a step size of 4.6 Å. The oligomers retained sufficient photoreactivity to undergo further oligomerization, either between themselves or upon adding fresh monomers, to generate longer oligomers. These oligomers are fully degradable as thermal reversibility allows for monomer recovery.

Introduction

The emergence of covalent organic frameworks (COFs),^{1–7} covalent organic cages,^{8–14} and other carbon-rich materials^{15–24} with fascinating properties has inspired chemists to explore simple synthetic methods useful for accessing such materials, starting with monomers encoded with functional information. Light-driven reactions, particularly polymerization, have proven to be efficient, mostly clean, environmentally friendly methods for producing functional materials.^{25–28} However, in homogeneous solutions, the persistent challenge of achieving stereoselectivity and specificity continues to exist, primarily due to the difficulty of controlling the reactivity of the excited state, thus leading to products that are challenging to distinguish and separate. This issue was partially addressed by employing noncovalent supramolecular interactions and performing photochemical reactions in the solid state, taking benefit of their preorganization.^{29–35} For example, the [4 + 4] photocycloaddition of anthracene derivatives, often exploited in supramolecular chemistry to create functional materials, is known to generate multiple regio- and stereoisomers in solution.^{36–43} The 9- and 10-substituted

anthracenes give two stereoisomers, *syn* and *anti*, of different proportions depending on the substitution. The number of isomers increases if the substitution is present in any other position.⁴⁴ However, recently, several anthracene-based macrocycles and cages were considered as starting building blocks for the preparation of highly ordered, stereoselective two-dimensional porous polymers and membranes in a single-crystal-to-single-crystal transformation.^{45–49} These studies primarily leverage monomer preorganization in a crystal or on a surface. Yet, attempts to achieve similar outcomes in solution have rarely been successful. A recent study employed time-resolved NMR spectroscopy to gain comprehensive insights into the polymerization of bis-anthracene monomers in solution but did not focus on isolating the resulting materials.⁵⁰ Control over the reactivity of the monomer would not only offer a method to regio- and stereoselectively produce polymers in solution without relying on the crystallization of the monomer in a preorganized fashion, but it would also provide opportunities for mechanistic studies by isolating intermediate oligomers, which form at the early stage of the reaction. Furthermore, oligomers may offer functional properties akin to those of polymers while offering better solubility, making them suitable for various solution-based techniques.^{51–53}

Herein, we explore the photochemical reaction of planar aromatic tetra-amide macrocycle **1** containing 1,8-diazaanthracene units in solution. Unlike anthracene, which typically yields mixtures,^{36–43} 1,8-diazaanthracenes (*i.e.*, pyrido-[3,2-*g*] quinolines) are known to produce exclusively the antiparallel photoproduct in solution to minimize dipolar repulsion (Fig. 1a) unless some constraints are applied to maintain the parallel arrangement of the reacting diazaanthracenes. In this case, the parallel photoproduct is quantitatively formed

^aBiomimetic Supramolecular Chemistry Laboratory, Department of Chemistry, School of Natural Sciences, Shiv Nadar Institution of Eminence Deemed to be University, Greater Noida, Uttar Pradesh 201314, India.

E-mail: bappaditya.gole@snu.edu.in

^bDepartment of Pharmacy, Ludwig-Maximilians-Universität München, Butenandtstr. 5–13, 81377 Munich, Germany

^cUniv. Bordeaux, CNRS, Bordeaux INP, CBMN, UMR 5248, F-33600 Pessac, France

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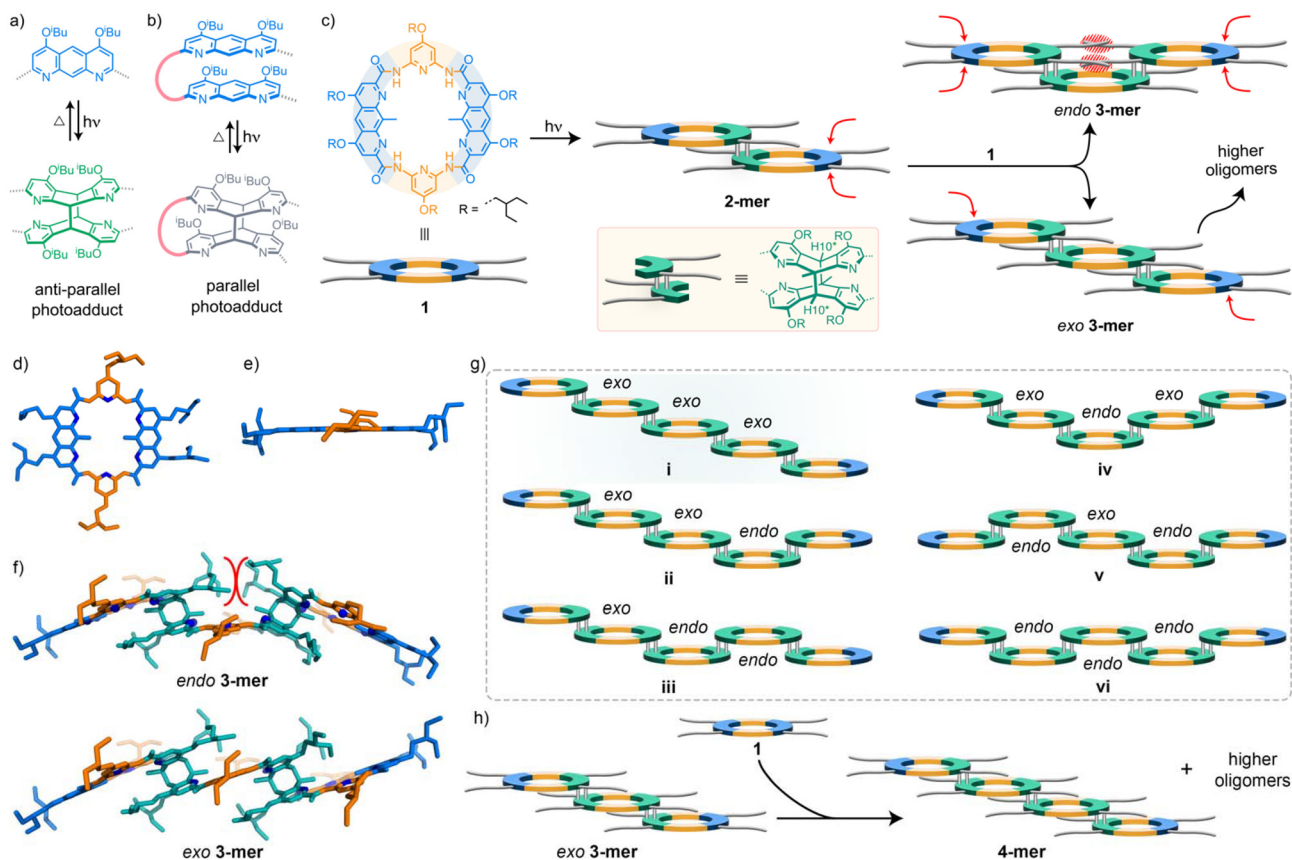


Fig. 1 Schematic illustration of the (a) intermolecular and (b) intramolecular [4 + 4] photocycloadditions of 1,8-diazaanthracene that give anti-parallel and parallel photoadducts, respectively. (c) Chemical structure of macrocycle **1** and the schematic representation of its intermolecular [4 + 4] cycloaddition. In principle, upon chain elongation, different stereoisomers may form depending on the relative orientation of the approaching monomers (red arrows). However, 2-ethyl butoxy solubilizing side chains have been chosen to produce the *exo*-isomer selectively. The red scribbles in the *endo* 3-mer highlight possible steric hindrance, which does not allow two monomers to approach a macrocycle from the same side. (d) Top and (e) side views of the geometry-optimized structure of **1** highlighting its planar structure. (f) Energy-minimized models (Merck Molecular Force Field static, MMFFs) of two possible stereoisomers, *endo* 3-mer and *exo* 3-mer. Steric hindrance is highlighted with red lines. (g) As a representative example, a 5-mer can have six possible stereoisomers: the all *exo*, i.e., isomer i, forms selectively due to the size of the side-chains. (h) Schematic illustration showing the chain growth mechanism, where a trimer, for example, can be photoirradiated in the presence of fresh monomers to produce higher oligomers.

(Fig. 1b).^{54–61} We show that macrocycle **1** undergoes intermolecular antiparallel [4 + 4] cycloaddition at both ends to create oligomers upon irradiation with UV light in chloroform (Fig. 1c). We have isolated oligomers of varying lengths with molecular weights of up to 10 kDa, allowing for their individual characterization. The photochemical reaction exclusively produced *exo*-[4 + 4] photoadducts and no *endo* isomer, thus providing a single isomer for each oligomer among multiple possibilities. Unlike conventional methods, which often rely on steric interactions near the reactive center to control, e.g., iso- or syndiotactic polymer architectures,^{62,63} the stereoselectivity in this system is governed by the manipulation of steric interactions at remote sites. The rigid structure of the macrocycle ensures that the resulting oligomers adopt a stiff, ladder-like architecture. Furthermore, we demonstrate that the shorter oligomers retain sufficient photoreactivity to undergo further oligomerization, either with themselves or with freshly added monomers, to generate longer oligomers (Fig. 1h).

Results and discussion

The shape-persistent tetra-amide macrocycle **1** was prepared from 1,8-diazaanthracene-2,7-dicarboxylic acid and 2,6-diaminopyridine derivatives in a single-step amide coupling reaction in 33% yield (Schemes S1–S3 and Fig. S1 and S2†). Geometry optimization by density functional theory (DFT) calculations revealed that **1** adopts a planar rigid structure suitable for photoproduct formation at the 1,8-diazaanthracene sites (Fig. 1d and e).

In addition to their role in solubilizing the macrocycles, we have identified the possibility for side chains in positions 4 and 5 of the diazaanthracene to control the stereoselectivity of the oligomers. For instance, when a monomer approaches a 2-mer, two stereoisomers may form: an *endo* 3-mer and an *exo* 3-mer (Fig. 1c). We reasoned that the formation of the *endo*-isomer could be prevented by using sufficiently hindering side chains that would cause steric clashes only in the *endo*



pathway, *i.e.* the transition state (TS) energy of the *endo* 3-mer is expected to be reasonably higher than that of the *exo* 3-mer, leading to the stereoselective formation of the *exo* isomer (Fig. S3†). The energy-minimized models of the two possible 3-mer stereoisomers bearing 2-ethyl butoxy side chains showed that the *exo* 3-mer is stabilized by 15 kJ mol⁻¹ compared to the *endo* 3-mer, which imposes a substantial strain on the central macrocycle to accommodate the bulky side chains (Fig. 1f). The energy-minimized structure showed that the strain on the central unit is reduced with smaller side chains, such as

methoxy or isobutoxy, which may alter the stereoselectivity (Fig. S3†). Furthermore, using smaller side chains would reduce the solubility drastically. We thus decided to use 2-ethyl butoxy side chains to both overcome solubility issues and achieve stereoselectivity. As illustrated in Fig. 1g, the 5-mer may exist as six possible stereoisomers. However, with our design strategy, only isomer **i** displaying four *exo*-bridges would be obtained. With increasing oligomer size, the number of possible stereoisomers increases; for example, it can be as high as 36 in the case of the octamer (Fig. S4†). Therefore,

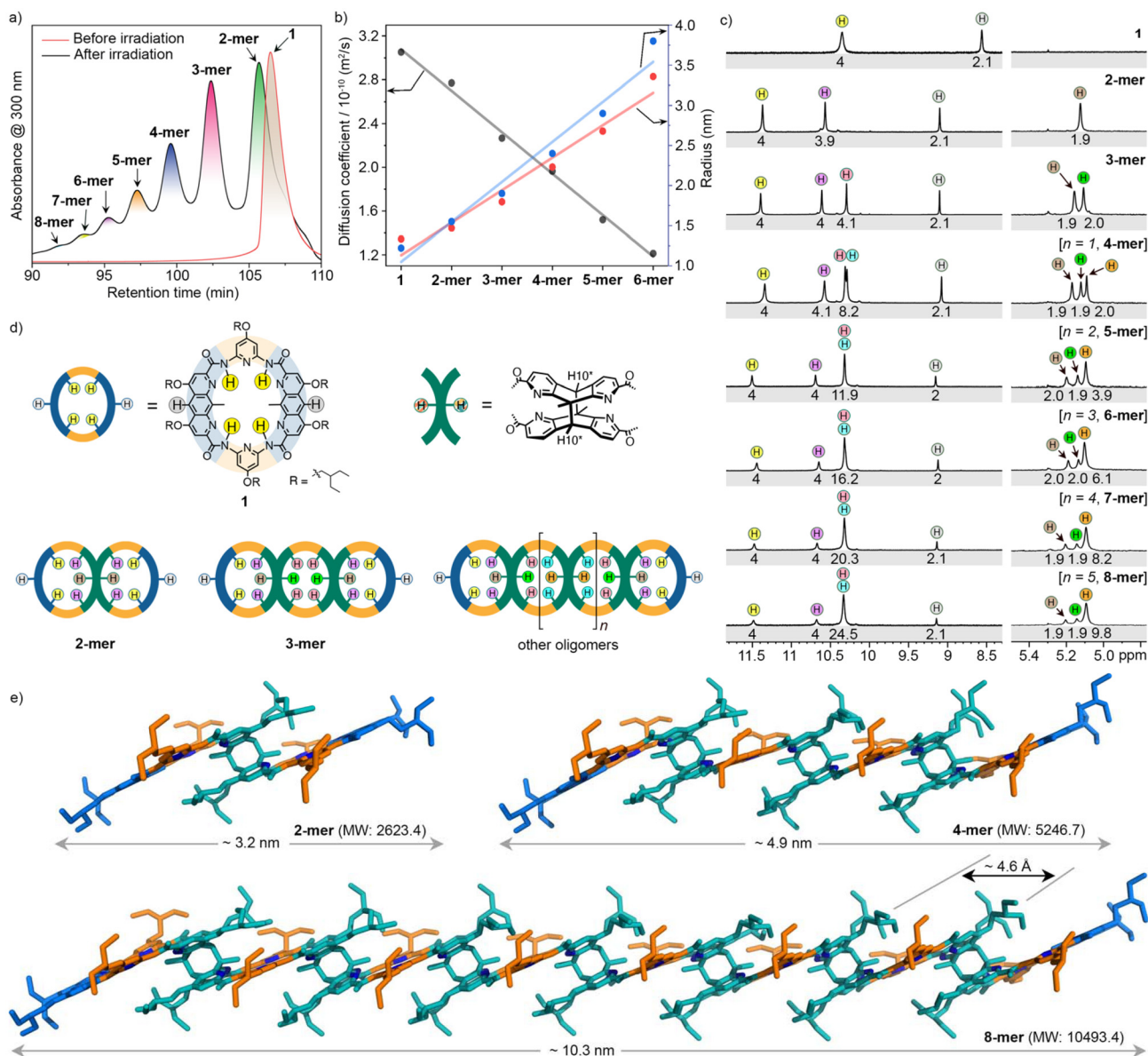


Fig. 2 (a) Recycling preparative GPC profiles after three cycles (CHCl₃, 25 °C, 7 mL min⁻¹) before and after photoirradiation of **1** in CHCl₃ showing a clear separation of oligomers of different sizes. (b) Diffusion coefficients (in grey) and corresponding solvodynamic radii (in red) of **1** and oligomers obtained from ¹H DOSY experiments. The calculated radius (in blue) of the geometry-optimized oligomers is given for comparison. (c) Selected areas of ¹H-NMR spectra (400 MHz, 298 K) in CDCl₃ of **1** and the oligomers with relative peak integration. The assigned protons are color-coded as in (d). 'n' represents the number of repeating units in the oligomers. n = 0 to 5 indicate oligomers 3-mer to 8-mer, respectively. (e) Side view of the energy-minimized molecular models of the 2-mer, 4-mer, and 8-mer using the MMFFs shown in tube representation. Hydrogen atoms are omitted for clarity. Length and molecular weight are indicated. Models for other oligomers are given in the ESI.†



creating only one isomer by simply adjusting side chains would be remarkable.

To assess the photo-oligomerization process, macrocycle **1** was irradiated with a broad wavelength range of 320–390 nm, matching well with its absorption maxima (Fig. S5†), for three hours under anaerobic conditions in chloroform (Scheme S4†). $^1\text{H-NMR}$ spectra were recorded at intervals, revealing significant changes compared to the initial spectrum and the emergence of several new signals (Fig. S6†). Notably, resonances in the range of 5.0–5.4 ppm, attributed to H10* (Fig. 2c), indicated the progress of $[4 + 4]$ cycloaddition.^{59–61} MALDI-TOF mass spectrometry confirmed the formation of higher oligomers (Fig. S7†), which were separated by recycling GPC (Fig. S8†). After three cycles, peaks all the way to the **8-mer** can be distinguished (Fig. 2a). The gradual decrease of peak intensity with increasing size suggests a lower reactivity as the oligomer size increases. One can observe that longer irradiation times lead to an increasing proportion of higher oligomers (Fig. S9†) but may cause thermal decomposition due to sample heating in our setup. However, this issue could be resolved by irradiating at lower temperatures. Each compound from the **2-mer** to the **8-mer** was isolated and identified using $^1\text{H-NMR}$ and mass spectrometry (Fig. S10–S23†).

All oligomeric species were found to be soluble in chloroform. $^1\text{H-DOSY}$ (diffusion ordered spectroscopy) NMR analysis (Fig. S24–S29†) of **1** and the **2–6-mers** showed that the oligomer diffusion coefficients decreased linearly with increasing their size (Fig. 2b). Using the Stokes–Einstein equation, we estimated their hydrodynamic radii to be 1.3, 1.5, 1.8, 2.1, 2.7, and 3.4 nm, respectively. This trend is in agreement with the theoretical radius of the oligomers (Fig. 2b). The well-resolved $^1\text{H-NMR}$ resonances in CDCl_3 (Fig. 2c) were consistent with the structures assigned based on mass spectrometry. In contrast to monomeric macrocycle **1**, which displays a single amide resonance, the **2-mer** showed two amide signals at 11.35 and 10.55 ppm, corresponding to terminal and central amide protons, respectively. Interestingly, the spectrum of the **3-mer** showed only one set of sharp signals, suggesting that only one of the two possible isomers had formed. For all the longer oligomers, three types of NH amide signals with varying intensities were observed. Based on symmetry, these signals can be assigned to terminal NHs (yellow, Fig. 2c and d), those neighboring the terminal NHs (magenta), and all the remaining central amides (pink and cyan). For the **4-mer**, two types of central NHs can be distinguished (pink and cyan), attesting to the sensitivity of NMR in distinguishing small differences in the environment. We thus infer that the single set of sharp resonances for all compounds reflects the presence of only one isomer, *i.e.*, other isomers would give rise to some distinct resonances should they be present. The chemical shift values of the amide signals remain relatively constant for the higher oligomers. Only the relative intensity of the central amide signal varies. This indicates that central amides are all in a similar environment, consistent with the oligomers remaining in a rigid structure. The signals in the 5–5.4 ppm region correspond to the H10* protons of diazaanthracenes

after the $[4 + 4]$ cycloaddition. The peak integration of each signal allowed us to confirm the oligomer length using the terminal resonance as a reference (Fig. 2c, yellow).

Since growing single crystals suitable for the X-ray analysis of the oligomers has so far eluded us, we used molecular modeling to gain structural insights (Fig. 2e, and Fig. S30–S36†). The models revealed rigid, ladder-shaped structures with an average plane-to-plane separation (*i.e.* step size) of 4.6 Å (Fig. S37†) and a length of up to 10 nm for the **8-mer**. The molecular lengths of the oligomers predicted by these models closely match the hydrodynamic radii derived from the DOSY experiments (Fig. 2b).

The oligomers are thermally quite stable up to 353 K with no noticeable changes over three days. However, beyond this temperature, they started decomposing into smaller oligomers.

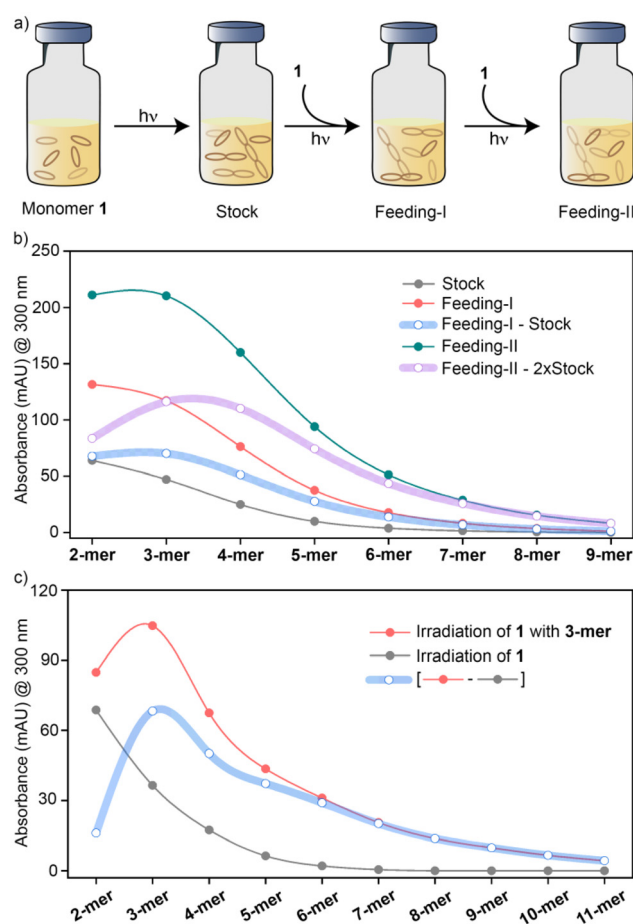


Fig. 3 (a) Schematic representation of multicycle oligomerization. Monomer **1** was initially irradiated to obtain a mixture of oligomers. Then, fresh monomers were added to obtain higher oligomers and enhance their proportions. (b) The plot summarizes the evolution of oligomers after every feeding step. The proportion of the oligomers is normalized against the initial irradiation step to indicate a significant enhancement in the oligomer concentration. (c) Evolution of oligomers when irradiation of monomers was performed in the presence of a **3-mer**. The normalized data with respect to the oligomers formed in the absence of the **3-mer** showed a significant role in forming higher oligomers. The lines connecting points are for visual guidance.



To achieve complete thermal reversibility, the **4-mer** was heated at 393 K in 1,1,2,2-tetrachloroethane over a period of 60 h (Fig. S38†). ¹H NMR analyses performed at intervals revealed the transient appearance of the **2-mer** and **3-mer**, thus hinting at a gradual degradation of the **4-mer** (Fig. S39†). Attempts to reverse the cycloaddition using light irradiation at 254 nm were ineffective.

To assess how a given oligomer could be further elongated and to decipher the mechanism of the formation of long oligomers, we conducted feeding experiments in which fresh macrocycle **1** was added at intervals during the reaction. Initially, **1** was irradiated in chloroform for 1 h to generate a mixture of oligomers. Subsequently, two feeding cycles were performed with the addition of fresh **1**, followed by irradiation. The evolution of the oligomer formation was monitored by GPC after each cycle (Fig. S40 and S41†), showing a significant increase in the proportion of higher oligomers (Fig. 3b), despite **1** being capable of independent oligomerization in every feeding cycle. The normalized data indicate the influence of the parent oligomers, *i.e.*, the oligomers present before fresh macrocycle monomers were added, in promoting the formation of higher oligomers. This is consistent with the oligomers reacting slowly with each other while they can still react with the monomer and undergo elongation. Consistently, we observed that performing the photo oligomerization of **1** in the presence of the **3-mer** facilitated the formation of higher oligomers, such as a **10-mer** and **11-mer** (Fig. S42 & S43†), which could not be achieved by irradiating **1** alone in the same timeframe. Moreover, we found that when we directly irradiated a preformed **2-mer** over one hour in chloroform, oligomers of the **2-mer**, such as the **4-mer**, **6-mer**, and **8-mer**, were formed in higher proportion, but odd-numbered oligomers also formed (Fig. S44–S46†), likely due to thermal reversibility caused by heat due to the irradiation. These experiments demonstrate that the oligomers are sufficiently photoreactive to undergo further oligomerization in the presence of monomers.

Conclusions

In conclusion, we have used a simple [4 + 4] cycloaddition reaction to photo-oligomerize a planar macrocycle containing light-sensitive diazaanthracene moieties. By properly choosing the solubilizing side chains, we were able to control the stereoselectivity to obtain discrete ladder-like oligomers out of many other possible isomers. Oligomers up to the **8-mer** were easily isolated by GPC and characterized by NMR and mass spectrometry. The ability to obtain large oligomers with molecular weights over 10 kDa in a single-step photoradiation is rare. We demonstrated that these oligomers remained sufficiently reactive to further react with fresh monomers to form even longer oligomers. The oligomers also exhibit thermal reversibility, returning to monomers along the same pathway. This approach holds promise for synthesizing other functional oligomers from different photoresponsive monomers.

Author contributions

B. G., Y. F., and I. H. conceived the project. J. M. K. planned and conducted the experiments. B. G. supervised the research, performed analysis, and wrote the original draft of the manuscript. Y. F and I. H. provided critical feedback during the whole process and edited the manuscript.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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