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Photochemical Unmasking of Polyyne Rotaxanes

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ABSTRACT: Bulky photo-labile masked alkyne equivalents (MAEs) are needed for the synthesis of polyyne polyrotaxanes, as insulated molecular wires and as stabilized forms of the linear polymeric allotrope of carbon, carbyne. We have synthesized a novel MAE based on phenanthrene and compared it with an indane-based MAE. Photochemical unmasking of model compounds was studied at different wavelengths (250 nm and 350 nm), and key products were identified by NMR spectroscopy and X-ray crystal-lography. UV irradiation at 250 nm leads to unmasking of both MAEs. Irradiation of the phenanthrene system at 350 nm results in quantitative dimerization *via* [2+2] cycloaddition to form a [3]-ladderane; irradiation of this ladderane at 250 nm generates a dihydrotriphenylene, which can be oxidized easily to a triphenylene. Irradiation of the indane-based MAE at 350 nm in the presence of traces of oxygen forms an endoperoxide and a bisepoxide. Both MAEs have been incorporated into rotaxanes *via* copper-mediated active metal template Glaser or Cadiot-Chodkiewicz coupling. The identity of the rotaxanes was confirmed by NMR spectroscopy and mass spectrometry. The phenanthrene rotaxane decomposes during attempted photochemical unmasking, whereas photolysis of the indane rotaxane results in unmasking of the polyyne thread to form a rotaxane with a chain of 16 *sp*-hybridized carbon atoms. This approach opens avenues towards the synthesis of encapsulated carbon allotropes.

INTRODUCTION

The predicted electronic and mechanical properties of carbyne, the linear *sp*-hybridized allotrope of carbon,^{1,2} have been debated for many years,^{3,4} providing a motivation for synthesizing this elusive carbon polymer. However, if carbyne could be prepared as an undiluted carbon allotrope, it is expected to be unstable and explosive, because acetylenic chains easily react with each other to form cross-linked materials,⁵⁻⁷ in a process that is highly exothermic.8 End-capped oligomeric analogues of carbyne, known as polyynes, have been prepared with chains of up to 44 sp-hybridized carbon atoms, stabilized by bulky terminal groups.^{9,10} This strategy is unlikely to be effective for stabilizing longer polyynes, because the influence of the end-groups diminishes as the chain gets longer. Polyynes have previously been stabilized by rotaxane formation,^{11,12} helical wrapping¹³ and co-crystallization with metal complexes.¹⁴ Very long polyynes, consisting of more than 6,000 carbon atoms, have been created and studied inside carbon nanotubes.¹⁵ However, this type of encapsulation largely screens the properties of the guest; for example, it is difficult to measure the charge transport, optical or mechanical properties of a polyvne inside a carbon nanotube, because all these aspects are dominated by the host nanotube. The most versatile strategy for creating stable forms of carbyne is probably to synthesize polyrotaxanes in which the carbon chain is threaded through many macrocycles to form an 'insulated molecular wire'.¹⁶ Threading also offers a strategy to stabilize the cyclic analogues of carbyne, cvclo[n] carbons,¹ by catenane formation, which would not be possible using encapsulation inside carbon nanotubes. Here we present an approach to the synthesis of polyyne rotaxanes using photo-labile masking groups. This chemistry opens an avenue towards carbyne polyrotaxanes and cyclocarbon catenanes.

Polyyne rotaxanes have been prepared previously using active metal templates.^{12,17-20} This strategy enables a single mac-

access to polyyne polyrotaxanes with many threaded macrocycles, unless temporary alkyne-masking groups can be used to prevent unthreading. Various masked alkyne equivalents (MAEs) have been developed, particularly for the synthesis of cvclo[n] carbons. MAEs are molecular subunits that can be transformed into alkynes by activation with light,²¹⁻²³ heat^{24,25} or chemical reagents²⁶⁻²⁸ (Scheme 1). Here, we explore the use of MAEs in the synthesis of polyyne rotaxanes. Our choice of MAEs is governed by the following factors: (i) the MAE must be chemically stable under the conditions of rotaxane formation; (ii) it should be bulky enough to act as a temporary stopper and prevent unthreading of the rotaxane; (iii) unmasking of the MAE must be possible in interlocked systems, and (iv) the MAE should be easy to synthesize. Photo-labile MAEs are appealing because they are expected to be stable under the conditions of rotaxane formation. Here, we investigate a new photo-labile MAE, based on the susceptibility of phenanthrene to undergo reversible photochemical [2+2]-cycloadditions to the double bond of the central ring (9,10-position). We compare this new MAE with an indane-based MAE developed by Tobe and coworkers.²¹ The photochemistry of both MAEs was investigated and some unexpected products were identified. Polyyne rotaxanes containing both types of MAEs were synthesized, but only the rotaxane with indane-based MAEs underwent clean photochemical unmasking.

rocycle to be threaded round a polyyne, but it does not provide

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Scheme 1. Examples of Masked Alkyne Equivalents.



RESULTS & DISCUSSION

Synthesis and testing of MAEs

The dichlorocyclobutene **1** is readily prepared from phenanthroline by photochemical addition of trichloroethylene followed by elimination of hydrogen chloride, as reported by Hacker and McOmie (Figure 1).²⁹ Sonogashira coupling was used to prepare the TIPS-protected model compound **2**. The crystal structure of **2** confirms the structure and gives an indication of its steric bulk (Figure 1c).³⁰ We compared this phenanthrene-based MAE with an indane-based MAE developed by Tobe and coworkers.²¹ A drawback of the indane group is its longer synthesis: indane dichloride **3** is prepared in eight steps from cyclopentanone (Figure 1b).^{21d} The TIPS-protected model compound **4** was prepared for comparison with **2**.

Irradiation of compound **2** as a solution in CDCl₃ (0.2 M concentration) with UV light (wavelength 250 nm) for 21 h leads to formation of phenanthrene in 80% yield, as illustrated by the ¹H NMR spectra in Figure 2. TIPS-triyne **5** is also formed in this reaction (Figure 3), although in poor yield (20% by NMR analysis) together with a complex mixture of products. A reference experiment showed that TIPS-triyne **5** is stable under the conditions of this photolysis, implying that the side products come from competing side reactions of **2**, rather than from decomposition of triyne **5**.

We also tested irradiation of **2** with UV light at longer wavelengths. Surprisingly, irradiation at 350 nm gave no phenanthrene; instead the [2+2] dimer, [3]-ladderane **6**, is formed quantitatively as a single diastereomer (as seen from the ¹H NMR spectra in Figure 4 and X-ray crystal structure in Figure 3b).³⁰ The geometry of **6** is similar to those of previously reported ladderanes,³¹ but steric congestion results in an unusually long bond length for the C-C bonds shared between two cyclobutane units (bond length 1.604(2) Å; see Supporting Information, Section S6 for a comparison of bond lengths from CSD and Section S7 for computational studies).



Figure 1. (a,b) Synthesis of MAEs models 2 and 4; (c) two orthogonal views of the structure of compound 2 in the crystal (thermal ellipsoids plotted at the 40% level).



Figure 2. ¹H NMR spectra (CDCl₃, 298 K, 400 MHz) of a solution of **2** (red peaks) before and after irradiation with UV light (250 nm, no purification). Phenanthrene (blue peaks) is the major product (80%) after irradiation for 21 h. Triyne **5** could be detected by ¹H and ¹³C NMR (yield 20% by ¹H NMR).

The unexpected outcome of the irradiation of **2** at 350 nm led to us to question whether [3]-ladderane **6** might be an intermediate in the cycloreversion observed on irradiation of **2** at 250 nm. This can be ruled out as dimer **6** does not yield the unmasked triyne **5** when irradiated with 250 nm UV light; instead it generates a different product **7** (50% yield) with only one phenanthrene unit expelled (Figure 3a). The formation of different products on irradiation at 250 and 350 nm must reflect the reactivity of different excited states. Compound **7** is also formed as a minor product (~10%) when **2** is irradiated with 250 nm UV light (peak at 3.77 ppm in Figure 2).

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Figure 3. Reactivity of phenanthrene MAE model 2. a) Irradiation of 2 at 250 nm or 350 nm, and irradiation of dimer 6 at 250 nm, showing a possible mechanism for the formation of dihydrotriphenylene 7 (R = TIPS). b) Two orthogonal views of the structure of 6 in the crystal. c) Structures of dihydrotriphenylene 7 and triphenylene 8. Thermal ellipsoids plotted at the 40% level for (b) and (c).



Figure 4. ¹H NMR spectra (C_6D_6 , 298 K, 400 MHz) of a solution of **2** (red peaks) before and after irradiation with UV light (350 nm, no purification). A [3]-ladderane **6** (green peaks) is formed quantitatively after irradiation for 14 h. (The outcome of photolysis in CDCl₃ is the same.)

The constitution of 7 was deduced from the fact that it is formed from 6 concomitantly with one equivalent of phenanthrene, and from the integration of the ¹H NMR signals. The crystal structure revealed that 7 has a surprising stereochemistry with a central *trans*-dihydrotriphenylene.³⁰ This suggests that a photochemical domino reaction takes place after phenanthrene is extruded by a [2+2]-cycloreversion to form intermediate A (Figure 3a). This cyclobutene appears to undergo a disrotatory 4π electrocyclic ring opening (photochemically allowed) to give a cyclohexadiene **B**, which then undergoes a conrotatory 6π electrocyclic ring opening (thermally allowed) to form a [10]annulene intermediate C, which recloses via a disrotatory 6π electrocyclic ring closure to yield 7. Alternatively, the central cyclobutane ring in A may undergo a photochemical [2+2]-cycloreversion to generate C in one step. A similar cycloreversion was observed in the formation of [12]annulene from a ladderane.³²

Intermediates A-C have not been observed but these mechanisms appear to be the most likely explanation for the generation of 7 from 6. The crystal structure has four molecules of 7 in the asymmetric unit, all with essentially the same molecular geometry: a representative molecule is shown in Figure 3c.

When exposed to air, dihydrotriphenylene 7 is slowly oxidized to triphenylene 8 in the course of weeks. This process can be accelerated by stirring a CH_2Cl_2 solution under an oxygen atmosphere for five days to achieve full oxidation. The structure of 8 was confirmed by NMR, MS and X-ray crystallography (Figure 3c).³⁰

Indane derivative 4 is a known compound,^{21d} but its photochemistry has not been reported. We found that irradiation with 250 nm UV light leads to rapid formation of indane and triyne 5 (both in 70% yield; Figure 5 and 6). Competition experiments show that 4 reacts about five times faster than 2 under identical photolysis conditions.



Figure 5. ¹H NMR spectra (CDCl₃, 298 K, 400 MHz) of a solution of **4** (red peaks) before and after irradiation with UV light (250 nm, no purification). Indane (blue peaks) is formed in 70% yield after irradiation for 1.3 h. The triyne **5** (formed in 70% yield) could be detected by ¹³C NMR.



Figure 6. a) Reactivity of indane MAE model **4**. b) Two orthogonal views of the structure of **10** in the crystal (thermal ellipsoids plotted at the 40% level).

In contrast to the clean photochemistry with 250 nm light, irradiation of indane 4 at 350 nm gives a complex mixture of products (Figures 6 and 7). Initially, a new set of 1 H resonances appears at 6.5 ppm and 4.6 ppm (green peaks

in Figure 7); this set of peaks disappears to be replaced by another set of peaks at 3.6 ppm and 3.2 ppm (purple peaks in Figure 7). Formation of indane is also observed, indicating that unmasking occurs to some extent as well. The two products, 9 and 10, were isolated and characterized (isolated yields of 9 and 10 after 3.5 h of irradiation 19% and 28%, respectively; 55% of 10 after 8 h of irradiation: ¹H NMR spectra shown in Figure 7b). The crystallographic structure determination of epoxide 10 (Figure 6b)³⁰ enabled us to deduce the structure of peroxide 9. The reactions are summarized in Figure 6a. Presumably, traces of oxygen are converted to singlet oxygen and undergo hetero-Diels-Alder reaction³³ with the diene moiety of 4. The formation of bis-epoxides from unsaturated endoperoxides has been reported previously.³⁴ As expected, products 9 and 10 are not formed when rigorously O₂-free solutions of 4 are irradiated. Tobe et al. reported the formation of a cyclooctatetraene and a semibullvalene (each in 7% yield, using a low-pressure mercury lamp in hexane) when irradiating TMSand TBDMS-analogues of 4^{21d} We also observed formation of these products, each in about 10% yield, when irradiating with 250 nm light, identified by their characteristic ¹H NMR spectra.

The crystal structure of **10** (Figure 6b) shows that the sixmembered cyclohexane-bisepoxide ring is remarkably flat (root-mean-squared deviation for 6 carbon atoms: 0.004 Å); this conformation has been reported previously in similar cyclohexane derivatives.³⁵ Comparing the crystal structures of **2** and **10** reveals that the geometry of the enediyne unit is unaffected by the decoration of the other side of the cyclobutene ring; bond lengths and bonding angles are essentially identical.

In summary, both the indane and the phenanthrene group are viable MAEs for masking polyynes. Unmasking is quicker and more efficient for the indane but the phenanthrene MAE is more accessible. The alkyne masked with indane 4 gives triyne 5 in 70% yield; the one masked with phenanthrene 2 gives 5 in 20% yield.

Rotaxane synthesis

With these encouraging results in hand, we synthesized interlocked structures based on the phenanthrene and indane MAEs. Bulky trityl stoppers were added to ensure that the macrocycle cannot slip off after a rotaxane is formed.³⁶ The synthesis of a phenanthrene rotaxane **16** and the corresponding reference thread **17** are shown in Schemes 2 and 3.

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Figure 7. a) ¹H NMR spectra (CDCl₃, 298 K, 400 MHz) of a solution of **4** (red peaks) before and after irradiation with UV light over 8 h. (350 nm, no purification). In addition to indane (blue peaks), two new species are formed (green and purple). b) ¹H NMR spectra (CDCl₃, 298 K, 400 MHz) of the purified compounds; the green peaks are assigned to peroxide **9** and purple peaks belong to bisepoxide **10**.

Scheme 2. Synthesis of phenanthrene-masked rotaxane 16.

Scheme 3. Synthesis of phenanthrene reference thread 17.



Synthesis of rotaxane 16 starts with desymmetrization of dichloride 1 *via* a Sonogashira coupling to TMS-acetylene to give chloride 13. In a subsequent Sonogashira coupling, diyne stopper 12 was installed to give stoppered masked phenanthrene 14 (Ar = 4-methylphenyl).

Stopper 12 was synthesized in one step from alcohol 11.³⁶ We found that it is necessary to separate the trityl unit and the MAE by two acetylene units to prevent a steric clash of the stoppers in the rotaxane forming step. A design based on a single acetylene spacer between stopper and MAE failed to give interlocked products (SI, Scheme S5).

After TMS-deprotection of 14 with $K_2CO_3/MeOH$, the phenanthrene rotaxane 16 was formed in 56% yield *via* coppermediated oxidative homocoupling using macrocycle 15.³⁷



The synthesis of indane rotaxane 27 proved more challenging than anticipated. Sonogashira couplings using TMSacetylene instead of TIPS-acetylene are less efficient on dichloride 3 (53% yield for TMS-acetylene, 100% for TIPSacetylene). Therefore, initial attempts focused on TIPS protected derivatives. We were not able to couple stopper 12 to the indane 18 (Scheme 4) and TIPS indane stopper 21 could only be obtained by Cadiot-Chodkiewicz coupling 12,38,39 to 20, the monodeprotected derivative of 4. Attempted deprotection of 21 to give 22 was not successful; a range of fluoride sources (TBAF, CsF, KF) led to rapid decomposition of 21 and analysis of the degradation products suggested that the trityl unit is cleaved from the molecule (Scheme S7). To overcome this problem, SiEt₃ (TES) was used as a protecting group, as it does not require fluoride for deprotection. Surprisingly, the Cadiot-Chodkiewicz coupling, which yielded the TIPS analogue 21, did not work on 23. It is puzzling that changing the silyl protecting group (for a smaller one) in a remote part of the molecule has such a drastic effect on reactivity (Scheme 4).

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Scheme 4. Unsuccessful approaches to the synthesis of compound 22.



The successful route to **24** proceeds *via* a low-yielding Sonogashira coupling (15%) to give **25** (Scheme 5). Subsequently a Sonogashira coupling gave **24** in 91% yield. In contrast to phenanthrene rotaxane **16**, we were unable to prepare indane rotaxane **27** *via* a copper-mediated oxidative homocoupling; only non-interlocked products were obtained. However, if indane stopper **24** is deprotected to **22** and converted to the corresponding alkynyl bromide **26**, rotaxane **27** can be formed in 47% yield (over three steps: TES-deprotection, bromination of half the material and Cadiot-Chodkiewicz coupling).^{12,39} Reference thread **29** was synthesized *via* a copper-catalyzed homocoupling (Scheme 6).

The ¹H NMR spectra of rotaxanes 16 and 27 are compared with those of the free threads 17 and 29, and macrocycle 15, in Figure 8. Compounds 16, 17, 27 and 29 exist as mixtures of meso and racemic diastereomers (SI, Scheme S8), causing splittings in the NMR resonances. The splittings in the signals of the phenanthroline macrocycle are more pronounced in 16, compared to 27; this can be attributed to the higher magnetic anisotropy and greater lateral extension of the phenanthrene group compared to the indane. In both rotaxanes, NOEs were observed between the aryl protons of the stopper (f and g in 16, e and f in 27) and proton A of macrocycle 15, indicating that the macrocycle can shuttle over the MAE units. The identity of rotaxanes was further confirmed by mass spectrometry; in contrast to the hydrocarbon precursors and dumbbells, all rotaxanes can be detected easily using ESI-MS (see SI for MS spectra).

Scheme 5. Synthesis of rotaxane 28.





Photochemical unmasking of polyyne rotaxanes

Scheme 6. Synthesis of dumbbell 30.



Figure 8. Comparison of the ¹H NMR spectra (CDCl₃, 298 K, 500 MHz) of phenanthrene thread **17**, phenanthrene rotaxane **16**, macrocycle **15**, indane rotaxane **27**, and indane thread **29**. Signal labeling according to Schemes 2 and 5. Signals from the macrocycle are highlighted in green.



Figure 9. Unmasking of rotaxanes **16** and **27**. a) ¹H NMR spectra (cyclohexane- d_{12} , 298 K, 400 MHz) of a solution of **27** before and after irradiation (250 nm) for 4 h. Green peaks are assigned to the macrocycle; red peaks to indane. Green peaks sharpen over time, indicating a higher symmetry. Unmasked octayne rotaxane **28** was isolated from the reaction mixture (34% yield). b) ¹H NMR spectra (cyclohexane- d_{12} , 298 K, 400 MHz) of a solution of **16** before and after irradiation (250 nm) for 12 h. Green peaks are assigned to the macrocycle; red peaks to phenanthrene. Phenanthrene is generated in the course of the reaction, but the macrocycle peaks broaden and lose intensity due to decomposition. No unmasked rotaxane **28** was isolated from the photolysis of **16**.

Sharpening of the macrocycle peaks does not occur on irradiation of phenanthrene rotaxane **16** (Figure 9b); all peaks belonging to the rotaxane broaden significantly and lose intensity, in spite of the release of phenanthrene. ESI-MS shows no new peaks, only the peak for starting material **16** (intensity decreases over time) was observed. No identifiable products could be isolated from photolysis of **16**.

As a reference, we synthesized the corresponding octayne dumbbell **30** by irradiating indane thread **29** with UV light (250 nm): the yield (31%) is similar to the unmasking of rotaxane **27** (Scheme 6). The ¹H NMR spectra of indane rotaxane **27**, unmasked rotaxane **28**, macrocycle **15** and unmasked thread **30** are compared in Figure 10. No strong NOEs are observed between the stopper units and the macrocycle of **28**, suggesting that the macrocycle resides near the center of the polyyne chain.

CONCLUSIONS

We have shown that MAEs can be successfully applied to make long polyyne rotaxanes. The unmasking reaction of two different MAEs was thoroughly studied and unexpected photochemical side products were identified by X-ray crystallography. The indane-based MAE developed by Tobe and coworkers can be unmasked yielding a polyyne rotaxane comprising eight -C=C- units, whereas the phenanthrene MAE proved to be unsuitable, due to competing side reactions during photochemical unmasking. This is the first demonstration that a polyyne can be released from a masked interlocked precursor. Our results indicate that encapsulated versions of carbyne and cyclocarbons should be accessible, but to achieve this objective, it will be necessary to use a smaller macrocycle (or increase the bulk of the MAE), so that the MAE groups act as temporary stoppers.



Figure 10. a) ¹H NMR spectra (CDCl₃, 298 K, 500 MHz) of masked indane rotaxane **27**, unmasked octayne rotaxane **28**, free macrocycle **15** and unmasked thread **30**. Signals numbered according to Scheme 5. The shifts of macrocycle **15** are hardly affected by the threaded dumbbell in rotaxane **28**.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsXXXXXXX.

Details of synthetic protocols, copies of NMR and MS spectra, crystallographic data (PDF).

Crystallographic data for compound 2 (2003495) (CIF)

Crystallographic data for compound 6 (2003496) (CIF)

Crystallographic data for compound 7 (2003497) (CIF)

Crystallographic data for compound **8** (2003498) (CIF)

Crystallographic data for compound **10** (2003499) (CIF)

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