Durham Research Online

Deposited in DRO:
02 November 2016

Version of attached file:
Published Version

Peer-review status of attached file:
Peer-reviewed

Citation for published item:
Danon, J. J. and Leigh, D. A. and McGonigal, P. R. and Ward, J. W. and Wu, J. (2016) 'Triply threaded [4]rotaxanes.', Journal of the American Chemical Society., 138 (38). pp. 12643-12647.

Further information on publisher's website:
http://dx.doi.org/10.1021/jacs.6b07733

Publisher's copyright statement:
ACS AuthorChoice - This is an open access article published under a Creative Commons Attribution (CC-BY) License, which permits unrestricted use, distribution and reproduction in any medium, provided the author and source are cited.

Additional information:

Use policy

The full-text may be used and/or reproduced, and given to third parties in any format or medium, without prior permission or charge, for personal research or study, educational, or not-for-profit purposes provided that:

- a full bibliographic reference is made to the original source
- a link is made to the metadata record in DRO
- the full-text is not changed in any way

The full-text must not be sold in any format or medium without the formal permission of the copyright holders.

Please consult the full DRO policy for further details.
Triply Threaded [4]Rotaxanes

Jonathan J. Danon,† David A. Leigh,‡,*,‡ Paul R. McGonigal,† John W. Ward,† and Jhenyi Wu‡

†School of Chemistry, University of Manchester, Oxford Road, Manchester M13 9PL, United Kingdom
‡School of Chemistry, University of Edinburgh, The King’s Buildings, West Mains Road, Edinburgh EH9 3JJ, United Kingdom

Supporting Information

ABSTRACT: [4]Rotaxanes featuring three axles threaded through a single ring have been prepared through active metal template synthesis. Nickel-catalyzed sp³→sp³ homocouplings of alkyl bromide “half-threads” through 37- and 38-membered 2,2′:6′,2″-terpyridyl macrocycles generate triply threaded [4]rotaxanes in up to 11% yield. An analogous 39-membered macrocycle produced no rotaxane products under similar conditions. The constitutions of the [4]rotaxanes were determined by NMR spectroscopy and mass spectrometry. Doubly threaded [3]rotaxanes were also obtained from the reactions but no [2]rotaxanes were isolated, suggesting that upon demetalation the axle of a singly threaded rotaxane can slip through a macrocycle that is sufficiently large to accommodate three threads.

INTRODUCTION

Although there are many examples of rotaxanes consisting of several macrocycles threaded onto a single molecular strand,1 rotaxanes with more than one axle passing through a ring remain rare.2,3 Doubly threaded [3]rotaxanes have been prepared1 by active metal template synthesis,5 exploiting the ability of the metal atom bound by the macrocycle to turn over and catalyze formation of a second axle through the ring. However, to date, no rotaxanes have been reported with more than two strands threaded through a macrocycle. This is likely due to a “Goldilocks requirement” of the macrocycle needing to be large enough to accommodate three (or more) molecular chains threaded through its cavity while also having to be sufficiently small to prevent the ring slipping over the stoppers of an individual axle.

We previously described the synthesis of doubly threaded [3]rotaxanes using the one-pot nickel-catalyzed sp³→sp³ reductive homocoupling of alkyl bromides2a through a 2,2′:6′,2″-terpyridyl macrocycle. In that system the macrocycle-nickel complex is able to turn over, allowing covalent capture of a second thread through a 35-membered ring. Here we report that modestly increasing the size of the macrocycle to a 37- or 38-membered ring enables a third axle to be assembled through the cavity, generating the first examples of triply threaded rotaxanes.

One might expect that a modest increase in the size of stoppers that are sufficiently bulky to form a [2]rotaxane should enable multiple threaded rotaxanes to be formed with a large macrocycle via successive covalent bond-forming reactions directed through the cavity. However, experiments1 and molecular modeling both show that even small increases in macrocycle cavity size require very significant expansion of the stoppers to prevent detethering of singly threaded [2]-

RESULTS AND DISCUSSION

A catalytically active macrocycle–Ni(0) complex of 1a was generated by the in situ reduction of a Ni(II) salt in N-methyl-2-pyrrolidone (NMP), to which was added 20 equiv of 2a in tetrahydrofuran (THF). The reaction mixture was heated at 60 °C for 18 h before washing with ammonial ethylenediaminetetraacetic acid disodium salt solution (Na₂EDTA–NH₃(aq)) to remove the metal ions (Scheme 1). Purification of the crude organic products by size-exclusion chromatography (SEC) led to the recovery of unreacted macrocycle 1a (44%) and the isolation6 of three products, identified by ¹H NMR (e.g., relative integration of axle-to-ring protons) and mass spectrometry (e.g., mass of molecular ion) as the non-interlocked thread 3a, doubly threaded [3]rotaxane 4a (19%) and triply threaded [4]rotaxane 5a (11%).

The absence of any singly threaded [2]rotaxane among the rotaxane products isolated from the active template reaction...
was initially surprising, but CPK models show that tris(‘Bu-phenyl)methylene stoppers can slip through the cavity of metal-free 1a. However, a singly threaded species must be an intermediate in the assembly of the more highly threaded structures, 4a and 5a. This is possible because the cavity of the nickel-coordinated macrocycle is considerably smaller than that of the metal-free ring. It appears that the nickel-complexed singly threaded [2]rotaxane intermediate is sufficiently long-lived to react with two more alkyl bromide molecules and catalyze the formation of a second axle to form the doubly threaded [3]rotaxane 4a before dethreading, either directly or through transient loss of the metal center (Figure 1). The 1H NMR spectra of rotaxanes 4a and 5a (Figure 2b,c) show several upfield chemical shifts with respect to that of their non-interlocked components, 1a and 3a (Figure 2a,d), consistent with face-on interactions of regions of one component with the aromatic rings of another. The shifts are

Figure 1. Proposed pathway for one-pot active template synthesis of triply threaded [4]rotaxanes, 5a–f. A Ni(0) center (purple) bound to the endotopic terpyridyl coordination site (red) of a macrocycle (blue) catalyzes the sp3−sp3 homocoupling reaction of two alkyl bromide building blocks through the cavity. The thread and macrocycle can dissociate, either by transient demetalation forming a larger cavity or through a slow dethreading process, or the coordinated Ni(0) can catalyze a second carbon–carbon bond-forming reaction between two more building blocks, to give a doubly threaded [3]rotaxane. The presence of a second thread prevents any further dethreading. A further Ni(0)-catalyzed homocoupling of thread building blocks generates the triply threaded [4]rotaxane 5a–f.

[3]rotaxane is kinetically stable and can, in turn, promote a third active template nickel-catalyzed homocoupling of thread-forming components 2a to form the triply threaded [4]rotaxane, 5a (Table 1, entry 1). The kinetic stability of the nickel-coordinated [2]rotaxane is also consistent with the recovery of significant quantities of macrocycle 1a after the reaction is worked up, despite the use of a large excess (20 equiv) of the thread-forming component. Much of the recovered macrocycle likely results from decomposition of the nickel-coordinated [2]rotaxane during the workup procedure, as <30% macrocycle was recovered in the previously reported active template reactions that use slightly smaller macrocycles and generate stable [2]rotaxanes.

The 1H NMR spectra of rotaxanes 4a and 5a (Figure 2b,c) show several upfield chemical shifts with respect to that of their non-interlocked components, 1a and 3a (Figure 2a,d), consistent with face-on interactions of regions of one component with the aromatic rings of another. The shifts are
generally more pronounced for the [4]rotaxane (Figure 2c), a consequence of the tight packing of the axles in the cavity resulting in them spending more time close to one another and to the macrocycle aromatic rings. A further indication of the increased steric crowding in the triply threaded [4]rotaxane is the broadening of a number of signals (e.g., H,M,I), likely due to restricted movement of the components relative to each other on the 1H NMR time scale.

The structure of the triply threaded [4]rotaxane 5a was modeled with Spartan’14. The semiempirical PM6 geometry minimized structure shows the terpyridyl group tilted almost orthogonally to the plane of the macrocycle, which maximizes the size of the cavity to accommodate the three threaded chains (Figure 3). In adopting this co-conformation, the stopper groups of the three chains are brought in close proximity to each other at both ends of the rotaxane. The protruding phenyl ring of the macrocycle is directed toward the aromatic rings of one of the stoppers, consistent with the upfield shifts of protons H_A and H_B after the third threading event (Figure 2b,c).

To probe the multiple threading process further, we investigated the influence of changes in both the axle length and macrocycle size on the active template rotaxane-forming reaction (Table 1). Increasing the distance between the bromine atom and the stopper of the axle-forming alkyl thread building block generally leads to decreased yields of the resulting [4]rotaxane (Table 1).

| entry | thread building block | [3]rotaxane 4 (%) yield | [4]rotaxane 5 (%) yield |
|-------|-----------------------|------------------------|------------------------|
| 1'    | 2a                    | 4a (19)                | 5a (11)                |
| 2'    | 2b                    | 4b (22)                | 5b (7)                 |
| 3'    | 2c                    | 4c (21)                | 5c (7)                 |
| 4'    | 2a                    | 4d (19)                | 5d (4)                 |
| 5'    | 2b                    | 4e (20)                | 5e (7)                 |
| 6'    | 2c                    | 4f (17)                | 5f (3)                 |

"20 equiv. \(^b\)Isolated yields with respect to amount of macrocycle employed. \(^c\)Performed with macrocycle 1a. \(^d\)Performed with macrocycle 1b. Reactions carried out under the conditions shown in Scheme 1; >95% conversion to homocoupled products was observed in each case.

Figure 2. \(^{1}H\) NMR spectra (600 MHz, CDCl\(_3\), 298 K) of (a) macrocycle 1a, (b) [3]rotaxane 4a, (c) [4]rotaxane 5a, and (d) free thread 3a. Assignments correspond to labeling in Scheme 1.

Figure 3. Spartan’14 semiempirical PM6 geometry-minimized structure of [4]rotaxane 5a. Hydrogen atoms not shown for clarity. (a) Viewed through the macrocycle cavity and (b) side-on.
bromides from a three-carbon chain in 2a to 6 and 12 carbons in 2b and 2c, respectively, had little effect on the yields of the corresponding [3]- (4b and 4c) and [4]rotaxanes (5b and 5c) (Table 1, entries 2 and 3). A comparison of the 1H NMR spectra of [4]rotaxanes 5a–c shows less shielding with increasing chain length (Supporting Information, Figure S2), indicating that the longer threads can position themselves to minimize unfavorable steric interactions between the bulky stoppers and the macrocycle. Nevertheless, the greater steric congestion from threaded chains close to the metal binding site in the synthesis of 5a does not significantly affect the ability of the nickel to catalyze the active template reaction.

Macrocycles 1b and 1c, containing one and two extra methylene groups in the southern hemisphere, respectively, were prepared to investigate the effect of a modest increase in cavity size on the ability to trap multiply threaded rotaxanes. Using the 38-membered ring macrocycle 1b, [3]rotaxanes (4d–f) and [4]rotaxanes (5d–f) were obtained in similar yields to those obtained using macrocycle 1a (Table 1, entries 4–6). The 1H NMR spectra of [4]rotaxanes 5a and 5d are very similar to each other (Supporting Information, Figure S3), although the downfield chemical shift of proton H3 (Δp = +0.09 ppm with the wider cavity) provides some indication that the 38-membered ring somewhat alleviates the close packing of the threads.

When macrocycle 1c was employed in the analogous active template reactions, however, no rotaxane products could be isolated. Apparently the cavity of the 39-membered ring is too small to accommodate the nickel to catalyze the active template reaction.

CONCLUSIONS

Rotaxanes with multiple axles threaded through the same ring are challenging to access, requiring a delicate balance of microcycle cavity size and conformation to stopper size, and a threading reaction that can tolerate increasing steric bulk at the site where the chains are joined through the ring. The one-pot active metal template nickel-catalyzed homocoupling of stoppered alkyl bromides of various lengths through 37- or 38-membered 2,2′,6,2′-terpyridyl macrocycles affords the first examples of tripod threaded [4]rotaxanes, together with doubly threaded [3]rotaxanes and the non-interlocked threads. Although the cavities of the macrocycles utilized are too large for 2|rotaxanes to be isolated using the stoppers employed, the nickel-coordinated macrocycles are small enough to form 2|rotaxane intermediates that can react further to form kinetically stable, multiply threaded, [3]- and [4]rotaxanes. We anticipate that this approach will be useful for the synthesis of rotaxanes featuring even more chains threaded through a single ring, by judicious choice of template, coupling reaction, macrocycle size and conformation, and stopper size and shape.

EXPERIMENTAL SECTION

General Experimental Procedure for the Active Metal Template Synthesis of [3]- and [4]Rotaxanes. To a solution of the terpyridine macrocycle (1a or 1b, 1 equiv) and NiCl2·6H2O (1 equiv) in NMP (2.5 mL) under an inert atmosphere of nitrogen was added activated Zn powder (20 equiv) and the resulting suspension stirred vigorously for 1 h, resulting in a color change from greenish yellow to deep purple. A solution of the stoppered bromide (2a, 2b, or 2c, 20 equiv) in THF (2.5 mL) was added and the resulting mixture stirred at 60 °C for 18 h. The reaction mixture was diluted with EtOAc (200 mL) and washed with a 17.5% aqueous solution of ammonia saturated with Na2EDTA (2 x 50 mL), H2O (3 x 50 mL) and brine (50 mL). The organic layer was dried over MgSO4, the solvent removed under reduced pressure and the resulting residue purified by size exclusion chromatography (CH2Cl2, as eluent, see Supporting Information, Figure S1) to yield the corresponding [3]- and [4]rotaxanes (4a–f and 5a–f, respectively) as colorless powders or clear films.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.6b07733.

Synthetic procedures and characterization data (PDF)

AUTHOR INFORMATION

Corresponding Author

david.leigh@manchester.ac.uk

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We thank the EPSRC for funding (EP/H021620/2), the EPSRC National Mass Spectrometry Centre (Swansea, U.K.) for high-resolution mass spectrometry and Miriam R. Wilson and Javier Jaramillo-Garcia for their assistance with the graphics and molecular modeling.

REFERENCES

(1) Xue, M.; Yang, Y.; Chi, X.; Yan, X.; Huang, F. Chem. Rev. 2015, 115, 7399–7501.

(2) (a) Klotz, E. J. F.; Claridge, T. D. W.; Anderson, H. L. J. Am. Chem. Soc. 2006, 128, 15374–15375. (b) Prikhod’ko, A. I.; Durola, F.; Sauvage, J.-P. J. Am. Chem. Soc. 2008, 130, 448–449. (c) Prikhod’ko, A. I.; Sauvage, J.-P. J. Am. Chem. Soc. 2009, 131, 6794–6807. (d) Lee, C.-F.; Leigh, D. A.; Pritchard, R. G.; Schultz, D.; Teut, S. J.; Timco, G. A.; Winpenny, R. E. P. Nature 2009, 458, 314–318. (e) Ackermann, D.; Schmidt, T. L.; Hannam, J. S.; Purohit, C. S.; Heckel, S. A.; Famulok, M. Nat. Nanotechnol. 2010, 5, 436–442. (f) Zhang, Z.-J.; Zhang, H.-Y.; Wang, H.; Liu, Y. Angew. Chem., Int. Ed. 2011, 50, 10834–10838.

(3) (a) Cheng, H. M.; Leigh, D. A.; Maffei, F.; McGonigal, P. R.; Slawin, A. M. Z. Wu, J. Am. Chem. Soc. 2011, 133, 12298–12303. (b) Yamashita, Y.; Mutoh, Y.; Yamasaki, R.; Kasama, T.; Saito, S. Chem. - Eur. J. 2015, 21, 2139–2145. (c) Hayashi, R.; Mutoh, Y.; Kasama, T.; Saito, S. J. Org. Chem. 2015, 80, 7536–7546. (d) Movsisyan, L. D.; Franz, M.; Hampel, F.; Thompson, A. L.; Tykynski, R. J.; Anderson, H. L. J. Am. Chem. Soc. 2016, 138, 1366–1376.

(4) (a) Aucagne, V.; Hänni, K. D.; Leigh, D. A.; Lusby, P. J.; Walker, D. B. J. Am. Chem. Soc. 2006, 128, 2186–2187. (b) Berná, J.; Crowley, J. D.; Goldup, S. M.; Hänni, K. D.; Lee, A.-L.; Leigh, D. A. Angew. Chem., Int. Ed. 2007, 46, 5709–5713. (c) Crowley, J. D.; Hänni, K. D.; Lee, A.-L.; Leigh, D. A. J. Am. Chem. Soc. 2007, 129, 12092–12093. (d) Goldup, S. M.; Leigh, D. A.; Lusby, P. J.; McBurney, R. T.; Slawin, A. M. Z. Angew. Chem. Int. Ed. 2008, 47, 3381–3384. (e) Berná, J.; Goldup, S. M.; Lee, A.-L.; Leigh, D. A.; Symes, M. D.; Teobald, G.; Zerbetto, F. Angew. Chem., Int. Ed. 2008, 47, 4392–4396. (f) Goldup, S. M.; Leigh, D. A.; Long, T.; McGonigal, P. R.; Symes, M. D.; Wu, J. J. Am. Chem. Soc. 2009, 131, 15924–15929. (g) Crowley, J. D.; Goldup, S. M.; Lee, A.-L.; Leigh, D. A.; McBurney, R. T.; Chem. Rev. 2009, 108, 1530–1541. (h) Crowley, J. D.; Hänni, K. D.; Leigh, D. A.; Slawin, A. M. J. Am. Chem. Soc. 2010, 132, 5309–5314. (i) Crowley, J. D.; Goldup, S. M.; Gowans, N. D.; Leigh, D. A.; Ronaldson, V. E.; Slawin, A. M. Z. J. Am. Chem. Soc. 2010, 132, 6243–6248. (j) Goldup, S. M.; Leigh, D. A.; McBurney, R. T.; McGonigal, P. R.; Plant, A. Chem. Sci. 2010, 1, 383–386. (k) Lahal, H.; Jobe, K.; Watkinson, M.; Goldup, S. M. Angew. Chem., Int. Ed. 2011, 50, 4151–4155. (l) Langton, M. I.;
(m) Weisbach, N.; Baranova, Z.; Gauthier, S.; Reibenspies, J. H.; Gladysz, J. A. *Chem. Commun.* 2012, 48, 7562–7564. (n) Lewandowski, B.; De Bo, G.; Ward, J. W.; Papmeyer, M.; Kuschel, S.; Aldegunde, M. J.; Gramlich, P. M. E.; Heckmann, D.; Goldup, S. M.; D’Souza, D. M.; Fernandes, A. E.; Leigh, D. A. *Science* 2013, 339, 189–193. (o) Winn, J.; Pinczewska, A.; Goldup, S. M. J. *Am. Chem. Soc.* 2013, 135, 13318–13321. (p) Bordoli, R. J.; Goldup, S. M. J. *Am. Chem. Soc.* 2014, 136, 4817–4820. (q) Noor, A.; Moratti, S. C.; Crowley, J. D. *Chem. Sci.* 2014, 5, 4283–4290. (r) Noor, A.; Lo, W. K. C.; Moratti, S. C.; Crowley, J. D. *Chem. Commun.* 2014, 50, 7044–7047. (s) Baranova, Z.; Amini, H.; Bhuvanesh, N.; Gladysz, J. A. *Organometallics* 2014, 33, 6746–6749. (t) De Bo, G.; Kuschel, S.; Leigh, D. A.; Lewandowski, B.; Papmeyer, M.; Ward, J. W. J. *Am. Chem. Soc.* 2014, 136, 5811–5814. (u) Hoekman, S.; Kitching, M. O.; Leigh, D. A.; Papmeyer, M.; Roke, D. J. *Am. Chem. Soc.* 2015, 137, 7656–7659. (v) Barat, R.; Legigan, T.; Tranoy-Opalinski, I.; Renoux, B.; Pérandeau, E.; Clarhaut, J.; Piolet, P.; Fernandes, A. E.; Aucagne, V.; Leigh, D. A.; Papot, S. *Chem. Sci.* 2015, 6, 2608–2613. (w) Franz, M.; Januszewski, J. A.; Wendinger, D.; Neiss, C.; Movsisyan, L. D.; Hampel, F.; Anderson, H. L.; Görling, A.; Tykwinski, R. R. *Angew. Chem., Int. Ed.* 2015, 54, 6645–6649. (x) Neal, E. A.; Goldup, S. M. *Chem. Sci.* 2015, 6, 2398–2404.

(5) (a) Saito, S.; Nakazono, K.; Takahashi, E. *J. Org. Chem.* 2006, 71, 7477–7480. (b) Saito, S.; Takahashi, E.; Wakatsuki, K.; Inoue, K.; Orikasa, T.; Sakai, K.; Yamasaki, R.; Mutoh, Y.; Kiyoshima, T. *J. Org. Chem.* 2013, 78, 3553–3560.

(6) The rotaxane yields reported are with respect to the macrocycle, the limiting reactant employed. The reactions typically proceeded with >95% conversion of the alkyl bromide building blocks, used in excess, to the combined homocoupled axle products (non-interlocked thread and [3]- and [4]rotaxanes).

(7) (a) *Spartan’14*; Wavefunction Inc., Irvine, CA, 2014. (b) Stewart, J. J. P. *J. Mol. Model.* 2007, 13, 1173–1213.