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REVIEW

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Cyclic polymers from alkynes: a review

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Cyclic polymers have applications across various fields, including material science, biomedicine, and inorganic chemistry. Cyclic polymers derived from alkyne monomers have expanded the application scope to include electronic materials and polyolefins. This review highlights recent advancements in the synthesis of cyclic polymers from both mono- and disubstituted alkynes. The aim is to provide a comprehensive overview of the synthetic methodologies and the application of cyclic polymers derived from alkynes. Additionally, this review will facilitate a comparative analysis of the advantages and limitations of various synthetic methods and describe opportunities for future development of novel catalytic systems to synthesize cyclic polymers from alkynes.

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1. Introduction

The increasing need for flexible electronic components has prompted extensive exploration into the synthesis of conjugated polymers. The unique combination of flexibility, electronic conductivity, and tuneable electrochemical properties make conjugated polymers ideal for advanced electronic applications. Poly(3-hexylthiophene) is an example of a conjugated polymer used in organic sensors and organic solar cells. Substituted polyacetylenes are also conjugated and exhibit electronic conductivity, electrochromism, and luminescence. Leveraging these properties, researchers use polyacetylenes for

(LEDs).²¹ Some polyacetylenes adopt a helical secondary structure,²² enabling applications in enantiomer resolution,²³ chirality sensing,²⁴ and circularly polarized luminescence.¹⁶ Most recently, and the subject of this review, conjugated polyacetylenes with a cyclic topology are now available, thus opening new opportunities to explore topological effects on all the aforementioned properties and applications. For example, discovered in our lab, cyclic polyacetylene (cPA) synthesized at any temperature has a predominantly *trans* configuration, whereas the linear forms initially with *cis* bonds. Additionally, films of cPA are more stretchable leading to a significant increase in conductivity compared to linear polyacetylene (lPA).

sensing cyanide,17 photoresists,20 and light emitting diodes

Cyclic and linear polymers with the same repeat unit and molar mass exhibit different thermal,²⁵ rheological,²⁶ crystallization,²⁷ and degradation²⁸ properties. For example, cyclic polyesters degrade slower than linear versions of identical molar mass,²⁸

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Synthesis of cyclic polymers from alkynes

a feature particularly advantageous for cyclic polyacetylenes given their tendency to degrade oxidatively.29 Nature also takes advantage of topology; cyclic DNA³⁰ and peptides^{31,32} exhibit increased thermal, chemical, and enzymatic stability.

The solid-state physical properties of cyclic and linear polymers are different. For example, the reduced free volume³³ created by a lack of chain ends in cyclic polymers result in higher glass transition temperatures (T_{σ}) than linear analogues of similar molar mass.^{25,34,35} Also, a lack of chain ends for cyclic polymers results in fewer entanglements leading to lower melt viscosities. 36 The higher T_g and lower melt viscosities of cyclic polymers suggest their promising application as thermoplastics with improved processing capability.

Amphiphilic block copolymers undergo self-assembly in both the solution and solid-state and given the decreased size of cyclic polymers their assemblies are smaller. Indeed, theoretical studies predict the spacing between cyclic block copolymers assemblies is 0.63 smaller.³⁷ Subsequent early experimental studies validated the theory revealing solid-state domain spacing 0.70-0.75 times smaller for assembled cyclic block copolymers compared to linear analogues,38,39 a result crucial to advancing block copolymer nanolithography. Recent studies suggest a cyclic topology has more impact on the self-assembly properties of polymers than previously thought.40

Despite the many studies illustrating the physical differences between linear and cyclic polymers many properties remain underexplored, for example conductivity in conjugated polymers41-43 with a cyclic topology. The synthesis of cyclic polymers in reasonable quantities and purity has always been the limiting factor to exploring the full suite of cyclic polymer properties. Ring closing of a linear polymeric precursor and ring expansion are the two most common methods for synthesizing cyclic polymers. Historically, catalysts used to polymerize alkynes do not provide control over the polymer end groups, thus preventing the synthesis of cyclic polyacetylenes via ring closure. Overcoming this synthetic limitation, catalysts can now directly synthesize conjugated cyclic polyacetylenes from alkynes according to Fig. 1. In this review, we highlight recent key developments in the synthesis of cyclic polyacetylenes from alkynes, their emerging applications, and the remaining challenges starting from the synthesis a pincer supported tungsten alkylidyne complex.

Cyclic polymers from monosubstituted alkynes

A breakthrough in cyclic polymer synthesis came in 2012 when we reported alkylidyne complex 1 initiates the polymerization of phenylacetylene.44 At the time, we were not aware of the cyclic topology of the resulting polyacetylenes. However, the clues to the hidden topology were there originally, including the isolation of tethered alkylidene intermediates 2 and 3 (ref. 44) and ring expanded metallacyclopentadienes 4 and 5 (Fig. 2).45 Eventually, in 2016 we reported the tethered alkylidene tungsten catalyst 6 initiates the polymerization of phenyl acetylene (1:10 000) to give cyclic poly(phenylacetylene) (cPPA)⁴⁶ in high yield (>94%) (Table 1, entry 1). Cyclic polyacetylenes from most alkyne monomers are now accessible using 6 including, aliphatic alkynes, gaseous alkynes (propyne and acetylene), and both electron poor and rich phenylacetylene derivatives. 34,35,47-49 The assignment of a cyclic topology for the polymers produced from 6 and alkynes comes from a variety of techniques; such as, atomic force microscopy (AFM),47,50 differential scanning calorimetry (DSC),25 gel permeation chromatography (GPC),48,51 and rheology.34,35 One of the exciting features of initiator 6 is its

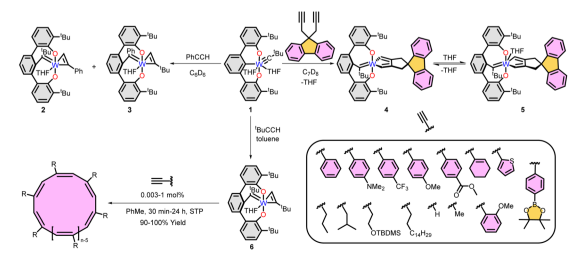


Fig. 2 Synthesis of ring expanded metallacyclopentadienes (top left), tethered alkylidene intermediates (top right), polymerization of alkynes and monomer scope (bottom).

Table 1 Polymerization of phenylacetylene

Entry	Monomer	Catalyst	[PhCCH]: [Cat]	Time	Yield (%)	M_n (kDa)	Đ
1	PA	6	10 000	15 min	94	30.8	2.2
2	PA	6	5000	15 min	96	69	1.8
3	PA	8	1000	12 h	94	62.9	1.6
4	PA	8	100	2 h	94	272	1.3
5	PA	10	1000	12 h	37	108	1.5
6	PA	10	1000	12 h	19	115	1.5
7	PA	11	500	12 h	59	48	4.1
8	PA	11	100	12 h	21	30	3.4
9	PA	12	50	12 h	16	32	3.6
10	PA	12	250	12 h	7	38	3.5

extreme reactivity toward alkynes, with measured activities of 180 000 000 g $\mathrm{mol_{cat}}^{-1}\,\mathrm{h}^{-1}.^{34}$ The cyclic polyacetylenes produced by 6 typically exhibit a predominantly cis (>90%) configuration with the notable exception of cyclic polyacetylene.

The mechanism of cyclic polymer formation remains unclear. Isolation or *in situ* detection of several metallacyclopentadienes^{45,52,53} lead to our initial proposal of a straightforward ring expansion polymerization (REP) where subsequent insertions of alkyne grow the cyclic polymer on the metal center. The role, if any, of the tethered alkylidene remains elusive.

Alkylidynes polymerize acetylenes to give linear polyacetylenes.⁵⁴ However, recent results by Mindiola and a patent suggest the possibility that cyclic polymers are also accessible from alkynes initiated by alkylidynes. In the patent, Exxon reports that (^tBuO)₃W≡C^tBu initiates the polymerization of phenylacetylene to give cyclic poly(phenylacetylene).⁵⁵ Evidence for the cyclic topology comes from DSC, GPC, and NMR data. Mindiola *et al.*, in 2024, reported the vanadium-alkylidyne 8 initiates the polymerization of phenylacetylene to yield cyclic poly(phenylacetylene) (with some linear impurities).⁵⁶ In earlier work,⁵⁷ they noted in passing that polymer forms when 9 was exposed to phenylacetylene but its topology was not interrogated. Revisiting those initial results revealed the cyclic

topology of PPA formed by **8**. Very good yields of polymer form within 12 h with a 1:1000 catalyst: monomer loading of **8** (Table 1, entry 3). Increasing the loading to 1:100 results in a corresponding faster reaction providing 94% in 2 h (Table 1, entry 4). Initial inquiries into the mechanism of polymerization revealed several key intermediates. The interplay between the noninnocent $dBDI^{2-}$ (ArNC(CH₃)CHC(CH₂)Nar, Ar = 2,6- i Pr₂C₆H₃) ligand is a key feature to access a cyclic topology. For example, intermediate **10**, bearing the protonated form of the $dBDI^{2-}$ ligand and a metallacyclohextriene was isolated and is active for the synthesis of cPPA. Although, the yield was low (19–37%) the cPPA was topologically pure (Table 1, entries 5 & 6).

Installing metal-carbon triple bonds within alkylidynes can be synthetically challenging. Circumventing the need to access an alkylidyne, Mindiola recently reported that treatment of Ti or V dihalide precursors with 1,3-dilithioallene yields metal-lacyclobuta-(2,3)-diene species capable of polymerizing phenylacetylene to yield cPPA. While the complexes are simple to prepare the yields of cyclic polyphenylacetylene are modest when using complexes 11 (21–59%) or 12 (7–16%), with relatively higher dispersity (3.5–4.1) compared to 8 and 10 (1.3–1.6) (Table 1, entries 7–10). The *cis/trans* content of cyclic poly(phenylacetylene) produced from catalysts 7–12 has not been evaluated.

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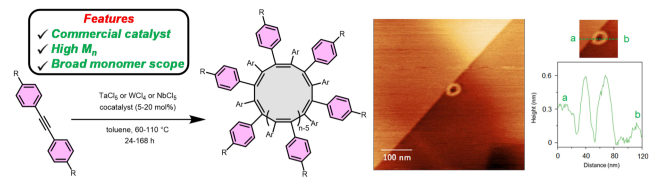


Fig. 3 Polymerization of disubstituted acetylenes using commercially available catalysts and AFM image of cyclic polydiphenylacetylene on HOPG. Reprinted with permission from Macromolecules. 2023, 56, 5873–5880. Copyright 2023 American Chemical Society.

3. Cyclic polymers from disubstituted alkynes

3.1 Polymerization of disubstituted alkynes

The polymerization of disubstituted alkynes was thoroughly studied. $^{59-63}$ However, the polymerization of disubstituted alkynes for cyclic polymer synthesis is receiving renewed attention. In 2021, Maeda *et al.* reported the polymerization of diphenylacetylenes at 80 °C with TaCl₅ pretreated with n Bu₄Sn in toluene to yield cyclic polymers (Fig. 3) with high *cis*-content. 64 The exciting discovery is the easy access to cyclic polyacetylenes from commercial reagents.

In subsequent work Maeda and Taniguchi et al. reveal NbCl₅ with PhSiH₃ also affords cyclic pretreated polydiphenylacetylenes from diphenylacetylenes substituted in either the meta or para positions in high yield and high molar mass.65 Additionally, the NbCl₅/PhSiH₃ catalysts system produces predominately cis-double bonds within the backbone. Moreover, using NbCl₅/PhSiH₃ the authors synthesized the first helical cyclic polyacetylenes by polymerizing monomers containing chiral side groups. These polymers form predominantly one-handed helices capable of circularly polarized luminescence. The high molar mass of the polymers appeared to drastically increase the induced circular dichroism intensities compared to previous reports.65

Expanding the monomer scope Maeda *et al.* reported a WCl₄ initiated polymerization of electron poor, rich, and neutral diphenylacetylenes. ⁶⁶ WCl₄ pretreated with Ph₄Sn and PPh₃ at 110 °C will polymerize diphenylacetylenes to produce cyclic polydiphenylacetylenes. AFM provided conclusive evidence for the cyclic topology of the polymers (Fig. 3).

3.2 Polymerization of cyclic alkynes

In 2016 Fisher *et al.* reported the ring closing of a linear polymer to give cyclic polyphenylene ethynylene during ring opening alkyne metathesis polymerization (ROAMP) as depicted in Fig. 4.⁶⁷ In that system the choice of catalyst dictates whether a linear or cyclic polymer forms. Specifically, the authors found that polymerizing 5,6,11,12-tetradehydrobenzo[a,e][8]annulene with alkylidyne 13 selective (for the chain end) backbiting led to the formation of cyclic phenylene ethynylenes (with minor

linear impurities) as confirmed by NMR and mass spectrometry (Fig. 4).

A common method to avoid linear impurities formed during ring closing of a linear polymer is to perform ring expansion of an already cyclic initiator. To this end just a few years later the first ring expansion alkyne metathesis polymerization (REAMP) of cyclic alkynes to yield cyclic polyalkynes was reported by Veige *et al.* Treating 3,8-didodecyloxy-5,6-dihydro-11,12-dihydrodibenzo[a,e]-[8]annulene (14), with the tethered alkylidyne initiator 15 in toluene produces cyclic poly(a-phenylene ethynylene) (**cPoPE**). Notably, the polymerization proceeds quickly (5 min) producing high yields (70%) of **cPoPE** with high molar mass (a_a > 100 kg mol⁻¹) but high dispersity (a = 4.2-9) (Fig. 5).

Using the methods described in this review, cyclic polymers are now accessible from almost any alkyne. Overall, the recent



Fig. 4 Synthesis of cyclic poly(o-phenylene ethylene) using ROAMP catalyst 13.

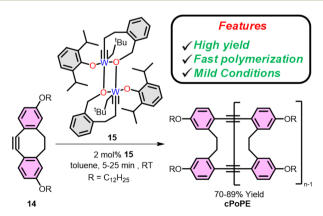


Fig. 5 Polymerization of 14 via REAMP using tethered alkylidyne 15.

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Fig. 6 Synthesis of cyclic poly(4-methyl-pentene) (also known as TPX™) and cyclic polypropylene using tethered tungsten alkylidene 6.

advances across all cyclic polymers derived from alkynes through catalyst design, scalability, and monomer scope are opening new applications for these interesting materials that exploit their lack of chain ends.

4. Applications of cyclic polyacetylenes

4.1 Precursor to cyclic polyolefins

The scalable synthesis of cyclic analogues of commercially relevant linear polymers is a longstanding challenge. Given the high activity of initiator 6 commercially relevant cyclic polymers are now accessible at scale. In pursuit of this goal Veige and Sumerlin *et al.* reported the synthesis of cyclic poly(propylene).

Complex 6 initiates the polymerization of propyne gas to form the unsaturated cyclic poly(propyne) and subsequent hydrogenation yields atactic cyclic poly(propylene) (cPP) (Fig. 6). cPP prepared with 6 has a higher glass transition temperature than previously reported linear atactic polypropylene. Investigation of the bulk rheological properties of cPP reveals a lack of a plateau modulus and a zero-shear viscosity ten times lower than linear poly(propylene) (IPP). Both properties suggest interesting potential of cyclic polyolefins as additives in commercial applications.

Poly(4-methyl-1-pentene) (or TPXTM) is a transparent thermoplastic used primarily in packaging and membranes. Treating 4-methyl-1-pentyne (4-MP) with 6 yields unsaturated cyclic poly(4-methyl-1-pentyne) and hydrogenation over Pd/C in cyclohexane yields atactic cyclic poly(4-methyl-1-pentyne)

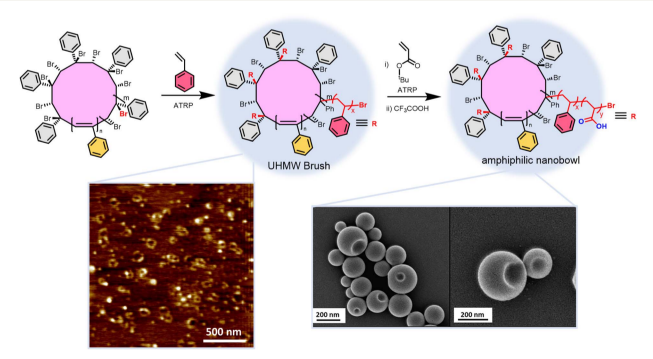


Fig. 7 Synthesis of ultra-high molecular weight bottle brushes *via* ATRP. AFM (bottom left) and SEM images (bottom right) of ultra-high molecular weight bottle brushes. Reprinted with permission from Macromolecules, 2020, 53, 9717–9724 (bottom left) and Macromolecules, 2022, 55, 7446–7453 (bottom right). Copyright 2020 and 2022 American Chemical Society.

Fig. 8 Synthesis of cyclic polyacetylene (left) and cyclic poly(acetylene-co-propylene) using tungsten tethered alkylidene 6.

(**cPMP**).³⁴ Again, **cPMP** exhibits a higher $T_{\rm g}$ (39 °C νs . 29 °C) with no plateau modulus and a zero-shear viscosity fifteen times less than a linear poly(4-methyl-1-pentyne) (**lPMP**) sample. Importantly, catalyst **6** initiates pentyne polymerization with an activity of 180 000 000 $\rm g_{polymer}$ mol_{cat}⁻¹ h⁻¹. Given its high activity initiator **6** has the potential to produce cyclic polyolefins at an industrially relevant scale.

4.2 Synthesis of cyclic brushes

Another application of cyclic polyacetylenes is as a precursor to ultra-high molar mass cyclic bottlebrush polymers *via* post-polymerization modification. Adding bromine to cyclic polyphenylacetylene brominates the backbone yielding a benzylic bromine capable of initiating atom transfer radical polymerization (ATRP) to grow cyclic brush polymers (Fig. 7). Butyl acrylate and styrene side chains grow from the macrocyclic initiator with 52–72% grafting efficiencies. Atomic force microscopy (AFM) interrogation of the bottlebrush polymers dispersed on a surface reveal the topology of the polymers produced from alkynes with 6.

Leveraging the broad monomer scope of grafting from via ATRP enables the synthesis of amphiphilic cyclic bottlebrushes from alkynes capable of self-assembling into porous spheres and nanobowls.68 Grafting from using styrene followed by chain extension using tert-butylacrylate provides the bottlebrush arms. Treating the cyclic bottlebrushes with trifluoroacetic acid in THF converts the tert-butylacrylate repeat units into acrylic acid. Dissolving the amphiphilic bottlebrush in THF forms spherical nano-assemblies with multiple pores. Dropwise addition of water to this solution converts the porous spheres to spheres containing only a single hole (nanobowls). Identical linear bottlebrushes only form the porous spherical nanoassemblies, indicating the topology can affect the selfassembly behaviour of amphiphilic copolymers. The unsaturated bonds of polyacetylenes are excellent sites to add functionality in post-polymerization modification; however, they also act as pathways for electron mobility making cyclic polyacetylenes semi-conducting polymers.

4.3 Synthesis of conductive cyclic polymers

Doping and irradiating polyacetylenes alters their conductivity. 5.7,42,69,70 The effect of a cyclic topology on conductivity is challenging to interrogate though some new insights are

emerging.⁷¹ Exposing a dilute solution of **6** to acetylene gas rapidly produces freestanding films of cyclic poly(acetylene) (**cPA**) (Fig. 8).⁴⁷ An important feature is **cPA** synthesized with **6** contains 99% *trans* double bonds regardless of the polymerization temperature, whereas linear poly(acetylene) (**IPA**) usually requires heating to convert *cis* to *trans*. The methodology allows for rapid synthesis under dilute conditions that temporarily generates soluble or homogenized **cPA**. Notoriously difficult to functionalize, cyclic brush polymers were generated from the temporarily soluble polymers using a similar bromination ATRP strategy as previously outlined.^{47,50} AFM imaging of **cPA** brush polymers add to the support of a cyclic topology.

Consistent with prior literature cyclic copolymers of acetylene and propyne exhibit significantly lower conductivities relative to **cPA**.⁷² Distortion of backbone planarity caused by the methyl substituent reduces the conductivity. However, simply stretching **cPA** either by hand or using dynamic mechanical analysis (DMA) drastically increases its conductivity.⁷³ By stretching I₂ doped **cPA** to 1.6 times its original length (Fig. 9) the conductivity of the sample increases by 5× while stretching **IPA** failed to produce similar results (Table 2). More stretchable films of **cPA**, due either to the

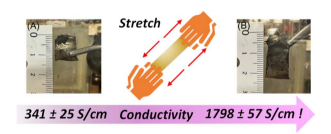


Fig. 9 Photographs of cPA films (A) before and (B) after manual stretching.

Table 2 Conductivities of I_2 doped cPA and IPA films before and after stretching

Sample	Composition	Conductivity (S cm ⁻¹)
Non-stretched cPA Hand-stretched cPA Non-stretched lPA Hand-stretched lPA	$(\mathrm{CHI_{0.34}})_n$ $(\mathrm{CHI_{0.31}})_n$ $(\mathrm{CHI_{0.27}})_n$ $(\mathrm{CHI_{0.28}})_n$	341 ± 25 1798 ± 57 524 ± 48 485 ± 65

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Fig. 10 Synthesis of a cyclic polymer gel by post polymerization modification of cPEPB followed by crosslinking with diglycerol. Reprinted with permission from Macromolecules, 2024, 57, 1779–1787. Copyright 2024 American Chemical Society.

cyclic topology or low crosslink density, lead to improved alignment of the polyacetylene fibrils and thus improved conductivity.

4.4 Synthesis of functional cyclic polymers

Catalyst 6 also polymerizes a variety of functional alkynes.⁴⁹ For example, 6 initiates the polymerization 4-ethynylanisole to produce cyclic poly(4-ethynylanisole).⁴⁸ Cleaving the methyl group using BBr₃ generates cyclic poly(4-ethynylphenol) (**cPEP-OH**). **cPEP-OH** is pH responsive; it was soluble at pH > 11 and insoluble at pH \leq 11.

Similarly, **6** initiates the polymerization of boron functionalized monomers to yield cyclic poly(4-ethynylphenylboronate esters) (**cPEPB**).²⁵ Using Suzuki coupling methods **cPEPB** is easily functionalized (92% by NMR). Treating boronate ester containing cyclic polymers with an excess of methylboronic acid in the presence of trifluoroacetic acid yields cyclic poly(4-ethynylphenylboronic acid) (**cPEPB-OH**) as depicted in Fig. 10. Dissolving **cPEPB-OH** in THF and treating with a diglycerol crosslinker produces gels. Gels produced from **cPEPB-OH** and linear **IPEPB-OH** have similar network stability, but the gel derived from the cyclic version has a higher storage modulus (*G'*).

5. Guidance for handling of cyclic polyacetylenes and cyclic polymer catalysts

Given the sensitivity of both the described catalysts and produced polymers it is important for those who wish to perform the synthesis of cyclic polyacetylenes to know how to properly handle both. Transition metal catalysts 1 and 6 are well known to be highly oxophilic^{49,74} making them highly sensitive to oxygen exposure. Thus, all steps of their synthesis and any polymerizations conducted using them must be performed in oxygen- and moisture-free environments. Despite its extreme reactivity it is also worth mentioning that catalyst 6 is highly stable in solution (months) and in the solid state (years) when properly stored (low temperature free of oxygen).

Less obviously, cyclic polyacetylenes degrade rapidly in oxygenated solutions^{49,75} and slowly in the solid state. Thus, precipitation should be performed without exposure to oxygen (into deoxygenated solution). Additionally, GPC and NMR analysis should be performed in the absence of oxygen which again is easily achieved by preparing the samples in an inert atmosphere

glovebox in appropriately airtight sample holders. The reader should also note that cyclic polyacetylenes are prone to electrocyclization upon extended dissolution in air free solvent at low temperatures.²⁹ Or in the solid state slowly at room temperature and quickly when exposed to elevated temperatures.⁷⁶

6. Conclusions

Thousands of reports over the past 60 years extoll the fascinating properties, chemistry, characterization, post-polymerization functionalization, imaging, stereochemistry, and application of linear polyacetylenes.⁷⁷⁻⁸⁰ New access to cyclic versions of linear polyacetylenes creates exciting opportunities to now investigate the role of topology. The ring expansion routes extolled herein are particularly exciting because they provide access to cyclic polymers free of functional groups in the polymer backbone used to produce comparable polymers via a ring-closing. Additional advantages of these methods over traditional ring closing methods include the possibility of eliminating the presence of linear polymer impurities typical of ring closing methods and eliminating the requirement of conducting the polymerization in highly dilute conditions. Combined, these advantages and the discovery of highly active catalysts 1 and 6 enable the large-scale synthesis of cyclic polymers. Access to large quantities of cyclic polyacetylenes will accelerate the discovery of new applications and will enable the properties of polyacetylenes to be re-examined but now with a cyclic topology.

In particular, access to cyclic polyolefins, such as cyclic polypropylene, provide an opportunity to exploit the inherent properties of cyclic polymers on commercially relevant materials. Catalyst 6 is central to new cyclic polymer discoveries since it possesses high polymerization activity, broad monomer scope, and operates under mild conditions if water and oxygen are absent. New catalyst systems continue to evolve. The simplicity of Maeda's *in situ* generated initiators provide easy access to disubstituted cyclic polyacetylenes. Broadening access further, Mindiola *et al.* demonstrates that alkylidynes in general are likely to play a vital role in the further development of cyclic polymer catalyst design.

Polymerization of monosubstituted alkynes with low catalyst loadings (0.003 mol%) using 6 is now possible. However, the polymerization of disubstituted alkynes requires 5 mol% of catalyst and 10–20 mol% additive. More active systems for the polymerization of disubstituted alkynes are desirable. Additionally, the polymerization mechanisms for the catalyst

systems reported thus far need more investigation to achieve ultimate control over cyclic polymer molecular weight, stereochemistry, dispersity, and monomer scope. Developing "living" catalysts with high initiation efficiency will allow control over both the molecular weight and dispersity of the produced polymers. Living catalysts for cyclic polymer synthesis will exponentially expand the field by enabling the synthesis of cyclic block copolymers.

Improvements in the polymerization of cyclic polymers from alkynes represent a major step towards realizing commercial applications. The scalable (decagram) synthesis of cyclic polyolefins from cyclic polyacetylenes *via* hydrogenation will enable experiments that probe the role of topology on commercially relevant polymers. In the near term the next steps will be to exploit the new access to large quantities of cyclic polymers from diverse alkyne monomers to further elucidate the role of a cyclic topology on the physical properties of polymers, composites, blends, thin films, nanostructured materials, crosslinked networks, vitrimers and many other advanced materials.

Data availability

Review

The data that support the findings of this review can be found in the cited references.

Author contributions

PTB composed the manuscript, curated the figures, tabulated the data, and created the artwork. ASV edited the manuscript.

Conflicts of interest

The authors declare the following competing financial interest(s): University of Florida, P. T. B. and A. S. V. have filed patents related to this work. A. S. V. is an equity stake holder in Oboro Labs Inc.

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