Communication

Post-Polymerization Modification of Ring Opening Metathesis Polymerization (ROMP)-Derived Materials Using Wittig Reactions

Ryan Duty and Christopher Hobbs*

Department of Chemistry, Sam Houston State University, Huntsville, TX 77341, USA; rsd015@shsu.edu
* Correspondence: chobbs@shsu.edu

Received: 14 May 2020; Accepted: 28 May 2020; Published: 29 May 2020

Abstract: This communication describes our recent efforts to utilize Wittig olefination reactions for the post-polymerization modification of polynorbornene derivatives prepared through ring opening metathesis polymerization (ROMP). Polymerizing α-bromo ester-containing norbornenes provides polymers that can undergo facile substitution with triphenylphosphine. The resulting polymeric phosphonium salt is then deprotonated to form an ylide that undergoes reaction with various aryl aldehydes in a one-pot fashion to yield the respective cinnamates. These materials can undergo further modification through photo-induced [2 + 2] cycloaddition cross-linking reactions.

Keywords: wittig reaction; ROMP; post-polymerization modification

1. Introduction

Post-polymerization modifications (PPMs) offer chemists the ability to change or fine-tune a macromolecule’s overall bulk properties. These reactions are typically carried out through the covalent modification of a polymer’s end groups, backbone, and/or pendant groups [1]. Although Hermann Staudinger receives much credit (deservedly so) for the development of “polymer-analogous reactions”, these types of transformations have been carried out for much longer [2]. However, recent decades have been witness to an explosion of literature examples describing various types of PPM reactions [3–5], including the modification of bio-based [6] polymers and plastic upcycling [7,8]. However, this renaissance is mostly attributable to the invention and development of the concepts of “click” and “green” chemistry [9,10], as well as controlled [11] “living” polymerization techniques, such as reversible addition/fragmentation chain transfer (RAFT) polymerization and atom transfer radical polymerization (ATRP) [12,13]. For example, Zhao et al. have recently shown that polymers containing Meldrum’s acid derivatives (prepared using RAFT) could undergo post-polymerization Knoevenagel reactions [14], while Sumerlin and coworkers showed that keto-enol tautomerization could be harnessed for the PPM of materials prepared from RAFT [15]. Bode’s laboratory recently showed that it was possible to utilize ATRP to prepare acyltrifluoroborate-containing polymers that could undergo post-synthetic modifications with functionalized amines [16].

In addition to RAFT and ATRP, ring opening metathesis polymerization (ROMP) [17]) has become one of the most widely utilized living polymerization techniques. This is thanks, in part, to the high functional group tolerance (and bench-stability) of Grubbs-type initiators that can easily facilitate the polymerization of high- and medium-strain cyclic monomers (i.e., norbornenes, cyclooctenes, etc.) for a number of applications [18]. Despite the high functional group tolerance of Grubbs initiators, there exist circumstances in which PPM reactions are required. For example, Nelson and coworkers recently described the use of Dewar heterocycles as monomers for ROMP [19]. Such species could be polymerized to provide polymeric -lactams that could undergo subsequent
modifications to yield water-soluble -amino acid polymers, functionalities that may be less well-tolerated in ROMP. Kiesling’s laboratory reported on the ROMP of oxazinone-based monomers. The resulting poly(oxazinone)s could undergo PPM with a variety of oxide ethers [20]. This same laboratory pioneered the use of “activated” N-hydroxsuccinimide (NHS) ester-containing ROMP materials, that could undergo PPMs to form biologically-active polymers [21]. Our lab has recently adopted this technique for the preparation of a polymer-supported DMAP catalyst [22]. Earlier examples describing the utility of multiple “click” reactions as tools for PPM of ROMP-based materials exist, as do those detailing PPM strategies to prepare polymer/protein conjugates [23–27].

For the last few years, our laboratory [28–33] (among others [34–36]) has been interested in the utilization of a “thio-bromo” click [37,38] reaction for carrying out PPMs. Of particular interest to us is the modification of polymeric materials prepared from ROMP [21]. The marriage of these two chemistries required the polymerization of norbornene/cyclooctene derivatives-decorated with electrophilic α-bromo ester moieties that could undergo PPMs through reactions with nucleophilic thiols. The utility of pendant alkyl bromides has been illustrated for other substitution reactions as well [39–45]. Following our initial reports, we reasoned that this functionality may serve as a convenient handle to introduce other useful functional groups, such as acrylates and cinnamates; these are α,β-unsaturated carbonyls that can undergo further modifications through various Michael reactions, as well as photo-facilitated [2 + 2] cycloadditions. Polymers decorated with acrylates and cinnamates have indeed been prepared through ROMP [46–49]. However, most of these rely on multistep monomer syntheses (≥ 2 steps) utilizing standard, substitution chemistry in a pre-polymerization strategy.

To this day, the Wittig reaction remains a highly efficient route toward the preparation of alkenes, including acrylate and cinnamate esters [50]. For this reason, it is a mainstay in sophomore organic chemistry courses. Although the Wittig and Wittig-like reactions have been used to prepare polymers and modify the termini of ROMP-based materials [51], it has not, to our knowledge, been utilized as PPM reaction for the modification of polymer-pendant groups, especially not with ROMP. The reason for its underutilization is, more-than-likely, the lack of atom-economy, a hallmark of “green chemistry”. This low atom economy is a consequence of the formation of a stoichiometric equivalent of a triphenylphosphine oxide which is, by no means, benign. For this reason, this reaction is often thought to be antithetical to the green chemistry philosophy (although catalytic variants have been developed [52]). Product purification only exacerbates this issue, as the removal of the byproduct often requires column chromatography, generating even more waste. However, this actually may not be as problematic for modifying polymers (especially on academic-lab scales), since polymer purification is commonly carried out by solvent precipitation. While not inherently “green”, solvent precipitations are relatively green, when compared to column chromatography (which can generate excessive amounts of solvent waste). Furthermore, this can offer a simpler way to isolate the phosphine oxide byproduct, which could presumably be reduced and recycled. With this in mind, we developed a straightforward method for modifying poly(norbornene) derivatives using Wittig reactions.

2. Results and Discussion

Our work began with the preparation of α-bromo ester-decorated norbornene 2 using procedures adapted from the literature [53]. This involved subjecting commercially available alcohol (mixture of endo and exo) to a reaction with bromoacetyl bromide in the presence of sodium bicarbonate for 2 h (Scheme 1). This led to the isolation of 2 in 65% yield. 2 was then subjected to ROMP in the presence of Grubbs 3rd generation initiator 3 [54] using [M]:[I] ratio of 250:1 in methylene chloride for 2 h. Subsequent quenching with ethyl vinyl ether (EVE) and precipitation into methanol led to the isolation of polymer 4 as gummy solid in 80% yield. Product formation was confirmed by the disappearance of the olefinic signals between 6.2 and 5.9 ppm and the appearance of broad signals between 5.48 and 5.07 ppm in the 1H NMR spectrum. Gel permeation chromatography (GPC) revealed that 4 possesses Mw and D values of 64,500 Da (Mw, theor. = 61,500 Da) and 1.03, respectively (see details of 1H NMR spectra and GPC chromatograms in supplementary materials).
In order to prepare the requisite polymeric ylide, we next subjected polymer 4 to a substitution reaction with triphenylphosphine in THF for 24 h. Phosphonium salt 5 was isolated in 71% yield by solvent precipitation into ether. Confirmation was obtained by \(^1\)H NMR, which showed a disappearance of the broad singlet at 3.85 ppm, corresponding to the \(\alpha\)-protons geminal to the bromide; this signal was shifted upfield and overlapped with the polymer’s olefinic signals. Phosphonium salt 5 was next subjected to reaction with sodium bicarbonate to generate a polymeric ylide (Scheme 2). Unfortunately, we found this ylide formation to be problematic, since it led to the formation of a completely insoluble gel through, presumably, unexpected cross-linking between the ylide and ester moieties [55].

We did find, however, that Wittig modifications could be carried out without the need to isolate the polymeric ylide. A biphasic, one-pot reaction of 5 (in methylene chloride), with a saturated, aqueous solution of NaHCO\(_3\) in the presence of excess benzaldehyde led to the formation of polymeric cinnamate ester 6 in 70% yield, after quenching with dilute acid (Scheme 3). Product formation was confirmed by \(^1\)H NMR spectroscopy, which showed the appearance of signals between 7.8–7.1 ppm and a doublet at 6.4 ppm, indicative of aryl and olefinic protons, respectively (Figure 1).
Figure 1. Photographs of crude, partial $^1$H NMR spectra of 5 (top) and 6 (bottom) showing the chemical shifts of the indicated protons for 6.

We next set out to determine the aldehyde scope of this reaction. As can be seen in Table 1, aldehydes containing various electron withdrawing groups were well tolerated, providing cinnamate products in moderate to good yields with good E/Z ratios (Table 1, entries 1–6). Heterocyclic aromatic aldehydes (e.g., furfural) could also be utilized (Table 1, entry 7). Unfortunately, this system is not without limitations; it was discovered that aliphatic aldehydes and aromatic aldehydes containing electron donating groups resulted in substantial cross-linking, as indicated by the complete gelation of the reaction mixture (Table 1, entries 8–10). This is probably caused by the lower electrophilicity of carbonyls containing electron donating groups.

Table 1. Aldehyde scope for Wittig reactions.  

| Entry | RCHO | Prd. | E/Z | Yield (%) |
|-------|------|------|-----|-----------|
| 1     |       | 6    | 18:1 | 70        |
| 2     |       | 7    | 17:1 | 95        |
| 3     |       | 8    | 7:1  | 65        |
| 4     |       | 9    | 12:1 | 83        |
| 5     |       | 10   | 14:1 | 90        |
| 6     |       | 11   | 7:1  | 89        |
| 7     |       | 12   | 5:1  | 65        |
Cinnamic esters (as well as other $\alpha,\beta$-unsaturated carbonyls) have been shown to undergo synthetically useful [2 + 2] cycloaddition reactions under UV light [46]. Gratifyingly, we found that the exposure of polymer 6 (in THF) to ca. 30 h of sunlight (albeit non-continuous) was sufficient to effect the formation of a completely insoluble material that was produced, presumably, through [2 + 2] cycloaddition cross-linking reactions (Figure 2).

Figure 2. (a) Cross-linking of 6 through [2 + 2] cycloaddition. (b) Photograph of THF solution of 6 before exposure to sunlight (left) and photograph after exposure to sunlight and decantation of excess solvent (right).

3. Conclusions

In conclusion, we have shown that a standard Wittig reaction can be used for the post-polymerization modification of materials prepared from ROMP. Polymerization of $\alpha$-bromo ester-containing monomers, followed by substitution with triphenylphosphine, led the way to carry out a one-pot reaction in which polymeric ylide formation and Wittig reaction could be carried out. This provided polymeric cinnamate esters containing electron withdrawing groups in moderate to good yield. Furthermore, these materials could be subjected to cross-linking reactions in the presence of sunlight, presumably occurring through [2 + 2] cycloadditions. Experiments designed to expand the scope of this process as well as exploring further PPM reactions utilizing Michael and Diels–Alder reactions are currently underway.

Supplementary Materials: The following are available online at www.mdpi.com/2073-4360/12/6/1247/s1,

$^1$H NMR spectra and GPC chromatograms.

Author Contributions: Conceptualization, C.E.H.; methodology, R.D.; writing—original draft preparation, C.E.H. and R.D.; writing—review and editing, C.E.H. and R.D.; funding acquisition, C.E.H. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the National Science Foundation, grant numbers 1744700 and 1847914 the Robert A. Welch Foundation, grant number X-0011.

Conflicts of Interest: The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.
References

1. Gauthier, M.A.; Gibson, M.I.; Klok, H.-A. Synthesis of functional polymers by post-polymerization modification. *Angew. Chemie Int. Ed.* 2009, 48, 48–58, doi:10.1002/anie.200801951.

2. Günay, K.A.; Theato, P.; Klok, H.-A. Standing on the shoulders of Hermann Staudinger: Post-polymerization modification from past to present. *J. Polym. Sci. Part A Polym. Chem.* 2013, 51, 1–28, doi:10.1002/pola.26333.

3. Liu, D.; Bielawski, C.W. Post-polymerization modification of poly(vinyl ethers): A Ru-catalyzed oxidative synthesis of poly(vinyl ether)s and poly(propenyl ether)s. *Polym. Chem.* 2016, 7, 63–68, doi:10.1039/C5PY01409C.

4. Coady, D.J.; Bielawski, C.W. N-Heterocyclic carbenes: Versatile reagents for postpolymerization modification. *Macromolecules* 2006, 39, 8895–8897, doi:10.1021/ma060230d.

5. Liu, D.; Bielawski, C.W. Synthesis of degradable poly[(ethylene glycol)-co-(glycolic acid)] via the post-polymerization oxyfunctionalization of poly(ethylene glycol). *Macromol. Rapid Commun.* 2016, 37, 1587–1592, doi:10.1002/marc.201600336.

6. Farmer, T.J.; Comerford, J.W.; Pellis, A.; Robert, T. Post-polymerization modification of bio-based polymers: Maximizing the high functionality of polymers derived from biomass. *Polym. Int.* 2018, 67, 775–789, doi:10.1002/pi.5573.

7. Williamson, J.B.; Na, C.G.; Johnson, R.R.; Daniel, W.F.M.; Alexanian, E.J.; Leibfarth, F.A. Chemoselective and regioselective functionalization of isocatic polypropylene: A mechanistic and structure–property study. *J. Am. Chem. Soc.* 2019, 141, 12815–12823, doi:10.1021/jacs.9b05799.

8. Lewis, S.E.; Wilhelmy, B.E.; Leibfarth, F.A. Organocatalytic C–H fluorooalkylation of commodity polymers. *Polym. Chem.* 2020, doi:10.1039/C9PY01884K.

9. Kolb, H.C.; Finn, M.G.; Sharpless, K.B. Click chemistry: diverse chemical function from a few good reactions. *Angew. Chemie Int. Ed.* 2001, 40, 2004–2021, doi:10.1002/1521-3773(20010601)40:11<2004::AID-ANIE20043-0.CO;2-5.

10. Anastas, P.T.; Constable, D.; Jimenez-Gonzalez, C. *Green Metrics*; Wiley-VCH: Weinheim, Germany, 2018.

11. Grubbs, R.B.; Grubbs, R.H. 50th anniversary perspective: living polymerization—Emphasizing the molecule in macromolecules. *Macromolecules* 2017, 50, 6979–6997, doi:10.1021/acs.macromol.7b01440.

12. Matyjaszewski, K. Discovery of the RAFT process and its impact on radical polymerization. *Macromolecules* 2020, 53, 495–497, doi:10.1021/acs.macromol.9b02054.

13. Beers, K.L. The first dive into the mechanism and kinetics of ATRP. *Macromolecules* 2020, 53, 1115–1118, doi:10.1021/acs.macromol.9b02460.

14. Yuan, L.; He, L.; Wang, Y.; Lang, X.; Yang, F.; Zhao, Y.; Zhao, H. Two- and three-component post-polymerization modifications based on Meldrum’s acid. *Macromolecules* 2020, 53, 3175–3181, doi:10.1021/acs.macromol.0c00482.

15. Easterling, C.P.; Coste, G.; Sanchez, J.E.; Fanucci, G.E.; Sumerlin, B.S. Post-polymerization modification of polymethacrylates enabled by keto–enol tautomerization. *Polym. Chem.* 2020, 11, 2955–2958, doi:10.1039/D0PY00383B.

16. Schauenburg, D.; Divandari, M.; Neumann, K.; Speigel, C.A.; Hackett, T.; Dzang, Y.-C.; Spencer, N.D.; Bode, J.W. Synthesis of polymers containing potassium acrylitrifluoroborates (KATs) and post-polymerization ligation and conjugation. *Angew. Chemie Int. Ed.* 2020, accepted.

17. Bielawski, C.W.; Grubbs, R.H. Living ring-opening metathesis polymerization. *Prog. Polym. Sci.* 2007, 32, 1–29, doi:10.1016/j.progpolysci.2006.08.006.

18. Grubbs, R.H., Wenzel, A.G., O’Leary, D.J., Khosravi, E. (Eds.) *Handbook of Metathesis*, 2nd ed.; Wiley-VCH: Weinheim, Germany, 2015.

19. Nistanaki, S.K.; Nelson, H.M. Dewar heterocycles as versatile monomers for ring-opening metathesis polymerization. *ACS Macro Lett.* 2020, 9, 731–735, doi:10.1021/acs.macrolett.0c00227.

20. Fishman, J.M.; Zwick, D.B.; Kruger, A.G.; Kiessling, L.L. Chemoselective, postpolymerization modification of bioactive, degradable polymers. *Biomacromolecules* 2019, 20, 1018–1027, doi:10.1021/acs.biomac.8b01631.

21. Strong, L.E.; Kiessling, L.L. A General synthetic route to defined, biologically active multivalent arrays. *J. Am. Chem. Soc.* 1999, 121, 6193–6196, doi:10.1021/ja990223t.

22. Fredlund, A.; Kothapalli, V.A.; Hobbs, C.E. Phase-selectively soluble polynorbornene as a catalyst support. *Polym. Chem.* 2017, 8, 516–519, doi:10.1039/C6PY02041K.

23. Schaefer, M.; Hanik, N.; Kilbinger, A.F.M. ROMP copolymers for orthogonal click functionalizations. *Macromolecules* 2012, 45, 6807–6818, doi:10.1021/ma301061z.

24. Van Hensbergen, J.A.; Burford, R.P.; Lowe, A.B. ROMP (co)polymers with pendant alkyne side groups:
Post-polymerization modification employing thiol–yne and CuAAC coupling chemistries. *Polym. Chem.* 2014, 5, 5339–5349, doi:10.1039/C4PY00604F.

25. Kim, K.O.; Kim, J.; Choi, T.-L. Controlled ring-opening metathesis polymerization of a monomer containing terminal alkyne and its versatile postpolymerization functionalization via click reaction. *Macromolecules* 2014, 47, 4525–4529, doi:10.1021/ma502052k.

26. Romulus, J.; Henssler, J.T.; Weck, M. Postpolymerization modification of block copolymers. *Macromolecules* 2014, 47, 5437–5449, doi:10.1021/ma5018109.

27. Wright, T.A.; Page, R.C.; Konkolewicz, D. Polymer conjugation of proteins as a synthetic post-translational modification to impact their stability and activity. *Polym. Chem.* 2019, 10, 434–454, doi:10.1039/C8PY01399C.

28. Hobbs, C.E.; Vasireddy, M. Combining ATRP and ROMP with thio-bromo, copper-catalyzed, and strain-promoted click reactions for brush copolymer synthesis starting from a single initiator/monomer/click partner. *Macromol. Chem. Phys.* 2019, 220, 1800497–1800502, doi:10.1002/macp.201800497.

29. Subnaik, S.I.; Hobbs, C.E. Flow-facilitated ring opening metathesis polymerization (ROMP) and post-polymerization modification reactions. *Polym. Chem.* 2019, 10, 4524–4528, doi:10.1039/C9PY00822E.

30. Ashlin, M.; Hobbs, C.E. Post-polymerization thiol substitutions facilitated by mechanochemistry. *Macromol. Chem. Phys.* 2019, 220, 1900350–1900354, doi:10.1002/macp.201900350.

31. Ashok Kotheapalli, V.; Shetty, M.; De Los Santos, C.; Hobbs, C.E. Thio-bromo “click”, post-polymerization strategy for functionalizing ring opening metathesis polymerization (ROMP)-derived materials. *J. Polym. Sci. Part A Polym. Chem.* 2016, 54, doi:10.1002/pola.27801.

32. Grubb, J.; Carosio, F.; Vasireddy, M.; Moncho, S.; Brothers, E.N.; Hobbs, C.E. Ring opening metathesis polymerization (ROMP) and thio-bromo “click” chemistry approach toward the preparation of flame-retardant polymers. *J. Polym. Sci. Part A Polym. Chem.* 2018, 56, 645–652, doi:10.1002/pola.28939.

33. Yao, Q.; Gutierrez, D.C.; Hoang, N.H.; Kim, D.; Wang, R.; Hobbs, C.; Zhu, L. Efficient codelivery of paclitaxel and curcumin by novel bottlebrush copolymer-based micelles. *Mol. Pharm.* 2017, 14, 2378–2389, doi:10.1021/acs.molpharmaceut.7b00278.

34. Kumar, S.; Deike, S.; Binder, W.H. One-pot synthesis of thermostressive amyloidogenic peptide–polymer conjugates via thio–bromo “click” reaction of RAFT polymers. *Macromol. Rapid Commun.* 2018, 39, 1700507, doi:10.1002/marc.201700507.

35. Döhler, D.; Kaiser, J.; Binder, W.H. Supramolecular H-bonded three-arm star polymers by efficient combination of RAFT polymerization and thio-bromo “click” reaction. *Polymer* 2017, 122, 148–158, doi:10.1016/j.polymer.2017.06.067.

36. Biewend, M.; Neumann, S.; Michael, P.; Binder, W.H. Synthesis of polymer-linked copper(i) bis(N-heterocyclic carbene) complexes of linear and chain extended architecture. *Polym. Chem.* 2019, 10, 1078–1088, doi:10.1039/C8PY01751D.

37. Rosen, B.M.; Lligadas, G.; Hahn, C.; Percec, V. Synthesis of dendrimers through divergent iterative thio-bromo “click” chemistry. *J. Polym. Sci. Part A Polym. Chem.* 2009, 47, 3931–3939, doi:10.1002/pola.23519.

38. Rosen, B.M.; Lligadas, G.; Hahn, C.; Percec, V. Synthesis of dendritic macromolecules through divergent iterative thio-bromo “click” chemistry and SET-LRP. *J. Polym. Sci. Part A Polym. Chem.* 2009, 47, 3940–3948, doi:10.1002/pola.23518.

39. Barlow, T.R.; Brendel, J.C.; Perrier, S. Poly(bromoethyl acrylate): A reactive precursor for the synthesis of functional RAFT materials. *Macromolecules* 2016, 49, 6203–6212, doi:10.1021/acs.macromol.6b00721.

40. Li, Z.-L.; Sun, L.; Ma, J.; Zeng, Z.; Jiang, H. Synthesis and post-polymerization modification of polynorbornene bearing dibromomaleimide side groups. *Polymer* 2016, 84, 336–342, doi:10.1016/j.polymer.2016.01.020.

41. Binder, W.H.; Kluger, C. Combining ring-opening metathesis polymerization (ROMP) with sharpless-type “click” reactions: an easy method for the preparation of side chain functionalized poly(oxynorbornenes). *Macromolecules* 2004, 37, 9321–9330, doi:10.1021/ma0480087.

42. Yang, S.K.; Week, M. Modular covalent multifunctionalization of copolymers. *Macromolecules* 2008, 41, 346–351, doi:10.1021/ma071052k.

43. Brendel, J.C.; Schmidt, M.M.; Hagen, G.; Moos, R.; Thelakkat, M. Controlled synthesis of water-soluble conjugated polyelectrolytes leading to excellent hole transport mobility. *Chem. Mater.* 2014, 26, 1992–1998, doi:10.1021/cm500500t.

44. Sugiyama, K.; Azuma, H.; Watanabe, T.; Ishizone, T.; Hirao, A. Anionic polymerization of 2-haloethyl methacrylates. *Polymer* 2003, 44, 4157–4164, doi:10.1016/S0032-3861(03)00350-1.

45. García-Loma, R.; Albéniz, A.C. Poly(ω-bromoalkynorbornenes-co-norbornene) by ROMP-hydrogenation:
A robust support amenable to post-polymerization functionalization. RSC Adv. 2015, 5, 70244–70254, doi:10.1039/C5RA15187B.

46. Wang, Y.; Noga, D.E.; Yoon, K.; Wojtowicz, A.M.; Lin, A.S.P.; García, A.J.; Collard, D.M.; Weck, M. Highly porous crosslinkable PLA-PNB block copolymer scaffolds. Adv. Funct. Mater. 2008, 18, 3638–3644, doi:10.1002/adfm.200800385.

47. Hreha, R.D.; Haldi, A.; Domercq, B.; Barlow, S.; Kippelen, B.; Marder, S.R. Synthesis of acrylate and norbornene polymers with pendant 2,7-bis(diarylamino)fluorene hole-transport groups. Tetrahedron 2004, 60, 7169–7176, doi:10.1016/j.tet.2004.06.069.

48. Lowe, A.B.; Liu, M.; van Hensbergen, J.A.; Burford, R.P. Combining ring-opening metathesis polymerization and thiol-ene coupling chemistries: Facile access to novel functional linear and nonlinear macromolecules. Macromol. Rapid Commun. 2014, 35, 391–404, doi:10.1002/marc.201300744.

49. Wu, Z.; Liu, L.; Cheng, P.; Fang, J.; Xu, T.; Chen, D. Reusable gold nanorod/liquid crystalline elastomer (GNR/LCE) composite films with UV-triggered dynamic crosslinks capable of micropatterning and NIR actuation. J. Mater. Chem. C 2019, 7, 14245–14254, doi:10.1039/C9TC04651H.

50. Rocha, D.H.A.; Pinto, D.C.G.A.; Silva, A.M.S. Applications of the wittig reaction on the synthesis of natural and natural-analogue heterocyclic compounds. Eur. J. Org. Chem. 2018, 2018, 2443–2457, doi:10.1002/ejoc.201800523.

51. Chen, Y.; Abdellatif, M.M.; Nomura, K. Olefin metathesis polymerization: Some recent developments in the precise polymerizations for synthesis of advanced materials (by ROMP, ADMET). Tetrahedron 2018, 74, 619–643, doi:10.1016/j.tet.2017.12.041.

52. Lao, Z.; Toy, P.H. Catalytic Wittig and aza-Wittig reactions. Beilstein J. Org. Chem. 2016, 12, 2577–2587.

53. Taylor, M.T.; Nelson, J.E.; Suero, M.G.; Gaunt, M.J. A protein functionalization platform based on selective reactions at methionine residues. Nature 2018, 562, 563–568, doi:10.1038/s41586-018-0608-y.

54. Love, J.A.; Morgan, J.P.; Trnka, T.M.; Grubbs, R.H. A practical and highly active ruthenium-based catalyst that effects the cross metathesis of acrylonitrile. Angew. Chemie Int. Ed. 2002, 41, 4035–4037, doi:10.1002/1521-3773(20021104)41:21<4035::AID-ANIE4035>3.0.CO;2-I.

55. Uijttewaal, A.P.; Jonkers, F.L.; Van der Gen, A. Reactions of esters with phosphorus ylides. 2. Mechanistic aspects. J. Org. Chem. 1978, 43, 3306–3311, doi:10.1021/jo00411a009.

© 2020 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).