Click Nucleophilic Conjugate Additions to Activated Alkynes: Exploring Thiol-yne, Amino-yne, and Hydroxyl-yne Reactions from (Bio)Organic to Polymer Chemistry

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ABSTRACT: The 1,4-conjugate addition reaction between activated alkynes or acetylenic Michael acceptors and nucleophiles (i.e., the nucleophilic Michael reaction) is a historically useful organic transformation. Despite its general utility, the efficiency and outcomes can vary widely and are often closely dependent upon specific reaction conditions. Nevertheless, with improvements in reaction design, including catalyst development and an expansion of the substrate scope to feature more electrophilic alkynes, many examples now present with features that are congruent with Click chemistry. Although several nucleophilic species can participate in these conjugate additions, ubiquitous nucleophiles such as thiols, amines, and alcohols are commonly employed and, consequently, among the most well developed. For many years, these conjugate additions were largely relegated to organic chemistry, but in the last few decades their use has expanded into other spheres such as biorganic chemistry and polymer chemistry. Within these fields, they have been particularly useful for bioconjugation reactions and step-growth polymerizations, respectively, due to their excellent efficiency, orthogonality, and ambient reactivity. The reaction is expected to feature in increasingly divergent application settings as it continues to emerge as a Click reaction.

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Special Issue: Click Chemistry
Received: October 2, 2020
1. INTRODUCTION

Alkynes that have electron withdrawing substituents in conjugation with the acetylenic moiety (i.e., activated alkynes) are tremendously useful synthetic intermediates as a result of their combination of good bench stability with high reactivity. Although these compounds can react via numerous pathways, including as dienophiles in cycloaddition reactions, they are most renowned for their interaction with nucleophiles in conjugate addition reactions. For alkynes in immediate conjugation with carbonyl moieties such as ketones (ynones), esters (propiolates), or amides (propiolamides), nucleophilic attack commonly furnishes a 1,2-addition product (at the carbonyl carbon) or the 1,4-addition product (at the acetylenic moiety) in addition to other more esoteric products (Figure 1). The former reaction is usually induced by hard or strong nucleophiles, including organolithiums or Grignard reagents, while the latter is famously more efficient and achieved using softer nucleophiles.

The 1,4-conjugate addition reaction, which is colloquially referred to as the “Michael addition” or “Michael reaction”, remains an extremely useful synthetic transformation. Although it was first established using carbon-centered nucleophiles in carbon−carbon bond forming reactions, the use of heteronucleophiles has continued to gain prominence as a consequence of their superior reaction efficacy and the general ubiquity of heteroatoms in organic compounds. Specifically, the heteronucleophilic 1,4-conjugate addition has been employed extensively for the construction of heterocycles and for highly efficient conjugation to biological compounds for bioorganic and medicinal chemistry applications. This especially applies to amine- and thiol-based nucleophiles which are abundant functionalities in many naturally occurring compounds, including amino acid residues of proteins or biopolymers such as polysaccharides. The high performance of nucleophilic conjugate additions has translated beyond organic transformations and into other disciplines, most notably polymer chemistry. In particular, it has stimulated increased interest in step-growth polymerizations to afford heteroatom-containing polymers of novel compositions, replacing inefficient condensation chemistries (such as acid-catalyzed esterification) which struggle to reach full conversion and lower the molecular weight of the resultant polymers. However, heteronucleophilic Michael addition reactions often afford very high molecular weight, well-defined step-growth polymers under comparatively mild reaction conditions and in some cases the polymerizations may proceed spontaneously at ambient temperature. These considerations are significant because polymer molecular weight is intimately connected to the performance of the end-material properties. More recently, Michael addition reactions have featured in postpolymerization conjugation reactions to afford functional materials and they have also been used to create high performance network polymers such as hydrogels as a consequence of excellent compatibility with aqueous environments.

Regardless of application setting and the overall utility of the transformation, reaction outcomes have historically been divergent, ranging from poor yields even at high temperatures to quantitative conversion at ambient temperatures without the use of a catalyst; the eventual outcome of the reaction is strongly reliant upon several aspects including electrophilicity of the activated alkyne, the strength of the nucleophile, solvent polarity, and the nature of the catalytic species in many cases. Even though some reactions are uncatalyzed, the majority are not, and judicious catalyst choice often has significant bearing on reaction efficiency and selectivity. In this Review, we have focused on reactions that are most congruent with classical Click chemistry criteria that were first outlined nearly 20 years ago. While some reported examples satisfy all (or most) Click requirements, others are less “Click-like” but are still highlighted because of either their historical significance in the development of this chemistry or to show where improvements are required to make the reactions more efficient and/or selective. Moreover, we have narrowed our focus to reactions that employ the most abundant and versatile nucleophiles, which includes thiols (thiol-yne), amines (amino-yne), and alcohols (hydroxyl-yne) because these have most successfully translated to other chemistry spheres outside of organic chemistry. Other heteroatom-centered nucleophiles (including boron, phosphorus, selenium, and tellurium) or dinucleophilic species (such as hydrazines, amidines, or hydroxylamines) are beyond the scope of this review because these are comparatively inefficient and/or require inert atmospheres, thus largely precluding them from being reasonably termed as Click reactions and hence inhibiting their widespread use beyond organic synthesis. In addition, they have been well documented in other recent reviews, and in this Review, we have focused on reactions that are most congruent with classical Click chemistry criteria that were first outlined nearly 20 years ago. While some reported examples satisfy all (or most) Click requirements, others are less “Click-like” but are still highlighted because of either their historical significance in the development of this chemistry or to show where improvements are required to make the reactions more efficient and/or selective. Finally, we made an effort to emphasize studies relating to polymeric materials and bioconjugate chemistry as these are rapidly expanding areas in which the heteronucleophilic Michael reaction is envisioned to be increasingly impactful as a consequence of its orthogonality, modularity, simple reaction requirements, and overall efficiency.

1.1. Substrate Overview

In this brief section, we will provide some context on the availability of substrates (nucleophiles and electrophilic alkynes) that commonly react in Michael reactions. It is by no means exhaustive, rather it is intended to provide a sense of the accessibility of frequently encountered substrates (Figure 2).

Because of their potent nucleophilicity, thiols are the most commonly used heteroatomic substrate in Click Michael reactions. Unfortunately, they lack relative abundance in commodity compounds and only naturally occur in a few amino acids. As a result, commercially sourced thiol-containing compounds can be relatively expensive in comparison to alcohols or amines. Still, many desired thiol substrates can be produced from alcohols, albeit in multistep schemes typically via a thioester intermediate or disulfide, but more direct catalytic pathways are also available. On the other hand, connected to the performance of the end-material properties. More recently, Michael addition reactions have featured in postpolymerization conjugation reactions to afford functional materials and they have also been used to create high performance network polymers such as hydrogels as a consequence of excellent compatibility with aqueous environments.
amines are abundant building blocks, particularly diamines due to their commercial importance in polyamides and polyurethanes. Finally, the hydroxyl group is among the most central functionalities in organic chemistry. However, alcohols are generally poorer nucleophiles compared to thiols and amines, often requiring unusual catalysts and/or conditions to react efficiently in conjugate additions.

In general, many of the electron deficient alkynes that are presented in this Review are isolated from readily accessible precursors using convenient synthetic techniques. Propiolates are arguably the most abundant substrate in associated literature. They are commonly synthesized using straightforward acid-catalyzed esterification (or analogous protocols such as the Steglich esterification for more challenging substrates) of the commodity chemical propionic acid or via simple O-alkylation of the carboxylate salt. It should be noted that the more reactive propiolyl chloride can also be esterified, however it is only transiently stable and can quickly decompose to hazardous byproducts. The synthesis of ynones is also very versatile, with many synthetic pathways available. Transition metal-catalyzed coupling between terminal alkynes and functional acyl chlorides enables a modular route to an expansive library of aromatic or alkyl ynones. Another prevalent method is the reaction between metalled alkynes and Weinreb–Nahm amides to yield desired ynone structures. However, a more exhaustive account of ynone synthesis is well described in another review. The synthesis of propiolamides can often be more challenging than propiolates because the condensation reaction between amines and propionic acid derivatives competes with the amine conjugate addition (i.e., amine-yne reaction) to the electron deficient alkyne. However, reasonable yields can be obtained via carefully controlled condensation reactions between propiolates and amines. Additionally, there are other methods that afford propiolamides such as acylation reactions employing carbamoyl chlorides, photocatalyzed decarboxylative ynonylation, hydrogenative coupling of terminal alkynes with formamides, or palladium-catalyzed multicomponent reactions among alkynes, isonitriles, and carboxylates. More esoteric electron deficient alkynes such as ethynylsulfones are conveniently isolated from sulfonyl chlorides, sulfonyl hydrazides, sulfinates, or via multicomponent reactions, for example, photocatalyzed coupling of sulfur dioxide and alkynyl bromides. While their use as a substrate in Michael reactions is comparatively rare next to other activated alkynes, they are still very reactive in reported examples and this provides ample opportunity for further work. Propiolonitriles are also susceptible to 1,4-conjugate additions from heteronucleophiles, however, they are not as common in this context due to a lack of functional modularity.

2. NUCLEOPHILIC THIOL-YNE REACTIONS
2.1. Overview and Historical Development

Soon after Arthur Michael’s seminal report on the conjugate addition of carbon nucleophiles to unsaturated substrates, Ruhemann et al. discovered the heteronucleophilic conjugate addition reaction between sodium thiophenate and propiolates to furnish α,β-unsaturated esters (Scheme 1). These initial explorations were conducted at ambient temperature with yields in excess of 80% using the sodium thiophenate nucleophile, but the reaction produced large exotherms that led to side products. Despite the use of strong bases needed to activate the nucleophiles, this study would lay the groundwork to eventually enable the nucleophilic thiol-yne Michael addition to become a versatile Click reaction in modern chemistry.

In the last several decades, the nucleophilic thiol-yne addition has advanced to a point where most, if not all, Click criteria are fulfilled. Quantitative conversions can be observed under ambient reaction conditions, which is undoubtedly aided by the high nucleophilicity of the thiolate anion and/or the use of highly electrophilic alkynes. Base catalysis has also been a key development in this area and further increased the already large thermodynamic driving force of the thiol-yne reaction, consequently producing exotherms which are frequently reported in the literature. But perhaps the most attractive Click feature is the expansive modularity of the thiol-yne conjugate addition. With numerous electron-withdrawing groups, β-carbon substituents, and building blocks containing thiols, the reaction can be exquisitely tailored for a desired outcome.

Stereospecificity is another key aspect of the Click umbrella and is pertinent to the thiol-yne nucleophilic reaction and its unsaturated products. Ruhemann’s initial reports did not indicate the stereochemical configuration of the reaction product from sodium thiophenate and phenylpropionate, but a later study suggested the products to be primarily Z-isomers. A firmer understanding for the observed reaction stereochromies arrived many decades later when the anti-addition rule (which largely applies for most conjugate additions) to the electron-deficient alkyne was formulated from a detailed study into the thiol-yne conjugate addition reaction by Truce et al. In this report, the addition of sodium p-toluenethiolate to numerous electrophilic alkynes in methanol established the Z-isomer as the dominant product, but occasionally the E-isomer can be prevalent in small quantities. The authors initially assumed that isomerization of the product was responsible for the observed small quantity of the E-isomer (and this is still sometimes a concern that can obfuscate modern reaction outcomes). Heating the thiol-yne Michael adducts to 50 °C in the presence of catalytic amounts of HCl or sodium p-
counterparts, are also reversible, undergoing retro-Michael thiol-yne Michael adducts, much like the well-known thiol-ene will be highlighted in many forthcoming examples. Finally, greater than 50 prevalence of isomerization (resulting from temperatures or alkaline environments. This topic has been a rising research focus in recent years as interest in dynamic covalent chemistries has significantly grown.

Nevertheless, analysis of other historical references indicating E/Z stereochemistry should be treated with caution as the prevalence of isomerization (resulting from temperatures greater than 50 °C, UV light or base catalysis) can obfuscate the actual obtained stereochemistry in the reaction products. Since the seminal works by Truce et al., the importance of controlling E/Z stereochemistry in unsaturated reaction products has become a central research theme. Through variation of base catalyst, temperature, and solvent conditions, E/Z isomerization can be effectively tuned, which will be highlighted in many forthcoming examples. Finally, thiol-yne Michael adducts, much like the well-known thiol-ene counterparts, are also reversible, undergoing retro-Michael addition reactions under various conditions such as high temperatures or alkaline environments. This topic has been a rising research focus in recent years as interest in dynamic covalent chemistries has significantly grown.

In addition to the nucleophilic pathway for the thiol-yne reaction, radical or thermally driven reactions can also ensue and complicate reaction selectivity and products. However, it should be noted that the thermally initiated thiol additions were also found to be inhibited in the presence of a radical trapping agent, which suggests that the thermal mechanism is likely facilitated via radical pathways. As it stands, the radical variant of the thiol-yne reaction has traditionally featured high conversions and good functional group tolerance which led to its growth in both organic and material chemistry alike. However, this method is inherently less controllable than the nucleophilic mechanism, primarily due to the facile thiol conjugate addition (higher rate than the initial addition) to the \( \alpha,\beta \)-unsaturated product when in the presence of radicals. As well as double addition side products, the regio- and stereochemistry of resulting adducts increases the number of possible combinations, and a simple radical-mediated thiol-yne addition may yield up to six distinct reaction products (Scheme 2). Together, these factors exclude the radical-based thiol-yne reaction from Click categorization, unlike the nucleophilic variant that is a focus of this Review.

**Table 1. Reaction between Sodium \( p \)-Toluenethiolate and Various Electrophilic Alkynes in Methanol and Subsequent Isomerization of Unsaturated Product with Sodium \( p \)-Toluenethiolate Unless Stated Otherwise**

<table>
<thead>
<tr>
<th>( R_1 )</th>
<th>Conversion (%)</th>
<th>Initial E/Z</th>
<th>Final E/Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>–CN</td>
<td>100</td>
<td>0/100</td>
<td>67/33</td>
</tr>
<tr>
<td>–SO(_2)C(_6)H(_4)-P</td>
<td>65</td>
<td>0/100</td>
<td>100/0</td>
</tr>
<tr>
<td>–C(_6)H(_4)NO(_2)-P</td>
<td>98</td>
<td>0/100</td>
<td>100/0</td>
</tr>
<tr>
<td>–CO(_2)Me</td>
<td>93</td>
<td>8/92</td>
<td>78/22</td>
</tr>
<tr>
<td>–COMe</td>
<td>93</td>
<td>18/82</td>
<td>78/22</td>
</tr>
<tr>
<td>–CONH(_2)</td>
<td>97</td>
<td>13/93</td>
<td>77/23</td>
</tr>
</tbody>
</table>

*Dilute HCl used as isomerization catalyst.*

toluenethiolate in methanol afforded higher quantities of the more stable \( E \)-isomer under these conditions. However, control experiments performed on the adduct formed from methyl propiolate and \( p \)-toluenethiolate at a range of temperatures (0–35 °C) with and without radical inhibitor indicated that both radical and thermal isomerization would not proceed under the initial reaction conditions for the thiol-yne Michael addition. Interestingly, the \( E \)-isomer was observed only in adducts containing a carbonyl moiety; it was postulated that the carbonyl group enabled the formation of a stable enol intermediate enabling small quantities of the more thermodynamically stable \( E \)-isomer to form directly.

Nevertheless, analysis of other historical references indicating \( E/Z \) stereochemistry should be treated with caution as the prevalence of isomerization (resulting from temperatures greater than 50 °C, UV light or base catalysis) can obfuscate the actual obtained stereochemistry in the reaction products. Since the seminal works by Truce et al., the importance of controlling \( E/Z \) stereochemistry in unsaturated reaction products has become a central research theme. Through variation of base catalyst, temperature, and solvent conditions, \( E/Z \) isomerization can be effectively tuned, which will be highlighted in many forthcoming examples. Finally, thiol-yne Michael adducts, much like the well-known thiol-ene counterparts, are also reversible, undergoing retro-Michael addition reactions under various conditions such as high temperatures or alkaline environments. This topic has been a rising research focus in recent years as interest in dynamic covalent chemistries has significantly grown.

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**Scheme 2. Radical Mediated Reaction of Terminal Alkynes and Monofunctional Thiyil Radicals with the Six Observable Products**

2.2. Trends in Reactivity

This straightforward conjugate addition reaction is now a multifaceted tool that has been utilized in a variety of settings, however, understanding the effects of reaction conditions (including solvent), additives, or catalysts and the nature of the reactants have been key to recent advances. In this section, we have classified each sub-section according to these primary reaction conditions, with the aim of demonstrating how these parameters can influence the reaction outcomes for the thiol-yne reaction and whether Click criteria (related to scope, efficiency, stereospecificity, and orthogonality) can be satisfied.

2.2.1. Substrate Effects. Thiols, and corresponding thiolate anions, are excellent nucleophiles as a consequence of the large atomic radius of sulfur that is able to stabilize charge effectively. With that said, the high nucleophilicity can lead to competing reactions with other functional groups present in the electrophile. Historic reactions of carbonyl-based electrophiles (propiolates and ynoles) with thiolate anions often led to competing 1,2-addition reactions. In modern examples, these byproducts are rarely reported due to milder conditions that frequently employ only catalytic amounts of base to enhance the nucleophilicity of the thiol. Thiols are notorious for their oxidation to form disulfide species, however, this formation is not kinetically competitive with the reaction rates for conjugate addition reactions and thus does significantly limit their utility in the presence of the open atmosphere which is an important requisite for Click reactions. It has also been noted that some phosphine catalysts...
(such as dimethylphenylphosphine) can even effectively reduce disulfide bonds under ambient conditions, thus further supporting excellent reaction efficacy under ambient atmosphere.69

A survey of the literature reveals there is a lack of fundamental studies that detail the effect of thiol structure (such as acidity or sterics) on the outcome of nucleophilic thiol-yne reactions. Nevertheless, some comparisons can be drawn from the analogous nucleophilic thiol-ene reaction which has been investigated in greater detail. A detailed mechanistic study of the nucleophilic thiol-ene reaction of hexyl acrylate with various thiols reported that the reaction rate (and overall conversion) was dependent upon the acidity of the thiol (Table 2).60 The addition with ethyl thioglycolate reached near quantitative conversion after 120 s, but the corresponding reaction with hexanethiol only achieved 19% conversion after 500 s. This relationship could be reasonably expected to hold with increasingly acidic thiols (thiophenols and thioacids) as well as correlate to the thiol-yne Michael addition. As it stands, no systemic study details the correlation and thioacids) as well as correlate to the thiol-yne Michael addition.

Table 2. Relationship of Thiol Acidity to Reaction Conversion and Rate for the Thiol-ene Michael Addition of Various Thiols to Hexylacrylate60

<table>
<thead>
<tr>
<th>Thiol</th>
<th>pKa</th>
<th>Conversion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-hexylamine</td>
<td>7.9</td>
<td>99</td>
</tr>
<tr>
<td>Thiol, 22 °C</td>
<td>9.3</td>
<td>61</td>
</tr>
<tr>
<td>C6H5O</td>
<td>10.6</td>
<td>19</td>
</tr>
</tbody>
</table>

There are a variety of acetylenic Michael acceptors that have been investigated for thiol-yne additions which include the following electron withdrawing groups (including ester, ketone, amide, sulfone, and cyano) in conjunction with the alkene unit. A noteworthy study investigated the reactivity differences (reaction rates and product stereochemistry) between propiolamides, propiolates, and ynoles in base-catalyzed conjugate addition reactions with thiol terminated peptides in aqueous acetonitrile solvent mixtures at ambient temperature (Table 3).67 Quantitative yields were observed for the alkyl and aryl terminal ynoles, while methylpropiolate (30%) and phenylpropiolamide (<5%) gave poorer yields. The rate of reaction followed the same trend and correlated to the electron-withdrawing ability of the functional group in conjugation with the thiol, where ketone > ester > amide. It was also noted that use of excess electrophile led to competing addition to the primary amine of alanine also present in the peptide substrate; this was only observed when using terminal ynoles (where R1 = Ph, R2 = H, and R3 = CH2CH2Ph, and R4 = H). In addition to studying the effect of the withdrawing group, the influence of β-carbon substitution was also investigated (Table 3). A lower yield is observed for ynoles that possess β-substituted n-butyl groups in neutral water/acetonitrile mixtures (<5%), however, quantitative yields are observed in pH = 8 buffered mixtures. The n-butyl groups seem to have little effect on the electronic nature of the ynone as it relates to reactivity and appear to only present a steric barrier for the thiolate addition.

The E/Z stereochemical outcome of the unsaturated product is also affected by the electronics of the Michael acceptor. Shiu et al. reacted a protected cysteine with various electrophilic alkenes in aqueous acetonitrile conditions (Table 4).67 Quantitative functionalization of the thiol moiety in buffered water/acetonitrile mixture (pH = 8) was achieved with all electrophiles, while the propiolamide failed to reach full conversion in unbuffered solution. Interestingly, the E/Z stereoselectivity was strongly correlated to the electron density of the acceptor (when the solvent was buffered to pH = 8), with the propiolamide affording the highest Z-content (∼95%) followed by the propiolate (∼89%) and the aryl ynone (83%), however, the alkyl ynone substrate yielded a more mixed stereochemical content (E/Z = 34/66). The same ratio (E/Z = 34/66) was observed for ynoles with β-substituted n-butyl, possibly indicating an equilibrium position for the stereochemistry. The overall trend in stereochemistry may be impacted by isomerization reactions, but it is more likely dependent upon stabilization of the enolate intermediates, where bond rotation to the more thermodynamically stable trans-isomer becomes more favorable as the electron-withdrawing ability increases (Scheme 3).68

The influence of the electron withdrawing group is not limited to the initial addition reaction but also to the stability of the Michael adduct itself. Thermally induced reversibility of the reaction has been reported for ynone65,66 and propiolate67

![Table 3. Conversion between a Thiol-Containing Peptide and Various Electrophilic Alkenes in Aqueous Acetonitrile Mixtures](https://doi.org/10.1021/acs.chemrev.0c01076)

<table>
<thead>
<tr>
<th>E/Z</th>
<th>R1</th>
<th>R2</th>
<th>H2O</th>
<th>pH = 8 buffer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NHP</td>
<td>H</td>
<td>20/80</td>
<td>5/95</td>
</tr>
<tr>
<td></td>
<td>OMe</td>
<td>H</td>
<td>66/34</td>
<td>11/89</td>
</tr>
<tr>
<td></td>
<td>Ph</td>
<td>H</td>
<td>25/75</td>
<td>17/83</td>
</tr>
<tr>
<td></td>
<td>C6H5</td>
<td>H</td>
<td>34/66</td>
<td>34/66</td>
</tr>
<tr>
<td></td>
<td>Ph</td>
<td>C6H5</td>
<td>20/80</td>
<td>34/66</td>
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<tr>
<td></td>
<td>C6H5</td>
<td>C6H5</td>
<td>66/34</td>
<td>34/66</td>
</tr>
</tbody>
</table>

![Table 4. Stereochemistry of Reaction Products between Protected Cysteine and Various Electrophilic Alkenes in pH Buffer and Aqueous Acetonitrile](https://doi.org/10.1021/acs.chemrev.0c01076)

<table>
<thead>
<tr>
<th>E/Z</th>
<th>R1</th>
<th>R2</th>
<th>H2O</th>
<th>pH = 8 buffer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NHP</td>
<td>H</td>
<td>20/80</td>
<td>5/95</td>
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<td></td>
<td>OMe</td>
<td>H</td>
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<td>11/89</td>
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<td>17/83</td>
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<td>H</td>
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<td>C6H5</td>
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<td>C6H5</td>
<td>C6H5</td>
<td>66/34</td>
<td>34/66</td>
</tr>
</tbody>
</table>
substrates as well as network materials using ynone monomers,\textsuperscript{30} but propiolamide–thiol adducts are irreversible under similar reaction conditions. The relative electron withdrawing strength of the functionalities can account for the discrepancy, at first glance, but further study is certainly warranted to wholly elucidate the relative reaction barriers and to screen other conceivable electron withdrawing groups (such as sulfones) for their reversibility.

Since the earliest studies, the efficiency of the reaction has shown little sensitivity to \( \beta \)-carbon substituents on the Michael acceptor. Acceptors bearing aromatic\textsuperscript{69,70} and alkyl\textsuperscript{71} groups have previously been shown to still react efficiently despite the steric barrier. For example, an \( n \)-butyl \( \beta \)-substituted substrate still reached near quantitative conversion, albeit at a notably slower rate than the unsubstituted analogue.\textsuperscript{67} Another study performed a \( N \)-methylmorpholine-catalyzed thiol conjugate addition to ethyl propiolate with varying \( \beta \)-functionality (Scheme 4).\textsuperscript{72} Using the terminal alkyne afforded the highest yield (91\%) with the ethyl ester \( \beta \)-substituted substrate producing slightly lower yield (86\%), followed by the \( n \)-butyl derivative (yield = 62\%). Any form of substitution on the \( \beta \)-position presents a steric barrier, which explains the lower yield in both latter cases. However, the additional ester group also activates the alkyne, making it “doubly” activated and thus more susceptible to nucleophilic attack, which can explain the increased yield for the ester substituent compared to the electron donating butyl group.

\subsection{2.2.2. Solvent Effects.} One of the greatest hallmarks of the thiol-yne Michael addition is that the suitable solvent scope is enormous and no additional solvent purification (e.g., drying or degassing) is usually required. Even in the presence of potentially nucleophilic hydroxy groups, such as in aqueous conditions or alcoholic solvents, the thiol-yne reaction can proceed cleanly with no hydroxyl-yne products.\textsuperscript{73} In one study, water was proposed to facilitate the reaction via hydrogen bonding activation of both carbonyl and thiol substrates because yields are significantly higher in water compared to neat conditions. The analogous thiol-ene reaction is also efficient in water without the need of an added catalyst species.\textsuperscript{74} The vast range of compatible solvents has led to several studies explicitly examining solvent effects on the Michael reaction. In most cases, solvent choice leads to a rather pronounced effect on both the rate of reaction and the stereochemistry of the \( \alpha \beta \)-unsaturated reaction products.

Generally, solvents with high polarity (i.e., large dielectric constant, including water) enhance the reaction rate and overall conversion, while less polar solvents retard the reactivity. A detailed investigation of solvent effects on the uncatalyzed, equimolar addition of thiophenol to ethyl propiolate was reported by Randive \textit{et al.} (Table 5).\textsuperscript{75} Polar solvents such as methanol, acetonitrile, or dimethylformamide, delivered high yields (>80\%) in less than 30 min, and reactions performed in water approached quantitative yields (98\%) in just 5 min. Reactions conducted in relatively nonpolar solvents (including toluene and \( n \)-hexane) proceed much slower and reached lower overall conversions. A moderate range of \( E/Z \) stereoisomer content (53\%–97\% \( Z \)-isomer) were also accessible by changing the reaction solvent; high polarity solvents (water, acetonitrile, and methanol) typically yielded products with high \( Z \)-isomer content with the notable exception of dimethylformamide.

Another study investigated solvent effects of the triethylamine-catalyzed addition of dodecanethiol to ethyl propiolate (Table 6).\textsuperscript{76} As previously noted, both the conversion and reaction rate were positively correlated to the dielectric constant of the solvent where dimethyl sulfoxide, acetonitrile, and acetone afforded quantitative conversions within 1 h. The stereochemistry of the reaction product was also discovered to be intimately tied to the solvent dielectric with product \( E/Z \) content ranging from 98/2 (benzene) to 22/78 (dimethyl sulfoxide). The authors posited that the higher polarity of the reaction medium more effectively stabilized the enolate intermediate to enable anti- over syn-addition during the
proton transfer step. On the other hand, the combination of a low polarity solvent with triethylamine could possibly form a hydrogen-bonded pair, so the syn-attack would be more favorable under those conditions.

As previously disclosed, the thiol-yne reaction can be conducted efficiently in water and/or aqueous mixtures, which is another hallmark of a Click reaction. Moreover, the pH of the aqueous reaction solvent can have a pronounced effect on the reaction rate and overall conversion. “Greener” recyclable solvents such as polyethylene glycol (PEG) have also shown promise for the thiol-yne Michael addition. For this uncatalyzed addition of thiophenol to ethyl 3-phenylpropiolate, yields near 90% were achieved at 60 °C, which falls short of the Click definition but could still be greatly improved by simply employing organobase catalysts. The steadfast efficiency under a wide range of reaction conditions, especially tolerance to aqueous environments, has vaulted the reaction into a similar class as its more popular thiol-ene counterpart and propelled it closer to biorthogonality.

### 2.2.3. Catalyst Effects

Although there are several examples of the thiol-yne Michael addition proceeding spontaneously, this is only achievable under narrowly defined reaction conditions such as in highly polar solvents and/or alkaline pH or by employing highly acidic thiol. However, employing a catalyst greatly broadens the scope of both operative Michael acceptors and thiols, which serves to improve the rate of reaction, total conversion, and stereoselectivity in many cases.

The rudimentary use of stoichiometric sodium to generate an isolated thiolate anion is effective, but it offers poor atom economy, more reaction steps, and ultimately does not fulfill Click criteria. However, the use of a catalytic quantity of amine as the nucleophilic thiol-yne reaction, which is another hallmark of a Click reaction. Moreover, the use of a catalytic quantity of amine as the nucleophile to form an allene enolate that can serve to concern that the use of these “nucleophilic” catalysts could form stable catalyst—Michael acceptor adducts to inhibit reactivity. However, this matter was largely settled in a thorough study that reported negligible adduct formation for phosphine- or tertiary amine-catalyzed reactions, even when the catalyst was employed in stoichiometric quantities.

There are several examples that compare different organobases as thiol-yne catalysts; one notable study investigated a range of amine and phosphine catalysts for the reaction between ethyl propiolate and 1-dodecanethiol in order to establish comparative catalytic performance on reaction rate, yield, and stereochemistry (Table 7).

### Table 7. Reaction of Ethyl Propiolate with Dodecanethiol Employing Different Organocatalysts at 0.1 M and Ambient Temperature

<table>
<thead>
<tr>
<th>catalyst</th>
<th>loading (mol %)</th>
<th>time (min)</th>
<th>conversion (%)</th>
<th>E/Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-hexylamine</td>
<td>10</td>
<td>60</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>isopropylamine</td>
<td>10</td>
<td>60</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>diisopropylamine</td>
<td>10</td>
<td>60</td>
<td>&lt;1</td>
<td>50/50</td>
</tr>
<tr>
<td>dicyclohexylamine</td>
<td>10</td>
<td>60</td>
<td>&lt;1</td>
<td>50/50</td>
</tr>
<tr>
<td>NEt₃</td>
<td>10</td>
<td>60</td>
<td>85</td>
<td>97/3</td>
</tr>
<tr>
<td>DIPEA</td>
<td>10</td>
<td>60</td>
<td>&lt;1</td>
<td>50/50</td>
</tr>
<tr>
<td>DBU</td>
<td>1</td>
<td>10</td>
<td>100</td>
<td>34/66</td>
</tr>
<tr>
<td>TBD</td>
<td>0.1</td>
<td>1</td>
<td>100</td>
<td>10/90</td>
</tr>
<tr>
<td>Ph₃PMe₂</td>
<td>1</td>
<td>10</td>
<td>100</td>
<td>50/50</td>
</tr>
<tr>
<td>Ph₃P</td>
<td>1</td>
<td>60</td>
<td>20</td>
<td>30/70</td>
</tr>
</tbody>
</table>

“DIPEA = disopropylethylamine. DBU = 1,8-diazabicyclo[5.4.0]-undec-7-ene. TBD = 1,5,7-triazabicyclo[4.4.0]dec-5-ene.

reaction could be retrieved in high yields from simple acid washing procedures to remove residual base catalyst. When primary amines such as hexylamine or isopropylamine were used, no conversion was observed because the catalyst was inactivated from competing addition of the amine to the electrophilic alkyne. Only trace amounts of products (reaction conversion <1%) were observed for the secondary amines (diisopropylamine and dicyclohexylamine), presumably as a consequence of similar addition byproducts. The relatively non-nucleophilic tertiary amine (triethylamine) offered a much-improved outcome (85% conversion in 60 min). Finally, the stronger amidine and guanidine bases, 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) and 1,5,7-triazabicyclo[4.4.0]dec-5-ene (TBD), were far more active, achieving quantitative conversions at low catalyst loadings (1 and 0.1 mol %, respectively) in just 10 min.

Trivalent phosphines have also shown promise for the nucleophilic thiol-yne reaction, shortly after they were discovered to be highly active catalysts in the thiol-ene...
reaction.\textsuperscript{85,86} Phosphine catalysts are widely assumed to follow the nucleophilic catalyst pathway (Scheme 5) on account that they lack the requisite basicity to effectively deprotonate many thiols; as such, their nucleophilicity is very important to overall catalytic efficiency.\textsuperscript{82,83} Hence, dimethylphenylphosphine was found to possess similar activity to the strong amidine bases, such as DBU, reaching quantitative conversion in 10 min.\textsuperscript{76} However, in the same study, triphenylphosphine was relatively ineffective as a catalyst (only 20% conversion in 60 min), likely as a consequence of its poor nucleophilicity. These results were corroborated in another report where tributylphosphine afforded quantitative yields in 15 min, but the triphenylphosphine-catalyzed reaction proceeded to only 17% conversion after 3 d using the same reaction conditions (ambient temperature in dichloromethane solvent).\textsuperscript{84}

Similar to the observed solvent effects on stereochemistry, the catalyst choice can also play an important role.\textsuperscript{76} All organobase-catalyzed thiol-yne reactions produced products with mixed stereochemistry or high Z-isomer content (up to 90%) from the thiol anti-addition across an alkyne, with the exception of triethylamine, which afforded high E selectivity (97%). It was postulated that these stark differences could be explained by relative basicity differences where the triethylamine could form an ion-pair with the thiol that would then add via a syn-addition to the alkyne moiety.\textsuperscript{57} Differences in the stereochemical outcome among other catalysts in the study could be explained by varying nucleophilicity and/or basicity that influenced the relative rates between both mechanistic pathways (e.g., anti vs syn-addition).

Double thiol addition to an alkyne to form the thiaoctenal species can also be observed when suitably strong bases are used in sufficient catalyst loadings and the stoichiometry of the thiol is adjusted. For example, employing TBD at 1 mol % loading in conjunction with 2 equiv of dodecanethiol (relative to ethyl propiolate) yields the corresponding thiaoctenal in near quantitative yield (>97%) at ambient temperature in just 10 min (Scheme 6).\textsuperscript{76} A similar reaction using tributylphosphine (at 20 mol % loading) with 2 equiv of benzyl mercaptan (relative to methyl propiolate) produces the thiaoctenal in 80% yield after 24 h at ambient temperature.\textsuperscript{88} In this example, the second addition of thiol proceeded approximately 1000 times slower, which implies that judicious choice of catalyst loading and/or reaction time can yield exclusively single addition products even when employing highly active nucleophilic catalysts. Finally, increasing the reaction temperature can also distinguish selectivity between single and double addition products. For example, the thiol-yne reaction in water at ambient temperature afforded single-substituted products, but increasing the reaction temperature to 80 °C yielded thiaoctenal products in suitable yields (67–81%).\textsuperscript{73}

Inorganic-based reagents, including BF\textsubscript{3},\textsuperscript{89} or NaH,\textsuperscript{90} have also been investigated to catalyze the thiol-yne reaction, however, the catalysts that arguably show greater promise are heterogeneous species that allow simple separation and potential recyclability. A study into the influence of common lab inorganic solids on the addition of 2-naphthalenethiol to methyl propiolate was undertaken by Kodomari \textit{et al.} (Table 8).\textsuperscript{91} Neutral alumina achieved a far higher conversion (96%) compared to reactions at all other conditions, including a control without any additive (19%), silica (20%), and even triethylamine (85%). Furthermore, when employing neutral alumina the Z-isomer was heavily preferred, while other catalysts were less stereoselective and afforded a mixture of E/Z isomers. However, although alumina is abundant, a vast quantity (>500 wt %) was employed, which falls short of the high atom economy requirement of a Click reaction. Finally, the reactions were efficient when using aromatic thiols, but alkyl thiols were not as reactive, only reaching <90% conversion after 48 h. Nevertheless, the use of a heterogeneous catalyst like alumina could still be attractive if it retained catalytic activity for batch recycling or flow chemistry processes.

The synthesis of dithiaoctenals have also been reported using heterogeneous catalysts.\textsuperscript{92} A more recent investigation found high surface area magnesium oxide could achieve near quantitative yields of the double addition product (98%) from the reaction of 1,3-propanedithiol to 4-phenyl-3-butyn-2-one.\textsuperscript{87} The reaction also showed a good tolerance when acceptors with \(\beta\)-substituents were employed, such as \(-\text{SiMe}_3\) (yield = 95%, 1 h) or \(-\text{C}_5\text{H}_{11}\) (yield = 98%, 3 h). As it stands, inorganic catalysts are less efficient than organic counterparts (often employed in at least stoichiometric quantities) and their future role in advancing the thiol-yne reaction will be contingent on the ability to leverage their heterogeneity, \textit{i.e.}, by physically recycling them.

Table 8. Reaction of Methyl Propiolate with 2-Naphthalenethiol Catalyzed by Various Inorganic Solids and Triethylamine in Diethyl Ether

<table>
<thead>
<tr>
<th>catalyst</th>
<th>conversion (%)</th>
<th>E/Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>none</td>
<td>19</td>
<td>50/50</td>
</tr>
<tr>
<td>Al\textsubscript{2}O\textsubscript{3}</td>
<td>96</td>
<td>10/90</td>
</tr>
<tr>
<td>SiO\textsubscript{2}</td>
<td>20</td>
<td>45/55</td>
</tr>
<tr>
<td>NEt\textsubscript{3}</td>
<td>85</td>
<td>58/42</td>
</tr>
</tbody>
</table>

2.3. Applications in Organic Synthesis

In addition to its high efficiency and good orthogonality, the ability to furnish multiple reactive functionalities (especially an electron deficient alkene) in the reaction products has allowed for widespread utility of the nucleophilic thiol-yne reaction in organic synthesis. This has most commonly been exploited in cascade heterocyclic syntheses, but it has also played a role in bioconjugate chemistry where orthogonal reactions can be utilized in tandem.

2.3.1. Heterocycle Synthesis. Versatile functionality, selectivity, and a large thermodynamic driving force are crucial to the synthesis of heterocyclic compounds, making the mild nucleophilic thiol-yne addition an attractive reaction in the synthesis of functional heterocycles. In particular the use of
commercially available “doubly activated” alkynes (e.g., diethyl acetylenedicarboxylate), which have gained increasing attention in organic chemistry, especially for the synthesis of sulfur-containing heterocycles. Additionally, the α,β-unsaturated products from the thiol-yne Michael addition have also been effectively exploited in subsequent intramolecular cyclization reactions to afford heterocyclic compounds. However, many of these heterocycle syntheses do not exhibit Click features, and thus this section will not cover every example; instead, the focus will be directed to more efficient and/or informative reports.

An early interesting reaction that demonstrated the versatility of the thiol-yne Michael addition centered on the attempted synthesis of functional benzothiazines from 2-aminothiophenol. Initially, the reaction was investigated using a doubly activated alkyne (dimethyl acetylenedicarboxylate) as the Michael acceptor, where the thiol Michael addition occurs before the in situ intramolecular amidation (Scheme 7-top). However, when using a singular activated alkyne (3-phenylpropionitrile), the Michael addition is selective for the thiol addition without the intramolecular amino Michael addition side reaction (Scheme 7, bottom). After this step, the substrate could be cyclized to yield benzothiazine and benzothiazole depending on the reaction conditions. This example highlights the orthogonal reactivity of the thiol over the amine in the Michael addition and should be an important consideration for reactions between complex substrates that may feature several functionalities.

Thiophenes are important building blocks in conjugated materials and medicinal chemistry, but the well-controlled synthesis of functionalized thiophene rings is a historical challenge. Variations of Fiesselmann’s thiophene synthesis exploit the functionality control of the Michael acceptor β-carbon substituent, the electronic nature of the alkyne activating group, and thiol nucleophile to effectively yield highly substituted thiophenes. In the seminal report, a propiolate or ynone was reacted with thiolglycolate to form a Michael adduct which then undergoes an intramolecular cyclization. Although the Michael addition reaction is high yielding when employing DBU as the catalyst, the subsequent cyclization is not, but it can be improved by changing the base catalyst from DBU to CsCO₃ to afford substituted thiophenes in good yields (Table 9).

The reaction is also famously insensitive to β-ynone substituents, and careful selection of the electron withdrawing groups has been exploited to afford thiophenes flanked by notoriously difficult-to-access functional groups. For example, amino-functionalized thiophene rings have been synthesized using Fiesselmann’s method by employing propionitrile acceptors, however, isolated yields are typically moderate (~70%) and sodium thiolate nucleophiles are generated instead of using catalytic bases. There has been a long-standing interest in synthesizing thiophene derivatives containing organoboron groups for further modification in cross-coupling applications. By installing a trifluoroborate salt on the β-position of an ynone reactant, the Fiesselmann route...
can afford a borate-substituted thiophene (Scheme 8). The initial Michael addition of the thioglycolate to the phenylpropynone borate salt proceeded quantitatively and an efficient subsequent cyclization afforded the thiophene compound in high yield (95%) under ambient conditions. However, the yield of this reaction is heavily contingent on reagents and conditions; for example, without K₂CO₃ or MeOH the reaction barely proceeds (<5% yield) and the yield fluctuates according to the nature of the ynone. Another representative example for the utility of the thiol-yne reaction is the synthesis of fluorinated thiophenes, which are challenging synthetic targets, but the Fiesselmann reaction can be employed to furnish heavily fluorinated thiophenes.

Longstanding limitations of the Fiesselmann synthetic pathway included the inability to functionalize each position independently and/or synthesize fully (tetra-) substituted thiophenes. However, these issues were simultaneously addressed by using pyridinium 1,4-zwitterionic thiolates as nucleophiles in the Fiesselmann thiophene synthesis. Nevertheless, these reactions do not fully satisfy Click requirements, as exemplified by their modest yields (60–75%) at 85 °C and the overall poor atom economy (due to the pyridine leaving group). Another important hallmark of Click chemistry is employing readily available reagents; a recent example sourced the low-cost and abundant mercaptoacetaldehyde dimer to synthesize thiophene (Table 10). Treating a functional ynone species with dimeric mercaptoacetaldehyde and stoichiometric triethylamine leads to a cyclic thioether intermediate possessing a hydroxyl group that can undergo subsequent dehydration via treatment with SiO₂ or HCl. Another advantage is the broad range of suitable ynones, however, the use of stoichiometric additives is not ideal, but further exploration of other organobases could optimize reaction conditions to be more compatible with Click requirements.

The combination of good orthogonality and rapid reaction rates has made the thiol-yne a popular choice for one-pot multistep syntheses. In one example, the ynone Michael acceptor was synthesized via Sonogashira coupling followed by in situ Michael addition and cyclization to afford a functional thiophene substrate from acyl chlorides (Table 11). The reaction demonstrated a good functional group tolerance for the α- and β-carbon substituents on the ynone, as exemplified by yields in excess of 90%. The same approach has also been used to make oligomeric thiophenes, which are known to exhibit aggregation induced emission (AIE). Despite good reaction orthogonality in this study, as evidenced by tolerance of metal species and diverse functionalities, the use of large quantities (up to equimolar amounts) of organobase catalysts fall short of the high atom economy that is crucial to satisfy Click criteria.

### Table 10. Organobase-Catalyzed Intermolecular Heterocyclization of Ynones with Dimeric 2-Mercaptoacetaldehyde

<table>
<thead>
<tr>
<th>R₁</th>
<th>R₂</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Me</td>
<td>Ph</td>
<td>75</td>
</tr>
<tr>
<td>Ph</td>
<td>n-butyl</td>
<td>91</td>
</tr>
<tr>
<td>Ph</td>
<td>SiMe₃</td>
<td>91</td>
</tr>
</tbody>
</table>

### Table 11. Tandem Reaction of Sonogashira Coupling and Thiol-yne Addition to Afford Trisubstituted Thiophenes

<table>
<thead>
<tr>
<th>R₁</th>
<th>R₂</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>p-MeOC₆H₄</td>
<td>Ph</td>
<td>97</td>
</tr>
<tr>
<td>p-NCC₆H₄</td>
<td>p-(CH₃)₂C₆H₄</td>
<td>91</td>
</tr>
<tr>
<td>2-thienyl</td>
<td>Ph</td>
<td>90</td>
</tr>
<tr>
<td>p-MeC₆H₄</td>
<td>ferrocenyl</td>
<td>91</td>
</tr>
</tbody>
</table>

#### 2.3.2. Bioconjugation Reactions.

Thiols are exemplary targeting moieties for bioconjugation reactions as a result of their high nucleophilicity compared to primary or secondary amino groups in peptides and related biomolecules. For example, the thiol of cysteine was determined to be 4 orders of magnitude more reactive as a nucleophile than any other amino acid functionality. It should be no surprise that the thiols in biological systems have been exploited via Michael addition reactions with electrophilic alkynes. An interesting application includes functionalizing free cysteine thiol groups in cotton with alkyne-containing reactive dyes to improve the staining efficiency. The reaction has even been successfully demonstrated in an analytical assay to quantify sulfur content in certain wines.

Historically, maleimide was the preferred Michael acceptor used for antibody–drug conjugates due to its superior reaction rate, however, stability issues arose due to the reversibility of the resulting Michael adduct because irreversible conjugation was required along with good targeting selectivity of the thiol. Wagner and co-workers screened multiple acetylenic electrophiles for rapid cysteine tagging in aqueous conditions (PBS:DMSO mixture) and observed good yields for 3-phenylpropionitrile (94%), N-phenylmaleimide (99%), N-phenylpropionamide (72%), and phenylpropynone (99%). Unlike the adducts formed from the ynone or maleimides, products from propionitrile substrates were indefinitely stable under the reaction conditions (Scheme 9).

#### Scheme 9. Irreversible Cysteine Tagging Using 3-Phenylpropionitrile

![Scheme 9](https://doi.org/10.1021/acs.chemrev.0c01076)
compared to the maleimide substrate, which was found to couple with multiple other amino acids (histidine, glycine, and serine) in non-negligible amounts (≥2.8%).

In medicinal applications, alkyne Michael acceptors have shown great promise in irreversible inhibition of protein targets. They show particular promise in the inhibition of Bruton’s tyrosine kinase (BTK): a protein linked to non-Hodgkin’s lymphoma. Acalabrutinib is an FDA approved drug for the inhibition of BTK that features a targeting propiolamide Michael acceptor that can bind with the protein substrate. Importantly, the drug has shown better targeting selectivity than the analogous drug candidate that contained an alkene Michael acceptor moiety, thus demonstrating the superior chemoselectivity of using acetylenic substrates.

Thiol-containing amino acids (such as cysteine or homocysteine) and small peptides featuring these units (such as glutathione) are important indicator molecules for assays related to disease diagnostics. However, it is crucial to be able to orthogonally discern among different thiol containing biomolecules, and this can be achieved by judicious choice of the alkyne Michael acceptor. In a particularly elegant example, a fluorophore with a flanking ynone functionality was first reacted with a thiol to generate a nonfluorescent complex. Cysteine or homocysteine was then reacted with the complex to release the thiol quenching molecule and generate a new fluorescent adduct that could be accurately quantified by fluorescence spectroscopy (Figure 3). This reaction relied on the reversible displacement of the thiol-ynone Michael adduct, which can be dynamic under alkaline pH so in this example the amino acid conjugation was performed at pH = 8.1 to ensure the efficient thiol displacement.

More recently, Cheng et al. designed a very simple and selective fluorescent tag based on a propiolamide electrophile that reacted preferentially with thiol nucleophiles over amines. The resultant thiol-Michael adduct itself is not fluorescent, however, upon an intramolecular cyclization with the neighboring amino group (which is not present in peptide-based thiols such as glutathione), the adduct becomes fluorescent (Figure 4).

2.3.3. Intermediates or Reagents. The thiol-yne Michael reaction has also been featured as a means to furnish valuable synthetic reagents and/or intermediates. For example, phosphonium ylides can be synthesized through the thiol-yne Michael addition of 2-aminothiophenol and di(methyl/ethyl) acetylene carboxylate in the presence of triphenylphosphine. Additionally, cyclic thioacetals are versatile umpolungs for carbonyl synthons, and they are smoothly produced from successive conjugate additions of propanedithiol or ethanediol thiol.

Efficient pathways for selective protection/deprotection of thiols remains challenging in organic synthesis. Vinylic Michael acceptors had previously been investigated as thiol protecting groups but did not perform well, especially during the deprotection step, however, the sulfone-based alkyne Michael acceptor, tosylacetylene, was demonstrated as a useful protecting group. Specifically, the Michael addition of tosylacetylene to thiols could be achieved in good to excellent yields (80–100%) at ambient temperature. The protecting group was also shown to have excellent reaction selectivity for thiols in the presence of alcohols, and the resulting adduct was stable in acidic and alkaline media, thus highlighting its utility. Finally, deprotection to regenerate the thiol was quantitatively achieved by reacting with excess pyrrolidinyl.

The deprotection of a thioacetic acid-propiolate Michael adduct with CsCO₃ has also been used as an efficient route to vinylic thiocetic acids including cysteine and homocysteine derivatives (Scheme 10). It is interesting to note that subsequent intermolecular reactions were not reported despite the nucleophilicity of the thiolate species. The thiol-yne reaction was also not quantitative (yield = 79%) in this example despite the low pKa of the thiol, but this can be rationalized by the fact that it is a secondary thiol and imidazole is a relatively poor base. Finally, the addition reactions of pyrrolecarboxdithioates to activated alkenes and alkynes was reported as a path to divinylsulfanes, albeit in low yields.
In a final example, the thiol-yne reaction between thiols and propiolates or propiolamides was employed for the synthesis of rotaxanes in chloroform using K₂CO₃ as an additive (150 mol %) (Figure 5). The addition reaction using only the acetylenic “axle” Michael acceptors (i.e., without the macrocycle) were successful for both the propiolate (24%) and propiolamide (53%). This was further translated to the reaction with the “wheel” to form the rotaxane in similar yields (19% for propiolate and 35% for propiolamide). Interestingly, the analogous thiol-ene reaction (i.e., employing an acrylate or acrylamide substrate) failed to form the targeted compounds. This example is also interesting because the propiolamide substrates afforded higher reaction yields even though the propiolate is normally accepted as the more reactive of the two moieties.

2.4. Applications in Polymer Chemistry

Soon after Click reactions gained prominence in organic chemistry, their use was investigated in polymer synthesis. One of the greatest success stories has been the thiol-ene reaction, including both nucleophilic and radical mechanisms, which has been used to synthesize step-growth polymers in high conversion with rapid kinetics and good orthogonality. Although the thiol-yne reaction is comparatively less developed in polymer chemistry, it offers not only many of the same advantages attributed to thiol-ene polymerizations but a few others as well. Principally, if the thiol-yne reaction proceeds via a nucleophilic pathway the product is an unsaturated polymer which provides opportunities for both postpolymerization functionalization as well as stereochemical structure−property manipulation. To date, much of the limited work on thiol-yne polymerizations has featured radical-based processes but these are inherently less selective (e.g., competition between single and double addition). Nevertheless, there are a few examples of nucleophilic thiol-yne polymerizations and its application in polymer chemistry is expanding rapidly.

2.4.1. Step-Growth Polymers. As a consequence of their exceptional reactivity, ynones were the first Michael acceptors to be polymerized with dithiols in a thiol-yne polyaddition. Bass and co-workers synthesized a series of disubstituted aryl internal ynones from the addition of lithium phenylacetylide to aryl dialdehydes. These diyne monomers were then reacted with dithiophenols in the presence of methylmorpholine to furnish unsaturated step-growth polymers with pendant phenyl groups. The synthesis of conjugated polymers can be challenging and is often restricted to organometallic cross-coupling reactions with a limited number of accessible building blocks. However, the straightforward synthesis of the diyne monomers and dithiophenols opens a library of conjugated structures where various stereochemistry, functionality, and diverse heterocycles can easily be incorporated. Unfortunately, these aromatic thiol-yne polymers were difficult to fully characterize as a result of their limited solubility, however, the polymers could be retrieved in high yields (>90%) by a simple precipitation procedure in methanol (Scheme 11).

Terminal aromatic ynone monomers have also been polymerized with dithiols and these reactions proceeded spontaneously at low temperatures (0−5 °C), thus highlighting the significant difference in reactivity between the internal and terminal ynone. Both types of aromatic polymers (from both internal and terminal ynone) were thermally stable and could be fabricated into useable materials using compression molding at high temperatures but this is somewhat surprising since
terminal ynone–thiol adducts have been investigated for their dynamic behavior. Further, another early study investigated the reaction of a thioacetal with a ynone monomer using a catalytic quantity of triethylamine to afford unsaturated polymers with a greater solubility than previously reported polymers from aliphatic dithiols.

More recently, propiolates have emerged as excellent monomers for thiol-yne step-growth polymerizations. The reaction of aromatic dithiols with aromatic dipropiolates (internal and terminal) proceeded to high conversion using a range of amine organobase catalysts (Scheme 12). Diphenylamine proved to be an active catalyst to afford a high molecular weight polymer (weight-average molar mass, $M_w = 30$ kDa) in 98% yield with simple isolation via precipitation. Very recently, it was discovered that the same reaction (varying the monomer only with an additional $\beta$-substituted methyl group) can proceed uncatalyzed in dimethylformamide at 60 °C to produce polymers with comparable molecular weights and isolated yields. In addition to their high thermal stability, these polymers possess exceptional optical transparency and refractive index (resulting from their high sulfur content), which makes them potentially useful in optics applications, however, there lacks critical analysis of the structure–property relationships. Some of the polymers also exhibit aggregation induced emission (AIE), where the polymers in the solution state are weakly emissive, but as they become less soluble, they aggregate and emit at a higher intensity. These AIE aromatic polymers have potential value for the detection of explosives.

Stereochemistry has been a key consideration for the thiol-yne Michael addition for decades, however, its importance was largely unheeded in polymer chemistry. More recently, there has been a push to rationally manipulate stereochemistry in order to influence bulk properties of polymers and materials. A series of dipropiolate monomers were synthesized from the Fischer esterification of propiolic acid and commercially available diols. These monomers were polymerized with commercial aliphatic dithiols using various solvents and organobases to modulate the $E/Z$-content in the polymer backbone (Figure 6a). Linear unsaturated polyesters were attained at very high molecular weight ($>100$ kDa) with excellent control of $Z$-isomer content (32~80%). The stereochemistry dictated the bulk mechanical properties where the high $Z$-content polymer was significantly tougher and stronger than the high $E$ polymer. This trend in mechanical strength was attributed to greater polymer chain entanglement and alignment of the $Z$ unit that led to increased crystallization; this is illustrated by wide-angle X-ray scattering.

Scheme 12. Base-Catalyzed Step-Growth Polyaddition of Aromatic Dithiol and Aromatic Dipropiolate

![Scheme 12 diagram]

Figure 6. Thiol-yne polymerization to afford stereocontrolled polyesters. (a) Reaction of dipropiolate with 1,6-hexanediol to afford unsaturated polyesters. (b) Wide-angle X-ray scattering (WAXS) indicating more order in higher cis content materials. (c) DSC thermograms showing crystallinity in high cis polymers. Reproduced under Creative Commons Attribution License from ref 144.
(WAXS) (Figure 6b) and endothermic transitions in the differential scanning calorimetry (DSC) thermograms (Figure 6c). The alkenes remaining in the unsaturated polymer were also reacted with sulfur (i.e., vulcanized) to create an elastic material which further exemplifies the utility of the thiol-yne reaction in polymer synthesis. An analogous series of thiol-yne polyester biomaterials also displayed independently tunable mechanical and degradation properties, which addresses a long-standing challenge in synthetic biomaterials, by simultaneously manipulating the Z-isomer content (stereocontrol) and succinate amount (compositional control) of the polymer.145 Previously, controlling one property involved complex changes to the material formulation that would then affect the former property. However, this approach could conveniently afford two materials with similar mechanical strength, but different degradation rates and vice versa. In general, thiol-yne polyesters display excellent mechanical strength, in addition to facile manipulation of bulk physical properties, which can be finely tuned according to alkene stereochemistry and/or monomer choice.

Another report described the synthesis of high molecular weight (>100 kDa) stereocontrolled unsaturated polyamides from dipropiolamide monomers (Figure 7a).24 The polyamides were extremely processable and malleable, relative to conventional nylons, and the stereochemistry once again influenced thermomechanical properties where the greater Z-content polymers displayed higher glass transition temperatures and greater physical stiffness than their isomeric analogues (Figure 7b-c). Although these trends corroborated the structure–property relationships observed for analogous polyesters, the differences among the properties were significantly less pronounced. This is likely due to the fact

Figure 7. Thiol-yne polymerization to afford stereocontrolled polyamides.24 (a) Reaction of dipropiolamide with 1,6-hexanedithiol to afford unsaturated polyamide. (b) Plot of Young’s modulus and Glass transition temperature against the E/Z content of the unsaturated polyamides. (c) Physical demonstration of excellent moldability compared to conventional nylons. Reproduced under Creative Commons Attribution License from ref 24.

Figure 8. Polyaddition of dithiol with doubly activated alkyne with dithiol using a TBD catalyst.148 Reproduced with permission from ref 148. Copyright 2019 American Chemical Society.
that the polyamides were amorphous at all E/Z ratios, whereas crystallinity was found to drive the changes to the bulk mechanical properties for the polyester materials. Nevertheless, the lack of crystallinity in the thiol-yne polyamides was beneficial to processing the materials, especially compared to conventional semicrystalline nylons that require extremely high temperatures to effectively process. Efforts to further improve stereoselectivity in this area toward 100% E or Z polymers continues to be a major aim and should afford even more divergent physical properties within respective polymer classes.

The nucleophilic thio-yne Michael reaction predominantly results in a single thiol-addition rather than a second conjugate addition to the unsaturated product, unless a strong base or excessive catalyst loadings are employed. This is often regarded as one of its prime advantages over the radical thiol-yne reaction, however, recent studies have exploited this second conjugate addition to synthesize linear thioacetic polymers. Endo and co-workers disclosed the efficient polyaddition of 1,4-bis(mercaptomethyl)benzene and methyl propiolate catalyzed by large quantities of tributylphosphine (50 mol %), but the barrier for second addition led to slow kinetics and poor overall conversions, only affording low molecular weight polythioacetics (<10 kDa). Nevertheless, thioacetic polymers have the potential to offer unique degradation pathways and optical properties due to their susceptibility to hydrolysis and high sulfur content. A similar system exploited β-substitution of ethyl propiolate to furnish a hydrophobic fluorinated polythioether. In another example, a “doubly” activated alkyne (dimethyl acetylenedicarboxylate) was reacted with several dithiols using the strong organobase, TBD, as the catalyst in chloroform (Figure 8). High molecular weight (>30 kDa) linear polymers were afforded in under 60 min and isolated by simple precipitation in acidified methanol. As a consequence of the high electrophilicity of the Michael acceptor and the strength of the organobase, the double addition was favored, resulting in linear polythioethers. A few other acetylenedicarboxylate esters (ethyl and tert-butyl) were also successfully polymerized with the potential to use other moieties to expand possible pendant functionality. Among the dithiols that were tested, a striking discovery found that both 1,2-ethanediol and 1,3-propanediol both favored intramolecular cyclization over propanization to afford the dithiane product. Importantly, these polymers were found to be dynamic and could be effectively depolymerized by the subsequent addition of monofunctional thiol and TBD. These studies perfectly highlight the scope of the thiol-yne Michael addition, and specifically how modular substrate design can provide access to a broad range of versatile polymers for different applications.

The thiol-yne Michael addition is also particularly suited for multicomponent tandem reactions because of its orthogonality. This strategy has been utilized by Tang and co-workers to create functional linear polymers (Scheme 13). Tandem Sonogashira coupling and conjugate addition reactions employing various combinations of acyl chlorides, phenylacetylenes, and thiols (where at least two of the species were disubstituted) formed linear polymers in a one-pot reaction. The high molecular weight linear polymers (>30 kDa) were retrieved by a filtration/precipitation protocol and isolated in near quantitative yields (>95%). The polymers showed varying UV absorption behavior and different refractive indexes, and this is likely to be related simply to the specific aryl functionalities that were employed. Nevertheless, the reaction illustrates the orthogonality of the Michael reaction with other catalysts and functionalities present. However, it should be noted that the Sonogashira coupling reaction is sensitive to atmospheric oxygen and therefore deviates from Click criteria. The rapid reaction rates and high efficiency of the thiol-yne Michael addition is also represented by its use in postpolymerization functionalization reactions. A polymer with propiolate-containing pendant groups was smoothly reacted with thiol in the presence of NEt₃ to afford the single addition thiol-Michael adduct. A subsequent second addition with a different thiol can be achieved using 1 equiv of monofunctional thiol and employing a stronger base such as TBD. Both reactions are observed to proceed to completion and are purified by precipitation to isolate the functionalized polymer in high yields (>90%). The same principle of stepwise thiol additions has also been applied to polyesters containing an activated internal alkyne along the polymer backbone achieving quantitative conversions when using TBD at 25 mol % loading.

2.4.2. Network Polymers. Hydrogels are important materials comprised of cross-linked (network) polymers that are swollen with aqueous media, with critical applications ranging from food to healthcare. Hydrogel synthesis has historically turned to natural polymers (such as alginates), but hydrophilic synthetic polymers are increasingly being used. In particular, the thiol-yne Michael reaction has proven to be highly useful for hydrogel synthesis due to its great efficiency.

Scheme 13. Multicomponent Reactions Combining Sonogashira Coupling and Thiol-yne Addition to Afford Various Linear Polymers

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**Polymers**

https://doi.org/10.1021/acs.chemrev.0c01076

`Chem. Rev. XXXX, XXX, XXX--XXX`

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and tolerance to aqueous environments. Multiarm polyethylene glycols (PEG) were functionalized with thiol and propiolate moieties via straightforward Fischer esterification and then reacted with one another in an aqueous solution to form robust hydrogels in under 30 min. This simple approach enabled tunable mechanical properties and mesh sizes by selecting three- or four-arm PEG units and varying the length of two-arm substrates (Figure 9). The same thiol-yne system has been exploited in conjunction with tetrazine-norbornene adducts to form interpenetrating dual network hydrogels. Both Click reactions proceeded orthogonally to form independent networks that were considerably tougher than either single network system.

Using a similar system, nonswellable hydrogels were also obtained, which is important for tissue engineering because uncontrolled swelling behavior can diminish mechanical performance of the material. A hydrogel based on the thiol-yne system still has residual electron deficient alkenes within the network; these can be further functionalized. As an example, an antimicrobial PEG-based thiol-yne hydrogel was formed via postsynthetic conjugation with a thiol-terminated antimicrobial peptide. Compatibility with biological systems has already been highlighted for the thiol-yne reaction, but this also enables the versatility of incorporating numerous natural polymers (such as alginate, chitosan, or hyaluronic acid) into synthetic hydrogels. Thiol-yne hydrogels that incorporate these natural polymers have exhibited self-healing behavior and high stretchability rendering them superior to hydrogels of similar composition (i.e., the same biopolymer) but not cross-linked with thiol-yne chemistry.

The reversibility of the second thiol addition in the thiol-yne reaction has been used to create reprocessable cross-linked materials. Similar to other dynamic Michael addition systems, the reversibility of the reaction is triggered by base or heat. This feature has been exploited to deliver self-healing injectable hydrogels by cross-linking multiarm PEG thiol with a small molecule alkyl ynone. The dynamic nature of the thiol-yne reaction was triggered by high pH buffer solution, enabling the hydrogels to undergo shear-thinning under stress, meaning they could be quickly injected using a syringe only to reform into a “solid” material after delivery (i.e., when the stress was removed). A similar thiol-yne double addition reaction was also employed to afford covalent adaptable networks (CANs) based on reversible cross-linking. By varying the substrates on the ynone monomers, the exchange rates of the second thiol addition were tuned by several orders of magnitude to create materials with diverse rheological properties that were important for their reprocessing behavior. Model reactions of the thiol exchange process showed that the presence of electron withdrawing groups on the aromatic unit increased the exchange rate of the thiol and ultimately reduced the requisite reprocessing temperature for the bulk materials (Figure 10).

3. AMINO-YNE REACTIONS

3.1. Overview and Historical Development

The amino-yne Michael reaction is broadly similar to the ubiquitous thiol-yne motif, where a primary or secondary amine attacks the β-carbon on an electron deficient alkyne, although differences lie in the catalysis because many amino-yne reactions proceed spontaneously (i.e., without an added catalyst) at ambient temperatures. As such, the efficiency of the reaction rivals, and in some cases surpasses, the thiol-yne Michael addition as a result of the excellent nucleophilicity of the amine functionality. Consequently, the reaction has been adopted in numerous materials and biological applications. Furthermore, the reactivity and versatility of the enamine Michael adduct itself has found good utility in organic synthesis.

The first amino-yne nucleophilic addition was reported in 1899 by Ruhemann and co-workers where the reactions of diethylamine (or piperidine) with diethylacetylene dicarboxylate and ethyl phenylpropionate were reported. The reactions were described as “violently exothermic” and proceeded in the absence of a catalyst at ambient temperature. This early work originally focused on employing secondary amine nucleophiles to avoid a second addition of the nucleophilic enamine Michael adduct. Subsequent work featured primary and multifunctional amines to induce
intramolecular condensation reactions after the initial conjugate addition to furnish heterocyclic compounds such as pyrones, pyridines, and cyclic amides. Other early work in the area also expanded the scope of Michael acceptors to propiolonitriles and ynones in equally high efficiency to the propiolates. It should be noted that nucleophilic amino-yne additions can also proceed without the requisite of an electron withdrawing functionality in conjugation with the alkyne, i.e., electronically neutral alkynes are sometimes reactive as well. The addition of amines and ammonia to nonactivated alkyl alkynes has been known for decades but typically requires harsh reaction conditions including high temperatures (>300 °C) and pressures or heavy metal catalysts. As such, these transformations are not discussed in further detail.

Modern organic synthesis has continued to focus on transition metal-catalyzed intramolecular (and intermolecular) cyclization reactions of amines to electron deficient alkynes (primarily ynones), and this area has been critically reviewed in recent years. Even though this section will feature overlap with some of this content, it will ultimately focus on highly efficient reactions presenting with Click features, emphasizing those with (near) quantitative yields, high selectivity, and/or simple reaction conditions. Finally, the section will be presented slightly differently than the thiol-yne reaction because there are not many conclusions to be made about reactivity trends according to reaction conditions and/or substrate scope. There will be a brief overview on reactivity, but subsequent parts of the section will be divided by catalyst class for organic transformations and then conclude with emerging applications in polymer and materials chemistry.

### 3.2. Trends in Reactivity

This convenient reaction has become a mainstay in modern organic chemistry, however, a detailed understanding of the effects of reaction conditions (including solvent, additives, and/or catalysts) and the structure of the reactants on reaction outcomes has not been systematically investigated. This is problematic because the efficiency of the reaction varies widely, sometimes proceeding spontaneously at ambient temperature or instead requiring the use of metal catalysts at elevated temperatures. Nevertheless, we will attempt to draw meaningful comparisons within and among reports, especially as related to the aforementioned reaction conditions.

The description of alkene stereochemistry in the amino-yne Michael adducts is notably less discussed in the literature because it is heavily contingent on internal hydrogen bonding and steric considerations rather than charge stabilization of the intermediate species as in the thiol-yne analogue. This means that there is often less opportunity to manipulate the stereochemical outcomes unless the amino nucleophile is substituted. Nevertheless, many conjugate additions of primary amines to acetylenic Michael acceptors typically select for the Z-isomer, especially when propiolates, propiolamides, and ynones are employed due to intramolecular hydrogen bonding with the adjacent carbonyl moiety, while secondary amines tend to be less stereoselective and sometimes favor the E-isomer (Scheme 14). This of course is not a universal characteristic, as the influence of the Michael acceptor can also change the stereochemistry and isomerization of the Michael adducts is possible, both of which can obfuscate stereochemical trends.

![Scheme 14. Generic Scheme Showing the 1,4-Conjugate Addition of 1° and 2° Amine Nucleophiles to Carbonyl-Containing Michael Acceptors](https://doi.org/10.1021/acs.chemrev.0c01076)

### 3.3. Applications in Organic Synthesis

#### 3.3.1. Metal-Catalyzed Reactions

The metal-catalyzed amino-yne reaction has been used to a great extent to synthesize N-containing heterocycles, primarily substituted pyroles or indoles, from precursors containing acetylenic units and amines. The first report of pyrrole synthesis via conjugate amino-yne addition was reported by Nozaki and co-workers in 1981 using a palladium catalyst to synthesize pyrroles in quantitative yields. Gabriele et al. expanded on this motif to form highly substituted pyrroles and quinolines at similar efficiency using a variety of palladium(II) and copper(II) catalysts (Scheme 15). The use of precious metal gold- and silver-based catalysts have also been employed with analogous reaction outcomes.

![Scheme 15. Transition Metal-Catalyzed Intramolecular Amino-yne Cyclization to Form Pyrroles](https://doi.org/10.1021/acs.chemrev.0c01076)

In 2010, Kievano et al. developed a one-pot quinoxaline synthesis (in yields of up to 92%) by Sonogashira coupling and subsequent intramolecular amino-yne coupling using a Pd/Cu catalyst at 70 °C. A palladium-catalyzed synthesis of isoindolines using Sonogashira coupling has also been reported with up to 94% yield in water at 50 °C. The copper(II)/triethylamine-catalyzed synthesis of pyrazoles in yields up to 99% was accomplished from the reaction of various yrones and substituted hydrazines.

These examples illustrate some of the more moderate reaction conditions (such as water as a solvent and/or low reaction temperatures) in the area and thus align more closely with Click criteria.

Six-membered nitrogen-containing heterocycles have also been intensely investigated using the amino-yne reaction, with a few examples reporting good efficiency. A one-pot strontium- and barium-catalyzed synthesis of isoquinolines by cascading amino-yne and amino-ene reactions was reported, achieving yields of up to 99% at a higher temperature (130 °C) which decreases its compatibility with Click criteria. The Gouault group described the total synthesis of (−)-epimyrtine, wherein one of the critical steps was a highly selective gold-catalyzed intramolecular amino-yne cyclization of an ynone to form a six-membered pyridone from a...
secondary amine (Scheme 17b). Although this reaction step was not quantitative, this is likely due to steric considerations from both the amino functionality and the ynone moiety.

Ribeci and co-workers used alkali metal salts or organo-bases to produce indoles via intramolecular cyclization with yields of up to 99% and 90%, respectively, however microwave heating at 200 °C diminishes the experimental simplicity. The Verma group later reported KOH to catalyze the conjugate addition of imidazole with electron deficient alkynes at yields of up to 84%. This example is particularly interesting because imidazole is sometimes employed as an organobase catalyst for other conjugate addition reactions and its nucleophilicity is comparatively low compared to other amine nucleophiles.

3.3.2. Organocatalyzed Reactions. Transition metal complexes are undoubtedly the catalyst of choice for amino-yne reactions, but organocatalyzed reactions have started to appear in the previous decade. Svete and co-workers developed a series of peptide syntheses employing N-methylmorpholine to catalyze the reaction of primary amines with ynone-terminated peptides at low temperatures (from 0−25 °C), affording vinylogous peptides in up to 92% isolated yields and with 100% E-selectivity or 87% Z-selectivity depending on the functional group. N-Methylimidazole has also been used to catalyze the synthesis of carboxylated pyroles from propiolates, achieving yields of up to 87% (Scheme 18). Proline-catalyzed activation of Michael acceptors is commonly encountered in organic synthesis, and this was employed to facilitate the reaction between an ynal (the aldehyde equivalent of an ynone) and an aniline derivative in quantitative yields at ambient conditions.

Analogous to thiol-yne reactions, trivalent phosphines have also shown efficacy as catalysts for the amino-yne Michael reaction. Kwon and co-workers have reported several phosphine-catalyzed nitrogen-containing heterocycles, synthesized in acetonitrile at ambient temperature and often in quantitative yields. Numerous benzannulated nitrogen heterocycles (such as indoles or benzimidazolines) were synthesized from simple dinucleophiles and acetylenic Michael acceptors by employing 1,3-bis(diphenylphosphino)propane at 20 mol % (Scheme 19a). Generally, the reactions were quite efficient (yields ranging from 82−96%), but trends regarding reaction outcomes according to electrophile (i.e., ynone vs propiolate) did not emerge. A one-pot synthesis of quinolines, with near quantitative yields observed for some substrates, was reported from the triphenylphosphine-catalyzed conjugate addition between o-tosylamidobenzaldehydes with several classes of activated acetylenes (Scheme 19b). Generally, the aryl ynones afforded the best yields (86−99%) followed by the alkyl ynone (88%) and alkyl propiolate (61%), which is not unexpected considering their relative electrophilicities. The efficiency is quite remarkable considering that triphenylphosphine is relatively non-nucleophilic, compared to trialkylphosphines, and a catalytic nucleophilic mechanism is presumably invoked for this case.

3.3.3. Uncatalyzed Reactions. Spontaneous amino-yne reactions are also documented in the literature, especially for reaction.
the highly nucleophilic primary or secondary alkyl amines. However, the extreme reactivity can sometimes lead to deleterious side reactions (such as “over-addition”), especially for primary amines that form a nucleophilic enamine Michael adduct. Together, these features detract from Click characteristics. Anilines are less nucleophilic than alkyl amines, but they have also been reported in spontaneous Michael addition reactions. En route to quinolines, an uncatalyzed, solvent-free conjugate addition of 2-aminobenzaldehyde with an alkyl ynone proceeded in 61% yield with predominantly Z-isomer formation (Scheme 20a). An iron-catalyzed one-pot method was further employed without isolating the intermediate enamine Michael adduct (or vinylogous amide) and afforded substituted quinolines in moderate to excellent yields ranging from 52−94% with aryl ynones being more efficient (Scheme 20b).

Recently, Tang and co-workers reported the uncatalyzed reaction of primary amines with a dual-activated alkyne to form maleimides in ethanol at ambient temperature with isolated yields up to 88%. This result is particularly encouraging because maleimides are usually formed from harsh condensation reactions requiring high temperatures and/or the use of catalysts. In 2010, the Sekiguchi group reported an uncatalyzed amino-disilyne reaction at cryogenic temperatures with a yield of 78%. The Michael addition reaction of various nucleophiles (anilines, amines, thiols, or thiophenol) and 4-hydroxy-2-alkynoates were investigated in water at ambient temperature under ultrasound sonication (Scheme 21). When amines were employed instead of thiols, a cascade conjugate addition/lactonization process afforded 4-amino-furan-2-one derivatives. The initial conjugate addition step was also most efficient when employing thiophenol derivatives, however primary amines also afforded good yields compared to alkyl thiols and anilines. Finally, the importance of water as the solvent was demonstrated and reactions conducted in acetonitrile required heating to 80 °C in order to obtain satisfactory yields.
A particularly powerful example of the amino-yne reaction is the efficient bioconjugation to native molecules without tedious prefunctionalization of the targets. An ynone substrate was quantitatively tagged onto several different biomolecules (including peptides, proteins, and polysaccharides) via the amino-yne reaction. Importantly, the reactions reached completion within minutes at ambient temperatures and also proceeded with high stereoselectivity (∼92% Z-isomer formation). In this study, the amino-yne reaction was more efficient than the analogous thiol-yne or hydroxyl-yne reactions, from reaction with thiols or alcohols, respectively, with the targeted biomolecules (Figure 11). The same group also reported the use of amino-yne Click reactions to efficiently immobilize proteins onto cell surfaces, representing a forward step toward biorthogonality.

In another contemporary example, pectin (which is an enzyme) was conjugated to an amino-functionalized polymer with up to 90% efficiency under ambient conditions and in the absence of a catalyst. Immobilization of the enzyme greatly improved its shelf life with 70% activity still observed after 60 d in addition to increased thermal and pH tolerances.

### 3.4. Applications in Polymer Chemistry

#### 3.4.1. Step-Growth Polymers

Just over 30 years ago, Bass and co-workers reported a new uncatalyzed step-growth polymerization between aromatic diynones and diamines to afford unsaturated nitrogen containing polymers. Although the polymers were isolated in near-quantitative yields, the reaction conditions required high temperatures of 100 °C. Soon after, Wilbur and Bonner extended this motif to aliphatic poly(enamine ketone)s with ≥100% Z-isomer selectivity at ambient temperatures, which brought the reaction more in line with Click criteria. More than two decades later, the polyaddition reaction was again investigated by Tang and co-workers, where they presented a copper-catalyzed amino-yne Click reaction using internal dipropiolate substrates to form a small library of moderate molecular weight (up to 12.8 kDa) polymers under a variety of reaction conditions (Scheme 22a). The use of CuCl was found to be catalytically superior to other salts, and interestingly, the Z-content decreased as the size of the halide increased (CuCl, E/Z = 12/88; CuBr, E/Z = 18/82; CuI, E/Z = 61/39). For the copper-catalyzed reaction: solvent effects on stereochemistry were also scrutinized where the Z-isomer within the unsaturated polymer ranged from 50% (dimethylformamide...
or DMSO) to 88% (tetrahydrofuran), however, the stereochemistry was not strongly correlated to solvent polarity, and the bulk reaction produced the highest stereoselectivity ($E/Z = 6/94$). The polymers were found to have good UV shielding efficiency below 400 nm and nearly perfect transmittance of visible light. The same group subsequently developed a room temperature, spontaneous version of this reaction in dichloromethane to afford high molecular weight polymers (up to 64 kDa) with quantitative conversion after only several hours (Scheme 22b). Disimilar to the metal-catalyzed protocol, this reaction was predominantly selective for the $E$-isomer, which better satisfies Click requirements. Interestingly, the addition of triethylamine afforded a polymer with more mixed stereochemistry, although isomerization to 100% $E$-content was observed when the reaction mixture was left at ambient temperatures for prolonged periods of time. Finally, these two examples also superbly demonstrate the importance of the Michael acceptor on reaction efficiency: the terminal propiolate proceeds without a catalyst at ambient temperature, whereas the former requires high temperatures and the use of a catalyst to yield only modest molecular weight polymers.

A variety of different polymer systems were reported soon after, including high molecular weight poly($\beta$-aminovinylsulfone)s with perfect stereoregularity (100% $E$) from ethynylsulfone monomers. This study is also interesting because it appears that the reaction is stereoselective for the $E$-isomer even when employing primary amine monomers, however, a postpolymerization isomerization to the thermodynamic product cannot be ruled out as a contributing factor. Another recent report described the spontaneous polymerization between aryl yrones and various 1° or 2° diamines and is similar to the substrates employed by Bass and co-workers in their seminal studies. Nevertheless, this work did improve the molecular weight of the resulting polymers (up to 49 kDa) and investigated the polymer stereochemistry where $Z$-isomeric polymers were obtained when employing 2° amines and $E$-isomers for 1° amines, in addition to high thermal stabilities ($T_{ds} > 300 \, ^\circ\text{C}$). In a related example, a multi-component tandem polymerization employing hydrazines, yrones, and terephthaloyl chloride yielded conjugated polypyrrazoles in high yields, all of which had high decomposition temperatures ($T_{ds} > 350 \, ^\circ\text{C}$) and possessed photoluminescent properties. Ni and co-workers have also used an amino-functionalized doxorubicin prodrug to end-cap an amino-yne polymer for drug delivery applications which proceeded at 35 $^\circ\text{C}$ in dichloromethane (Scheme 23).

Recent trends in the field have pivoted away from the use of chlorinated solvents to more benign solvents such as tetrahydrofuran or water (in vivo) and even solvent-free, which are more congruent with Click reaction principles.

Finally, an interesting example featured a sequence-controlled polymer by employing sequential tandem amino-yne/thiol-ene reactions (Figure 12). Amine-terminated sequence-controlled oligomers were first obtained by exploiting the orthogonality between the two nucleophiles, i.e., amine vs thiol monomers, before a subsequent step-growth polymerization with a dipropiolate monomer to afford high molecular weight polymers.
weight (up to ∼60 kDa) materials in dimethylformamide at ambient temperature (up to 82% yield). Although this example loses some Click characteristics, it does showcase good modularity and applicability for the Michael reaction.

3.4.2. Network Polymers. As a consequence of its exceptional efficiency and compatibility with aqueous environments, the amino-yne reaction has recently been employed for the synthesis of hydrogels. Injectable and pH responsive hydrogels based on cross-linking the free amino groups within the biopolymer chitosan provided a platform for tunable drug release behavior (Figure 13). Importantly, the amino-yne cross-linking reaction proceeded spontaneously under physiological conditions which is critical for tissue engineering applications. Langer and co-workers also utilized the amino-yne reaction to afford synthetic hydrogels based on propiolate terminated multiarm PEG precursors. The study was also thorough in its investigation of the substitution pattern for the multifunctional amine nucleophiles. Interestingly, the use of 2° amines efficiently formed gels under all conditions, while 1° amines resulted in gelation at only a high solids content (30 wt %). The greater efficiency of 2° amines vs 1° amines is similar to the trend observed in previously referenced reports for step-growth polymers. Nevertheless, the gels displayed good physical properties that could enhance their biocompatibility including fracture stresses ranging from 16 to 65 kPa, compressive modulus of 6 to 27 kPa, and the ability to swell up to 300%.

The reaction has also been employed to form hyperbranched polymer architectures. In one report, the polymerization of a fluorescent amine monomer was used to probe the aggregation behavior of amphiphilic amino-yne polymers as they self-assembled into nanoparticles. In another interesting study, the amino-yne reaction was used as a junction point to create a miktoarm copolymer. A secondary amine terminated with two alcohols was quantitatively reacted with a propiolate-PEG substrate to afford a difunctional PEG macromonomer. This species was then used to initiate the ring-opening polymerization (ROP) of caprolactone to afford an amphiphilic copolymer that formed nanostructured materials with various morphologies that depended on the polymer arm lengths and reaction conditions. The biorthogonality of the amino-yne reaction, combined with its impressive versatility, should enable innovative biomaterial applications.

4. HYDROXYL-YNE REACTIONS

4.1. Overview and Historical Development

Alcohols have also been widely employed as nucleophiles for 1,4-conjugate addition reactions with activated alkynes in what is now commonly known as the hydroxyl-yne reaction. In fact, some of the first heteronuclear conjugate addition reactions by Ruhemann in 1900 also screened phenolates to furnish α,β-unsaturated esters (Scheme 24).


At first glance, the hydroxyl-yne reaction appears principally analogous to the thiol or amine variants, however, a distinguishing feature is the reduced acidity (relative to the thiol) and nucleophilicity (compared to either) of the hydroxyl species. The effect of this characteristic is that hydroxyl-yne reactions do not proceed spontaneously and demand the addition of a catalyst and/or base to suitably activate the hydroxyl moiety. The net effect of these harsher reaction requirements means that the orthogonality of the hydroxyl-yne reaction can also suffer. Nevertheless, the reaction can still proceed efficiently under certain conditions and there are examples that satisfy Click characteristics. Finally, alcohols are one of the most ubiquitous functional groups in organic chemistry so this balances the comparative inefficiency relative to less abundant heteronucleophiles and also provides palpable motivation for continued progress in the field. However, as a result of the aforementioned drawback of hydroxyl-yne reactions the overall lesser number of Click reactions is still perceptible, especially compared to thiol-yne or amino-yne reactions. Similar to the amino-yne section, we will start with a brief overview on reactivity and then categorize the reactions according to catalyst choice and conclude with examples in polymer chemistry.

4.2. Trends in Reactivity

There are just a handful of studies that investigate structure-reactivity relationships for hydroxyl-yne reactions. The most noteworthy study concluded that the organobase-catalyzed reaction of primary alcohols and propiolates can proceed efficiently (i.e., quantitative conversion in <2 h at ambient temperature), however when secondary and tertiary alcohols are employed as nucleophiles, the reactions are comparatively sluggish and rarely reach completion. Unfortunately, other reaction factors such as the nature of the acetylenic acceptor or solvent effects were not examined, although both are known to be important factors related to reaction outcomes in other conjugate addition reactions. Finally, the authors described the products as primarily Z-isomers, although specific values were not reported in this example.

In general, the hydroxyl-yne reaction is often reported to generally favor the E-isomer over the Z-isomer in many cases. Similarly to the thiol-yne variant, the stereochemical reaction outcome shows dependence on both the solvent and catalyst choice. A detailed study by Inanaga et al. investigated the
effects of catalyst and solvent on the addition reaction between benzyl alcohol and methyl propiolate (Table 12).\textsuperscript{82} When the

![Scheme 25. Palladium-Catalyzed Intramolecular Hydroxyl-yne Cycloadditions to Form Either 1,3-Dihydroisobenzofurans or Isochromenes According to Solvent Selection\textsuperscript{221}](https://doi.org/10.1021/acs.chemrev.0c01076)

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<td>480</td>
<td>86</td>
<td>75/25</td>
</tr>
<tr>
<td>PPh\textsubscript{3}</td>
<td>CH\textsubscript{3}Cl\textsubscript{2}</td>
<td>480</td>
<td>85</td>
<td>83/17</td>
</tr>
<tr>
<td>PPh\textsubscript{3}</td>
<td>THF\textsuperscript{a}</td>
<td>480</td>
<td>62</td>
<td>75/25</td>
</tr>
<tr>
<td>PPh\textsubscript{3}</td>
<td>MeCN\textsuperscript{b}</td>
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<td>83/17</td>
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</tr>
<tr>
<td>PBu\textsubscript{3}</td>
<td>C\textsubscript{6}H\textsubscript{6}</td>
<td>10</td>
<td>83</td>
<td>100/0</td>
</tr>
<tr>
<td>PBu\textsubscript{3}</td>
<td>CH\textsubscript{3}Cl\textsubscript{2}</td>
<td>10</td>
<td>98</td>
<td>100/0</td>
</tr>
<tr>
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<td>THF</td>
<td>10</td>
<td>98</td>
<td>100/0</td>
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<tr>
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<td>MeCN</td>
<td>10 &gt;98</td>
<td>99/1</td>
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</tr>
<tr>
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<tr>
<td>P(O\text{Me})\textsubscript{3}</td>
<td>MeCN</td>
<td>480</td>
<td></td>
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</table>

\textsuperscript{a}THF = tetrahydrofuran. \textsuperscript{b}MeCN = acetonitrile.

Table 12. Screening the Hydroxyl-yne Reaction between Benzyl Alcohol and Methyl Propiolate Using Various Solvents and Different Trivalent Phosphine Catalysts\textsuperscript{42}

reaction was catalyzed by triphenylphosphine, the quantity of E-isomer in the product only fluctuates slightly (75–83%) despite large differences in solvent polarity (e.g., benzene vs acetonitrile). These solvent effects are dramatically smaller than those reported for the thiol-yne reaction.\textsuperscript{76} Moreover, when the reaction is catalyzed by tributylphosphine, the solvent effect is nullified and reaction is stereoselective for the E-isomer. Together the surveyed reaction conditions only provide products ranging from 75–100% E-content. Although this demonstrates a smaller range compared to the thiol-yne reaction, some examples are perfectly stereoselective which satisfies an important Click requirement.

### 4.3. Applications in Organic Synthesis

The most popular use of the hydroxyl-yne reaction has been to synthesize heterocyclic compounds, notably furans. Nozaki and co-workers reported the first furan syntheses via intramolecular hydroxyl-yne reactions\textsuperscript{218} and this has been extensively reviewed in the 1990s.\textsuperscript{219} There are a few recent, excellent summaries that describe the chemistry of ynones and reports therein have suitably captured work in this area from the last few decades.\textsuperscript{4,5} Instead, we will largely spotlight reactions that are excellently summarized in several reviews\textsuperscript{226} so they will not be discussed further.

The nucleophilic hydroxylation of nonactivated acetylenes has also afforded various nitrogen- or oxygen-based heterocycles using gold\textsuperscript{230–234} and copper-catalyzed\textsuperscript{176,177,235} approaches. In particular, the reactions catalyzed by gold species have primarily involved intramolecular cyclization of enyne alcohols and can often be high yielding depending on the functional groups (Scheme 27).

![Scheme 26. Gold-Catalyzed Tandem Conjugate Additions to Form Substituted Furans\textsuperscript{225}](https://doi.org/10.1021/acs.chemrev.0c01076)

The pioneering work on the gold-catalyzed hydroxy-yne addition was accomplished by Larock and co-workers with a tandem conjugate addition reaction where the intermediate enolate attacked the internal alkyne as the nucleophile using AuCl\textsubscript{3} as the catalyst, affording heavily substituted furans (up to 90% isolated yields) under ambient conditions in only 1 h (Scheme 26).\textsuperscript{225} Many comparable gold-catalyzed furan syntheses have been reported using this general approach and are excellently summarized in several reviews\textsuperscript{226–229} so they will not be discussed further.

![Scheme 27. Metal-Catalyzed Intramolecular Hydroxyl-yne Cycloadditions Form Unsaturated Heterocycles\textsuperscript{46}](https://doi.org/10.1021/acs.chemrev.0c01076)

(a) Gold-catalyzed cascade reactions to form bridged heterocycles;\textsuperscript{233} (b) copper-catalyzed synthesis of pyroles and furans.\textsuperscript{275}
4.3.2. Organocatalyzed Reactions. Organocatalysts have become increasingly popular for hydroxyl-yne transformations in recent years. Nucleophilic phosphine catalysts were already mentioned as efficient catalysts for the hydroxyl-yne reaction, however, amine catalysts have also been screened. At 20 mol % catalyst loading, 1,4-diazabicyclo[2.2.2]octane (DABCO) afforded quantitative conversion (along with tributylphosphine) for the 1,4-conjugate addition of octanol to methyl propionate in just 2 h. Interestingly, other amines performed more poorly, including 4-dimethylaminopyridine (DMAP) (26%), triethylamine (38%), and N-methylmorpholine (85%).
Regardless of the catalyst and overall reaction efficiency, the E-isomer was still formed exclusively.

Proline- and pyrrolidine-based catalysts have become very useful after their first report in 2006 by Gouverneur and co-workers, affording yields of up to 90% and E/Z ratios of >99/1. The Taran group later reported use of a trivalent phosphine to catalyze the ambient temperature intermolecular reaction between an azido-alcohol and an ynone to afford 1,3-oxazines or 1,4-oxazepines, dependent upon the substituents (aryl vs alkyl) of the alcohol substrate, in up to 98% yield (Scheme 28).

Several studies have reported proline-catalyzed reaction of ynones and ynals with phenols in quantitative yields as a room-temperature synthesis of chromene derivatives (Scheme 29). Pyrrolidine was employed in a more recent report to catalyze the tandem Michael addition of ortho-substituted aminophenols to ynals at yields of up to 95%. Another study also described the efficient bioconjugation of polysaccharides with ynone substrates by employing DMAP as a catalyst. The authors also examined the stereochemistry of the conjugates, reporting 100% E-isomer stereoconfiguration for the reaction products.

### 4.4. Applications in Polymer Chemistry

In the 1990s, Endo and co-workers disclosed several reports describing polyaddition reactions that satisfy most Click criteria by reacting a variety of diols and electron-deficient diacetylenes. Specifically, the authors employed tributylphosphine to catalyze the polymerizations at ambient temperature in tetrahydrofuran, affording a large library of moderate molecular weight (up to 20 kDa) polymers in excellent yields (up to 94%). More recently, the Tang group reported DMAP as a catalyst for the step-growth polymerization between diphenols (such as bisphenol A) and diynones to produce stereoregular E-isomeric polymers with high molecular weights (up to 35 kDa). On the other hand, the same group also reported polymers with mixed stereo-configurations (~50/50 E/Z) and high thermal stability (T_d > 320 °C) when hydroxyl-yne polymerizations were catalyzed by a phosphazene species. However, because different alkyne monomers (aryl acetylenes vs ynones) were employed, it is difficult to draw meaningful conclusions between the
studies. Very recently, 1,4-diazabicyclo[2.2.2]octane (DABCO) was used to catalyze hydroxyl-yne polymerizations at ambient temperature using propiolate monomers and various diols producing polymers with $T_d > 300 \, ^\circ\text{C}$ and $T_g = 15\sim50 \, ^\circ\text{C}$, indicating most are glassy plastics at ambient temperature.\textsuperscript{246} When aliphatic diols were reacted, the resulting polymers primarily contained E-isomeric units but stereoselectivity diminished when employing phenolic monomers. Although there are only a handful of reports that feature alcohol nucleophiles in Michael addition polymerizations, the monomer scope is already extensive and exemplifies its versatility (Scheme 30). There is also one example of a network polymer where a 4-armed hydroxy-terminated polycaprolactone macromonomer was reacted with bis-propiolate monomers at ambient temperature to afford luminescent gels (Figure 14).\textsuperscript{247} A functionalization reaction of hydroxypropylcellulose (HPC) with an ynone substrate catalyzed by DMAP proceeded at ambient temperature in good yield (Scheme 31).\textsuperscript{299}

5. SUMMARY AND OUTLOOK

Nucleophilic conjugate addition reactions of heteronucleophiles to activated acetylenes, chiefly the thiol-yne, amino-yne, and hydroxyl-yne reactions, have long served as efficient transformations in organic synthesis and more recently within polymer science where their usage has prompted the development of many emergent and innovative materials. In many settings, the Michael reactions can satisfy Click criteria by exhibiting wide scope and good selectivity (orthogonal chemical reactivity, regioselectivity, and stereoselectivity) under mild temperatures and ambient atmosphere. The ability to perform efficient reactions under ambient conditions is one of the greatest hallmarks of the reaction and truly distinguishes it from metal-catalyzed cross-coupling transformations in organic synthesis and the inefficient condensation chemistries currently employed for the synthesis of some commodity polymers.

Nevertheless, reaction requirements and outcomes can vary widely among similar reactions even by simply changing the nucleophile, alkyne, solvent, or catalyst. To date, the field is lacking more fundamental studies that could help establish universal concepts, especially those related to the influence of reaction conditions on stereochemistry, kinetics, and reversibility. By undertaking systematic studies to assess structure-reactivity relationships, this should unlock important insights in order to establish more uniform design principles for the Michael reaction. This could be especially beneficial to applications in bioconjugate chemistry, where reliable protocols are critical to the clinical translation of technologies. It is envisioned that future developments in the field will ultimately seek to optimize conditions (especially relating to catalyst development) to enable a more generalized approach and rationally obtain Click reactivity on demand.

Selective modulation of product stereochemistry by judicious choice of catalyst is a unique advantage compared to many other Click reactions, especially the corresponding thiol-yne reaction. However, although the stereoselectivity can be rationally tuned through careful choice of base or solvent, thus far stereopure reaction products have remained rare. This presents the opportunity to develop more stereoselective catalysts, including other organocatalysts or single-site transition metal catalysts, which are both areas in need of development. Achieving improvements to stereoselectivity will push this family of reactions closer to satisfying complete Click criteria. Stereocontrol will also be critical to the effective translation to a broader range of applications. This is especially important for polymeric materials, where thiol-yne polymers have demonstrated superb structure-property manipulation just by simply changing the stereochemistry of the alkene in the polymer backbone. From a fundamental viewpoint, this is very exciting because it could enable the creation of a material portfolio where polymers are compositionally similar but feature a range of divergent physical properties. By reacting the same monomers and only adjusting the stereochemical content of the polymer, the material behavior could vary between hard/soft or elastic/plastic, which provides an exceptionally powerful approach to synthesizing next-generation, functional materials.

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Notes

The authors declare no competing financial interest.

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Connor J. Stubbs is currently studying his Ph.D. under the supervision of Prof. Andrew Dove at the University of Birmingham. Previously, he graduated with a MChem degree from the University of Hull in 2016. His research interests range from leveraging backbone stereochemistry to manipulate bulk polymer properties and designing chemically recyclable polymeric materials.

Matthew J. Price is a postgraduate researcher under Professor Andrew Dove in the School of Chemistry at the University of Birmingham.
Prior to this, he studied at the University of Cambridge, completing a Master’s in Natural Science including research work on polymer postfunctionalization and supramolecular cage synthesis. His research now focuses on the depolymerization of ‘real-world’ polymer wastes.

Andrew P. Dove is a Professor of Chemistry in the School of Chemistry at the University of Birmingham. Andrew undertook postdoctoral research first under the guidance of Prof. Robert M. Waymouth at Stanford University, California, and then as a CIPMA postdoctoral fellow at IBM, San Jose, California, under the supervision of Dr. James L. Hedrick and Prof. Robert M. Waymouth. Andrew returned to the UK to take up a RCUK Fellowship in Nanotechnology at the University of Warwick in 2005 and moved to the University of Birmingham in 2018. His research focusses on the development and application of sustainable polymers and degradable polymeric biomaterials.

ACKNOWLEDGMENTS

A.P.D., C.J.S. and J.C.W. acknowledge funding from the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation programme under grant agreement No. 681559. M.J.P. thanks The University of Birmingham Global PhD Scholarship for financial support.

REFERENCES


(47) Lowe, A. B. Thiol-yne ‘Click’ coupling Chemistry and Recent Applications in Polymer and Materials Synthesis and Modification. Polymer 2014, 55, 5517−5549.


(64) Yao, B.; Mei, J.; Li, J.; Wang, J.; Wu, H.; Sun, J. Z.; Qin, A.; Tang, B. Z. Catalyst-Free Thiol-Yne Click polymerization: A Powerful and Facile Tool for Preparation of Functional Poly(vinylsulfide)s. Macromolecules 2014, 47, 1325−1333.


Tributylphosphine-catalyzed Reaction of Ethanethiol with Alkynyl Acetylenes Bearing Electron-Withdrawing Groups with Thiols.

Stereochemical Control in the Addition of Thiols to p-Toluenesulfonylacetylene. Synthesis of disubstituted Thiophenes and Oligothiophenes.

Addition of Thiols to p-Toluenesulfo


Substituted Acrylates by Regioselective and Stereoselective Addition of Thiols and Amines to an Alkynyl Ester in Water.

Conjugated Addition of Thiols to Dithianes from Acetylenic Ketones.

Addition of Thiols to Methyl Propiolate.

Comparison of 13 Homogeneous Thiol-X Reactions.


Addition of Thiols to Activated Unsaturated Compounds.


Addition of Thiols to Electrophilic Unsaturated Compounds.


(140) Chuo, Y. Conjugated Polymer Synthesis: Methods and Reactions; Wiley-VCH Verlag, 2010.


