Phosphorus-Based Catalysis
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ABSTRACT: Phosphorus-based organocatalysis encompasses several subfields that have undergone rapid growth in recent years. This Outlook gives an overview of its various aspects. In particular, we highlight key advances in three topics: nucleophilic phosphine catalysis, organophosphorus catalysis to bypass phosphine oxide waste, and organophosphorus compound-mediated single electron transfer processes. We briefly summarize five additional topics: chiral phosphoric acid catalysis, phosphine oxide Lewis base catalysis, iminophosphorane super base catalysis, phosphonium salt phase transfer catalysis, and frustrated Lewis pair catalysis. Although it is not catalytic in nature, we also discuss novel discoveries that are emerging in phosphorus(V) ligand coupling. We conclude with some ideas about the future of organophosphorus catalysis.

1. INTRODUCTION
The study and application of organic phosphorus compounds in organic synthesis have a long history that dates back to the Middle Ages.1 Scientific investigations into these types of compounds, especially from a synthetic perspective, have unveiled the rich chemistry of organophosphorus compounds. In this realm, Michaelis and Arbusov, as well as other chemists, were the pioneers who laid the foundations of what we know as phosphorus chemistry today.1 One significant feature of organic phosphorus compounds is that they can be used as ligands, catalysts, or reagents to facilitate various organic reactions.2,3 When asked about the roles of phosphorus compounds in organic synthesis, however, a majority of organic chemists might offer stereotypical answers: ligands in transition-metal-promoted reactions, reagents for Wittig and Mitsunobu reactions, and organocatalysts for reactions that corresponding nitrogen counterparts also facilitate [e.g., the Morita–Baylis–Hillman (MBH) reaction]. In truth, phosphorus-based organocatalysis is a unique, multifaceted field, with many promising research opportunities and applications. For example, chiral phosphoric acid catalysis4 and phosphine oxide Lewis base catalysis5 play important roles in asymmetric organocatalysis. Frustrated Lewis pairs (FLPs), combinations of Lewis acids and Lewis bases, are important species for the activation of small molecules.6 As a green sustainable tool in organic synthesis, phosphonium salt-based phase transfer catalysis is garnering increasing attention.7 Bifunctional phosphonium catalysts have especially great potential in asymmetric synthesis. Chiral iminophosphorane super base catalysts have demonstrated their usefulness in the deprotonation of high-pKa pronucleophiles for enantioselective transformations.8 An abundance of structural and stereo-
has been made in the field. In this Outlook, we highlight the key features of three such topics: nucleophilic phosphine catalysis, organophosphorus catalysis to bypass phosphate oxide waste, and organophosphorus compound-mediated SET processes. We also provide an overview of several other important topics related to phosphorus-based catalysis in the Miscellaneous Catalyses section. We do not go into too much detail regarding these miscellaneous topics but instead draw the reader’s attention to several comprehensive reviews. Moreover, interesting and useful chemistry is emerging in phosphorus-based reactions. For example, phosphorus(V) ligand coupling, although not catalytic in phosphorus, is becoming a method complementary to transition-metal-catalyzed coupling for the functionalization of heterocycles; we discuss these processes briefly. Although this Outlook is not intended to provide a comprehensive review, we are aiming to give an overview of phosphorus-based catalysis, point out the key developments in the past two years, and proffer our perspective about the future of phosphorus-based catalysis, hoping to inspire researchers to consider the rich chemistry of organophosphorus compounds in their own endeavors.

2. NUCLEOPHILIC PHOSPHINE CATALYSIS

Nucleophilic phosphine catalysis typically operates through initial addition of a tertiary phosphine to an electron-deficient multiple bond, generating a reactive zwitterionic species. Various substrates, including alkenes, allenes, alkynes, and MBH alcohol derivatives (MBHADs), can be involved in this type of activation. The generated reactive zwitterionic intermediates can undergo further reactions with a variety of nucleophiles and electrophiles, or a combination of both. Nucleophilic phosphines also play a special role in the activation of C=N, N=N, strained C=C, or strained C=X (e.g., C=O, C=N) bonds. The broad substrate scope, mild reaction conditions, and diverse reaction modes have established nucleophilic phosphine catalysis as a powerful tool in organic synthesis. Since our group’s comprehensive review of phosphine organocatalysis in 2018, more than 80 new reports of nucleophilic phosphine catalysis have been published. A survey of the progress in nucleophilic phosphine catalysis for the past two years has revealed some interesting features. The main thrusts of the research continue to be the discovery of new patterns of annulation, asymmetric variants of previously reported reactions, and investigations into known reactions using substrates that are much more elaborate than those reported originally. In this Outlook, we classify these reactions of nucleophilic phosphine catalysis based on the nature of their electrophiles with different types of activated multiple bonds—specifically, alkenes, allenes, alkynes, and MBHADs. We further categorize these reactions in terms of the second reaction partner—namely, nucleophiles, electrophiles, and their combinations. In this section, we discuss some recent representative examples of nucleophilic phosphine catalysis to illustrate its unique features and rich chemistry.

2.1. Nucleophilic Phosphine Catalysis of Alkenes

The nucleophilic phosphine catalysis of alkenes involves the addition of phosphine to electron-deficient alkenes 2–1 to form Horner’s zwitterionic adduct 2–2, which has multifarious modes of reaction depending on its reaction partners (Scheme 1). As such, reactions of the zwitterion 2–2 with a pronucleophile (NuH), an electrophile (E), or a hybrid of both can lead to various synthetically useful scaffolds. Recent reports on the reactions of alkenes with pronucleophiles have been limited to the

\[
\text{NuH} \stackrel{\text{PR}_3}{\text{E}} \stackrel{\text{EWG}}{\text{E}} \text{EWG}
\]

phosphine-catalyzed Michael addition, in which pronucleophiles, such as phosphine oxides and TMSCN, have been introduced to produce $\beta$-phosphoryl- and $\beta$-cyano-carbonyls, respectively. Among the reactions of the zwitterionic adduct with electrophiles, the MBH and Rauhut–Currier (RC) reaction are regularly mentioned as representative classical examples. The broad scope and general applicability of these two reactions for the construction of acyclic and carbo- and heterocyclic compounds have undoubtedly contributed to the significant attention that they have garnered from synthetic chemists. Recent progress in these two reactions has focused mainly on using elaborated alkenes to construct complex polycyclic structures in an enantioselective manner. In an elegant example, in 2019, Vicario and co-workers reported a catalytic enantioselective transannular MBH reaction to afford various fused bicyclic compounds containing stereogenic tertiary alcohol moieties in excellent yields and enantioselectivities (Scheme 2). In their proposed model 2–6 for stereoeinduction, hydrogen bonding interactions between the NH functional groups of the bifunctional phosphine catalyst and the electrophilic ketone unit of the substrate were crucial to impart the high enantioselectivity. To further demonstrate its utility, they adopted this new method in the first asymmetric synthesis of the sesquiterpene natural product $\gamma$-gurjunene. Specifically, when the racemic ketone 2–7 was subjected to the conditions, kinetic resolution occurred to afford the desired bicyclic adduct 2–8 in 42% yield and 90% ee, paving the way for the total synthesis of $\gamma$-gurjunene. In another recent example of reactions between the zwitterionic adduct and electrophiles, Lin and co-workers reported the direct $\beta$-acylation of 4-arylidene pyrazolones and 5-arylidene thiazolones with acyl chlorides via phosphine catalysis. When an electrophile–nucleophile is reacted with an alkene, annulation usually occurs. For example, Guo and co-workers reported an efficient annulation between $\gamma$-sulfonamido-enones and sulfamate-derived cyclic imines to furnish multicyclic imidazolidine derivatives.
Tertiary phosphines add to electrophiles even in the presence of electron-deficient alkenes. A recent paper by Zhang and co-workers unveiled a phosphine-catalyzed vicinal difunctionalization of α,β-enones with TMSN₃ (Scheme 3). In this unique transformation, the reaction was initiated by the addition of 1,4-bis(diphenylphosphino)butane (dpbb) to TMSN₃, to form a phosphazide zwitterion 2−12. Michael addition of this intermediate followed by intramolecular cyclization generated 2−13, whose decomposition formed the β-amino α-diazo carbonyl compounds 2−11. To slow down the decomposition of the phosphazide zwitterionic intermediate 2−12 to the iminophosphorane, the bulky TMS group was intentionally selected. A bifunctional chiral phosphine catalyst 2−14 promoted the asymmetric version of this novel transformation, affording moderate enantioselectivities. In a related study, the same group reported a phosphine-catalyzed [3 + 2] cycloaddition of α-diazoacetates and β-trifluoromethyl enones to access 4-(trifluoromethyl)pyrazolines.

2.2. Nucleophilic Phosphine Catalysis of Allenes. Phosphine catalysis of allenes is the most active area of nucleophilic catalysis, due to the diverse modes of reactivity displayed by allenes. In particular, highly functionalized allenes that deviate from the generic versions used in classic reactions often result in unforeseen reactivity patterns. Novel reactions continue to be invented, often accompanied by their asymmetric variants mediated by chiral phosphines, reflective of the potpourri of quality chiral phosphines made available in the past decade, specifically for nucleophilic organocatalysis. The phosphine catalysis of allenes reported in the past two years is presented here in three sections, categorized based on the nature of the second starting material: nucleophile, electrophile, or electrophile—nucleophile.

2.2.1. Nucleophilic Phosphine Catalysis of Allenes with Nucleophiles. Reactions of Allenes with Nucleophiles. Umpolung addition is the most prevalent mode of reaction between an electron-deficient allene and a nucleophile in the presence of a tertiary phosphine catalyst. Among the umpolung additions of allenes, β'- and γ-umpolung additions are the most commonly observed reaction modes (Scheme 4). For allenoates lacking an alkyl substituent at the α-position, γ-umpolung addition may occur; α-alkyl-substituted allenoates undergo β'-umpolung addition. Less common is the Michael addition onto the β-carbon of allenoates, as has been demonstrated with α-fluoro-β-ketoamide nucleophiles. Enantioselective γ-umpolung additions of allenes with nitrogen-centered nucleophiles have continued to be studied. The pronucleophile, either mono or double remote 1,7-addition products can be obtained. Mechanistically, the ε-umpolung addition can be viewed as a vinyllogous γ-umpolung addition triggered by the γ-vinyl substituent on the allenoate. Nucleophilic addition of a phosphine catalyst to the γ-vinyl allenoate 2−15 would form the zwitterion 2−18. Subsequent deprotonation of the pronucleophile and nucleophilic addition at the remote ε-carbon atom, followed by proton transfer and β-elimination of the catalyst, would afford the product 2−16. Among the series of phosphines tested, the bifunctional phophine 2−17 was the best agent. Huang et al. reasoned that the reaction rate was accelerated as a result of additional interactions between the multifunctional phosphine and the substrates.

Reactions of Allenes with Dinucleophiles and Trinucleophiles. While simple nucleophiles are known to promote umpolung addition of allenes, a distinctive [m + n] mode of annulation can occur when dinucleophiles or trinucleophiles are introduced. Furthermore, when combined with multifunctionalized allenes, these nucleophiles can result in several unique modes of annulation. For example, phosphine-catalyzed [3 + 2] annulations of the dinucleophile 2-sulfonamidomalonate with δ-acetoxy allenoates have allowed access to pyrrolines; when vinyl allenoate was reacted with trinucleophiles, complex tetracyclic structures were constructed with the formation of two new rings. Along with those novel annulation processes, new patterns of [4 + 1] annulation to form 5-membered carbo- or heterocycles have been discovered. In an elegant example, Huang and co-workers reported a novel phosphine-catalyzed [4 + 1] cyclization of allenyl imides with methyl ketimines and enamines (Scheme 6). This method provided efficient access to cyclopentenoyl enamines and imines. The phosphazinvinyl methyl ketene 2−21 was proposed as the key 1,4-(bis)electrophile intermediate, which reacted with the nucleophiles 2−22 and 2−24 to afford the cyclopentenones.
Notably, changing the traditional allenoates to allenyl imides resulted in unforeseen reactivity patterns.

2.2.2. Nucleophilic Phosphine Catalysis of Allenes with Electrophiles. Unlike the addition reactions of nucleophiles with allenes, the majority of reactions of allenes with electrophiles forge two new bonds to afford cyclic scaffolds. A diverse range of electrophiles (e.g., alkenes, imines, ketones/aldehydes, aziridines, and azomethine imines) can be employed, rendering phosphine-catalyzed reaction of allenes with electrophiles a powerful tool in organic synthesis.

Lu’s Allene–Alkene/Imine [3 + 2] Annulation. While annulations of allenes with ketones, aziridines, azomethine imines, troponones, and other reaction partners have continued to be explored in this area, Lu’s [3 + 2] annulation has been the most thoroughly studied and adopted for a variety of synthetic applications. Researchers in the field have continued to find new ways to reinvent Lu’s allene–alkene/imine [3 + 2] annulation. Whereas engagement of previously unreported allenes remains a dominant activity, significant attention has also been paid to the preparation of enantioenriched cyclopentenes and pyrrolidines. In 2019, Lu and co-workers reported a phosphine-catalyzed enantioselective [3 + 2] annulation between allenes and isoindigos, resulting in spirooxindoles having two vicinal quaternary stereogenic centers (Scheme 7). The threonine-derived bifunctional phosphine catalyzed the [3 + 2] annulation to afford dimeric spirocyclic bisindoline motifs in high yields and excellent enantiomeric excesses. Notably, the product 2–28 was transformed into the bisoxindole 2–30, which acted as a common intermediate for the bisoxindoles 2–31 [a known precursor of the natural products (−)-diprytophenalene and (−)-WIN 64821] and 2–32 [a known precursor of (−)-chimonanthine and (−)-folicanthine].

Several examples of allene–nitroolefin [3 + 2] annulations have been reported. In particular, the engagement of 3-nitroindoles, 3-nitrobenzofurans, and 3-nitrobenzothiophenes resulting in the dearomative functionalization of electron-deficient aromatic rings demonstrates that phosphine organo-catalysis is potentially a platform that can be complementary to metal-catalyzed dearomative reactions (Scheme 8). Two elegant examples have appeared from the Zhang and Lu groups, who demonstrated the asymmetric dearomative functionalization of electron-deficient allenoates 2–33 with 3-nitroindoles 2–34 catalyzed by the bifunctional phosphines 2–36 and 2–37, respectively. Compared with the preparation of enantioenriched cyclopentenes, examples of enantioselective allene–imine [3 + 2] annulation are rare. To overcome this deficiency, Kwon and co-workers prepared the novel P-stereogenic bicyclic chiral phosphine CarvoPhos from the natural product carvone and applied it to the asymmetric syntheses of a bevy of pyrrolines, including the antiarrhythmogenic small molecule efsevin.
In the context of allene–electrophile [4 + 2] annulation, δ-acetoxy allenoates have been good precursors for the synthesis of benzene rings. In a striking example, Tong and co-workers found that the phosphonium diene intermediate formed by addition of the phosphine to δ-acetoxyallenoates could undergo [4 + 2] annulation with 2-hydroxyquinone derivatives to form naphthoquinones 2–40 (Scheme 9). The efficient and stereoselective synthesis of axially chiral naphthaquinone atropisomers was facilitated by the cooperative effects of the tertiary phosphate and tertiary amine in the bifunctional chiral phosphine NUSIOC-Phos, promoted the reactions between 2-noates acted as novel fi

Scheme 9. Synthesis of Aryl-Naphthaquinone Atropisomers

Allene–Electrophile Multicycle Annulation. When allenes and/or electrophiles having additional functionalities (e.g., conjugated C=CC bonds, a leaving group, or a pronucleophile) are employed, never-before-seen reactions, especially multicycle annihilations, may ensue to give complex polycyclic structures. For example, when γ-benzylallenoates were treated with (E)-2-(3-aryllallylidene)malononitriles in the presence of a phosphine catalyst, multicycle annulation occurred to give bicyclic[3,3,0]-octene derivatives. In another example, a C2-symmetric chiral phosphine, NUSIOC-Phos, promoted the reactions between pyrrolidine-2,3-diones and cyclic N-sulfonyl imines could also generate functionalized benzenes.

Scheme 10. Domino Annulation of δ-Sulfonamidoallenoates to Access Hydroisoquinolines Derivatives

δ-Umpolung Addition. Given that most reactions between allenes and electrophiles have been cyclizations, Xu’s report of a catalytic asymmetric dienylation of para-quinone methides (p-QM) 2–48 with γ-benzylallenoate 2–47 through δ-umpolung addition is noteworthy (Scheme 11). Various optically pure diarylmethanes 2–49 were obtained in moderate to good yields through this method. They proposed that the phosphonium dienolate intermediate 2–51 underwent proton transfer to afford the vinylogous ylide 2–52. Subsequent 1,6-conjugate addition to the p-QM 2–48, followed by a series of proton transfers and β-elimination of the catalyst 2–50, afforded the product 2–49.

2.2.3. Nucleophilic Phosphine Catalysis of Allenes with Electrophile–Nucleophiles. When allenes react with a molecule containing both electrophilic and nucleophilic groups in the presence of catalytic phosphine, annihilations typically occur. Representative examples of nucleophile–electrophiles include salicyaldehyde and salicylamine. Recently, Zhou and co-workers introduced an additional pronucleophile, malonate, into the salicylaldehyde structure to give the dinucleophile 2–51, followed by a series of proton transfers and β-elimination of the catalyst 2–50, afforded the product 2–49.

Scheme 11. δ-Umpolung Addition

2.3. Nucleophilic Phosphine Catalysis of Alkynes. A variety of pronucleophiles, electrophiles, and nucleophile–electrophiles can be involved in the phosphine catalysis of alkynes. Although phosphine catalysis with alkynes has received less attention than that with allenes, reports of known...
reactions with previously unexplored substrates, as well as discoveries of novel reactions, have continued to appear. Among these transformations, Michael addition and $\alpha$-umpolung addition have been the two major reaction pathways when nucleophiles are present (Scheme 13). In one recent example, Salin and co-workers reported a bishydrophosphorylation of electron-deficient alkynes through nucleophilic phosphine catalysis.\textsuperscript{36a} In another example, Zhang and co-workers demonstrated a chiral bifunctional phosphine-promoted double Michael addition of a tryptamine-derived oxindole dinucleophile onto ynones.\textsuperscript{36b} Contemporarily, the Wang group reported the efficient synthesis of (E)-2-nitromethylcinnamates through phosphine catalysis of alkynes (Scheme 14).\textsuperscript{37a} In this transformation, initial phosphine-catalyzed $\alpha$-umpolung addition of the alkyne 2−62 with the nitromethane 2−61 formed the 1,2-disubstituted alkene intermediate 2−64. Subsequent phosphine-promoted 1,3-rearrangement of the nitro group generated the more stable trisubstituted alkene 2−63.

Anticarboboration, silaboration, and diboration of alkynoates can be realized using phosphine catalysis.\textsuperscript{38} A recent example by Santos and co-workers involved PBu$_3$ catalyzing the transphosphinoboration of internal alkynes 2−66 (Scheme 15).\textsuperscript{37b} The reaction was initiated through $\alpha$-umpolung addition of the alkyne 2−68 to form the ylide 2−71. Ensuing proton transfer, followed by an aldol reaction, generated the annulation product benzo[b]azepin-3-one 2−70. 2.4. Nucleophilic Phosphine Catalysis of MBHADs. Together with alkenes, allenes, and alkynes, the MBHADs are also excellent substrates for nucleophilic phosphine catalysis. The MBHADs 2−73 are typically activated by a tertiary phosphine through SN$_2$′ addition to give an active phosphonium intermediate 2−74, which undergoes various reactions when combined with nucleophiles, electrophiles, and nucleophile−electrophiles (Scheme 17). When a nucleophile is present, phosphine catalysis of the MBHAD will involve a unique S$_N$2′−S$_N$2′ process leading to apparent S$_N$2 allylation. As is the case with allenoates, MBHADs can provide three-carbon units in annulation reactions when treated with electrophiles. The known MBHAD−alkene or MBHAD−azo [3 + 2] annulations continue to be reported for asymmetric versions with more elaborate substrates.\textsuperscript{40} MBHADs undergo the [4 + 1] annulations when mixed with enone/enimine/diene electro-
philes in the presence of a phosphine catalyst. Several recent reports have demonstrated the incorporation of various electrophiles, including 2-enoylpyridine (oxide)s, ortho-quinone methides, and α,β-unsaturated imines.41

Alongside these developments, several new reactions have been discovered using more elaborate substrates.42 In one such example, the Huang group reported the domino [3 + 3] cyclization of MBH carbonates 2−75 with nucleophile−electrophile para-quinamines 2−76 to access hydroquinolines 2−77 (Scheme 18).42a The reaction was initiated by the SN2−SN2′ process to afford an allylic amination intermediate 2−78. Subsequent addition with the phosphine, followed by proton transfer, formed the ylide 2−80. Intramolecular Michael addition and elimination of the phosphine generated the hydroquinoline product 2−77. In another example, the same group disclosed a novel phosphine-catalyzed annulation of an MBHAD with perfluoroalkylimidoyl chloride as the electrophile to construct the perfluoroalkylated benzazepines 2−83 (Scheme 19).42b Here, the phosphonium species generated from the MBH carbonate and the phosphine was deprotonated by base; it then added to the fluorinated imidoyl chloride to give the intermediate 2−84. Cyclization of the deprotonated 2−84 and subsequent enamine-to-imine tautomerization afforded the perfluoroalkyl-substituted benzazepines. Notably, the perfluoroalkylimidoyl chlorides provided four atoms of the benzazepine products.

Conventionally, MBH alcohols have been converted to their derivatives, including acetates, carbonates, and halides. A recent report by Guo and co-workers revealed that phosphine catalysis enables the direct reaction of MBH alcohols with azomethine imines to access bicyclic (epoxymethano)-pyrazolo[5,1-b]-quinazolines (Scheme 20).42c They proposed that the MBH alcohol 2−88 was activated through reaction with the phosphine to form the zwitterionic intermediate 2−90. The alkoxide of 2−90 underwent sequential nucleophilic addition to the electron-deficient azomethine imine 2−87, intramolecular Mannich-type reaction, and SN2 displacement of the catalyst to form the bridged tetracycle 2−89.

2.5. Miscellaneous Nucleophilic Phosphine Catalysis.
In addition to activated C—C multiple bonds, tertiary phosphine catalysts can also add to C=O and N=O groups and strained-ring electrophiles, resulting in a diverse array of reactions. Several recent examples are noteworthy. For example, the Behrouz and Rad group reported an ultrasound-promoted PPh3-catalyzed three-component condensation of benzil, urea, and aldehyde for the synthesis of 2,4,5-trisubstituted imidazoles.43 Recent progress in phosphine catalysis has unveiled several unique rearrangements of strained cyclopropanes.44 The Xu group demonstrated that phosphine-catalyzed rearrangement of vinylcyclopropylketones can provide access to cycloheptenones.44a Subsequently, the Li and Xu group reported the synthesis of tri- and tetrasubstituted furans, and trisubstituted dienones, through phosphine-catalyzed rearrangements of alkylidenecyclopropylketones (ACPs, Scheme 21).44b The development of this reaction was inspired by a known palladium-catalyzed rearrangement of ACPs.45 In this new study, three types of substrate-controlled rearrangement of alkylidenecyclopropylketones occurred to afford three different product types. The results could be explained by considering the allylic phosphonium zwitterions intermediates 2−101, 2−102, and 2−103 that formed upon nucleophilic addition of the phosphine to the ACPs. When R1 was an alkyl group, and R2 was an electron-withdrawing group, an Sn2 process gave trisubstituted furan 2−96; when both R1 and R2 were aryl groups, an

Scheme 18. Synthesis of Hydroquinolines

Scheme 19. Synthesis of Perfluoroalkylated Benzazepines

Scheme 20. Phosphine-Catalyzed Reactions of Unmodified MBH Alcohols

Scheme 21. Phosphine-Catalyzed Rearrangements of Alkylidenecyclopropylketones

ACS Cent. Sci. 2021, 7, 536−558

https://dx.doi.org/10.1021/acscentsci.0c01493
\textit{S}_{2,2}' \textit{ pathway formed tetrasubstituted furan 2–98; when } R^1 \text{ was a primary alkyl group } (\text{RCH}_2) \text{ and } R^2 \text{ was an aryl substituent, the intermediate } 2–103 \text{ underwent elimination and the alkene isomerization to afford the trisubstituted dieneone } 2–100. 

\textbf{2.6. Summary.} Nucleophilic phosphine catalysis has remained one of the most vigorous areas of research in the field of phosphorus-based organocatalysis. It continues to expand its horizons to the assembly of carbo- and heterocycles through new modes of reaction. Recent progress in this area has shared some common features. Unforeseen reactivity patterns continue to be discovered when highly functionalized substrates, deviating from generic versions, are used in classical reactions (e.g., Schemes 5 and 6). Enantioselective variants of the classical nucleophilic phosphine catalysis processes continue to be reported (e.g., Schemes 7 and 8). Brand new reactions continue to be uncovered, often accompanied by asymmetric variants when using chiral phosphines (e.g., Schemes 9 and 10). The application of nucleophilic phosphine catalysis is a promising approach for the syntheses of functional molecules (e.g., Schemes 2 and 7). In addition, the zwitterionic adducts generated in situ from nucleophilic addition of a phosphine to electron-deficient multiple bonds can be efficient basic catalysts for asymmetric Mannich-type reactions, Strecker reactions, Michael additions, and aldol reactions. 46 In a recent example, Wang and co-workers revealed that the zwitterion 2–109 generated from the multifunctional chiral phosphine 2–107 and the allenolate 2–108 was highly efficient in the asymmetric decarboxylative Mannich reactions between the cyclic ketimines 2–104 and the \( \beta \)-keto acids 2–105 (Scheme 22). 37 Notably, the cyclization reactions are driven by the formation of strong P(V)\( =\)O double bonds. The formation of a phosphine oxide as waste is, however, an issue that has troubled chemists for a long time. Although the issue can be partly solved through immobilization and precipitation of the phosphine oxide, 48 catalytic cycling of the phosphorus species would be more atom-economic and environmentally friendly. 49 Two strategies, namely, redox-neutral and redox-driven phosphine oxide catalysis, are commonly adopted to recycle phosphorus species in such reactions (Scheme 23). 49,50 The redox-neutral catalysis

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Scheme_22.png}
\caption{Phosphonium Zwitterion-Catalyzed Decarboxylative Mannich-Type Reaction}
\end{figure}

\textbf{Scheme 23. Redox-Neutral and Redox-Driven Phosphine Oxide Catalysis}

requires in situ activation of the phosphine oxide to form an active P(V) intermediate, which will further react with substrates and regenerate phosphine(V) oxide. In the redox-driven catalysis, P(III) usually reacts with substrate 3–1 to form an active P(V) intermediate, which will further react with the substrate 3–2 to give the product and the phosphine oxide. To turn over the phosphine oxide to P(III), stoichiometric reductant is needed. The harsh conditions required for the activation or reduction of the inert P(V) double bond, however, have largely plagued the development of catalytic processes. In the past decade, significant advances have been made in this field. 49 Examples include the development of catalytic Wittig-type reactions using arsine or telluride ylides, 51 gas evolution (N\(_2\), CO\(_2\)) to assist the redox-neutral catalytic process, 49,50,52 and the use of silanes as reductants in the redox-driven catalytic cycle. 49,53 Several new advances have emerged since our review in 2018, 49 and we highlight them below.

\textbf{3.1. Redox-Neutral Phosphine Oxide Catalysis.} In redox-neutral phosphine oxide catalysis, the phosphine oxide is converted directly into another reactive P(V) species. For example, treatment of a phosphine oxide with an isocynate or oxazolyl chloride produces iminophosphorane 52 or chlorophosphonium chloride 50 intermediates that have been used in the aza-Wittig or Appel reaction and in the dehydration of oximes to nitriles. The use of this mode of phosphine oxide catalysis, although known since 1962, has lagged behind the thriving activities of redox-derived catalysis. 54 Recently, however, in what could be considered one of the most exciting innovations of late, Denton and co-workers developed a redox-neutral organocatalytic Mitsunobu reaction (Scheme 24). 54 The Mitsunobu reaction is commonly used for the nucleophilic substitution of primary and secondary alcohols with inversion of their chiral centers in a stereospecific manner. Despite its usefulness, the requirement of stoichiometric amounts of reagents (PPh\(_3\) \_ azo oxidant) and the generation of undesired waste have undermined this powerful transformation. Although several attempts have been made to develop a redox-driven organocatalyzed Mitsunobu reaction, a stoichiometric amount of reductant has been required to recycle the phosphine oxide, and a stoichiometric amount of oxidant has been required to regenerate the azo oxidant in cases where a substoichiometric amount of the azo species is used. 59 In Denton’s study, a
phosphine oxide catalyst 3–5 was employed without any other additives. Notably, the previously required azo oxidant is not necessary here, because the active dihydrobenzooxaphospho-

Scheme 25. Reductive C–N Cross-Coupling between
Nitroarenes and Boronic Acids

3–11. The second deoxygenation began with addition of the phosphete to the nitrosoarene to form the betaine intermediate 3–16, which underwent 1,2-migration to give the desired product with concomitant generation of the phosphete oxide 3–11. The Radosevich group recently demonstrated that nitromethane is amenable to the reaction, realizing methylation of arylboronic acids.61c When the nitroarenes have different ortho substituents, sequential reductive coupling and cyclization could occur efficiently to yield various heterocycles (Scheme 26).62 A tandem coupling/acylation would generate oxindoles 3–18 and quinoline-diones 3–19, while a tandem coupling/condensation would form indoles 3–20 and benzimidazoles 3–21.

Scheme 26. Synthesis of Azaheterocycles through Reductive Coupling between Nitroarenes and Boronic Acids

One of the reactions that is mediated by a phosphine, but not widely known, is the cross-coupling between nitroarenes and boronic acids for the synthesis of di(hetero)arylamines, reported by Csáky.60 In 2018, the Radosevich group disclosed a deoxygenative C–N crosscoupling between nitroarenes and boronic acids (Scheme 25).61a,b While transition-metal-mediated C–N couplings are used widely, the P(III)-catalyzed reaction is a complementary main group re-oxidation method. In particular, this transformation tolerates various functional groups, including a series of heterocycles and halogen substituents. Radosevich et al. proposed a two-stage deoxygenation sequence for the reductive C–N coupling. Initially, [3 + 1] cheletropic addition of the nitrobenzene 3–8 and the phosphete 3–12 generated the intermediate 3–13, decomposition of which furnished nitrosoarene and the oxide

3–11. The second deoxygenation began with addition of the phosphete to the nitrosoarene to form the betaine intermediate 3–16, which underwent 1,2-migration to give the desired product with concomitant generation of the phosphete oxide 3–11. The Radosevich group recently demonstrated that nitromethane is amenable to the reaction, realizing methylation of arylboronic acids.61c When the nitroarenes have different ortho substituents, sequential reductive coupling and cyclization could occur efficiently to yield various heterocycles (Scheme 26).62 A tandem coupling/acylation would generate oxindoles 3–18 and quinoline-diones 3–19, while a tandem coupling/condensation would form indoles 3–20 and benzimidazoles 3–21.

Scheme 26. Synthesis of Azaheterocycles through Reductive Coupling between Nitroarenes and Boronic Acids

Compared with the oxidation of S(II) species into valuable sulfoxides and sulfones, the deoxygenative transformation of sulfur-containing compounds is less recognized as synthetically useful. The Sharpless group had previously demonstrated that trialkylphosphite could reduce sulfenyl chlorides to corresponding sulfenyl chlorides and, subsequently, to sulphenyl chlorides, and that the sulphenyl chloride could be captured by nucleophiles (e.g., alcohols) to form sulfinate esters.63a The final step in their proposed mechanism for the sequential deoxygenation of sulfonyl chloride was the reaction of the sulfenyl chloride with trialkylphosphite to form trialkyl(arylthiol)phosphonium chloride, which underwent a Michaelis–Arbuzov reaction to produce the corresponding phosphorothiolate and alkyl chloride.63b Recently, Radosevich and co-workers reported that the phosphete oxide 3–11 was capable of catalyzing electrophilic sulfonylations through deoxygenation of sulfonyl
chlorides with indoles as the nucleophile (Scheme 27). The reaction is suitable for a diverse range of indoles and sulfonyl chlorides, providing access to indole derivatives. For this 2-fold deoxygenative sulfonylation, 3−25 was proposed as the key intermediate responsible for the electrophilic substitution.

Again, the P(V)/P(III) cycle was driven by phenylsilane-mediated reduction of P(III) oxide to P(V). Halophosphonium cations have proven to be effective for deoxygenation in both the Appel reaction and amide formation. This behavior inspired the development of deoxygenative sulfenylation, which collapsed to the product 3−11 and phenylsilane was used to replace the previously employed stoichiometric phosphine. The reduction selectively targeted the activated double bonds without affecting other functional groups susceptible to hydrogenation. Remarkably, water was tolerated by the silane-promoted P(III)/P(V) redox cycling.

Scheme 27. Electrophilic Sulfenylation through Deoxygenation of Sulfonyl Chlorides

![Scheme 27. Electrophilic Sulfenylation through Deoxygenation of Sulfonyl Chlorides](image)

The annulation was amenable to both aniline and benzylamine for the construction of six- and seven-membered rings. The key to the success of this sequential annulation is the generation of the halophosphonium cation 3−29 from the in situ-generated phosphetane and the oxidant diethyl bromomaldehyde (DEBM). Activation of a carboxylic acid with the phosphetane cation, followed by coupling, resulted in the first dehydration to form an amide intermediate. The amide bond was further activated by the phosphetane species, with the ensuing S₈Ar cyclodehydration generating the desired heterocycles.

Alkene reduction is one of the most fundamental reactions in organic synthesis. While transition metal catalysis dominates the field, organocatalysis approaches may complement the former by providing unique selectivity and functional group compatibility. Indeed, several successful examples of phosphine-mediated reduction of activated double and triple bonds have been reported. These reductions have involved addition of the phosphine to the double or triple bond to form the zwitterion 3−32 and a subsequent reaction with water to form the phosphorane 3−33, which collapsed to the product 3−31 and the phosphate oxide (Scheme 29). A stoichiometric quantity of the phosphate was, however, required, producing undesired phosphate oxide waste. Last year, the Werner group reported the first example of a phosphine oxide-catalyzed alkene reduction through silane-mediated P(III)/P(V) redox cycling. In this transformation, a combination of the strained phosphetane oxide 3−11 and phenylsilane was used to replace the previously employed stoichiometric phosphine. The reduction selectively targeted the activated double bonds without affecting other functional groups susceptible to hydrogenation. Remarkably, water was tolerated by the silane-promoted P(III)/P(V) redox cycling.

3.3. Asymmetric Phosphine Oxide Catalysis. Although chiral phosphines have been recognized as reagents for asymmetric Wittig olefination and Staudinger−aza-Wittig reactions, explorations in this field have been limited by the noneconomical nature of those reactions. Today, significant advances in phosphine oxide catalysis have given an impetus for renewed interest in this area, such that enantioselective processes driven by the P(III)/P(V) redox cycling of chiral phosphine oxides are emerging as powerful tools for the synthesis of chiral molecules. In particular, desymmetrization of meso compounds is a valuable strategy for accessing molecules with high optical purity. In 2014, Werner et al. reported the first enantioselective catalytic Wittig reaction through desymmetrization of 2-(3-bromo-2-oxopropyl)-2-methylcyclopentane-1,3-dione with catalytic Me-DuPhos (3−37) and phenylsilane as the terminal reductant. They found that Me-DuPhos worked efficiently for asymmetric induction with up to 90% ee, but with less than 10% yield. Three years later, the Christmann group reinvestigated this reaction and found that the desymmetrizing Wittig reaction could provide high enantiomeric excess (92−96% ee) and moderate yields (Scheme 30). They found that premature debromination of the triketone, caused by impurities in the starting materials, might have been responsible for the low yield reported initially. Although high temperature (150 °C) was required for the catalytic intramolecular Wittig reaction, good

Scheme 29. Reduction of Activated Alkenes through P(III)/P(V) Redox Cycling Catalysis

![Scheme 29. Reduction of Activated Alkenes through P(III)/P(V) Redox Cycling Catalysis](image)

Scheme 30. Synthesis of ent-Dichrocephone A and ent-Dichrocephone B

![Scheme 30. Synthesis of ent-Dichrocephone A and ent-Dichrocephone B](image)
enantioselectivities were obtained when using Me-DuPhos. The optically pure bicyclic enone products were further elaborated into various carbocyclic propellanes through sequential Michael addition with Lipshutz’s cyanocuprates and ring-closing metathesis.\textsuperscript{70b} The propellane 3–39 was applied to the first asymmetric syntheses of ent-dichrocephone A and ent-dichrocephone B.\textsuperscript{70c}

Although good selectivity was obtained in some cases for the asymmetric Wittig olefination at high temperature, mild room-temperature reactions would be more desirable. The inert nature of P(V)═O poses a challenge for its silane-mediated reduction to a trivalent phosphine at ambient temperature. Nevertheless, continued studies in this field have unveiled some factors for lowering the energy barrier of the silane-mediated phosphine oxide reduction, such as the use of the strained small-ring phosphetane oxide 3–11 as the catalyst. Interestingly, strained phosphine oxides can be reduced more readily than unstrained ones. A recent computational study explained why strained phosphetanes and bridged [2.2.1] bicyclic phosphines are privileged structures in redox-driven phosphine oxide catalysis.\textsuperscript{71} One reason is that one of the C—P—C angles of these phosphines is close to that in the proposed transition state for the intramolecular hydride delivery in the (phosphoniooxy)silicate(IV) adduct formed during the phosphine oxide and the silane. Studies have also demonstrated that both acidic and basic additives can facilitate the silane-mediated phosphine oxide reduction.\textsuperscript{72} Consolidating these lessons, in 2019, Kwon and co-workers developed a catalytic asymmetric Staudinger−aza-Wittig reaction for the synthesis of chiral cyclic imines 3–44 (Scheme 31).\textsuperscript{73} In this reaction, the commercially available \( P\)−chiral [2.2.1] bicyclic endo-phenyl-HypPhos 3–45 was used to facilitate desymmetrization of the two C═O groups of various 1,3-diketones and to form enantioenriched iminoketones containing a chiral quaternary center. The strained nature of the catalyst, in conjunction with the 2-nitrobenzoic acid additive, enabled room-temperature redox cycling mediated by phenylsilane.

In another recent report, the Voituriez group documented an efficient method to access enantioenriched (trifluoromethyl)cyclobutenes 3–48 through an asymmetric \( \alpha\)-umpolung addition−Wittig olefination cascade between the dione 3–46 and the acetylenedicarboxylate 3–47 (Scheme 32).\textsuperscript{74} The catalyst employed here, \( \varepsilon\text{-anisyl-HypPhos} (3–49)\), is also a strained [2.2.1] bicyclic phosphine oxide, which, along with the phosphoric acid additive, facilitated mild P(III)/P(V)═O redox cycling. This tandem reaction represents another powerful example of phosphine oxide redox cycling as a tool for the synthesis of attractive chiral molecules.

3.4. Summary. Although significant progress has been made, challenges remain in the area of phosphine oxide catalysis. Most redox-driven phosphine oxide catalysis has been based on silane-mediated in situ reduction. The need for a stoichiometric amount of reductant raises economic and environmental concerns.\textsuperscript{75} A recent study by Sevov and co-workers revealed that the triphenylphosphine oxide byproduct of the Wittig olefination can be transformed directly into \( \text{PPh}_3\) through electroreduction (Scheme 33).\textsuperscript{76} Revitalized adaptation of such unconventional technologies may open new doors for redox-driven phosphine oxide catalysis. Conversely, as demonstrated by Denton and co-workers in their redox-neutral organocatalytic Mitsunobu reaction, designing novel catalysts may also be a promising avenue for green chemistry. Although catalytic versions of those phosphine oxide-driven reactions are appealing, unless their selectivities and yields are comparable with those of the original versions, their utility will be limited. In this regard, there is a great need to develop mild methods for reduction or activation of phosphine oxides. Accordingly, asymmetric phosphine oxide catalysis is another appealing field benefiting from ongoing progress in phosphine oxide catalysis. Although much needs to be done, phosphine oxide catalysis finds itself at the threshold of a bright future.

4. PHOSPHORUS-MEDIATED RADICAL PROCESSES

Organophosphorus compound-mediated SET processes have received much less attention than their nitrogen counterparts,\textsuperscript{77} despite trivalent phosphine radical cations and phosphoranyl radicals having been recognized as reaction intermediates as early as the 1950s.\textsuperscript{78} Progress in this emerging field has established that organophosphorus compounds can act as radical initiators, catalysts, and radical precursors in various SET transformations for the synthesis of important structural motifs.\textsuperscript{79} With the rapid development of photocatalysis in recent years, the use of organophosphorus compounds as radical precursors, especially for the purpose of deoxygenation or desulfurization, is becoming established as a reliable method.\textsuperscript{79} Several examples of organophosphorus compound-mediated radical processes are discussed below.
4.1. Use of Substoichiometric Phosphine. Electron donor—acceptor (EDA) complexes, also known as charge-transfer complexes (CTCs), are typically formed from electron-rich and electron-poor compounds and possess distinct properties differing from those of their parent compounds, enabling some unique transformations. In particular, even when the parent compounds cannot absorb visible light, their EDA complexes can, facilitating useful transformations. In step with recent attention paid to the study of EDA complexes, especially in the area of photoredox chemistry, the use of phosphorus compounds, which are good electron donors for forming EDA complexes, is gathering momentum. In 1990, the Huang group discovered that a catalytic amount of PPh₃ could promote the perfluoroalkylation of simple alkenes with perfluoroalkyl iodides through a radical chain reaction (Scheme 34). In this transformation, an EAD complex 4→4, formed from PPh₃ and perfluoriodine 4→1, would decompose to the radical intermediate R. Addition of R to an alkene generates the alkyl radical intermediate 4→5. The reaction of 4→5 with perfluoroiodine forms the product 4→3 and regenerates R for radical chain propagation. While subsequent examinations of this reaction were conducted under more or less the same thermal conditions as those in Huang’s original study, Czekelius and co-workers found in 2019 that light also facilitated activation of the EAD complex, allowing the reaction to occur at close to ambient temperature (Scheme 34). A systematic study revealed that Bu₄P and light from a blue LED (461 nm) were optimal for promoting this radical chain reaction.

In the EDA complex-initiated radical processes discussed above, the substrates are involved in the EDA complexes and incorporated into the final products. This strategy requires careful selection of substrates to match the electronic properties of the phosphines, limiting their scope and, consequently, the utility of this strategy. A more general and effective strategy, however, may involve a CTC catalyst that can transfer an electron to a substrate to assist the radical generation and later accept an electron from the late-stage intermediate to complete the catalytic cycle. This strategy was realized by the Fu group in 2019. Although most photoredox catalysts involve precious metal complexes or organic dyes, they demonstrated a new type of photocatalytic decarboxylative alkylation using the combination of PPh₃ and sodium iodide (Scheme 35). This novel alkylation was realized with several types of alkyl radical precursors (decarboxylative alkylation via redox-active esters, deaminative alkylation via Katritzky’s N-alkylpyridinium salts, and trifluoromethylation with Togni’s reagent) and radical acceptors (e.g., silyl enol ethers, activated heterocycles, and 1,1-disubstituted alkenes). In the proposed mechanism, Fu et al. suggested that the CTC 4→10 is formed initially through Coulombic interactions between PPh₃, sodium iodide, and the N-(alkylcarbonyl)phthalimide 4→8. SET from iodide to the phthalimide moiety then prompts decomposition and concomitant generation of an alkyl radical, which adds to the radical acceptor. Finally, the transient radical cation complex 4→12 oxidizes the amine radical cation 4→15 to the product 4→16 and turns back to the PPh₃-NaI complex 4→10. A chiral phosphoric acid could enable enantioselective Minisci-type reactions with high levels of stereoselectivity. Interestingly, the blue LED light (456 nm) used here is very similar to that (461 nm) used in the perfluoroalkylation of alkenes (Scheme 34), suggesting that related complexes might have formed.

In a related study, the He, Xue, and Chen groups reported a radical C–H arylation of oxazoles with aryl iodides. They proposed that the complex D formed from Cs₂CO₃ and 1,1'-bis(diphenylphosphino)ferrocene (dpff) could serve as a strong electron donor, reducing aryliodine to form the radical anion aryliodine complex 4→20 and further initiating a radical chain reaction (Scheme 36). The radical 4→20 would undergo C–I bond cleavage to form the aryl radical 4→21. Addition of 4→21 to

![Scheme 34. Phosphine-Initiated Perfluoroalkylation of Alkenes](image)

![Scheme 35. Photocatalytic Decarboxylative Alkylations Mediated by PPh₃ and NaI](image)

![Scheme 36. dpff/Cs₂CO₃-Mediated Oxazole C–H Arylation](image)
oxazoles, followed by deprotonation, would form the radical anion 4–24. Reduction of the aryl iodide by the radical anion 4–24 would form the product 4–25 and regenerate the radical anion 4–20 for radical chain propagation. The radical anion 4–24 could also reduce D(+)1 to D and complete the redox cycle. DFT calculations indicated, however, that dissociating CsCO3− from D(+)1 to generate a potential electron acceptor (dppe)3+ is thermodynamically challenging. The authors favored the radical chain reaction pathway, while not ruling out the redox cycle.

In addition to complexing with other compounds to form catalysts for SET processes, phosphorus compounds themselves can also act as electron shuttles in redox cycles. In 2015, Tan and Liu employed Togni reagent II as the source of trimethylphosphine for the phosphine-catalyzed bistriﬂuoromethylation of enamides (Scheme 37).86a The reaction involved multiple SET processes, with mechanistic studies indicating that the catalytic cycle was driven by redox cycling of trivalent phosphine and a phosphorus-centered radical cation. Initially, the phosphine 4–28 was oxidized by the Togni reagent II through SET to form the CF3 radical and the phosphorus-centered radical cation 4–30. Addition of the CF3 radical to the alkene, followed by a 1,5-hydride shift, generated the α-amidobenzyl radical 4–33. Subsequent oxidation by the Togni reagent II and deprotonation formed the enamide 4–35. Addition of the CF3 radical to 4–35 afforded the radical intermediate 4–36, which was further oxidized by the phosphorus radical cation and deprotonated to give the product 4–38. The SET oxidation also recycled the phosphine catalyst 4–28. This method was further developed by the authors to allow concomitant trifluoromethylation of the alkene and subsequent remote alcohol or amine α-C–H activation to afford valuable trifluoromethyl-substituted ketones and aldehydes.86b

4.2. Use of Stoichiometric Phosphine. In addition to their roles as radical initiators and catalysts as described above, trivalent phosphorus(III) compounds are increasingly being used as radical precursors for SET transformations.8 Although, at present, stoichiometric reagents (e.g., ylide or phosphine) are required, the byproducts formed in the deoxygenation or desulﬁtization process may be recycled in situ, as we have discussed above regarding phosphine oxide catalysis. Mechanistically, fragmentation of phosphoranyl radicals, through either α- or β-scission, is an efﬁcient means of forming radicals for further transformation. The mechanisms of formation of phosphoranyl radicals can be distinguished by whether they involve addition of an external radical to a phosphine or addition of a nucleophile to a phosphorus-centered radical cation, and also by whether light is involved in the formation of the phosphoranyl radicals. Because this emerging ﬁeld has been summarized in two recent reviews,9 we discuss only selected examples in this Outlook, including the α- and β-scissions of phosphoranyl radicals, light-associated and non-light-associated SET processes, as well as two modes of phosphoranyl radical generation.

Miura and co-workers reported recently the photoredox-driven fragmentation of phosphoranyl radicals through α-scission to generate active radical species.9 They disclosed a practical and efﬁcient synergistic photoredox approach for the preparation of elongated esters from alkenes and stabilized phosphorus ylides under mild conditions (Scheme 38). The phosphoranyl radical intermediate generated through [IrIII]*-mediated reduction of the phosphonium salt 4–42 further fragmented into PPh3 and the radical 4–44 for subsequent addition to alkenes. Notably, the phosphoranyl radical was generated through neither of the methods described above.

The Zhu and Xie group provided an example of photoredox-driven fragmentation of phosphoranyl radicals through β-scission in their efﬁcient approach to ketones through the deoxygenation of aromatic carboxylic acids using synergistic photoredox catalysis in the presence of PPh3 (Scheme 39).86a

The reaction was initiated through oxidation of PPh3 to its radical cation, mediated by a photoexcited catalyst. The radical cation coupled with the carboxylate anion to generate the phosphoranyl radical intermediate 4–46, the β-scission of which generated the acyl radical 4–47. The ﬁnal product was formed through radical addition to an alkene. In addition, corresponding deuterated aldehydes could be formed by the acyl radical intermediate abstracting a deuterium atom from D2O.86b
Concurrently, the Doyle group reported the deoxygenation of alcohols and carboxylic acids 4−48 through a similar phosphine-promoted photoredox-catalysis reaction.90,91

Whereas the examples above involving phosphorus compound-mediated SET were driven by photocatalysis, phosphoranyl radical intermediates can also be formed under thermal conditions. In 2018, the Schmidt group reported an intermolecular anti-Markovnikov hydroamination of alkenes using triethylphosphate and N-hydroxyphthalimide (Scheme 40).89 The reaction was initiated by hydrogen atom abstraction from N-hydroxyphthalimide to produce PhthNO•, which added to triethylphosphite to form the phosphoranyl radical intermediate 4−54. Subsequent β-fragmentation generated the radical PhthN•, which added to the alkene 4−51 to generate the radical 4−56. Abstraction of a hydrogen atom from 4−50 formed the anti-Markovnikov alkene hydroamination product 4−52 and regenerated PhthNO•.

### 4.3. Summary

Organophosphorus compound-promoted SET reactions have reactivity complementary to that of nucleophilic phosphine catalysis. In nucleophilic phosphine catalysis, the substrates are limited to electron-deficient alkenes, whereas inert alkenes (as demonstrated in Schemes 34 and 37−40) and C−H bonds (as demonstrated in Schemes 35−37) can be activated in SET processes. Special attention may be given to the EDA complexes of phosphorus compounds, which are sensitive to light and can either initiate or catalyze SET processes. Exploration of the EDA complexes of phosphorus compound should unveil rich chemistry in this underdeveloped area. A special type of EDA complex, the FLP, can be a powerful tool in the activation of small molecules. Although a two-electron-transfer mechanism is generally recognized, recent reports have indicated that SET processes might also play important roles in FLP chemistry. An interesting study by Stephan and co-workers unveiled that a frustrated radical pair Ph(Mes)P•/[E(C6F5)3] (E: B/Al) could be formed through a SET process, while the analogous Bu3P pair behaved differently without undergoing such a process. This finding, as well as others, provides evidence that the structural features of tertiary phosphines greatly influence the stability of phosphorus-centered radical cations and can even alter the reaction pathway. Moreover, a tertiarly phosphine itself can function as an electron shuttle in a redox-catalytic cycle. Organophosphorus compound-promoted SET processes, therefore, have huge potential for applications in photochemistry and electrochemistry. The limited number of reports of organophosphorus compound-catalyzed SET reactions may be due to the short half-lives of trivalent phosphine radical cation species. Designing phosphines or their complexes with structural frameworks of suitable electronic potential may result in longer half-lives for the open-shell species and benefit this area. The unique role of organophosphorus compounds to generate radical intermediates through SET processes is also appealing.

### 5. MISCELLANEOUS CATALYSTS

In addition to nucleophilic phosphine catalysis, catalysis to bypass phosphine oxide waste, and phosphorus-mediated radical processes, organophosphorus compounds are used in multitudes of catalytic processes, including phosphonium salt phase transfer catalysis (PTC), iminophosphorane super base catalysis, asymmetric phosphine oxide Lewis base catalysis, FLP catalysis, and chiral phosphoric acid catalysis. PTC is considered a powerful and green sustainable tool for addressing synthetic challenges. In this context, phosphonium-containing organocatalysts, especially bifunctional catalysts, are emerging as powerful phase transfer catalysts for various asymmetric transformations (e.g., alkylation, Michael addition, Darzens reaction, Mannich reaction, and Strecker reaction). Deprotonation of pronucleophiles with chiral Bronsted bases (e.g., cinchona alkaloids and derivatives) to induce enantioselective reactions via ion pairs is an established reliable method for realizing various enantiodifferentiating transformations. Although powerful, the application of this method to high-pKₐ pronucleophiles has been limited. The availability of phosphorus-based super bases [e.g., P-spiro chiral triaminomorphosphorane, chiral bis( guanidino)iminophosphorane, and bifunctional iminophosphorane], with values of pKₐ ranging from 22 to 32.9, has opened a new door in this area, realizing various challenging reactions that are difficult to perform using cinchona-based catalysts. Another type of phosphorus-related base catalysis is asymmetric phosphine oxide Lewis base catalysis. Generally, a polarized phosphine oxide bond coordinates to chiral phosphines to form pentavalent silicon complexes, thereby enhancing the electrophilicity of the silicon species to activate electrophiles (e.g., carboxyls, epoxides, and imines) for further transformation. In the presence of chiral phosphine oxides (e.g., chiral binaphthyl dioxides and chiral phosphoramides), a variety of asymmetric transformations (e.g., allylation, aldol, hydrophosphonylation, and reduction of unsaturated compounds) can be promoted.

Alongside base catalyses, phosphorus-based Bronsted acid catalysis has also become established as a reliable and efficient method to form a variety of C−C, C−H, and C−X bonds enantioselectively. Since the introduction of chiral phosphoric acid into catalytic asymmetric reactions in 2004, the following years have witnessed tremendous progress, resulting in the development of a variety of chiral acids (e.g., BINOL derivatives, TADDOL derivatives, VAPOL derivatives, and spiro phosphoric acids). Today, phosphorus-based Bronsted acid catalysis remains an active area of inquiry. Finally, FLPs, formed from combinations of Lewis acids and Lewis bases, have been used in the development of main group element catalysis processes for the activation of small molecules (e.g., H₂, CO₂, CO, SO₂). In this context, various phosphines have played important roles in combination with Lewis acids [e.g., B(C₆F₅)₃, Al(C₆F₅)₃] to form active FLPs. Although such studies constitute a large genre of the field of organophosphorus-based catalysis, we cannot cover them in this Outlook, due to limited space; therefore, we direct the reader’s attention to recent reviews on each topic.
Although the scope of organophosphorus-based catalysis is large, most of its topics are “young” in that they have been established as fields of systematic inquiry only within the last 20 years. For example, redox-driven phosphine oxide catalysis began to gain significant interest in 2009 after O’Brien published the catalytic Wittig reaction;\(^\text{93}\) the phosphorus compound-mediated SET process has been attracting more attention recently as a result of the rapid development of photoredox chemistry;\(^\text{94}\) FLP chemistry became popular only after Stephan reported, in 2006, that H\(_2\) can be activated by an FLP.\(^\text{95}\) Notably, one of the original functions of chiral phosphoric acid, first reported in 1971, was as a stoichiometric reagent for the resolution of racemic amines.\(^\text{94}\) The rich chemistry of phosphorus compounds will enable more interesting reactions to be added to the toolbox of organic synthesis. One lesson we can take here is that new chemistry is emerging and that previous stoichiometric reactions may turn into catalytic ones in due course. Although not catalytic in its organophosphorus reagents, phosphorus(V) ligand coupling has been gaining renewed interest recently. We discuss these so-called “contractive coupling” reactions briefly in the next section.

### 6. PHOSPHORUS(V) LIGAND COUPLING

Like the other main group elements below the second row of the periodic table (e.g., S, I, Si), phosphorus can adopt hypervalent states featuring an expanded valence-shell, tending to extrude an electron pair to form a more stable octet.\(^\text{93}\) The collapse of the hypervalent species results in various transformations, one of which is ligand coupling.\(^\text{96}\) The notion of “ligand coupling” was first introduced by Oae to describe the exclusion of two ligands from hypervalent species to form a stable octet and the coupling product.\(^\text{96,97}\) It was also termed “contractive coupling” by Newkome earlier in describing a similar process for the synthesis of biheteroaryls.\(^\text{98}\) While the concept has been applied to various main group elements,\(^\text{97}\) recent progress has demonstrated that P(V)-mediated ligand coupling can be a powerful method for heterocycle functionalization.\(^\text{98}\) Many heterocycles, including pyridines, diazines, and benzoazoles, are important pharmacophores. While transition-metal-mediated functionalization of aromatic rings and electrophilic aromatic substitution are very well established, functionalization of these heterocycles remains a big challenge in terms of site-selectivity and functional group compatibility.\(^\text{100}\) In this context, phosphorus ligand coupling has found itself very useful in medicinal chemistry. Although a stoichiometric amount of reagent is required at present for phosphorus ligand coupling, the contractive coupling process mimics reductive elimination in transition-metal-catalyzed coupling (Scheme 41). Recently, the main group element-mediated two-electron redox catalysis has demonstrated its potential in coupling reactions.\(^\text{101}\)

The Radosevich group reported a T-shaped P(III) compound as the catalyst for transfer hydrogenation of azobenzenes,\(^\text{101b}\) and Cornella and co-workers unveiled a novel Bi complex that facilitates the fluorination of aryloboronic esters through redox catalysis.\(^\text{101c,d}\)

Such progress sheds light on the potential for catalytic phosphorus-mediated ligand coupling. Below, we discuss some recent advances in the area of phosphorus ligand coupling.

The phenomenon of coupling between two ligands has been observed frequently at the phosphorus center.\(^\text{97,98,102,103}\) For example, phosphine oxides or phosphonium salts bearing two or three 2-pyridyl groups can undergo contractive ligand coupling to afford bipyrindines.\(^\text{98,102c,e}\) The introduction of the heterocycles at the phosphorus center was, however, the most challenging step in those reports. In Newkome’s study, phenylpyridylphosphine (oxide)s were prepared through reactions between diliathiophosphinephosphate and halopyridines or through addition of 2-lithiopyridines to dichlorophenylphosphine.\(^\text{98}\) The success of ligand coupling requires a reliable method to access the hypervalent pentacoordinated phosphorus. In the case of phosphine oxides, addition of organometallic reagents (e.g., alkoxides, Grignard reagents, and organolithium reagents) can provide ready access to hypervalent pentacoordinated phosphoranes.\(^\text{96,97}\) For example, Newkome and Hager reported that heating phenyl(2-pyridyl)phosphine oxides at 100 °C in the presence of EtONa afforded 2,2′-bipyridines in 50–60% yield.\(^\text{98}\) Phosphonium salts containing two 2-pyridyl groups display similar behavior. Oae and Uchida demonstrated that treatment of benzyltri(2-pyridyl)phosphonium bromide in water or alcohol in the presence or absence of acid could result in ligand coupling.\(^\text{102b−e}\) They also found that when tri(2-pyridyl)phosphine oxide was treated with nucleophilic organometallic reagents, both the coupling products of the two pyridine ligands and the coupling products of the pyridine ligand and the incoming nucleophile were formed.\(^\text{102b}\) When using benzyl magnesium chloride as the nucleophile, 2-benzylpyridine was formed as the major product.\(^\text{102b}\) Despite these advances, ligand coupling of heterocycles at the phosphorus center has found limited use because no general and mild conditions have been established to access related phosphonium salts bearing heteroaryls.

In the pioneering study reported by the Anders group, pyridine was transformed into a pyridylphosphonium salt in a site-selective and efficient manner through sequential addition of TfO, PPh\(_3\), and Et\(_3\)N.\(^\text{104}\) In one of the limited examples of the application of this finding, they showed that addition of NaN\(_3\) as the nucleophile to the pyridylphosphonium salt resulted in coupling of the azido and pyridine groups, followed by a Staudinger reaction to give the iminophosphorane.\(^\text{104a}\) The potential of this methodology in ligand coupling remained unexplored until recently, when McNally and co-workers investigated the site-selective functionalization of pyridines (Scheme 42).\(^\text{99a}\) They found that, in most cases, the original organic base (Et\(_3\)N) gave good results when forming phosphonium salts, but DBU was superior in some situations. The reaction temperature was also a critical factor of this protocol, with some substrates requiring low temperatures and some others needing higher temperatures. The purification of the phosphonium salts was performed through simple precipitation, without chromatography. The transformation displayed good regioselectivity, with the 4-substituted product being the major product. When the 4-position was blocked, 2-substituted products formed. Various functionalities were tolerated; specifically, pyridines with halogen substituents were introduced smoothly into the phosphonium salt. Addition of sodium alkoxide to the phosphonium salt produced the

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**Scheme 41. Transition-Metal-Catalyzed Coupling and Phosphorus(V) Ligand Coupling**

![Scheme 41. Transition-Metal-Catalyzed Coupling and Phosphorus(V) Ligand Coupling](https://dx.doi.org/10.1021/acscentsci.0c01493)
alkoxylated pyridine. Based on previous reports,96–98 McNally and co-workers proposed the formation of the pentavalent alkoxyporphorane intermediate 6–5, followed by contractive ligand coupling to construct the C–O bond. They could not, however, rule out the nucleophilic aromatic substitution (S_NAr) pathway. The reactions displayed good functional group tolerance and site-selectivity. For example, both nicotine and loratadine were transformed into phosphonium salts and reacted with sodium alkoxide to afford the alkoxylated products in good yields. Furthermore, sodium thiolate, sodium azide, and 2-benzothiazol-2-yl-triphenylphosphonium triates previously, they did not fully study the scope and application of these salts.104b

In 2018, McNally and co-workers used phosphonium salts as a platform for the synthesis of heterobiaryls through contractive C–C coupling via P(V) intermediates (Scheme 43).99c Facing the challenge of preparing di(azaaryl)phosphonium salts under mild conditions, they developed a unique route, without using heteroaryl halides or a lithium reagent. The key to the introduction of two heterocycles to the phosphorus center was the use of the fragmentable phosphine 6–12. Numerous examples of the reaction proceeded with synthetically useful overall yields (17–34% over three steps). In addition, the synthesis of complex heterobiaryls is possible, including the use of bioactive molecules as coupling partners. The McNally group has also provided further advances in phosphorus(V) ligand coupling.99d–g

Although the phenomenon of phosphorus(V) ligand coupling was observed as early as 1948,102a its potential in heterocycle functionalization was unveiled only recently. Contractive phosphorus(V) coupling is a coupling method for heterocycle functionalization that is complementary to transition-metal-mediated reactions in terms of selectivity and functional group tolerance. In transition-metal-catalyzed cross-coupling, redox cycling between the n and n + 2 oxidation states, through oxidative addition and reductive elimination, is crucial for catalysis. The phosphorus contractive coupling resembles reductive elimination, albeit in a noniterative manner. In the three-stage coupling sequence designed by McNally and co-workers, they realized a combination of stoichiometric oxidative addition and reductive elimination. In light of recent progress in the field of main group element redox catalysis, one cannot help but look forward to inventions that will enable P(III)/P(V) redox catalysis for phosphorus ligand coupling.

7. CONCLUSION AND OUTLOOK
Phosphorus-based organocatalysis is a vast field. This brief Overview of various topics in phosphorus-based catalysis highlights key recent advances in the field, with emphasis on four topics: nucleophilic phosphine catalysis, phosphine oxide catalysis, phosphorus-mediated SET processes, and phosphorus(V) ligand coupling. Below, we provide a summary and outlook for each topic.

(1) In this well-established but still active area, the present focus of nucleophilic phosphine catalysis lies in the development of new patterns of annulations, along with asymmetric variants of known reactions. Unforeseen reactions continue to be discovered when employing highly engineered substrates. The application of nucleophilic catalysis in the synthesis of functional molecules remains promising. Moreover, nucleophilic addition of phosphines to electron-deficient C–C multiple bonds promotes the in situ formation of zwitterionic adducts that can act as efficient base catalysts for various asymmetric transformations. With continued progress, we expect to see the different reactivity patterns of
nucleophilic phosphine catalysis merging with transition metal catalysis and photoredox catalysis.

(2) Reactions driven by phosphine oxide formation remain highly popular among synthetic chemists. Catalytic, economical, and environmentally benign versions of these reactions will undoubtedly have a huge impact on synthesis in both academic and industrial settings. Recent advances have unveiled insights into how to improve the in situ activation of strong P(V)=O bonds. Multiple studies have revealed that a strained phosphacycle scaffold can facilitate the in situ reduction of phosphine oxides during redox-driven catalysis. Denton and co-workers found that a carefully designed phosphine oxide catalyst could realize additive-free phosphine oxide catalysis in a redox-neutral setting. Collectively, these findings signify that the design of phosphine (oxide)s with appropriate skeletons will be pivotal in moving the field forward. As demonstrated by Sevov and co-workers, new technologies, including electrochemistry and photoredox chemistry, can also add impetus to the in situ recycling of phosphine oxides. Alongside these advances, asymmetric phosphine oxide catalysis is also appealing for preparing highly valuable chiral molecules. Ultimately, we wish to see catalytic versions of traditional phosphine oxide-generating reactions applied in both academia and industry.

(3) At present, organic phosphorus compound-mediated SET processes remain undeveloped. Although limited in scope, this area is complementary to nucleophilic phosphine catalysis in terms of its mechanisms and functional group tolerance. While nucleophilic phosphine catalysis has been limited to electron-deficient multiple bonds, phosphorus radical catalysis can activate inert alkenes and C–H bonds. The design of phosphines or their complexes with structural frameworks of suitable electronic potential may lead to the open-shell species having longer half-lives, allowing rapid advances in this area. Several recent accounts have revealed that the EDA complexes of phosphines with RLi, NaI, or Cs2CO3 have a good SET ability, especially in the presence of visible light, potentially inspiring more discoveries in this field. Phosphorus-based reagents also facilitate the generation of active radical intermediates through deoxygenation or desulfurization, especially under photoredox catalysis. For the latter, challenges remain in the requirement for stoichiometric phosphorus reagent and the generation of waste. These issues may be addressed through recycling of the phosphine oxide, as we have discussed in Section 3. With recent developments in photochemistry and electrochemistry, we envision phosphorus compound-mediated SET processes to be a promising area.

(4) Phosphorus ligand coupling displays the versatility of phosphorus compounds, and it can serve as a complementary method for transition-metal-mediated coupling for heterocyclic functionalization. Although such reactions remain stoichiometric at present, their reductive elimination steps emulate those of transition-metal-promoted reactions. Together with other main group elements (e.g., S, I, Bi), the realization of catalytic P(V) ligand coupling may be on the horizon.

(5) Finally, several additional topics—phosphonium salt PTCs, iminophosphorane super base catalysis, asymmetric phosphine oxide Lewis base catalysis, FLP catalysis, and chiral phosphoric acid catalysis—are also important aspects of phosphorus-based organocatalysis. With the rich chemistry of organophosphorus compounds, this area continues to progress with the development of novel transformations. Phosphorus-based catalysis is at the threshold of a bright future. With continued invention, we anticipate phosphorus-based catalysis to have many more applications in the chemical and pharmaceutical industries.

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The authors declare no competing financial interest.

Acknowledgments

Financial support by the NIH (R01GM071779) is acknowledged.

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