Decarboxylative cross-coupling reactions for P(O)–C bond formation

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Phosphorus-containing compounds are one of the most important classes of organic compounds, which have wide applications in organic chemistry, medicinal chemistry, agricultural chemistry, and materials chemistry. In particular, organophosphorus compounds bearing a P(O)–C bond have attracted significant attention in recent decades due to their widespread biological and pharmacological activities. In this review, we will highlight the most important developments in the construction of P(O)–C bonds through decarboxylative C–P cross-coupling reactions. The literature has been surveyed from 2011 to May 2018.

1. Introduction

Organophosphorus compounds are among the most important classes of organic compounds, which are diversely found in natural products,1 pharmaceutical drugs,2 insecticides,3 and ligands.4 In particular, P(O)–C bond-containing compounds have attracted significant attention of medicinal chemists and pharmacologists in recent years owing to their broad spectrum of pharmacological profiles.5 Several currently marketed drugs contain this important structural unit, which is used for the treatment of various kinds of diseases. For example (Fig. 1), fosinopril 1 with the brand name Monopril is a phosphinate-containing angiotensin-converting enzyme inhibitor marketed worldwide for the treatment of hypertension (high blood pressure) and some types of chronic heart failures.6 Adefovir 2, with trade names Preveon and Hepsera, is a promising antiviral drug that is used to treat chronic hepatitis B; the drug works by blocking reverse transcriptase, an enzyme crucial for the hepatitis B virus to reproduce in the body.7 Alendronic acid 3 (Fosamax) is a bisphosphonate drug used to treat osteoporosis and several other bone diseases.8 As a consequence, the construction of P(O)–C bonds is an essential issue and has gained much attention from the research community interested in chemical synthesis.

The most general and straightforward methods for the construction of P(O)–C bonds involve either C–P cross-coupling of organic halides or triflates with P(O)H compounds (Fig. 2, route a)9 or direct dehydrogenative C–P cross-coupling between

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R–H compounds and P(O)H compounds (Fig. 2, route b). However, each of these methods suffers from various inherent drawbacks, and the most important of these drawbacks are as follows: (i) most of the organic halides are environmentally toxic and liberate hazardous halogen by-products; (ii) triflates are unstable, expensive, and liberate potentially genotoxic trillic acid; and (iii) the selective activation of a specific C–H bond poses a particular challenge. An alternative and efficient protocol for the formation of P(O)–C bonds involves decarboxylative C–P cross-coupling between carboxylic acids and P(O)H compounds (Fig. 2, route c). To the best of our knowledge, a comprehensive review on decarboxylative C–P cross-coupling reactions has not been reported in the literature so far. In continuation of our recently published reviews on metal-catalyzed cross-coupling reactions and the green synthesis of inorganic polymers, synthesis of inorganic complexes and bioinorganic chemistry.

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In this review, we will highlight the most important developments of decarboxylative C–P cross-coupling reactions from 2011 to May 2018. It is worth mentioning that a special emphasis is placed on the mechanistic aspects of the reactions.

2. Coupling of alkenyl acids with P(O)H compounds

The first synthesis of alkenylphosphorus compounds through transition-metal-catalyzed decarboxylative coupling reactions of alkenyl acids with P(O)H compounds was reported by Hu and co-workers in 2011; they treated various cinnamic acids with hydrogen phosphoryl compounds in the presence of 10 mol% of low-cost, commercially-available Cu$_2$O as the catalyst, 10 mol% of 1,10-phenanthroline as the ligand and 3.0 equiv. of Ag$_2$O as the additive in NMP. The reactions were carried out at 120 °C for 6 h and provided the expected alkenylphosphorus compounds in moderate to excellent yields (Scheme 1). It should be mentioned that all three kinds of P(O)–H compounds (H-phosphinates, H-phosphonates, and secondary phosphine oxides) were applicable to this coupling reaction. The decreasing order of the reactivities of hydrogen phosphoryl compounds with cinnamic acids under these reaction conditions was secondary phosphine oxides > H-phosphinates > H-phosphonates. Interestingly, the electronic character of the substituents in the phenyl ring periphery of cinnamic acids had negligible effect on the rate of the reaction. However, steric hindrance led to lower yields. A plausible mechanism that explains this transformation is depicted in Scheme 2. Initially, P(O)–H compound reacted with the copper/phen catalyst to form the intermediate A. Meanwhile, the reaction of alkenyl acids with Ag$_2$O produced intermediate B through a decarboxylation process. This intermediate B upon transmetalation...
A formed the organocopper intermediate C. Finally, reductive elimination of intermediate C afforded the observed alkenylphosphorus compounds 6 along with the initial copper species.

Next, Gao’s group demonstrated for the first time the usefulness of nickel catalysts for decarboxylative coupling of alkenyl acids with P(O)-H compounds. Thus, in the presence of Ni(dppf)Cl$_2$ as the catalyst and Ag$_2$O as the additive in DMSO at 120 °C, coupling of cinnamic acids 7 with hydrogen phosphoryl compounds 8 afforded the corresponding (E)-1-alkenylphosphorus compounds 9 in moderate to high yields (Scheme 3). A variety of sensitive functional groups including NO$_2$, NMe$_2$, CO$_2$Me, OMe, CF$_3$, F, and Cl were tolerated by the reaction conditions employed. This made further derivatization of the products possible. It is noted that other nickel catalysts such as Ni(OAc)$_2$, Ni(acac)$_2$, NiBr$_2$, and NiCl$_2$ also promoted this coupling reaction; however, lower yields were observed. Based on density functional theory (DFT) calculations and experimental data, the authors found that in this coupling reaction, the phosphine ligand exhibited better catalytic performance than the nitrogen ligand in the reductive elimination step owing to stronger nucleophilicity and larger size.

In a related investigation, Tang and Wang along with their co-workers found that the treatment of alkenyl acids 10 with P(O)-H compounds 11 (phosphine oxides and H-phosphonates) in the presence of Cu(OAc)$_2$·H$_2$O/K$_2$S$_2$O$_8$/MeCN combination as a catalytic system afforded the corresponding (E)-1-alkenylphosphorus compounds 12 in moderate to high yields with excellent stereoselectivities (Scheme 4). Some important points regarding the reactions are listed below: (i) aliphatic alkenyl acids failed to enter the reaction; (ii) depending on the electronic effects of the substituents on the aryl ring of cinnamic acids, substrates with electron-donating groups gave higher yields than those with electron-withdrawing groups; and (iii) phosphine oxides gave lower yields of desired products compared with H-phosphonates. It is noteworthy that this catalytic system was also successfully applied in the cross-coupling of various P(O)-H compounds with styrenes and β-nitrostyrene. A plausible mechanism for this transformation starts with the formation of the phosphonyl radical A by oxidation of the starting phosphonate 11 with in situ generated Cu$^{II}$ species [Cu(CH$_3$CN)$_n$]$^{2+}$; its nucleophilic attack on the alkenyl acid 10 led to the formation of radical B, which could be converted into a more stable conformation C. Next, the oxidation of this intermediate C by the Cu$^{II}$ species gave the carbocation intermediate D, which finally afforded the final product by a ligand-exchange-decarboxylation sequence (path A). In another possibility (path B), intermediate C gave complex D through ligand exchange, followed by decarboxylation to afford the corresponding (E)-1-alkenylphosphorus product (Scheme 5).

Very recently, Liu and co-workers described efficient transition-metal-free phosphorylation of alkenyl acid derivatives with P(O)H compounds via radical-promoted decarboxylation under mild conditions. They carefully tested several oxidants, bases, additives and solvents, and the system of TBPB/K$_2$HPO$_4$/KI/DMSO was found to be superior. Under optimized conditions, various aromatic and heteroaromatic alkenyl acids 13, including those exhibiting substitution on the C=C bond,
and symmetrical secondary phosphine oxides 14 reacted to give good yields of the corresponding products 15 (Scheme 6). In addition to mild reaction conditions and excellent functional group tolerance (F, Cl, Br, CN, CO$_2$H, CO$_2$Me, OCOMe, OMe, and SMe), the major advantage of this method is the formation of (E)-1-alkenylphosphorus compounds 15 on a gram-scale. The authors also showed the application of this procedure for the high yielding syntheses of the estrone derivative 17, a steroidal progestin used in clinic, with estrogenic activity (Scheme 7). They assumed that the mechanism of this coupling reaction involved the initial formation of tert-butoxyl and benzoyloxyl radicals via decomposition of TBPB, which was promoted with the assistance of iodide anions. Next, the abstraction of a hydrogen atom from H(O)PR$_1$R$_2$ 14 by a tert-butoxyl or benzoyloxyl radical afforded the phosphorus radical A, which underwent radical addition to the cinnamic acid anion B (generated from the reaction of the starting acid 13 with a base) to provide the radical intermediate C. Finally, decarboxylation of intermediate C followed by single-electron oxidation of the radical anion afforded the expected products 15 (Scheme 8).

3. Coupling of alkynyl acids with P(O)H compounds

In 2011, Liang and Yang along with their co-workers reported the first example of decarboxylative C–P cross-coupling of alkynyl acids with P(O)H compounds.$^{15}$ They showed that electron-rich alkynyl acids 18 underwent rapid C–P cross-coupling with various hydrogen phosphoryl compounds 19 in the presence of Cu$_2$O/Pd(acac)$_2$/1,10-phenanthroline/PPh$_3$/AgOAc combination as a catalytic system in NMP at 120 °C. The corresponding alkynylphosphorus compounds 20 were obtained in moderate to high yields (Scheme 9). The reaction is noteworthy since both alkyl and aryl alkynyl carboxylic acids and all three kinds of P(O)–H compounds were tolerated. It was noted that the presence of both copper and palladium catalysts is essential in this reaction; unsatisfactory yields of the expected products were obtained in the absence of any of these catalysts. The proposed mechanism by the authors is illustrated in Scheme 10. First, a Cu$^{II}$–phosphine-complex A was formed via the reaction of P(O)H compound 19 with the copper/phen catalyst. In parallel, decarboxylation of alkynyl acid 18 by silver gave alkynyl–silver species B. The in situ generated Pd$^{II}$ complex (by oxidation of Pd$^{0}$ with silver) transmetallated with the alkynyl–silver species B to generate the alkynyl–palladium intermediate C. The transmetallation of this highly active intermediate C with the Cu$^{II}$-complex A then gave the intermediate D. Finally, reductive elimination of intermediate D resulted in the formation of the expected alkynylphosphorus compounds 20. Later, Gao’s group showed that alkynyl phosphonate derivatives were also formed from the corresponding alkynyl acids and diisopropyl phosphonate by a simple process employing Ni(NO$_3$)$_2$·6H$_2$O as the catalyst and Ag$_2$CO$_3$ as the oxidant in DMSO at 100 °C. However, the yields of the desired products were low (36–45% for three examples).$^{16}$

Scheme 5  Mechanistic proposal for the reactions in Scheme 4.
In 2014, Li and co-workers reported that the combination of \( \text{Cu(OAc)} \cdot \text{H}_2\text{O}/1,10\text{-phenanthroline} \) could efficiently catalyze the decarboxylative cross-coupling of alkynyl acids 21 with dialkyl H-phosphonates 22 under relatively mild conditions in the most significant green solvent water (Scheme 11). Various aryl alkynyl acids and several dialkyl H-phosphonates were well tolerated in this catalytic system with moderate to high yields (40–88%). However, alkyl alkynyl acids failed to participate in this reaction. Interestingly, other copper salts [e.g., \( \text{CuCl}_2 \), \( \text{Cu(OTf)}_2 \), and \( \text{CuSO}_4 \cdot 5\text{H}_2\text{O} \)] were also found to promote the reaction albeit with reduced efficiencies. According to the mechanism proposed by the authors (Scheme 12), this reaction consists of the following key steps: (i) initial formation of active copper(II) intermediate \( A \) via coordination of 1,10-phenanthroline to \( \text{Cu(OAc)}_2 \); (ii) ligand exchange between intermediate \( A \) and alkynyl acid 21 to afford the copper(II) intermediate \( B \); (iii) decarboxylative reaction of intermediate \( B \) to give the copper(II) intermediate \( C \); (iv) reaction of the intermediate \( C \) with a phosphonate anion (generated from H-phosphonate 22 and \( \text{K}_3\text{PO}_4 \)) to form the copper(II) intermediate \( D \) and (v) reductive elimination of intermediate \( D \) to give the desired products 23.

At almost the same time, Hu, Gao, and Zhao demonstrated elegant and innovative copper-catalyzed decarboxylative cross-coupling of alkynyl acids with hydrogen phosphoryl compounds for the formation of alkenylphosphorus compounds. Thus, in the presence of a catalytic amount of \( \text{CuCl} \) (10 mol%) in DMF under nitrogen atmosphere, cross-coupling of aryl alkynyl acids 24 with H-phosphine oxides 25 afforded the corresponding \( (E) \)-alkenylphosphine oxide derivatives 26 in moderate to almost quantitative yields (Scheme 13).

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**Scheme 6** Transition-metal-free phosphorylation of alkenyl acid derivatives with P(O)H compounds reported by Liu.

**Scheme 7** Synthesis of estrone derivative 17 through a decarboxylative C–P cross-coupling reaction.
The reaction could also be conducted successfully on a gram scale. However, the reaction did not give good yields with alkyl and heteroaryl alkynyl acids. The proposed mechanism of the reaction by the authors is depicted in Scheme 14. It consists of the following key steps: (i) initial formation of the alkynyl copper intermediate \( A \) via decarboxylation of alkynyl acid \( 24 \) with \( \text{CuCl} \) and the generation of one molecule of \( \text{CO}_2 \) and one molecule of \( \text{HCl} \) at the same time; (ii) coordination of \( \text{P(O)H} \) compound \( 25 \) (in the form of trivalent phosphine oxide \( 25^0 \)) to the intermediate \( A \) to form the intermediate \( B \); (iii) migration of a proton from the oxygen atom to the \( \text{C sp} \) atom in the intermediate \( B \) to give a three-membered-ring transition state \( C \); (iv) conversion of intermediate \( C \) (in the form of four-centered transition state \( C^0 \)) to a thermodynamically more stable

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**Scheme 8**  Mechanism that accounts for the formation of (E)-1-alkenylphosphorus compounds 15.

**Scheme 9**  Cu/Pd-catalyzed decarboxylative C–P cross-coupling of alkynyl acids 18 with P(O)H compounds 19.

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**Scheme 10**  Mechanism proposed to explain the synthesis of alkynylphosphorus compounds 20.
Scheme 11  Decarboxylative cross-coupling of alkynyl acids 21 with dialkyl H-phosphonates 22 mediated by Cu(OAc)\(_2\)-H\(_2\)O.

Scheme 12  Proposed mechanistic pathways for the formation of alkynylphosphorus compounds 23.

Scheme 13  Cu-catalyzed decarboxylative cross-coupling of alkynyl acids 24 with H-phosphine oxides 25 for the stereoselective synthesis of (E)-alkenylphosphine oxides 26.
alkenyl copper intermediate D with the E configuration; and (v) protonolysis of intermediate D with alkynyl acid 24 or HCl to generate the desired product 26.

Recently, the same research team reported an interesting silver-mediated cascade decarboxylative C–P cross-coupling and annulation reaction of alkynyl acids 27 with diarylphosphine oxides 28 to form structurally sophisticated 2-phosphinobenzo[b]phosphole oxide frameworks 29 (Scheme 15).21 The reaction was performed under an argon atmosphere using AgOAc as the oxidant in DMF; it tolerated a number of functional groups (e.g., OMe, OCF₃, F, Cl, Br, CN, CHO, COMe, CO₂H, and CO₂Et) and generally provided highly substituted 2-phosphinobenzo[b]phosphole oxides 29 in moderate to high yields. According to mechanistic studies, this reaction proceeded through the formation of an arylethynyl silver intermediate A from the decarboxylative reaction of alkynyl acid 27 and AgOAc under heat, following the coordination of this intermediate A with 28 (in the form of the trivalent phosphine oxide 28') to give the intermediate B, which then underwent silver-mediated oxidative cross-coupling to produce the critical intermediate C via two single-electron oxidations. Subsequently, the addition of the diphenylphosphonyl radical (generated from 28) to the intermediate C afforded the alkenyl radical intermediate D, which underwent intramolecular addition to the phenyl ring at the ortho position of the phosphorus atom to afford the intermediate E. Then, the oxidation of this intermediate by AgOAc along with the removal of a proton led to the final product 29. In another possibility, the alkenyl radical D attacked the phenyl ring at the carbon position directly attached to the phosphorus atom to form the intermediate F, followed by C–P bond cleavage to give the phosphorus radical G. Next, the addition of G to the phenyl ring generated the intermediate E, which after oxidation, afforded 2-phosphinobenzo[b]phosphole oxide frameworks 29 (Scheme 16).

![Scheme 14](image_url)  Mechanistic explanation of the synthesis of (E)-alkenylphosphine oxide derivatives 26.

![Scheme 15](image_url)  Ag-mediated cascade reaction of arylpropioelic acids 27 with diarylphosphine oxides 28 developed by Gao.
4. Coupling of benzoic acids with P(O)H compounds

Decarboxylative C–P cross-coupling of benzoic acids with P(O)H compounds has been scarcely studied; in fact, only one example of such a reaction has been reported so far. In this report, Xiao and co-workers showed that the treatment of ortho-nitrobenzoic acids 30 with H-phosphonates 31 in the presence of a bimetallic palladium/silver catalytic system [Pd(OAc)\(_2\)/Ag\(_2\)CO\(_3\)] in DMF resulted in the formation of functionalized aryl phosphonates 32 in low to good yields (Scheme 17). However, meta- and para-substituted benzoic acids were inert to this coupling reaction. The results showed that the reaction was not very efficient and general since it had a number of limitations and exceptions. Thus, the development of a novel and efficient catalytic system that improves the efficiency of this interesting coupling reaction is highly desirable. According to the mechanism proposed by the authors (Scheme 18), this reaction starts with the reaction of silver salt with benzoic acid 30, leading to the formation of the silver benzoate species A. Subsequently, decarboxylation of this intermediate yields an aryl-Ag species B; its transmetalation with Pd\(^{2+}\)X\(_2\) forms an aryl-Pd intermediate C along with regeneration of the Ag\(^+\) salt. Next, ligand exchange of intermediate C with H-phosphonate 31 generates the Pd\(^{2+}\) complex D. Finally, reductive elimination of D affords the observed phosphonation product 32 and Pd\(^0\). Pd\(^0\) is then oxidized to Pd\(^{2+}\) to close the catalytic cycle.

5. Coupling of amino acids with P(O)H compounds

After the pioneering study by Hu and co-workers on decarboxylative coupling of N-benzyl-protected proline with diphenylphosphine oxide employing CuCl as the catalyst, the first
general report on the synthesis of biologically important tertiary amino phosphorus compounds 36 through three-component decarboxylative coupling of natural \( \alpha \)-amino acids 33 with aromatic aldehydes 34 and \( P(\text{O})\text{H} \) compounds 35 was published in 2011 by Yang et al.\(^{23} \) They carefully studied the effects of reaction variables such as catalyst, additive, solvent, and temperature and found that performing the reaction in toluene at 130 °C using CuI/DIPEA (N,N-diisopropylethylamine) combination as the catalytic system was the optimum reaction condition. Various \( \alpha \)-amino acids, H-phosphonates, secondary phosphine oxides, and both electron-poor and electron-rich aldehydes reacted well under the optimized reaction conditions to produce the corresponding tertiary amino phosphorus compounds 36 in moderate to quantitative yields (Scheme 19). The proposed mechanism by the authors is depicted in Scheme 20. First, an oxazolidin-5-one intermediate A was formed via condensation of an \( \alpha \)-amino acid 33 with an aldehyde 34; this intermediate underwent decarboxylation to generate the azomethine ylide B. Finally, the addition of \( P(\text{O})\text{H} \) compound 35 (in the form of the trivalent phosphine oxide 35' ) to this dipole B gave the expected product 36.

In a closely related investigation, Kaboudin’s group showed that many \( N \)-benzyl-2-(di(ethyl)phosphoryl)pyrrolidine derivatives 40 were successfully formed from the reaction of proline 37.

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**Scheme 18** Plausible mechanism for the formation of aryl phosphonates 32.

**Scheme 19** Cu-catalyzed three-component decarboxylative coupling of natural \( \alpha \)-amino acids 33 with aromatic aldehydes 34 and \( P(\text{O})\text{H} \) compounds 35.

**Scheme 20** Mechanism that accounts for the formation of tertiary amino phosphorus compounds 36.
with aldehydes 38 and diethyl phosphate 39 in the absence of any transition metal catalyst by simple heating in toluene (Scheme 21). Various aromatic, heteroaromatic, and aliphatic aldehydes were used to establish the general applicability of the method. The results demonstrated that the aromatic aldehydes afforded better yields compared to the aliphatic aldehydes. However, *ortho*-substituted benzaldehydes failed to enter this reaction.

6. Oxidative decarboxylative coupling of carboxylic acids with P(O)H compounds

6.1. Alkynyl acids

The possibility of metal-catalyzed oxidative decarboxylative coupling of alkynyl acids with P(O)H compounds was first demonstrated by Song and his research team in 2015, and they synthesized a series of β-ketophosphonates 43 from the reaction of propiolate acids 41 with H-phosphonates 42 in the presence of only 5 mol% of CuOTf as the catalyst, 10 mol% of FeCl3 as the co-catalyst, and 1.0 equiv. of Et3N as the base in DMSO under oxygen atmosphere (balloon). This Cu/Fe-cocatalyzed reaction tolerated both aryl as well as heteroaryl alkynyl acids and various symmetrical H-phosphonates and provided β-ketophosphonates 43 in fair to good yields (Scheme 22). However, aliphatic alkynyl acids did not function well under these reaction conditions. It should be mentioned that this catalytic system was also successfully applied in the oxyphosphorylation of terminal alkynes with P(O)H compounds. The mechanistic pathway of this oxidative decarboxylative coupling reaction is shown in Scheme 23, and it involved the initial formation of the dialkyl phosphonate cation radical A through the transfer of a single electron from iron(II) species to H-phosphonate 42 in the presence of molecular oxygen. Next, protonation of this cation radical A in the presence of a base gave the dialkyl phosphonyl radical B, which then attacked the triple bond of the alkynyl acid 41 to produce the radical intermediate C. Subsequently, the reaction of this radical with Cu(II)−(OOH) species (generated from Cu(I) under dioxygen atmosphere) gave the hydroperoxide species D. Finally, the removal of oxygen from D afforded the observed β-ketophosphonates 43 through the intermediate E.

Concurrently, Zhang and co-workers presented efficient copper-catalyzed oxidative decarboxylative coupling of alkynyl acids 44 with H-phosphine oxides 45 to form β-ketophosphine oxide derivatives 46. The results of this investigation revealed that the optimum condition for this reaction was the combination of CuSO4·5H2O (10 mol%), TBHP (2 equiv.), and NH3·H2O (25%) as the catalytic system using acetonitrile as the solvent at 60 °C. Under optimized conditions, the reaction tolerated various symmetrical and unsymmetrical H-phosphine oxides and a range of aromatic and heteroaromatic alkynyl acids and gave the expected products in moderate to excellent yields (Scheme 24). However, aliphatic alkynyl acids failed to enter the reaction. Mechanistically, this reaction was proposed to be initiated by the formation of tert-butoxy and hydroxyl radicals via decomposition of TBPB in the presence of Cu(II). Then, the abstraction of a hydrogen atom from H-phosphine oxide 45 by a tert-butoxy radical afforded the phosphorus radical A. Meanwhile, the reaction of alkynyl acid 44 with Cu(II) generated a salt of cupric carboxylate B. Subsequently, the addition of the phosphorus radical A to the α-position of the triple bond of the cupric carboxylate B gave the radical intermediate C, which was ultimately trapped by a hydroxyl radical to provide the intermediate D. The formation of the alkynyl copper...
Scheme 23  Mechanistic proposal for the reaction in Scheme 22.

Scheme 24  CuSO$_4$·5H$_2$O-catalyzed oxidative decarboxylative coupling of alkynyl acids 44 with H-phosphine oxides 45 into $\beta$-ketophosphine oxides 46.

Scheme 25  Plausible mechanism for the formation of $\beta$-ketophosphine oxides 46.
intermediate E occurred next, followed by its protonolysis to give the intermediate F. Finally, isomerization of F led to the formation of the desired product 46 (Scheme 25).

The synthesis of β-oxophosphine oxides in moderate to good yields (40–73%) was also reported by Zeng and co-workers through the reaction of corresponding alkynyl acids with H-phosphine oxides using Ag$_2$CO$_3$ as an efficient catalyst in ethanol at room temperature and air conditions.$^{27}$ Other silver salts such as Ag$_2$O, AgOAc, and AgOTf were also found to promote this reaction, albeit with lower yields. Under optimized conditions, both dialkyl and diaryl phosphine oxides were well tolerated. The reaction could also be conducted successfully on a gram scale.

### 6.2. Alkenyl acids

In 2015, an innovative and efficient synthetic protocol was reported by Zhou and co-workers for the oxidative decarboxylative coupling of various alkenyl acids 47 with H-phosphonates 48 to afford a diverse range of β-ketophosphonates 49 using CuOTf/FeCl$_3$/Et$_3$N combination as the catalytic system in DMSO (Scheme 26).$^{28}$ The reactions were performed under air and tolerated a series of sensitive functional groups including fluoro, chloro, bromo, methoxy, and cyano functionalities; the target products were obtained in moderate to good yields (49–79%). The authors showed that both H-phosphinate oxides and phosphine oxides were also compatible in this transformation.

### Scheme 26

**Cu/Fe-catalyzed oxidative decarboxylative coupling of various alkenyl acids 47 with H-phosphonates 48.**

[[Chemical structure]]

### Scheme 27

Chen’s synthesis of β-keto phosphorus compounds 52.

### Scheme 28

Plausible mechanism for the formation of β-keto phosphorus compounds 52.
and gave desired products in moderate yields. Moreover, they successfully extended this chemistry to the synthesis of a variety of functionalized β-ketophosphonates via a domino Knoevenagel-decarboxylation–oxygenphosphorylation sequence from aromatic aldehydes and H-phosphonates.

The synthesis of β-keto phosphorus compounds 52 in good to excellent yields (up to 92%) was also reported by Chen’s research team through the oxidative decarboxylative coupling reaction of cinnamic acids 50 with P(O)H compounds 51 by employing CuSO4·5H2O as an efficient catalyst and air oxygen as an environmentally benign oxidant (Scheme 27). Solvent has a dramatic role in this reaction; among the common solvents (e.g., THF, DCE, DMF, DMSO, MeCN, EtOH, and 1,4-dioxane), acetonitrile was the most effective solvent. Interestingly, the other solvents were completely ineffective. The authors proposed a mechanism for this decarboxylation–oxygenphosphorylation reaction in which an acetonitrile-soluble complex [Cu(MeCN)][Cu][4] A, generated from the copper(n) ion and MeCN, underwent a reaction with air oxygen to form a copper-active-oxygen complex B, which then quickly attacked the P(O)H compound 51 to produce the hydroperoxide complex C as well as the phosphonyl radical D. Subsequently, the addition of radical D to cinnamic acid 50 afforded the radical E, which in the presence of the hydroperoxide complex C yielded a hydroperoxide intermediate F. Next, dehydration of this intermediate F gave the intermediate G. Finally, energetically favorable decarboxylation of intermediate G afforded the final products 52 (Scheme 28).

7. Summary and conclusion

Carbon–phosphorus bond construction represents a key step in the synthesis of various medicines, insecticides, ligands, and modified nucleosides and nucleotides. In particular, P(O)–C bond formation reactions have attracted much attention owing to their presence in a number of commercialized drugs such as fosinopril, adefovir, alendronic acid, cidofovir, and brincidofovir. General methods to construct the P(O)–C bond involve transition-metal-catalyzed cross-coupling of organic halides, triflates, or tosylates with P(O)H compounds or the nucleophilic substitution reaction of organometallic reagents with electrophilic P-reagents. Despite significant progress, these methods still suffer from some limitations such as the use of metal reagents, expensive and unstable triflates, environmentally toxic organic halides, and poor tolerance of functional groups. In the last seven years, the decarboxylative C–P cross-coupling reaction between carboxylic acids and P(O)H compounds has emerged as one of the most reliable and robust tools for the construction of P(O)–C bonds.

As illustrated, all three kinds of P(O)–H compounds (H-phosphinates, H-phosphonates, and secondary phosphine oxides) were applicable to this novel coupling reaction. Interestingly, most of the reactions covered in this review could be scaled up to provide multigram quantities of the desired coupling products. These results showed the potential applications of these reactions in industry. Despite all these successes, many challenges still remain to be overcome: (i) the number of reported examples in this interesting research arena is limited. There is still a further need to study the scope and limitations of this approach for the formation of P(O)–C bonds; (ii) most of the reported decarboxylative C–P cross-coupling reactions are performed at high temperatures; thus, the exploration of milder processes is highly desirable, and (iii) other reactions such as coupling of allylic and benzylic carboxylic acids with various P(O)–H compounds should be explored. We conclude this review by hoping that it will stimulate researchers to carry out further research in this domain.

Conflicts of interest

There are no conflicts to declare.

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