Light-Promoted Organic Transformations Utilizing Carbon-Based Gas Molecules as Feedstocks

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Carbon-based gas molecules are readily available feedstocks and are widely used in industry as building blocks or fuels. However, their application in the synthesis of fine chemicals has been hampered due to operational complexity, poor reaction efficiency and selectivity. Recent development of photoredox-promoted transformations using such gaseous reagents has received considerable attention from the synthetic community. In this review, efforts in developing light-promoted organic transformations using carbon-based natural gases as C1 or C2 feedstocks and to overcome the associated challenges are briefly summarized.

1. Introduction

Carbon-based gas molecules are abundant and readily available feedstocks for use in chemical synthesis. Carbon monoxide (CO) and carbon dioxide (CO2) are substantially produced from the combustion of carbon-based substrates, and ethylene, acetylene, ethane and methane are among the most common carbon-based natural gases employed in chemical industry as C1 or C2 building blocks for chemical synthesis. For example, carbon dioxide has been widely used in the synthesis of polycarbonates and cyclic carbonates and for the production of urea through the Bosch-Meiser process.[1] Due to the ease of access to these gases, synthetic strategies have been developed to synthesize value-added fine chemicals from these simple and abundant feedstock compounds.[2] In this context, the development of light-promoted transformations utilizing carbon-based natural gases has gained momentum and is briefly summarized in this review.

2. Carbon Monoxide

Carbon monoxide is an inexpensive and abundant single-carbon source that is used in various carbonylation reactions. Early 20th century studies of the Fisher Tropsch synthesis and hydroformylation reactions have led to a better understanding of carbonylation reactions.[3] These reactions are now widely used in industries to produce valuable fine chemicals. For instance, the industrial production of acetic acid is through the carbonylation of methanol employing the Monsanto or Cativa process.[4] However, precautions are necessary due to the high toxicity of CO when used as the starting material.

Carbonylation using CO as the carbonyl source provides an efficient and atom-economic way to install a variety of carbonyl-based functional groups. A useful method is the radical carbonylation. However, this method did not receive attention until the early 1990s, when the groups of Bakac, Goldman and Ryu independently reported kinetic studies of radical carbonylation.[5] Since then, a series of radical-based carbonylation transformations, including light-mediated radical carbonylation reactions, have been developed.

2.1. Light-Promoted Radical Carbonylation in the Absence of Photosensitizers

In 1939, Faltings reported that acetone was produced when ethane and CO were irradiated with UV light.[6] Following Faltings’ discovery, Ryu et al. demonstrated a light-promoted, atom-transfer-carbonylative three-component coupling reaction between α-phenylselenides, terminal alkynes, and CO to afford a variety of adducts (Scheme 1).[7] A proposed mechanism involved the selenide undergoing homolysis to generate radical species. Subsequent radical addition of alkyl radicals 3 produces an alkyl radical 4, which undergoes a second radical addition with CO to yield an acyl radical 5. A group transfer of the phenylselenenyl group leads to the formation of the acyl selenide 6. Various products (6a–6f) could be synthesized in this manner. Notably, a sophisticated reactor system is required in light-promoted carbonylation with high pressure CO (Figure 1). The reaction was conducted in a stainless steel autoclave with quartz glass windows, which enables chemical transformations with high pressure gaseous reagents under light-irradiation.

Based on the fact that the C–I bond in an alkyl iodide could be homolytically cleaved to generate an alkyl radical under light irradiation,[8] the Ryu group reported a light-mediated catalyst-free esterification of alkyl iodides with alcohols (Scheme 2A).[9] The endothermic transfer of iodine from the alkyl iodide to an acyl radical was aided by the addition of a base. Moreover, the same protocol could be...
employed for the amidation of alkyl iodides using amines as the nucleophiles (Scheme 2B).[10] Syntheses of ¹¹C radiolabeled alkyl acids and related derivatives have also been achieved utilizing this protocol.[11] The same group extended this strategy to carbonylation of aryl iodides irradiated by a 500 W xenon (Xe) lamp.[12]

### 2.2. Light-Promoted C–H Carbonylation through Hydrogen Atom Transfer

An early discovery of C–H carbonylation was made in 1991 by Crabtree et al., who observed the formation of aldehydes from alkyl radicals generated by mercury photosensitization in the presence of CO.[13] Subsequently, the Goldman group demonstrated the synthesis of aldehydes by carbonylation of cyclohexane, catalyzed by aryl ketones under UV light.[5b] Mechanistically, the aromatic ketone catalyst is photo-excited to generate an excited ketone species 16. A hydrogen atom transfer (HAT) process between cyclohexane and 16 forms a ketyl radical 17 and cyclohexyl radical 18, which then traps CO to form an acyl radical 19. A reverse HAT process between 19 and 17 produces the aldehyde product and regenerates the aromatic ketone.

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catalyst (Scheme 3). In 1995, the Hill group reported a tetrabutylammonium decatungstate (TBADT)-catalyzed alkane carbonylation under 1 atm CO, irradiated with a 550 W Hg lamp (>280 nm). A TON of 54 was achieved for this reaction using cyclohexane as the substrate.\[14\] Ryu and co-workers realized a three-component coupling reaction involving alkanes, CO at high pressure (80 atm), and electron deficient alkenes, employing TBADT as a catalyst and a xenon (Xe) lamp light source.\[15\] They envisioned that TBADT would be photo-excited to generate an excited polyoxotungstate anion \([W_{10}O_{32}]^{4–}\), which promotes the formation of a carbon-centered radical 23 through an HAT process with an alkane (Scheme 4). Subsequent radical addition to CO delivers an acyl radical 24, which is trapped by an electron-deficient alkene to selectively form a radical adduct 25. A second HAT process could transfer the hydrogen atom from the reduced tungstate anion to the radical adduct 25 to form the desired product 22 and regenerate the TBADT catalyst. In addition, Ryu et al. showed that β-carbon carbonylation of cyclopentanones and aliphatic nitriles could be achieved with a similar protocol to yield β-alkylated ketones.\[16\] Alternatively, acyl hydrazide compounds could be obtained by trapping the acyl radical using diisopropyl azodicarboxylate.\[17\]

2.3. Recent Development in Photoredox-Promoted Carbonylation

The widely utilized iridium (Ir)- and ruthenium (Ru)-based photosensitizers have showed excellent catalytic activity in carbonylation reactions. In 2015, the Xiao group developed a visible-light-promoted decarboxylative carbonylation reaction between aliphatic carboxylic acids and benzodioxolone-substituted alkynes catalyzed by Ir[dF(CF3)ppy]2(dtbbpy)PF6, generating yrones in good to excellent yields (Scheme 5).\[18\] It was proposed that the reaction is initiated by a single-electron oxidation of the carboxylic anion 26 by the excited state of the iridium(III) photocatalyst to generate an alkyl radical 29. Subsequent radical addition of 29 to CO gives the acyl radical 30, which undergoes a second radical addition to the alkyne 27 to yield

Scheme 2. Photo-induced carbonylation of alkyl iodides.

Scheme 3. Aryl ketone catalyzed carbonylation of cyclohexane under UV light.
a vinyl radical 31. An elimination of a benzodioxolone (BI) radical produces the desired ynone 28. In 2018, Bousquet et al. developed a carboxylation reaction of aryldiazonium salts employing H₂O as a nucleophile and Ru(bpy)₃Cl₂ as the photocatalyst.[19] The Polyzos group reported a visible-light-mediated Ir(ppy)₃(dbtbpy)PF₆-catalyzed annulative alkoxy-carbonylation of alkyl-tethered aryldiazonium salts to yield 2,3-dihydrobenzofurans products in a continuous flow system.[20] In 2019, the Xiao group reported an efficient copper-catalyzed radical aminocarbonylation of cycloketone oxime esters in the presence of CO and amines under visible-light irradiation.[21] Mechanistically, the authors proposed a visible light-driven Cuᴵ/Cuᴵᴵ/Cuᴵᴵᴵ-based catalytic cycle (Scheme 6). The in situ generated copper(I) complex 36 reacts with phenylamine to form a new copper(I) complex 37, which after light irradiation forms the photoexcited copper complex 38. A subsequent SET-mediated reduction of 39 by 38 affords an iminyl radical 40 and copper(II) complex 41 (path a). Alternatively, 40 and 41 could be generated by reduction of 39 by 37 (path b). The iminyl radical 40 would then undergo a selective β-C-C bond scission to give a cyanoalkyl radical 42, which couples with 41 to form a high-valent CuᴵᴵІ complex 43. Subsequent coordination and insertion of CO to 43 would lead to the formation of acylcopper intermediate 44 or 45. Reductive elimination of the acylcopper complex produces cyanoalkylated amide 46 and regenerates the Cuᴵ 36.

Organic dyes have been widely used as photoredox catalysts to replace the costly Ir and Ru noble metals, and have exhibited excellent photo-catalytic activities.[22] In this context, the Xiao group reported a visible-light-promoted alkoxy-carbonylation reaction of aryldiazonium salts under 80 atm of CO, using fluorescein as the photocatalyst (Scheme 7A).[23] A wide range of substituted aryldiazonium salts and alcohols are well tolerated in this reaction. The photocatalytic carboxylation protocol can be used with chiral alcohols to obtain enantiomerically pure products (49c, 49f).

Based on control experiments, a plausible mechanism for this reaction was proposed as shown in Scheme 7B. The catalytic cycle begins with a single electron reduction of aryldiazonium salt by the light-excited photocatalyst to
generate a phenyl radical $50$ and a fluorescein radical cation. The phenyl radical $50$ subsequently reacts with CO to yield an acyl radical $51$, which is oxidized by the fluorescein radical cation to give benzyldieneoxonium ion $52$. Finally, a nucleophilic attack by an alcohol to the benzyldieneoxonium ion $52$ yields the desired ester product $49\text{a}$.

In 2015, Wangelin and co-workers demonstrated that the same transformation could also proceed employing eosin Y sodium salts under green light irradiation.$[24]$ Later, Gu and co-workers reported an eosin Y-catalyzed aryl ketone and indol-3-yl aryl ketone synthesis using aryl diazonium salts and heteroarenes.$[25]$ In 2019, Fensterbank and co-workers demonstrated a three-component carbonylation reaction between organosilicates, CO and alkenes (Scheme 8)$.[26]$ A wide range of alkyl silicates and electron-withdrawing substituents on the alkenes are tolerated in the reaction, yielding products in moderate to good yields. Interestingly, a non-carbonylated product $55\text{t}$ was obtained when the reaction was conducted using disulfonylethene. The strong radical accepting character of the sulfonyl group appeared to lead to the direct addition of the cyclohexyl radical to the Michael acceptor without carbonylation. It was also demonstrated that when allylsulfones were employed, $\beta\gamma$-unsaturated ketone products were obtained through the elimination of a phenyl sulfinate radical.$[26\text{a}]$ Recently, the same group expanded this method to the synthesis of amides in the presence of amines and CCl$_4$ (as a halogen donor).$[27]$ The reaction was thought to be initiated through photo excitation of the photocatalyst, 1,2,3,5-tetrakis(carbazol-9-yl)-4,6-dicyanobenzene (4CzIPN) to the excited state 4CzIPN*, which can oxidize the silicate to yield an alkyl radical $56$ and silicon cation along with a reduced 4CzIPN$\text{C}/\text{C}_0$ (Scheme 9). The resulting alkyl radical $56$ is then trapped by CO to form an acyl radical $57$, which can undergo a second radical addition to the alkene to generate the radical adduct $58$. The adduct $58$ is then reduced by 4CzIPN$\text{C}/\text{C}_0$ to yield a carbanion $59$, which is protonated to form the desired product $55$. In the alkyl sulfonate reaction, the eliminated phenyl sulfinate radical is reduced to regenerate the photocatalyst.

### 2.4. Light-Promoted Transition Metal-Catalyzed Carbonylation

In the 1980s, the groups of Watanabe and Suzuki independently observed that the reaction efficiency of transition metal-catalyzed carbonylation of alkyl iodides could be improved under light irradiation.$[28]$ They proposed that CO would insert into the alkyl-metal species to yield an acyl-metal species. This was further extensively evaluated by Ryu et al., who discovered that Pd complexes could accelerate the photo-induced atom-transfer-carbonylation (ATC) reaction of a wide range of alkyl iodides.$[29]$ For example, the addition of Pd complexes led to improved yields (from 54% to 87%) and shortened reaction time (from 50 h to 14 or 16 h) in the ATC reaction between 1-iodooctane, CO, and ethanol (Scheme 10A)$.[30]$ They proposed that the radical initiation step may involve a single electron transfer (SET) process from Pd$^0$ to cleave the C$\text{I}/\text{C}_0$ bond in an alkyl iodide $60$, generating a Pd$^+$ species and an alkyl radical $62$. The resulting radical $62$ could then undergo a radical addition to carbon monoxide to form an acyl radical $63$, which can couple with Pd$^2$-I to give a Pd$^3$ complex $64$. Subsequent metathesis of $64$ with ethanol and reductive elimination produced an ethyl ester $61$ (Scheme 10B). Importantly, terminal alkynes, boronic acids, amine and alkenes were also found to be able to participate in the three-component reactions to form the corresponding carbonyl products.$[31]$ In addition, this strategy was applied to the synthesis of the (−)-hinokinin precursor and dihydrocapsaicin.$[32]$ Interestingly, cyclized double car-
borylation products were obtained in good yields when 4-alkenyl iodides were used in the reactions. By light irradiation of a Pd catalyst, the authors also achieved four-component coupling reactions between iodoalkanes, alkenes, alcohols, and CO (Scheme 10C). It is noteworthy that alkyl chloride was tolerated (68b) and perfluorohexyl iodide proceeded effectively to furnish ester 68c.

Notwithstanding the efforts devoted to enabling carbon-cylation with CO, limitations such as the requirement for high pressure CO and harsh conditions, narrow substrate scopes still exist, and prevent the further development and application of these methods. The observation by the Ryu group[29] that the efficiency of palladium-promoted carbonylation could be improved by light, inspired Arndtsen et al. to develop highly efficient palladium-catalyzed photo-carbonylation reactions.[33] They questioned whether visible light could drive both the oxidative addition and the reductive elimination in one catalytic cycle. Their detailed reaction optimization and mechanistic studies suggested that light-induced single-electron transfer from Pd0 to aryl iodide, a radical-induced oxidative addition, was followed by formation of acyl radicals and a radical-involved reductive elimination. This breakthrough in the activation mode in light-induced CO carbonylation allowed carbonylation to proceed under ambient conditions, low pressure of CO, and with challenging aryl or alkyl halides and nucleophiles, generating valuable carbonyl derivatives such as acid chlorides, esters, amides, or ketones (Scheme 11). Both aryl- and alkyl-iodides and bromides are reactive toward carbonylation. Heterocycles (71g, 71i, 71o, 71r, 71s, 71u) sterically hindered alkyl halide (71t) and steroidal halide (71v) were all well-tolerated.

A plausible, experiment-based mechanism for this reaction was proposed (Scheme 12). Light-induced single-electron transfer from Pd0 complex 72 to aryl iodide leads to the formation of Pd1 complex 73 and an aryl radical 74. Subsequent oxidative addition of the putative Pd1 complex 73 with CO and the aryl radical 74 leads to the formation of Pd2 complex 75. Notably, this proposed radical-induced oxidative addition pathway with aryl bromides and alkyl iodides is not feasible under thermal conditions, demonstrating the unique role of visible light in this oxidative addition. Subsequent halide exchange produces a Pd complex 76. Light excitation of 76 would presumably lead to the formation of acyl radical 78, which after chlorine abstraction from 77 affords an acid chloride 79. Importantly, the light irradiation reduces the reductive elimination to low energy barriers.
In summary, studies on light-promoted radical carboxylation with CO have been briefly summarized. Radicals generated by light-induced bond homolysis, HAT processes, or redox processes can react with CO to form transient acyl radicals with or without the assistance of transition-metals, enabling a straightforward strategy for construction of a wide range of carboxylated compounds. The requirement however for high pressure CO to avoid decarbonylation, in comparison to conventional transition-metal-catalyzed carboxylation, remains a barrier to its broad application. A promising direction to achieve mild and practical carboxylation is to merge light activation with transition-metal catalysis.

3. Carbon Dioxide

Carbon dioxide (CO₂) is an attractive and renewable C₁ feedstock due to its high natural abundance, low cost and lack of toxicity. A number of strategies, such as transition metal-catalyzed and light-mediated carboxylation, have been developed to address its thermodynamic stability and kinetic inertness and convert CO₂ into carboxylic acids and derivatives.[34] Compared with conventional methods, the light-mediated systems are more environmentally friendly and can bypass the need for excess metallic reagents or harsh conditions. Pioneering studies in light-driven CO₂ fixation were first seen in the twentieth century, but the reported systems suffered from narrow substrate scopes and poor reaction efficiency and selectivity.[35] Major breakthroughs have been made in recent years, especially in the area of photocarboxylation.

3.1. UV Light-Promoted Carboxylation Using Carbon Dioxide

3.1.1. UV Light-Promoted Carboxylation of C(sp³)–H bonds

Direct C–H carboxylation through activation of the C–H bond is an atom- and step-economic process for synthesizing carboxylic acids and their derivatives. Although there has been impressive progress in transition-metal-catalyzed CO₂ functionalization of C(sp³)–H bonds,[36] direct carboxylation of unactivated C(sp³)–H bonds remains challenging. Numerous light-promoted carboxylation reactions of C(sp³)–H have recently been developed to tackle this challenge. In 2015, Murakami and co-workers disclosed the first carboxylation of benzyl C(sp³)–H bonds of ortho-alkylphenyl ketones promoted by UV or sunlight (Scheme 13).[37] A variety of aryl compounds (including an aryl aldehyde 81d) were well tolerated with moderate to good yields. It was proposed that the excited diaryl carbonyl group could abstract a hydrogen atom from the o-methyl group to yield a biradical species 84. This biradical species will presumably undergo rearrangement and endergonic isomerization to generate a highly reactive o-quinone methide 85 through a Norrish type II pathway. A subsequent [4+2] cycloaddition between the E-isomer of 85 and CO₂ affords a six-membered cycloadduct 86, which undergoes a ring-opening to provide the desired acid 87. The Z-isomer, on the other hand, would undergo a 1,5-hydrogen atom shift to regenerate the starting material. It is worth noting that sunlight can effectively promote this transformation.

In 2016, Murakami et al. reported a photo-induced carboxylation reaction of allylic C–H bonds using CO₂ and a combination of an aryl ketone type photocatalyst and a copper carboxylation catalyst under UV light irradiation (Scheme 14).[38] Both acyclic and cyclic alkenes were tolerated in the reaction, providing the corresponding carboxylic acids. However, it was observed that unsymmetrical alkenes containing two distinct types of C–H bonds at the allylic positions afforded mixtures of isomers (89d and 89e).

Mechanistically, photocatalyst 90 is excited by UV light to generate an excited state 91, inducing an intermolecular HAT process between 91 and the allylic substrate to generate a transient radical pair species 92 (Scheme 15). This radical pair would presumably undergo a radical-radical coupling reaction, producing the tertiary homoallylic alcohol intermediate 93. Subsequent deprotonation of 94 by copper tert-butoxide and C–C bond-cleaving reaction through β-carbon elimination yield the allylcooper species 95. A nucleophilic addition of 95 to CO₂ generates 96, which undergoes ligand...
exchange with potassium tert-butoxide to afford product 97 as the potassium salt, and regenerates the copper catalyst 98.

In 2016, Jamison and co-workers developed a novel continuous-flow method using 

\[
\text{CO}_2 + \text{amines} \rightarrow \text{products}
\]

under UV light irradiation (Scheme 16A). A wide range of benzyl amines with different functionalities were tolerated in the reaction, affording a variety of amino acids 100a–100g in good yields. High regioselectivity favoring the benzylic carboxylation was observed and non-benzylic amines with less activated \(\alpha\)-amino \(C\)/\(C_0\) bonds, such as \(N\)-cyclohexylpiperidine, selectively yielded an amino acid product 100h with moderate yields. The free amino acid could be obtained by cleavage of the 4-piperidone protecting group under neutral conditions, employing a polymer-supported amine scavenger such as JandaJel-NH_2 (Scheme 16B).

It was proposed that the excited singlet state of \(p\)-terphenyl 102 is generated by irradiation of \(p\)-terphenyl with UV light (Scheme 17). A subsequent SET between 102 and the tertiary amine 99 produces the strongly reducing \(p\)-terphenyl radical anion 103 and the corresponding amine radical cation 105. The radical anion 103 then donates an electron to \(\text{CO}_2\) to form a \(\text{CO}_2\) radical anion 104. In the meantime, deprotonation of amine radical cation 105 gives a neutral amine radical 106, which after a radical-radical coupling with \(\text{CO}_2\) radical anion 104 and subsequent protonation affords the carboxylic acid 100.

In 2019, Murakami and co-workers reported a UV light-promoted carboxylation of benzylic and aliphatic \(C\)/\(C_0\) bonds via ketone/Ni dual catalysis. Various substituted benzene derivatives were carboxylated at the benzylic positions (Scheme 18A). This methodology was also effective in the carboxylation of simple alkanes such as cyclohexane, cyclopentane, and pentane, although a mixture of three regioisomers was generated when \(n\)-pentane was the alkane substrate (Scheme 18B).
In the proposed mechanism, a ketone is excited by UV light (Scheme 19). The excited ketone $112$ is able to abstract the H atom from benzylic or aliphatic C(sp$^3$)/C$^0$H bonds producing a ketyl radical $113$ and a benzylic or aliphatic radical $115$. The ketyl radical $113$ is deprotonated by tBuO$^-$ to give a ketyl radical anion $114$, which is able to reduce the Ni$^{II}$ precursor to Ni$^0$. The benzylic radical or aliphatic radical formed ($115$) is captured by Ni$^0$ to form a Ni$^0$ complex $116$. Insertion of CO$_2$ into C-Ni$^0$ gives a Ni$^0$ carboxylate species $117$, which is reduced by ketyl radical anion to produce the carboxylate and regenerate the Ni$^0$ catalyst.

3.1.2 UV Light-Promoted Carboxylation of Alkenes with Carbon Dioxide

Continuous-flow β-selective hydrocarboxylation of styrenes was demonstrated by Jamison et al., employing 1,2,2,6,6-penta-methylpiperidine (PMP) as a reductant under UV light irradiation (Scheme 20). In this catalytic system, the photoexcited $p$-terphenyl catalyst ($E^0 = –2.63$ V vs. SCE in DMF) undergoes a SET with PMP to yield a strongly reducing $p$-terphenyl radical anion and a PMP radical cation.$^{[41]}$ CO$_2$ is reduced to the corresponding radical anion ($E^0 = –2.21$ V vs. SCE in DMF) by the $p$-terphenyl radical anion $122$. The CO$_2$ radical anion $104$ then attacks the β-position of the styrene, producing a stable benzylic radical $120$. The benzylic radical $120$ can then be reduced by the photocatalytic cycle to yield

![Scheme 18.](image1)

![Scheme 19.](image2)

![Scheme 20.](image3)
a carbanion 121, which is protonated to yield the desired dicarboxylated product. Notably, the reaction tolerates both \(\alpha\) - and \(\beta\)-substituted styrenes selectively providing the \(\beta\)-carboxylated products 119 in moderate yields. The reaction time required to complete this transformation can be reduced to 8 min by taking advantage of the continuous-flow apparatus.

3.2. Visible-Light-Promoted Carboxylation with Carbon Dioxide

3.2.1. Visible-Light-Promoted Carboxylation of C–Halogen Bonds and Their Equivalents

The carboxylation of C–halogen bonds provides a facile and straightforward method with which to introduce carboxylate functionalities. Early studies by Yamamoto et al. in the field of transition-metal-mediated carboxylation demonstrated that PhNi(L)Br (L = bipyridine) can react with CO\(_2\) to generate benzoic acids in moderate yields.[42] Significant advances were made by Martin and co-workers in 2009,[43] who reported the first catalytic carboxylation of alkyl bromides.[44] However, a stoichiometric amount of organometallic reductants was required in this transformation. In 2017, Iwasawa and co-workers reported the first example of visible-light-mediated carboxylation of aryl halides with CO\(_2\) using catalytic amount of Pd(OAc)\(_2\) and Ir(ppy)\(_2\)(dtbbpy)PF\(_6\) as a photoredox catalyst and \(N,N\)-diisopropylethylamine (iPr\(_2\)NEt) as a stoichiometric reductant, thereby overcoming the need for stoichiometric organometallic reductants (Scheme 21).[45] Various arylbromides (123a–123e) and chlorides (123f–123j) with different steric and electronic environments and heteroatoms could be used in this protocol to afford carboxylation products 124 with moderate to excellent yields.

It was proposed that the reaction is initiated by the reductive quenching of the excited photocatalyst by iPr\(_2\)NEt to yield an amine radical cation and reduced Ir\(_{\text{III}}\) species (Scheme 22). Subsequent reduction of the Pd\(_{\text{II}}\) complex by Ir\(_{\text{II}}\) via single electron reduction generates the active Pd\(_{\text{II}}\) species, which undergoes an oxidative addition with aryl halides 123 to yield a Pd\(_{\text{II}}\) complex 125. The insertion of CO\(_2\) into the Ar-Pd bond in 125 may occur before or after the single electron reduction of Pd\(_{\text{II}}\) to produce either 126 or 128. Both proposed pathways however lead to the formation of Pd\(_{\text{II}}\) carboxylate species 127, which undergoes a further single electron reduction to obtain the product and regenerate the Pd\(_{\text{II}}\) species.

König and co-workers later reported the carboxylation of aromatic and aliphatic bromides and triflates with CO\(_2\) (Scheme 23).[46] In this reaction, Ni was used as the transition metal catalyst and 4CzIPN was employed as the photocatalyst. Hantzsch ester (HEH) was employed as a reductant while K\(_2\)CO\(_3\) served as the CO\(_2\) source. The reaction displayed a broad substrate scope. Aryl- and alkyl- bromides and aryl triflates performed well in this reaction, generating carboxylic acids 131 in moderate to excellent yields. Notably, primary alkyl bromides, including tert-butoxycarbonyl (Boc) protected amines, can react effectively to yield the corresponding acid products 131h. In addition, unactivated cyclic alkyl bromides can be converted to the corresponding acid products 131i in moderate yield by adding one equivalent of cesium fluoride (CsF).

Similarly, the groups of Iwasawa and Jana separately described the carboxylation of aryl triflates with CO\(_2\) by using Pd/Ir dual catalytic systems under visible-light irradiation.[47] Carboxylation of alkanyl triflates was also achieved under Iwasawa’s conditions with a relatively short reaction time.

In 2019, the Feng group demonstrated the selective carboxylation of gem-difluoroalkanes through photoredox/Pd dual catalysis (Scheme 24).[48] A variety of arylidifluoroalkanes were evaluated to furnish predominantly Z-\(\alpha\)-fluoro-
crylic acids 133 in moderate to good yields, while alkyldifluoroalkenes failed to couple with CO2. The reaction displayed good functional group compatibility, tolerating dibenzofuran (133d), benzothiophene (133f) and indole (133q) moieties.

Mechanistically, the reaction is initiated by a SET between iPr2NEt and the photo-excited photocatalyst to produce the reduced photocatalyst IrII, which reduces the gem-difluoroalkene 132 to yield fluoroalkenyl radical 134 (Scheme 25). The carbon radical 134 is intercepted by Pd0 to form a putative PdI species 135, representing a rare example in which a transition metal is employed to intercept the fluorine-containing carbon radical to form a versatile organometallic species. Subsequent CO2 coordination and migratory insertion with 136 affords a carboxyl-PdI complex 137. A SET between 137 and IrII photocatalyst regenerates Pd0 and yields the carboxylate 138.

Yu and co-workers recently developed a dearomative arylcarboxylation of indoles with CO2 which selectively produces indoline-3-carboxylic acids by a successive single-electron transfer (SSET) strategy (Scheme 26).[49] Under transition-metal catalysis, the dearomative reductive coupling of indoles with two electrophiles is challenged by the low reactivity of the aromatic carbon-carbon double bond. It was proposed that these challenges could be overcome by using a radical relay SSET strategy. The authors envisioned that the reduction of the carbon-halide bond by visible-light-induced single-electron transfer would lead to a highly reactive aryl radical which would then undergo a selective intramolecular radical addition to the C2/C3 double bond in indoles instead of CO2. Subsequent SET would reduce the benzylic radical to a benzylic anion, which after trapping with CO2 and protonation would produce the arylcarboxylation product. Indeed, after reaction optimization, this transformation exhibited a broad substrate scope and high chemo- and diastereoselectivity. Substrates bearing functional groups such as bromo (140e), fluoro (140g) and thiophene (140i) were well tolerated (Scheme 26A). Notably, an all-carbon quaternary center could be formed to deliver product 140d with high diastereoselectivity. For unactivated aryl halides (Scheme 26B), with modified conditions, various aryl bromides
and iodides could proceed the reaction effectively, albeit with reduced diastereoselectivity.

In light of these experiments, a possible mechanism was proposed (Scheme 27). Radical anion 4CzIPN$^-$/C0 and the radical cation DIPEA$^+$ are generated by reductive quenching of the photo-excited 4CzIPN ($E_{1/2} [4CzIPN*/4CzIPN] = +1.35 \text{ V vs. SCE in MeCN}$) by DIPEA ($E_{1/2} \text{ox} = +0.63 \text{ V vs. SCE in DMF}$). An arylbromide $139$ is reduced by a single electron transfer from 4CzIPN$^-$/C0 to generate the radical anion $143$. Subsequent fragmentation of $143$ releases a bromide anion to give an aryl radical $144$, which undergoes an intramolecular radical addition to the C2$^-$/C0$^+$C3 double bond of indole to deliver the benzylic radical $145$. A second single electron transfer from 4CzIPN$^-$/C0 to $145$ leads to the formation of a carbon anion $146$, which undergoes nucleophilic addition to CO2 followed by protonation to deliver the desired dearomative arylcarboxylation product.

The Yu group disclosed the first external reductant-free visible-light-driven carboxylations of organic tetraalkyl ammonium salts with CO2 (Scheme 28).$^{[50]}$ The cross-electrophile coupling tolerates primary, secondary and tertiary C$^-$/C0$^+$N bonds and a variety of functional groups, giving $\alpha$-substituted arylacetic acids $148$ in moderate to good yields. TEMPO trapping experiments indicated the involvement of radicals. In addition, a control experiment with D2O as the additive demonstrated the HAT with solvent (DMF) and the formation of a benzylic anion intermediate. Moreover, the reaction with deuterated substrates indicated that the amine radical cation deprotonation and benzylic anion protonation occurs during the reaction process. Stern–Volmer luminescence studies and electrochemical analysis suggested that the iridium catalyst might be quenched by NMe3 generated in situ from the ammonium salt or an $\alpha$-amino radical. Moreover, the presumed benzyl radical ($E_{1/2} \text{red} = 1.43 \text{ V vs. SCE}$) might undergo further reduction by Ir$^{III}$-catalyst ($E_{1/2} \text{red (IrIII/IrII)} = 1.51 \text{ V vs. SCE}$) to afford a benzylic carbon anion. The subsequent nucleophilic attack of the carbon anion to CO2
and protonation gave the carboxylic acid products. However, the authors could not exclude other alternative pathways, including the coupling between benzyl radicals and ketyl radicals.

### 3.2.2. Visible-Light-Promoted Carboxylation of Alkenes with Carbon Dioxide

Pioneering studies from Hoberg et al. demonstrated the light-mediated formation of five-membered nicklacycles from Ni$^0$, alkenes and CO$_2$. Encouraged by Hoberg’s work,[3] the Rovis group reported the first Ni-catalyzed hydrocarboxylation of aryl alkenes using a stoichiometric amount of ZnEt$_2$ as the reductant.[3] Subsequently, several other groups demonstrated the use of various complexes of transition metals such as Fe, Rh and Ru as suitable catalysts for hydrocarboxylation of olefins.[33] In 2017, Iwasawa and co-workers disclosed the first visible-light-driven catalytic hydrocarboxylation of alkenes with CO$_2$ using a combination of a Rh complex as a carboxylation catalyst and a [Ru(bpy)$_3$]$^{2+}$ photocatalyst in the presence of a tertiary amine (Scheme 29).[54] The substrate scope was limited to only electron-deficient styrenes, which yielded products 152 in moderate yields. Mechanistically, the authors were able to prove that the reductant, iPr$_2$NEt, could serve as a proton source to form the key Rh-hydride intermediate under photoredox conditions. Control experiments revealed that the photoredox catalyst, light and the tertiary amine were all essential for the carboxylation reaction. A luminescence quenching study showed that a triplet-triplet energy transfer process could be involved in the carboxylation. The key catalytic species, the Rh$^1$ complex 153 undergoes hydrometallation and nuclophilic carboxylation to yield the Rh$^2$ carboxylate complex 155. Complex 155 then undergoes electron and proton transfers enabled by the photoredox catalyst to generate a Rh$^{III}$ dihydride 156, which delivers the carboxylic acid in the presence of a base and regenerates complex 153.

König and co-workers developed a regioselective hydrocarboxylation of aryl alkenes using CO$_2$ via the synergistic merger of a photoredox catalyst with a nickel catalyst (Scheme 30).[55] The regioselectivity of this reaction is dependent on the choice of the ligand. The use of neocuproine as ligand yielded the Markovnikov products 158 (branched:linear = 100:1) (Scheme 30A), while the use of 1,4-bis(diphenylphosphino)butane (dpbb) as ligand promoted the formation of anti-Markovnikov products 159 (Scheme 30B).

#### A) Markovnikov hydrocarboxylation

- **CO$_2$ (balloon)**
- 4CzIPN (1 mol%)
- L$_{NBR}$ (l = neocuproine)(10 mol%)
- K$_2$CO$_3$ (1.0 equiv), DMF, blue LEDS, r.t., 24 h

**Selected substrates**

<table>
<thead>
<tr>
<th>R = H</th>
<th>158a, 72%</th>
</tr>
</thead>
<tbody>
<tr>
<td>R = OMe</td>
<td>158b, 42%</td>
</tr>
<tr>
<td>R = COOMe</td>
<td>158c, 81%</td>
</tr>
</tbody>
</table>

#### B) Anti-Markovnikov hydrocarboxylation

- **CO$_2$ (balloon)**
- 4CzIPN (1 mol%) N$_2$B$_2$P$_2$ (10 mol%)
- dpbb (20 mol%), LiOCC(CH$_3$_2)$_2$ (1.0 equiv), 4 Å MS (50 mg)

**Selected substrates**

<table>
<thead>
<tr>
<th>R = H</th>
<th>159a, 44% (b = 99.2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R = OMe</td>
<td>159b, 43% (b = 99.2)</td>
</tr>
</tbody>
</table>

**Scheme 30.** Visible-light-driven Ni-catalyzed regioselective hydrocarboxylation of aryl alkenes.

Based on the proposed mechanism (Scheme 31), the active Ni$^0$ [E$_{1/2}^{red}$ (Ni$^3$/Ni) = −1.2 V vs. SCE] species is generated from Ni$^0$ in situ via two single electron transfer steps under photocatalytic reductive conditions [4CzIPN$^-$ (E$_{1/2}^{red}$ = −1.21 V vs. SCE)]. When neocuproine is used as the ligand, protonation by the Hantzsch ester radical cation produced a Ni hydride complex 160. Ni hydride 160 then undergoes hydrometallation with styrene, yielding an organonickel complex 161. Reduction of the complex followed by the insertion of CO$_2$ leads to the generation of a nickel carboxylate intermediate 163. Further reduction of 163 gives the Markovnikov product. On the other hand, when dpbb is employed, a five-membered nicklacycle 166 is generated.
through the formation of complex 165. The nickelacycle 166 can subsequently be reduced to yield a NiI complex 167. Finally, single electron reduction of 167 affords the anti-Markovnikov product and regenerates the Ni0 catalyst.

Compared to hydrocarboxylation of alkenes, difunctionalization of alkenes using CO2 is more challenging and provides products with higher structural diversity. In this context, Martin and co-workers reported the first example of visible-light-promoted, photoredox-catalysis assisted difunctionalization of alkenes with CO2 (Scheme 32).[56] In their study, the authors demonstrated a redox-neutral intermolecular dicarbofunctionalization of styrenes with CO2 and simple radical precursors, such as sulfonates including the Langlois reagent (CF3SO2Na), trifluoroborates and oxalates without requiring stoichiometric reductants. Notably, the merits of the trifluoromethycarboxylation were demonstrated by its tolerance of wide range of sensitive functional groups, including tosylates (171e) and boronic esters (171g). Mechanistically, the reaction is thought to proceed through a radical generated by single electron reduction with the excited photocatalyst Ir(ppy)2(dpbbpy)PF6 [E1/2^red(IrIII/IrII) = −1.51 V vs. SCE in MeCN] adding to the vinyl arene 172 to form a benzylic radical 173. Another single electron reduction of radical 173 would generate a benzylic anion, which finally traps CO2 to yield the carboxylate 174 (Scheme 33). Li’s group utilized perfluoroalkyl iodides as radical precursors to achieve a visible-light-driven carboxylative cyclization of allyl amines to give perfluoroalkylated oxazolidinones in the presence of CO2 and a base.[57] Yu et al. further expanded the reaction scope to difluoroalkyl bromides assisted by photoredox catalysis.[58] In 2019, Cheng’s group reported a palladium-catalyzed radical carboxylative cyclization of 2-(1-arylvinyl)anilines to generate 1,4-dihydro-2H-3,1-benzoxazin-2-ones using alkyl bromides and CO2 under visible-light irradiation.[59]

A visible-light-promoted, iron-catalyzed thiocarboxylation of styrenes and acrylates with CO2 was reported in 2017 by the Yu group (Scheme 34).[60] A range of styrenes and acrylates could be employed in this system to give useful β-thioacids 177 in good yields with high regio-, chemo- and diastereo-selectivity. The precise mechanism was not fully clear, but the authors proposed that CO2 radical anion and alkyl radicals might be involved in the catalytic cycle to achieve this unique regioselectivity.

Scheme 31. Proposed mechanism of visible light driven Ni-catalyzed regioselective hydrocarboxylation of aryalkenes.

Scheme 32. Dicarbofunctionalization of styrenes with CO2.

Scheme 33. Proposed mechanism of dicarbofunctionalization of styrenes with CO2.

Scheme 34. Thiocarboxylation of styrenes and acrylates.
Wu and co-workers developed a three-component metal-free silacarboxylation and carbocarboxylation between electron-deficient alkenes, CO₂ and either silanes or C(sp³)-H alkanes through a synergistic combination of photoredox and HAT catalysis (Scheme 35).[61] The silacarboxylation tolerates a variety of functional groups including cyano (179a), ketone (179b), carboxylic acid (179d) and amide (179e) groups, producing β-silacarboxylic acids in moderate to good yields.

The synthetic application of this protocol was also demonstrated in the late-stage elaboration of a biologically relevant complex molecule 179h. Common silanes such as tripropylsilane, triisopropylsilane, dimethylbenzylsilane, dimethyltert-butylsilane, and dimethylphenylsilane were viable in this transformation. The strategy was also successfully extended to carbocarboxylation, where excellent functional group compatibility was illustrated, giving products such as γ-amino acids (180a, 180b) which are important structural motifs in many pharmaceutically important compounds. Notably, aryl ketones could be used as electrophiles in a three-component coupling reaction with silanes and alkenes to produce γ-silyl alcohols 182.

The mechanism was proposed to begin with the photoactivation of 4CzIPN to 4CzIPN* (Scheme 36). 4CzIPN* would then be reductively quenched by 3-acetoxyquinuclidine, yielding reduced 4CzIPN and a radical cation intermediate 183. This radical cation 183 promotes a HAT from Si–H or C–H to generate a silyl or carbon radical 185 and a quinuclidinium cation 184. The generated silyl or carbon radical 185 adds to the alkene to yield radical intermediate 186 (~0.77 V vs. SCE for p-CN benzyl radical analogue), which is then reduced by the 4CzIPN⁻ (E_{1/2}^{red} = –1.21 V vs. SCE in MeCN) to generate an anion 187. The anion 187 subsequently undergoes nucleophilic addition to carbon dioxide and protonation to yield the desired silacarboxylation or carbocarboxylation product.

Inspired by the recent efforts in the development of alkene difunctionalization, Li and co-workers developed a visible-light-driven reductive carboarylation of styrenes with CO₂ and aryl halides in a regioselective manner using the readily available and inexpensive HCO₂K as the terminal reductant (Scheme 37).[62] This method provided an atom-economic and direct access to valuable hydrocinnamic acid derivatives 190. The reaction exhibits a broad substrate scope of aryliodides, bromides, with good yields and excellent regioselectivity (Scheme 37A). Moreover, a variety of (hetero)aryl moieties were well tolerated, affording products 193 with moderate to good yields (Scheme 37B). Less reactive aryl chlorides were effective in this reaction and the substrate scope can also be extended to alkyl halides. In addition, substituted styrenes were more effective than α,β-unsubstituted styrene derivatives, while aliphatic alkenes were unreactive. By employing ¹³C labeling, it was found that 74% of ¹³C incorporation was from CO₂ gas with the rest of the CO₂ source being derived from HCO₂K or K₂CO₃.

A plausible mechanism was proposed based on all the experimental results and literature precedents (Scheme 38). The photo-excited *IrIII is produced upon blue light irradiation, which is subsequently reductively quenched by 1,4-diazabicyclo[2.2.2]octane (DABCO) to give IrII and a radical cation of DABCO. A hydrogen atom transfer from HCO₂K promoted by DABCO radical cation gives a CO₂ radical anion, which reduces aryl halide to an aryl radical 194. Subsequent regioselective addition of this aryl radical to alkene affords a benzylic radical 195, which is further reduced by IrII to a benzylic anion 196. Nucleophilic addition of 196 to CO₂ produces the carboxylate 197, which is methylated to deliver an ester product.
Yu and co-workers expanded the scope of the visible-light-mediated difunctionalization of alkenes to phosphono-carboxylation with enamides and acrylates (Scheme 39). A range of aryl enamides were difunctionalized, providing various \( \beta \)-phosphono-\( \alpha \)-amino acids bearing a quaternary carbon center with moderate to excellent yields (Scheme 39A). However, the hydrophosphinylation product instead of the desired carboxylative product was obtained when an alkyl enamide was used. Phosphonocarboxylation of various acrylates is shown in Scheme 39B, although hydrophosphinylation of electron-deficient alkenes with phosphine oxides and phosphites (H-P(O)) compounds via nucleophilic addition has been well documented. Notably, an enantioenriched H-P(O) compound derived from a (4R,5R)-taddol derivative was used for an enantioselective transformation with CO\(_2\), which resulted in poor diastereoselectivity (Scheme 39C). Preliminary mechanistic studies revealed an \( \alpha \)-amino benzylic anionic species as the key intermediate and the involvement of a reductive quenching photoredox catalytic cycle.

More recently, Xi and co-workers reported a visible-light-promoted carbocarboxylation of styrenes using CO\(_2\) and amines catalyzed by photocatalyst 4CzIPN to access \( \gamma \)-amino acids 205 and 208 (Scheme 40). Mechanistically, photoexcited 4CzIPN is reduced by aniline via SET to give a radical cation intermediate, which is deprotonated to

**Scheme 39.** Phosphonocarboxylation of alkenes with carbon dioxide via visible-light promoted photoredox catalysis.

Yu and co-workers expanded the scope of the visible-light-mediated difunctionalization of alkenes to phosphono-carboxylation with enamides and acrylates (Scheme 39). A range of aryl enamides were difunctionalized, providing various \( \beta \)-phosphono-\( \alpha \)-amino acids bearing a quaternary carbon center with moderate to excellent yields (Scheme 39A). However, the hydrophosphinylation product instead of the desired carboxylative product was obtained when an alkyl enamide was used. Phosphonocarboxylation of various acrylates is shown in Scheme 39B, although hydrophosphinylation of electron-deficient alkenes with phosphine oxides and phosphites (H-P(O)) compounds via nucleophilic addition has been well documented. Notably, an enantioenriched H-P(O) compound derived from a (4R,5R)-taddol derivative was used for an enantioselective transformation with CO\(_2\), which resulted in poor diastereoselectivity (Scheme 39C). Preliminary mechanistic studies revealed an \( \alpha \)-amino benzylic anionic species as the key intermediate and the involvement of a reductive quenching photoredox catalytic cycle.

More recently, Xi and co-workers reported a visible-light-promoted carbocarboxylation of styrenes using CO\(_2\) and amines catalyzed by photocatalyst 4CzIPN to access \( \gamma \)-amino acids 205 and 208 (Scheme 40). Mechanistically, photoexcited 4CzIPN is reduced by aniline via SET to give a radical cation intermediate 210, which is deprotonated to
form an α-amino carbon radical species 212. A subsequent radical addition of 212 to styrene generates a γ-amino benzylic radical 213, which is reduced by the reduced 4CzIPN radical anion via a SET to afford a γ-amino benzylic carbanion 214, proposed as a lithium chelated species. A subsequent nucleophilic addition of 214 to CO₂ or LiCl activated CO₂ and protonation provides the desired γ-amino acid (Scheme 41).

More recently, König and co-workers reported a redox-neutral photocatalytic C(sp²)-H carboxylation of styrenes and (hetero)arenes. Their protocol provides a direct access to trans-cinnamic acid derivatives, in contrast to the existing light-mediated hydrocarboxylation of styrenes (Scheme 42A). The substrates bearing electron-donating groups (217c–217e) or weak electron-withdrawing groups (217f) were able to give corresponding product in good yield. When stronger electron-withdrawing group was applied (217g), a lower yield was observed. Moreover, 4-cyanostyrene (217h) was not suitable for this reaction. A possible mechanism has been proposed based on a series of mechanistic investigation, (Scheme 42C). The photocatalyst TMAH is deprotonated by cesium carbonate to form an anionic species TMA⁻, which under visible light irradiation generates a strongly-reducing excited anion *TMA⁻ [E(TMA⁻/*TMA⁻) = −2.92 V vs. SCE]. Although a direct reduction of CO₂ by *TMA⁻ is thermodynamically feasible, the luminescence lifetime of *TMA⁻ in the CO₂-saturated solution of DMSO remained almost unchanged in the Stern–Volmer experiment, which indicates a kinetic barrier preventing the formation of CO₂ radical anion. On the other hand, the luminescence lifetime of *TMA⁻ decreased obviously in the presence of styrene, demonstrating that a SET between *TMA⁻ and styrene initiates the reaction. Then a radical anion 219 traps CO₂ to give a radical carboxylate 220. Oxidation of 220 by TMA via SET generates

Scheme 40. Photoredox-catalyzed dicarbofunctionalization of styrenes with amines and CO₂.

Scheme 41. Proposed reaction mechanism for the synthesis of γ-amino acid.

Scheme 42. Redox-neutral photocatalytic C(sp²)-H carboxylation of styrenes and (hetero)arenes.
a benzylic cation followed by a deprotonated elimination to give the carboxylate product. According to the proposed mechanism, electron-deficient styrenes lead to reduced yield since electron-withdrawing groups can stabilize the radical anion intermediate and reduce its nucleophilicity to CO₂.

Notably, this protocol can be extended to a series of arenes to achieve directly C(sp²)-H carboxylation with CO₂, including naphthalene derivatives and heteroaromatic compounds (Scheme 42B).

3.2.3. Visible-Light-Promoted Carboxylation of Alkynes with Carbon Dioxide

Substituted alkenes are important structural motifs found in pharmaceuticals, agrochemicals and biologically active natural products and are also versatile building blocks in organic synthesis. Access of alkenes through the direct functionalization of alkynes represents a straightforward pathway. Although several groups have reported transition-metal-catalyzed alkyne hydrocarboxylation, the molecular scaffolds compatible with these protocols were limited to linear acrylic acids and 2-pyrones. Furthermore, stoichiometric amounts of reductants, such as Et₂Zn, silanes, Mn, or Zn, were required. The first breakthrough that overcame these limitations was reported by Wu, Zhao, and co-workers, who developed the first visible-light-driven hydrocarboxylation and carbocarboxylation of alkynes using CO₂ in a Ir/Co dual catalysis system. Both aryl and aliphatic substituted internal alkynes are good candidates for this transformation (Scheme 43). In general, the regioselectivity of this reaction is dependent on the relative steric environment of the unsymmetrical alkene, where the CO₂ insertion happens adjacent to the less sterically hindered substituent. Notably, it was observed that aryl bromides, which were usually incompatible with metal reductants, can be tolerated in this protocol generating products such as 223d. Interestingly, 2-pyrones were generated when terminal alkynes were employed, representing a rare case of synthesis of 2-pyrones via intermolecular [2+2+2] cycloaddition between two alkynes and CO₂.

Upon the introduction of an ancillary substituent on the aryl ring, various products, such as coumarins, 2-quinolones and 2-benzoxepinones, could be accessed directly through a one-pot alkyne hydrocarboxylation/alkene isomerization/cyclization sequence (Scheme 44). Here, the Ir photocatalyst plays two roles. First, it promotes the single electron transfer in alkylene hydrocarboxylation and second, energy transfer in the following alkene isomerization. Notably, this method could also generate seven-membered heterocycles by switching phenyl ether substituted alkynes to benzyl ether substituted alkynes, highlighting its utility in accessing heterocycles with different ring sizes. Based on the hypothesis that a five-membered cobaltacycle intermediate derived from the Co species, alkynes and CO₂ might be involved in the process, the authors strategically incorporated an electrophilic substituent in the alkylene substrate to trigger a kinetically favored intramolecular addition of the cobaltacycle intermediate, realizing alkylene difunctionalization. Indeed, γ-hydroxybutenolides and could be obtained using CO₂ with orthoester substituted aryl alkynes under the photoredox/cobalt dual catalytic conditions (Scheme 45), serving as a novel method to access these interesting structural motifs, which can be found in natural products such as ianthellidone G, manoalide and petrosaspongiolide M, and in pharmaceutically important agents.

Mechanistically, the transformation is initiated by a reductive quenching of visible light-excited IrIII with Pr₂NET to yield a reduced IrI species and an amine radical cation. The IrI species [Ir(ppy)_2(Ne)I] = −1.51 V vs. SCE in MeCN is then oxidized by a CoII species (E(1/2) = −0.72 V vs. SCE in MeCN) to regenerate IrIII and yield a CoI species. CoI couples with CO₂ and alkynes to form the key five-membered cobaltacycle intermediate (Scheme 46). Protonolysis of the CoIII cobaltacycle intermediate may undergo photoreduction to regenerate CoII. ZnBr₂ aids in this process.
by transmetallation to furnish a carboxylate 237. An Ir-mediated triplet-triplet energy transfer process followed by an acid-mediated intramolecular cyclization delivers the desired product 226. In the case of terminal alkynes, migratory insertion of another alkyne molecule into the five-membered ring 234 would yield a seven-membered cobaltacycle 236, which then undergoes reductive elimination to yield 2-pyrones 224. 

\[ \text{Scheme 45. Synthesis of } \gamma\text{-hydroxybutenolides from ortho-ester substituted aryl alkynes and } \text{CO}_2 \text{ driven by light irradiation.} \]

\[ \text{Scheme 46. Proposed mechanism of visible light driven Co-catalyzed alkyne hydrocarboxylation and } [2+2+2+2] \text{ cycloaddition with } \text{CO}_2. \]

Zn complex 241. Complex 241 underwent two rounds of tautomeration to yield \( \gamma \)-hydroxybutenolides 230.

3.2.4. Visible-Light-Promoted Carboxylation of Enamides and Imines with Carbon Dioxide

While the utilization of \( \text{CO}_2 \) as an abundant and reusable raw material has attracted significant interest, the construction of amino acids utilizing \( \text{CO}_2 \) as a C1 source represents one particularly intriguing but challenging transformation. There is an impetus to develop new methods for the catalytic hydrocarboxylation of readily available enamides or imines for this important family of bioactive compounds.

The first catalytic, transition metal-free light-promoted hydro-carboxylation of enamides and imines was pioneered by the Yu group in 2018, who used 1 atm of \( \text{CO}_2 \) to generate important \( \alpha, \alpha \)-disubstituted \( \alpha \)-amino acids (Scheme 48). [69]

Through the generation of \( \alpha \)-amino carbanions achieved by visible-light-driven reduction of enamides, they were able to overcome the inherent nucleophilicity at the \( \beta \)-position of enamides, leading to selective formation of \( \alpha, \alpha \)-amino acids. A wide range of aryl enamides (Scheme 48A) and aryl imines (Scheme 48B) could be applied using iPr2NEt as a reductant. Subsequently, the same group realized a visible-light-mediated phosphonocarboxylation of enamides as described in Scheme 39A.

Walsh and co-workers reported a visible light-mediated, Ir(ppy)2(bpy)PF6-catalyzed hydrocarboxylation of non-protected aryl imines with N,N-dicyclohexylmethylamine (Cy2NMe) as a sacrificial electron donor for the synthesis of \( \alpha, \alpha \)-disubstituted \( \alpha \)-amino acids (Scheme 49). [70] This method has good functional group compatibility, tolerating aryl chlorides (248d), aryl(hetero)cycles (248b, 248e) and allyl groups (248c) to provide these \( \alpha \)-amino acids with moderate to excellent yields.

The authors proposed that the coordination of an amine radical cation and imines forms a complex 249 which contains

\[ \text{Scheme 47. Proposed mechanism of light mediated } \gamma\text{-hydroxybutenolide synthesis from ortho-ester substituted aryl alkenes and } \text{CO}_2. \]
a 2-center-3-electron bond, facilitating the reduction by Ir\(\text{II}\) to form the radical anion intermediate (250/251). Ir\(\text{III}\) is regenerated in the process (Scheme 50). The highly reactive N-radical 251 is quickly quenched by the amine radical cation via HAT to form the \(\alpha\)-amino carbanion intermediate 252. Intermediate 252 acts as a strong nucleophile and reacts with \(\text{CO}_2\) to give the adduct 253, which after methylation affords the ester product. A key factor in controlling the reactivity is the resonance form of 251, which has a greater spin density on the nitrogen (0.37) than on the diaryl substituted carbon (0.18), indicating that diaryl substituted carbon carries more anionic characters.

3.2.5. Visible-Light-Promoted Carboxylation of N-sulfonylhydrazones

Recently, the König group reported a photo-Wolff–Kishner type thiocarboxylation of N-tosylhydrazones assisted by Ir\(\text{III}\)-based photoredox catalysis (Scheme 51). 255. A series of aromatic aldehyde-derived N-tosylhydrazones (256a–256f, 256m–256p, 256r, 256s, 256v, 256y, 256z, 256w, 256x, 256k),
256m–256n) and ketone-derived N-tosylhydrazones (256g–256l, 256o–256p) bearing various functional groups proved to be effective substrates. Increasing the steric hindrance at the α-position of the N-tosylhydrazones failed to decrease the product yield (256q–256s). Notably, the thiocarboxylation of a substrate 256t derived from an aliphatic ketone was also achieved at a low temperature (0°C). The scope of thiols was also evaluated. Thiophenols with electron-donating (256v) or electron-withdrawing groups (256u, 256w) were well tolerated and aliphatic thiols (256x–256z) were also able to generate the desired products in moderate yields.

The authors proposed a plausible mechanism (Scheme 52) based on a series of spectroscopic investigations and control experiments. This reaction is initiated by a SET between thiophenolate and the excited photoredox catalyst. N-tosylhydrazone is added by the resulting sulfur radical to generate the aminyl radical species 259. Then, a functionalized diazene intermediate 260 is formed by the fragmentation of this radical species 259, and an arenesulfonyl radical is produced simultaneously. Intermediate 260 subsequently undergoes a Wolff–Kishner type nitrogen extrusion process to give a sulfur carbanion 261 which attacks CO₂ to produce the thiocarboxylated product. The photoredox catalytic cycle is turned over by another SET between the arenesulfonyl radical and IrII.

Scheme 52. Proposed mechanism of photo-Wolff–Kishner type thiocarboxylation.

3.2.6. Visible-Light-Promoted Carboxylation of C(sp³)/C₀H bonds

There has been much development in the activation of C(sp³)/C₀H bond, but the direct carboxylation of C(sp³)/C₀H bonds remains limited to acidic C(sp³)/C₀H bonds, or requires harsh conditions and UV-light. In 2019, Martin and König collaborated to demonstrate a selective C(sp³)/C₀H carboxylation reaction catalyzed by the merger of photoredox and Ni catalysis. The selectivity of the C–H bond carboxylation was controlled through an alkyl bromide-induced Ni-chain walking process. The choice of ligand had a major impact on the selectivity. Subtle changes in substituents on carbons adjacent to the nitrogen atoms in the ligand can lead to a marked decrease in reactivity, and ligand L₁ appeared to be the optimal ligand for both reactivity and selectivity. The use of desiccants led to lower yields, suggesting that water played a significant role in the reaction. The carboxylation at benzylic sp³ C–H bond showed excellent selectivity and functional group tolerance, although products (263) were obtained in moderate yields (Scheme 53).

Scheme 53. Substrate scope of benzylic C(sp³)/H carboxylation.

In addition to benzylic C(sp³)/H bonds, the reaction is also compatible with non-activated alkanes, and the highest efficiency was achieved by using L₂ (Scheme 54). The reaction was found to tolerate a variety of functional groups, such as esters, nitriles (265g), ketones (265k), alkyl chlorides (265f), amides (265j) and indoles (265h). Most
substrates showed high selectivity for the desired linear products over the branched products.

Through combined computational and experimental studies, the authors proposed a plausible mechanism that begins with the reduction of Ni$^{II}$ to Ni$^{0}$ (Scheme 55). Ni$^{0}$ then undergoes an oxidative addition to the alkyl bromide to produce an alkyl-Ni$^{II}$ complex ($\text{266}$). DFT calculations suggested that complexes $\text{266}–\text{268}$ have similar energies and exist in rapid equilibrium via facile β-H elimination from cationic Ni$^{II}$ intermediates. It was also suggested by DFT calculations that CO$_2$ insertion was less favorable for the Ni$^{II}$ complex, thus supporting CO$_2$ insertion through a Ni$^{I}$ complex. Insertion of CO$_2$ into the Ni–C bond of Ni$^{I}$ species $\text{269}$ yields a Ni-carboxylate $\text{270}$. Ni-carboxylate $\text{270}$ then undergoes a SET to produce the carboxylate $\text{271}$ and regenerate Ni$^{0}$.

Another breakthrough in the field of visible light-mediated direct carboxylation of sp$^3$–C–H bonds was achieved by König and co-workers when they presented a metal-free approach to the carboxylation of the benzylic C(sp$^3$)–H bonds through an indirect HAT process, mediated by a triisopropylsilane thiol HAT catalyst.[74] The reaction shows a broad substrate scope of benzylic type compounds, including electron-withdrawing and electron-donating substituents, as well as cyclic and heterocyclic compounds (Scheme 56). Notably, in the case where there is more than one benzylic C–H site in the substrate, monocarboxylated acids ($\text{273a}–\text{273w}$) are formed exclusively. The applicability and efficiency of this protocol was demonstrated through the synthesis of several drug molecules containing the structural motif of 2-phenylpropionic acid, such as fenoprofen ($\text{274}$), flurbiprofen ($\text{275}$), naproxen ($\text{276}$) and ibuprofen ($\text{277}$) (Scheme 57). In the case of ibuprofen, where two sets of benzylic protons are present, a selective formation of ibuprofen $\text{277}$ over the alternative product $\text{278}$ was observed. The high chemoselectivity can be presumably attributed to the steric environment adjacent to the benzylic protons, where the carboxylation leading to ibuprofen was less sterically hindered and thus favored.

During the mechanistic investigation of the photocatalytic decarboxylation, it was observed that a cyano (CN) group on 4CzIPN was replaced by an ethylbenzene moiety. This formed 2,3,4,6-tetra(9H-carbazol-9-yl)-5-(1-phenylethyl)benzonitrile (4CzPEBN), which was proposed as the active photocatalyst to catalyze the reaction. 4CzPEBN is excited by light $[E_{1/2}^\text{PC/PC}^* = +1.19 \text{ V vs. SCE}]$, and then the thiol HAT catalyst ($E_{1/2}^{\text{thiol}/\text{HAT}} = +0.28 \text{ V vs. SCE}$)[75] is oxidized, yielding a reduced 4CzPEBN$^–$ (Scheme 58). The oxidized thiol
catalyst then loses a proton to yield a thiol radical (BDE S-H = 88.2 kcal mol⁻¹). The thiol radical abstracts a hydrogen from the benzylic C–H bonds (BDE C-H = 85.4 kcal mol⁻¹)⁷⁵ to yield a benzylic radical 279, regenerating the thiol catalyst. The benzylic radical (E₁/₂ red = 1.60 V vs. SCE for the phenylethyl radical)⁷⁶ is then reduced by 4CzPEBN⁻ [E₄/₃ (PC/PC) = −1.69 V vs. SCE] to form a carbanion 280 and regenerate the photocatalyst. The anion 280 is then captured by CO₂, and is protonated to yield the carboxylic acid 273a.

The recent development of light-driven carboxylation utilizing CO₂ as a C₁ source to access value-added chemicals has been briefly summarized in this section. These works represent novel activation modes complementary to existing transition-metal catalyst approaches and have inspired new perspectives for utilizing CO₂ in carboxylation, which can produce useful compounds for the agrochemical and pharmaceutical industries. Considering the rapid development in photocatalysis, we foresee that photo-mediated carboxylation using CO₂ will continue to progress rapidly to achieve challenging and previously inaccessible transformations, enabling greener and more sustainable protocols for fine chemical synthesis using CO₂.

4. Gaseous Alkenes and Alkynes

Gaseous alkenes and alkynes, such as ethylene, propene, 1,3-butadiene and acetylene, are important industrial feedstock chemicals for the synthesis of value-added chemicals. However, most light-promoted reactions involving such unsaturated gaseous compounds are limited to photo-cycloadditions.⁷⁷ In 2017, the Wu group developed a “stop-flow” micro-tubing (SFMT) reactor enabling efficient, safe and convenient screening of photo-mediated transformations involving the aforementioned gases (Figure 2).⁷⁸ The SFMT reactor platform is a design adapted from continuous flow systems with the addition of batch elements. Instead of allowing the reaction mixture to flow through the tubing continuously, the SFMT platform allows the flow to be stopped at will. The platform differs from earlier approaches in that the SFMT is based on a “switch in-and-out” approach using shut-off valves on each end of the micro-tubing reactor. This means that the SFMT system can be used as an independent module, in which the reactor can be filled at a pre-determined pressure maintained by a back-pressure regulator (BPR) with both valves closed and disconnected from the system. Multiple isolated reactors can be placed in parallel under the determined reaction conditions until the desired reaction time has been achieved. Thus, each of the sealed micro-tubing reactors behaves like a high-pressure reactor. The SFMT reactor may possess a poorer mixing efficiency compared to stirring in batch reactors, or circular flow patterns from Taylor flow, but the excellent gas-liquid interfacial contact in the SFMT reactor can still result in good reaction efficiency. An added advantage is that, compared to continuous flow techniques, reaction screening with SFMT reactors can save time by parallel screening. The residence time is not limited by the reactor size and can be held for as long as necessary.

SFMT reactors provide a simple and effective platform for photosynthesis using gaseous reagents, especially at high pressures. Wu and co-workers were able to achieve the photocatalytic vinylation of fluorinated aryl bromides 282 using acetylene gas in moderate to high yields assisted by the
SFMT reactors (Scheme 59). In stark contrast, only trace amounts (< 5%) of vinylated products were detected in conventional batch reactors. The gases acetylene escaped the reaction solution at 60 °C in conventional batch reactors, while the SFMT reactor can keep the gas within the reaction solution even at high temperatures.

Wu and co-workers developed a divergent synthesis of 1,2-diarylethanones, 1,4-diarylbutanes, 2,3-diarylbutanes, and ethyl-arenes from a common aryl halide and ethylene through the synergistic combination of photoredox and Ni catalysis (Scheme 60). By varying the photocatalysts, nickel catalysts, reaction temperature, and ethylene pressure, various products such as reductive Heck products, one or two-ethylene insertion linear products, and two-ethylene insertion branched products could be obtained in good yields and high selectivity. This is the first report of divergent synthesis achieved by different catalytic pathways accessed through modulating the oxidation state of organometallic catalysts by the use of photoredox catalysts. Notably, the SFMT reactor was employed to keep ethylene in the reaction solution at high temperature and high pressure to deliver the two-ethylene insertion branched products.

The proposed catalytic cycle (Scheme 61, catalytic cycle A) for the synthesis of 1,2-diarylethane begins with formation of Ru(bpy)₃Cl₂-promoted Ni I. Subsequent oxidative addition of aryl halides to the Ni I intermediate forms an aryl-Ni III species. Migratory insertion of ethylene into aryl-Ni III followed by aryl transfer by a transmetallation between an Ni III species and the aryl-Ni III species forms an aryl alkyl Ni III species. Then a rapid reductive elimination would yield product and regenerate the active Ni I catalyst. The cationic NiIII species generated during the transmetallation process would be reduced by a Ru-based photoredox cycle to regenerate NiI. For the synthesis of 1,4-diarylbutanes (catalytic cycle B), the catalytic cycle starts with the oxidative addition of aryl halides to Ni 0 reduced by Ir-based photoredox cycle to form the Ni II species. Ethylene migratory insertion into leads to the formation of alkyl NiII complex. Transmetallation between two molecules of complex followed by reductive elimination produces and Ni 0. For the catalytic generation of 2,3-diarylbutanes (catalytic cycle C), the alkyl Ni II complex undergoes a β-hydride elimination triggered by high temperature to yield a Ni hydride species. Insertion of into styrene forms a benzyl Ni intermediate. Subsequent transmetallation between two molecules of followed by reductive elimination affords and regenerates Ni 0. The reductive Heck-type product could be obtained by quenching the intermediate with water.

### 5. Gaseous Alkanes

The use of gaseous alkanes as inexpensive feedstocks in synthetic transformations is very attractive from an economic standpoint. However, it is often challenging to activate these gaseous alkanes with their strong C(sp³)–H bonds, poor...
solubility in most solvents and high ionization energies. Furthermore, conventional methods to activate such inert C–H bonds suffer from the lack of chemoselectivity as harsh conditions are often needed, and could also lead to functionalization of solvents or over-functionalization of products. To overcome the limitations, elegant transition-metal-catalyzed activation of gaseous alkanes has been reported in the past decades. More recently, photoredox catalysis as a new activation mode has been realized. This technology avoids the need for high reaction temperature and directing groups required for the transition-metal-catalyzed C(sp)–H activation.

In 2018, Zuo and co-workers developed a selective, photoredox-promoted functionalization of gaseous alkanes process through a synergistic merger of ligand-to-metal charge transfer (LMCT) and HAT catalysis. By utilizing inexpensive cerium salts and alcohols as catalysts, the amination of gaseous alkanes (Scheme 62), and the alkylation and arylation of methane and ethane were realized (Scheme 63). Notably, the amination of methane (5000 kPa) and ethane (101 kPa) achieved high turnover numbers of 2900 and 9700 respectively. To demonstrate the robustness of this method, the large-scale application of the reaction was investigated using continuous flow micro-tubing reactors. Remarkably, at a pressure of 1500 kPa for ethane and a flow rate of 0.75 mL min⁻¹ for the liquid solution stream, the amination product was obtained in 90% yield with a residence time of only 6 min. This translates to a production throughput of 2 mmol h⁻¹. On the other hand, the amination of methane only afforded 15% yield due to the pressure limit of commercial flow reactors (1800 kPa). In addition, methylation and ethylation of electron-deficient alkenes and arylation of methane and ethane were also achieved with moderate to high efficiency (Scheme 63).

The authors proposed that simple alcohols and a Ce IV salt would react in situ to generate a Ce IV-alkoxy complex which would undergo photo-induced LMCT to generate a high energy electrophilic alkoxy radical and a reduced Ce III species (Scheme 64). The alkoxy radical would then abstract a hydrogen atom from the alkane, generating an alkyl radical which readily couples with di-tert-butyl azodicarboxylate (DBAD, 307) to form a new C–N bond and an N-centered radical. This radical would undergo single-electron reduction by the reduced cerium catalyst to regenerate Ce IV and also deliver the anionic form of the desired product. The desired product is obtained by final protonation.

Energy-electrophilic alkoxy radical and a reduced Ce III species (Scheme 64). The alkoxy radical would then abstract a hydrogen atom from the alkane, generating an alkyl radical which readily couples with di-tert-butyl azodicarboxylate (DBAD, 307) to form a new C–N bond and an N-centered radical. This radical would undergo single-electron reduction by the reduced cerium catalyst to regenerate Ce IV and also deliver the anionic form of the desired product. The desired product is obtained by final protonation.

The Wu group has reported C–H alkylation and allylation reactions using hydrochloric acid (BDE = 103 kcal mol⁻¹) as the HAT catalyst precursor together with Mes-Acr⁺ photocatalyst assisted by SFMT reactors (Scheme 65). The reported reactions are irreproducible or even inaccessible in conventional batch reactors. Synthetically challenging ethane...
alkylation (Scheme 65) was achieved with strong Michael acceptors, giving alkylated products with moderate to good yields and good functional group tolerance. However, reactions with other alkenes bearing unsaturated ketones or esters were sluggish, presumably due to the fact that the radical adducts formed were not sufficiently strong oxidizing agents to turn over the photoredox catalytic cycle. Gram-scale synthesis using a 27.2 mL tubing reactor was achieved. This example showcased the SFMT reactor for efficient photo synthesis using gas at high pressures and temperatures. It is worth noting that this reaction is probably not suitable for continuous-flow synthesis, as it requires more than 24 h to go to completion.

In the proposed mechanism, the excited state of Mes-Acr\(^+\) could readily oxidize the chloride anion to a chlorine radical, which subsequently abstracts a hydrogen atom from the C(sp\(^3\))-H partner (ethane) to produce an ethyl radical \(\text{311}\). Nucleophilic addition of radical \(\text{311}\) to benzylidenemalononitrile would lead to radical adduct \(\text{312}\) which is then reduced by Mes-Acr radical to produce anion \(\text{313}\), which after protonation yields the ethylation product.

More recently, No/lc and co-workers reported a C(sp\(^3\))/C0\(\text{H}\) functionalization of methane, ethane, propane, and isobutane in continuous-flow reactors under 365 nm LED irradiation using decatungstate as the photocatalyst (Scheme 67).\(^{[85]}\)

Through the use of a photo hydrogen atom transfer catalyst TBA-DT in flow, they were able to avoid the use of harsh reaction conditions and obtain hydroalkylated adducts at room temperature in good isolated yields and high selectivity. Significantly, the utilization of flow chemistry provided key advantages over conventional batch reactions that facilitated the successful execution of this transformation. Homogenous irradiation of an entire reaction mixture can be achieved in microflow reactors and allows for efficient generation of alkyl radicals. The low solubility of these gaseous alkanes in organic solvents hampered the development of reactions in batch conditions. Here, the back-pressure regulators in flow allowed high pressure reaction conditions and can force the gaseous alkanes to dissolve in the organic solvents, allowing efficient mix of reagents. Moreover, high pressure reactions in flow are safe and scalable. The authors demonstrated that isobutane (316a–316d), propane (316e–316h) and ethane (316i–316l) can be efficiently functionalized by a variety of activated olefins including benzylidene-malononitriles, methyl trans-\(\alpha\)-cyanocinnamate, triethyl ethylene-tricarboxylate, \(N\)-phenylmaleimide and 3-methylene-2-norbornanone with high selectivity. Interestingly, high endo selectivity was observed in the formation of 316d and 316h, consistent with a hydrogen back-
donation from the less hindered side of the norbornane moiety. The reaction with methane shows narrower scope and forced conditions are required. Solvent functionalization was observed but could be circumvented by using deuterated acetonitrile.

A tentative mechanism was proposed (Scheme 68). Photocexcitation of decatungstate produces a singlet excited state of decatungstate, which rapidly relaxes to the actual reactive state wO. A subsequent hydrogen atom transfer from gaseous alkane to wO forms the carbon-centered radical species 317. A radical addition of 317 to olefin 315 yields the adduct 318, which abstracts a hydrogen atom from $\text{H}^+\left[\text{W}_{10}\text{O}_{32}\right]^{4-}$ to form the product 316.

Scheme 68. Proposed mechanism of the decatungstate catalyzed C-(sp$^3$)-H functionalizations of light hydrocarbons.

7. Summary and Outlook

This review has summarized the use of small gas molecules, including carbon monoxide, carbon dioxide, gaseous alkenes, alkynes and alkanes, as feedstocks in the synthesis of fine chemicals under light-irradiation. These gaseous reagents are abundant, inexpensive and readily available carbon-based feedstocks, and are therefore ideal feedstocks in organic synthesis. However, their utilization in synthesis of fine chemicals has been hampered by the difficulty in their handling and is mainly limited to transition-metal catalysis. Their utilization in photochemistry has recently witnessed a remarkable increase of attention in academia. We attribute this to two important reasons. The first reason is the emergence of photo-organic synthesis, which provides various novel activation modes to access to previously inaccessible transformations. The second reason is the advances of engineered reaction equipment for simple, convenient, and efficient usage of these gaseous reagents under light-irradiation.

The merits of continuous-flow reactors, including the improved light penetration, the excellent surface to volume ratio, significantly enhanced safety, and easy scaling up, offer a great degree of operational flexibility in the handling of gaseous reagents. The development of stop-flow micro-tubing (SFMT) reactors which combines the advantages of both batch reactors and flow reactors, provides an excellent platform for light-promoted reaction screening using gaseous reagents, especially at high temperatures and pressures.

Carbon monoxide has been used in UV light-promoted reactions since the 1930s. UV light was used to homolytically cleave chemical bonds to generate radicals which were trapped by CO to form an acyl radical, a key intermediate in carbylation reactions. Recent development in visible-light-promoted carbylation reactions involved the use of inexpensive organic dyes, such as fluorescein, eosin Y and 4CzIPN. Photosensitizers based on iridium and ruthenium have also demonstrated excellent catalytic activities. While the carbylation reactions have demonstrated a sustainable and efficient strategy for construction of a wide range of carbylated products, the high pressure required for the reactions continues to limit the applicability of the strategy. Recent development of transition-metal catalysis assisted by light activation provides a promising direction for incorporation of CO into complex molecules under mild conditions.

Transition-metal catalysis was widely used to overcome the thermodynamic stability and kinetic inertness of carbon dioxide. Photoredox catalytic systems recently have proven to be more environmentally friendly as they can bypass the need for harsh conditions or stoichiometric amounts of metallic reagents. The methods presented in this review are novel activation modes and are complementary to existing approaches. However, there is an absence of stereoselective transformations for the synthesis of enantioenriched acids and derivatives. Future efforts may be aimed at enantioselective transformations with CO$_2$. A promising strategy is to merge photoredox catalysis with transition-metal catalysts equipped with chiral ligands.

Gaseous alkenes and alkynes are important industrial feedstocks in the synthesis of value-added chemicals. The stop-flow micro-tubing reactor is developed to enable more efficient, safe and convenient screening of photo-mediated transformations using these flammable gases. The gases reagents can be sealed in the reaction solutions even at high temperature under photo-irradiation to achieve high efficiency. However, the lack of scaffold diversity of these gaseous alkenes and alkynes may limit their broad usage in the synthesis of complex molecular scaffolds.

Employing gaseous alkenes as synthetic feedstock is attractive from an economic standpoint, but it is usually difficult to activate these gaseous alkenes due to their strong C(sp$^3$)--H bonds. The harsh conditions required to activate such bonds would often lead to solvent functionalization or over-functionalization of products. Several recent reports on photo-mediated hydrogen atom transfer catalysis have led to promising results enabling functionalization of these inactive alkenes under mild conditions. The transformations can be potentially scaled up by taking advantage of the continuous-flow reactors. However, the efficient functionalization of methane, which is the most appealing but challenging substrate, is still difficult to achieve.

The synthesis of value-added fine chemicals using gaseous reagents under photo conditions has been considered difficult-to-handle in the past. However, as shown in this review,
these reagents can now be safely and efficiently processed through photo-irradiated transformations to enable novel opportunities and applications in organic synthesis. The rapid development of photoredox and hydrogen atom transfer catalysis, together with the advanced micro-tubing reactors, offers great momentum to develop green and sustainable synthetic protocols for photosynthesis using gaseous reagents. The innovation will continue to grow, impact synthetic chemistry, and bring about great benefits for both pharmaceutical and chemical industries in the future.

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Conflict of interest

The authors declare no conflict of interest.


Light-Promoted Organic Transformations Utilizing Carbon-Based Gas Molecules as Feedstocks

Photo-mediated fixation of carbon-based gas molecules: C–C bond formation utilizing carbon-based small gas molecules, including CO, CO₂, acetylene, ethylene, ethane and methane, as feedstocks is attractive but challenging. The emergence of photoredox chemistry and the development of engineered flow technology have provided vast opportunities for gas conversions, which has gained tremendous momentum over the past decade.