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## Carbon Dioxide as a Carbon Source in Organic Transformation: Carbon-Carbon Bond Forming Reactions by Transition-Metal Catalysts

### Yasushi Tsuji\* and Tetsuaki Fujihara

Recent carbon–carbon bond forming reactions of carbon dioxide with alkenes, alkynes, dienes, aryl zinc compounds, aryl boronic esters, aryl halides, arenes having acidic C–H bonds are reviewed in which transition-metal catalysts play an important role.

#### 10 1. Introduction

Carbon dioxide (CO<sub>2</sub>) is an ideal carbon source owing to its abundance, low cost, nontoxicity, and good potential as a renewable source.1 However, it is not easy to activate such a thermodynamically and kinetically stable material. Therefore, 15 catalysts must play an important role to develop useful transformations utilizing this attractive and environmentally friendly raw material. In this review article, recent progress on utilization of CO2 in carbon-carbon bond forming reactions is surveyed. Such survey will offer basic idea to employ CO2 as an 20 attractive carbon source in organic transformations. There have been several excellent reviews on a similar subject.<sup>2</sup> On the other hand, carbon-oxygen bond and carbon-hydrogen bond forming reactions of CO2 are also important for syntheses of organic carbonates (including CO<sub>2</sub>/epoxide copolymerization), formic 25 acid derivatives, and methanol. However, these reactions are not covered in the present article, since several useful reviews are available.3d-g, 4e-h

# 2. Stoichiometric reaction relevant to catalysis with $CO_2$

#### 30 2.1 Transition metal complexes bearing CO2

In order to develop efficient catalytic reactions, fundamental behaviours of  $CO_2$  in the presence of a transition metal complex must be understood. As the first isolated and characterized metal  $CO_2$  complex, Aresta et al. prepared  $[Ni(CO_2)(PCy_3)_2]$  ( $Cy = \frac{1}{2}$  cyclohexyl) complex in 1975 by treating  $[Ni(PCy_3)_3]$  or  $[\{Ni(PCy_3)_2\}_2N_2]$  with  $CO_2$  in toluene. (Scheme 1). The  $CO_2$  coordinates to the electron rich Ni(0) centre as  $\eta^2$  form. This finding suggested that  $CO_2$  could be activated on a metal centre. More relevant to catalytic reactions of  $CO_2$  Hoberg et al. found

**Scheme 1** Synthesis of Ni(0)  $\eta^2$ -CO<sub>2</sub> complex.

$$[Ni(cdt)] + CO_2 + H_3CC = CCH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

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$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = CH_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = TT_1 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = TT_2 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = TT_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C = TT_3 + tmeda \xrightarrow{\text{rt, 48 h}} (tmeda) Ni$$

$$H_3C =$$

**Scheme 2** Stoichiometric reaction of Ni(0) complex in the presence of 2-butyne and ethylene: cdt = 1,5,9-cyclododecatriene, tmeda = N,N,N',N'-tetramethylethylenediamine, dbu = 1,8-diazabicyclo[5.4.0]undec-7-ene

that  $CO_2$  reacts with alkenes or alkynes in the presence of a stoichiometric amount of Ni(0) complexes to afford oxanickelacycles (Scheme 2).<sup>6</sup> Even the metal complexes were used in stoichiometric amounts, useful carbonyl compounds were obtained as products. Furthermore, a wide variety of complexes bearing  $CO_2$  in various coordination styles ( $\eta^1$ ,  $\eta^2$ ,  $\mu^2$ -  $\eta^2$ ,  $\mu^2$ -  $\eta^3$ , etc.) have been isolated and characterized to date.<sup>7</sup>

#### 2.2 Seminal catalytic reactions of CO<sub>2</sub>

A series of pioneering works for catalytic transformation of  $CO_2$  with unsaturated compounds were carried out by Inoue et al. in 1976–1979. Sa-c First, they reacted 1,3-dienes under a  $CO_2$  pressure of 50 atm in the presence of a catalytic amount of  $Pd(dppe)_2$  (dppe =  $Ph_2(CH_2)_2PPh_2$ ) at 120 °C (Scheme 3a). They found the lactone formed from  $CO_2$  in a complex mixture of various products. In the presence of  $Ni(cod)_2$ —dppb (cod = 1,5-

**Scheme** 3. Early catalytic reactions of hydrocarbon unsaturates with CO<sub>2</sub>. dppe = Ph<sub>2</sub>P(CH<sub>2</sub>)<sub>2</sub>PPh<sub>2</sub>, dppb = Ph<sub>2</sub>P(CH<sub>2</sub>)<sub>4</sub>PPh<sub>2</sub>, cod = 1,5-cyclooctadiene, dba = dibenzylideneacetone

cyclooctadiene, dppb = Ph<sub>2</sub>P(CH<sub>2</sub>)<sub>4</sub>PPh<sub>2</sub>) as a catalyst at 120 °C, two terminal alkynes reacted with CO<sub>2</sub> (50 atm) to afford lactones in only low yields (Scheme 3b). <sup>8b</sup> On the other hands, with the same catalyst system, internal alkynes provided the corresponding products in much higher yields (Scheme 3c). <sup>8c</sup> As for alkenes, Pd(0) catalyzed reaction of methylenecyclopropanes with CO<sub>2</sub> (40 atm) provided lactones successfully in good yields (Scheme 3c). <sup>8d</sup> In these reactions, yields of the products from CO<sub>2</sub> were not so satisfactory. However, these results clearly indicated that such catalytic reactions are very useful and promising.

# 3. Carbon-carbon bond forming reaction with CO<sub>2</sub>

Carbon-carbon bond forming reactions are quite important to develop a wide variety of organic transformations employing CO<sub>2</sub>. The most classical methods to realize the C-C bond forming reaction with CO<sub>2</sub> are the reaction with Grignard and organolithium reagents. Highly nucleophilic these reagents react with CO<sub>2</sub> without a catalyst. However, in these reactions, many useful functionalities such as ketones, esters, and nitriles are not tolerant. Thus, in order to develop transformations compatible with various functionalities, much less nucleophilic substrates must be employed. In such cases, catalysts are important and indispensable for successful C-C bond formation reactions with 25 CO<sub>2</sub>.

### 3.1 Reactions of Zinc compounds with CO<sub>2</sub>

In place of Grignard and lithium reagents, less nucleophilic

with cat or without cat,

Scheme 4 Carboxylation reactions of organozinc and -borane compounds using  $CO_2$ 

87%

73%

83%

82%

Boc

organozinc were utilized with CO<sub>2</sub> (Scheme 4a), since a wide variety of functionalities are compatible with organozinc compounds. Yorimitsu and Oshima reported that alkylzinc iodide–LiCl readily react with CO<sub>2</sub> in THF in the presence of Ni(acac)<sub>2</sub>–PCy<sub>3</sub> as a catalyst to afford the corresponding carboxylic acids. <sup>9a</sup> They found that the addition of LiCl was essential for the carboxylation. They claimed that the Ni catalyst is indispensable for alkylzinc compounds, while PhZnI–LiCl reacted with CO<sub>2</sub> in THF without the Ni catalyst to afford benzoic acid in 36% yield. Independently, Dong et al reported that carboxylation of arylzinc bromides with CO<sub>2</sub> proceeded in the presence of a similar Ni and Pd catalysts bearing PCy<sub>3</sub> as a ligand. <sup>9b</sup> They reported that no carboxylation occurred without the transition metal catalyst. On the other hand, Kondo et al found that transition metal catalysts are not necessary for these

reactions, but only 2.8 equiv of LiCl in DMF is sufficient for the reaction. 9c Activities of organozinc compounds seem to be varied considerably depending on reaction conditions, and some reactions really proceed without a transition metal catalyst.

#### 5 3.2 Reactions of organoboronic esters with CO<sub>2</sub>

Organoboron compounds are easily prepared by various methods and are highly useful substrates in a variety of organic synthesis. Iwasawa et al. reported that aryl- and alkenylboronic esters react with CO<sub>2</sub> in the presence of [Rh(OH)(cod)]<sub>2</sub> (3

**Scheme 5** Possible catalytic cycles and isolation of catalytic species in Scheme 4c

**Scheme 6** Formal reductive carboxylation of terminal alkenes via hydroboration

97%

**Table 1** Nickel-mediated 1,4-double-carboxylation of 1,3-diene

1,3-diene	product	yield (%)
	CO <sub>2</sub> H CO <sub>2</sub> H	68
F <sub>3</sub> C	CO <sub>2</sub> H CO <sub>2</sub> H	75
F	CO <sub>2</sub> H	73
	HO <sub>2</sub> C - CO <sub>2</sub> H	50

mol%), dppp (Ph<sub>2</sub>P(CH<sub>2</sub>)<sub>4</sub>PPh<sub>2</sub>, 7 mol%), and CsF (3 equiv) to provide the corresponding carboxylic acids in high yields (Scheme 4b). <sup>10a</sup> In the reaction, ester part of the boronic esters affected the carboxylation considerably. Actually, only 2,2-dimethyl-1,3-propanediolate ester (Scheme 4b) afforded the carboxylation products in high yields, while other propan1,3-diolate, ethylene glycolate, pinacolate, and catecholate gave almost no products. On the other hand, insertion of CO<sub>2</sub> into carbon-Cu bonds has been known. <sup>11</sup> Hou et al. reported that the same carboxylation of organoboronic esters are possible with N-20 heterocyclic carbene copper catalysts, in which the same effect of the ester moiety was observed (Scheme 4c). <sup>10b</sup> Importantly, most copper catalytic species (2 and 3 in Scheme 5) were isolated in

ray crystal structure analysis. The mechanism shown in Scheme 5 was supported by DFT calculations indicating insertion of  $CO_2$  into the Cu-Ar bond is the rate-determining step in which nucleophilic attack of the aryl moiety on  $CO_2$  affords the new C – C bond. <sup>10c</sup> Iwasawa et al. reported similar carboxylation by a

high yields (91% and 92%) and fully characterized including X-

copper catalyst. 10d
Sawamura et al. (Scheme 6a) 12a and Hou et al. (Scheme 6b) 12b independently found that the copper catalyzed carboxylation of alkylboranes were carried out via hydroboration of terminal alkenes with 9-borabicyclo[3.3.1]nonane (9-BBN-H). The overall process represents a useful reductive carboxylation of terminal

#### 3.3 Carboxylations of dienes and alkenes

35 alkenes with CO<sub>2</sub>.

Mori et al. developed carboxylation of 1,3-dienes and allenes with CO<sub>2</sub> via allylnickel intermediates.<sup>13</sup> When 1,3-dienes were reacted with 1 equiv of Ni(cod)<sub>2</sub> and 2 equiv of dbu under CO<sub>2</sub>,

**Table 2** Nickel-mediated arylative mono-carboxylation of 1,3-diene

1,3-diene 
$$\begin{array}{c} CO_2 \text{ (1 atm)} \\ Ni(cod)_2 \text{ (1 equiv)} \\ DBU \text{ (2 equiv)} \\ THF 0 °C, 4-7 \text{ h} \end{array} \begin{array}{c} \text{"ArZn" (5 equiv)} \\ 0 °C, 2 \text{ h} \\ \text{then } CH_2N_2 \end{array} \end{array} \begin{array}{c} \text{product} \\ \text{yield (%)} \\ \end{array}$$

1,3-diene  $\begin{array}{c} \text{"ArZn"} \end{array} \begin{array}{c} \text{product} \\ \text{Ph} \end{array} \begin{array}{c} \text{Vield (%)} \\ \text{Ph} \end{array} \begin{array}{c} \text{Ph} \end{array} \begin{array}{c} \text{CO}_2\text{Me} \\ \text{Ph} \end{array} \begin{array}{c} \text{Ph} \end{array} \begin{array}{c} \text{Ph} \\ \text{Ph} \end{array} \begin{array}{c} \text{CO}_2\text{Me} \\ \text{A1 : 59} \end{array} \begin{array}{c} \text{A5} \\ \text{CO}_2\text{Me} \end{array} \begin{array}{c} \text{A6} \\ \text{A7} \\ \text{A7} \end{array} \begin{array}{c} \text{CO}_2\text{Me} \\ \text{A1 : 59} \end{array} \begin{array}{c} \text{CO}_2\text{Me} \\ \text{A2 : 57} \end{array} \begin{array}{c} \text{A8} \\ \text{CO}_2\text{Me} \end{array} \begin{array}{c} \text{A8} \\ \text{A3 : 57} \end{array} \begin{array}{c} \text{CO}_2\text{Me} \\ \text{A3 : 57} \end{array} \begin{array}{c} \text{A3} \\ \text{CO}_2\text{Me} \end{array} \begin{array}{c} \text{A6} \\ \text{A7} \\ \text{A3 : 57} \end{array} \begin{array}{c} \text{CO}_2\text{Me} \\ \text{A3 : 57} \end{array} \begin{array}{c} \text{A6} \\ \text{A7} \\ \text{A7} \\ \text{A8} \\ \text{A8} \\ \text{A9} \\ \text{A9}$ 

and then further with 5 equiv of Me<sub>2</sub>Zn, 1,4-double-carboxylation products were afforded in good yields (Table 1). 13a Although the mechanism of the stoichiometric reaction was not clear, the two CO<sub>2</sub> molecules, but no Me moieties, were incorporated in the 5 products. In contrast, when aryl zinc reagents (ArZn) instead of Me<sub>2</sub>Zn were employed in the similar stoichiometric reaction, proceeded mono-carboxylation (Table Furthermore, by using bis-1,3-dienes as the substrates, the catalytic reaction provided ring-closing carboxylated products 10 regio- and stereoselectively in high yields (Scheme 7a). 13b The asymmetric carbonylative cyclization were realized under similar reaction conditions with chiral phosphine ligands (Scheme 7b). 13c Mori et al. also found Ni-catalyzed double-carboxylation of  $CO_2^{13d}$ with trimethylsilylallene Ni-promoted 15 carboxylation/cyclization cascade reaction of allenyl aldehydes.13e In the former case, the silyl moieties were indispensable for the double carboxylation.

Iwasawa et al. reported hydrocarboxylation of allenes with  $CO_2$  catalyzed by Pd pincer catalyst (Table 3). <sup>14a</sup> AlEt<sub>3</sub> was <sup>20</sup> employed as a reducing agent and a hydrido source after the  $\beta$ -hydride elimination of the resulting ethyl palladium intermediate (Scheme 8). With a similar Pd catalyst system, one-to-one coupling of 1,3-dienes with  $CO_2$  occurred (Table 4). <sup>14b</sup>

**Scheme 7** Nickel-catalyzed ring-closing carboxylation of bis-1,3-dienes

**Scheme 8** Catalytic cycle of Pd-catalyzed hydrocarboxylation of allenes

**Table 3** Pd-catalyzed hydrocarboxylation of allenes

allene	yield of product
$R^{1}$ =Me, $R^{2}$ =Me	95%
$R^{1}$ =Me, $R^{2}$ =Ph	73%
$R^{1}$ =Me, $R^{2}$ = -(CH <sub>2</sub> ) <sub>2</sub> CH=CMe <sub>2</sub>	85%

$$X = -(CH_2)_2 - 72\%$$
 $= C(OCH_2)_2$ 
 $= C=O$ 
 $= CHPh$ 
 $X = -(CH_2)_2$ 
 $= C=O$ 
 $= CHPh$ 
 $= CO_2H$ 
 $= CO_2H$ 

cat: 
$$\begin{array}{c} \text{OTf} \\ \text{Ph}_2 \text{P} - \text{Pd} - \text{PPh}_2 \\ \text{Si} \\ \text{Me} \end{array}$$

**Table 4** Pd-catalyzed hydrocarboxylation of 1,3-diene: one-to-one coupling with CO<sub>2</sub>

1,3-dienes	products	yields (%)
Me Ph	Me COOMe	79
Ph	COOMe	92
Boc N	Boc N	79
	O COOH	81

<sup>&</sup>lt;sup>a</sup> The same Pd complex in Table 3 was used as a catalyst

Scheme 9 Ni-catalyzed reductive carboxylation of styrenes using  $CO_2$ 

(a) 
$$()_n$$
 Et  $()_n$  Et

(b) 
$$R^1$$
  $R^2$   $R^2$   $R^1$   $R^2$   $R^2$   $R^1$   $R^2$   $R^2$   $R^2$   $R^2$   $R^1$   $R^2$   $R^2$ 

**Scheme 10** Carboxylation of  $\alpha$ , $\omega$ -divines with CO<sub>2</sub>

As for alkenes, Rovis et al. found reductive carboxylation of styrene derivatives with  $Et_2Zn$  and  $CO_2$  (Scheme 9). They proposed an insertion on Ni-H into carbon-carbon double bonds of styrenes followed by insertion of  $CO_2$ , rather than the Hoberg's metallacycle shown in Scheme 2.

#### 3.4 Carboxylation of alkynes

Pioneering works for catalytic carboxylation of alkynes were carried out by Inoue et al. as shown in Scheme 3b,c. <sup>8b,c</sup> Tsuda and Saegusa found that the reactions of  $\alpha,\omega$ -diynes under pressurized CO<sub>2</sub> (50 atm) in the presence of Ni-phosphane catalyst proceeded smoothly and afforded bicyclic  $\alpha$ -pyrones in good yields (Scheme 10a). <sup>16a</sup> In this transformation, IPr ligand in place of phosphanes is much more effective and provided the similar carboxylated products in high yields under 1 atm of CO<sub>2</sub> (Scheme 10b). <sup>16b</sup>

On the other hand, Yamamoto et al. reported that terminal alkynes reacted with 1 equiv of Ni(cod)<sub>2</sub> and 2 equiv of dbu

Scheme 11 Carboxylation of terminal and TMS acetylenes

under CO<sub>2</sub> (1 atm) at 0 °C in a highly regio- and chemoselective manner (Scheme 11a).  $^{17a}$  Similar carboxylation reaction with 1 equiv of Ni(cod)<sub>2</sub> was also carried out using bidentate amidine ligands.  $^{17b}$  When alkyl or aryl zinc reagent was added into the 5 Yamaoto's system, alkylative and arylative carboxylation of alkynes proceeded (Table 5).  $^{17c,e}$  Interestingly, BuZn reagent afforded the butylated product (entry 3), while EtZn gave hydrocarboxylated product via  $\beta$ -hydride elimination (entry 4). This alkylative carboxylation could be carried out catalytically when trimethylsilyl acetylenes were employed as substrates

**Table 5** Carboxylation of alkyne with  $Ni(cod)_2$  and organozinc reagents

BnO		Ni(co	(1 equiv) od) <sub>2</sub> (1 equiv) (2 equiv) 0 °C, 1 h	"RZn" 0 °C, 2 h	product
entry	"RZn"	equiv	yield (%) <sup>a</sup>	product	
1	PhZnCl	2	69	BnO	CO <sub>2</sub> H
2	PhCH <sub>2</sub> ZnCl	1.5	81	BnO	∕CO <sub>2</sub> H
3	BuZnl	3	68	BnO	CO₂H
4	Et <sub>2</sub> Zn	1.5	9 <sup>p</sup>	BnO	CO₂H

<sup>&</sup>lt;sup>a</sup> Isolated yield based on Ni(cod)<sub>2</sub>

**Table 6** Cu-catalyzed hydrocarboxylation of alkyne using CO<sub>2</sub> and hydrosilanes

<sup>a</sup> Conditions A: IMesCuF (1–2 mol%) in 1,4-dioxane at 100 °C for 12 h. Conditions B: CIIPrCuF (2.5 mol%) in hexane at 70 °C for 14 h. Conditions C: IPrCuF (2.5 mol%) in 1,4-dioxane at 100 °C for 14 h.

(Scheme 11b).17d

In order to utilize CO<sub>2</sub> in C-C bond formations, reducing reagents have been indispensable in the reactions. To date, highly reactive and pyrophoric reagents such as Et<sub>2</sub>Zn<sup>9,15,17a,d,e</sup> and <sup>15</sup> Et<sub>3</sub>Al<sup>14</sup> were employed in stoichiometric and catalytic reactions. We recently reported that hydrosilanes, more mild and easy-to-handle reducing reagents, can be used in Cu catalyzed hydrocarboxylation of alkynes. <sup>18a</sup> Various alkynes were hydrocarboxylated efficiently using HSi(OEt)<sub>3</sub> in good yields <sup>20</sup> (Table 6). In the reactions, HSi(OEt)<sub>3</sub> could be replaced with polymethylhydrosiloxane (PMHS), which is a cheap by-product of the silicon industry. Ma et al. found that the same reaction proceeded with Ni(cod)<sub>2</sub>-CsF as a catalyst and Et<sub>2</sub>Zn as a reducing reagent. <sup>18b</sup> Ma also reported methylative carboxylation <sup>25</sup> of homopropargylic alcohols. <sup>18c</sup>

Inoue et al. found that terminal alkynes reacted with

Table 7 Au-catalyzed carboxylation of aromatic C-H bonds

(a) 
$$R = H + CO_2 + n - C_6H_{13}Br$$
  
 $5 \text{ mmol} \quad 1 \text{ atm} \quad 10 \text{ mmol}$ 

Cul (0.2 mmol)

 $K_2CO_3 (30 \text{ mmol})$ 
in DMA
 $100 \,^{\circ}C$ , 4h
 $V_2CO_3 (30 \text{ mmol})$ 
 $V_3CO_3 (30 \text{ mmol})$ 
 $V_3CO_$ 

**Scheme 12** Carboxylation of terminal alkynes with CO<sub>2</sub> with and without bromoalkanes

bromoalkanes under  $CO_2$  in the presence of a catalytic amount of  $CuI.^{19a}$  Insertion of  $CO_2$  into the acetylenic C-H bonds to give alkyl 2-alkynoates at 100 °C in good yields (Scheme 12a). CuBr, AgI, or AgNO<sub>3</sub> can be utilized as the catalyst successfully. This 5 reaction was applied to one-pot synthesis of arylnaphthalene lactones. <sup>19b</sup> Recently, IPrCuCl was found to be a more efficient

and the reaction proceeded even at lower temperature (60 °C). <sup>19c</sup> Furthermore, insertion of  $CO_2$  into the C-H bond proceeded without bromoalkanes in the presence of a  $Cu^{20a,b}$  (Scheme 12b) <sup>10</sup> or  $Ag^{20c}$  catalyst. The same reactions could be carried out without the transition metal catalysts at 120–160 °C, but 2 equiv of  $Cs_2CO_3$  was indispensable. <sup>20d</sup>

#### 3.5 Carboxylation of aryl halides

15 As mentioned earlier, the carboxylation reactions of arylzinc (section 2.1) and arylboronic esters (section 2.2) with CO<sub>2</sub> have been studied intensively, since in these reactions various functionalities that are compatible unlike with Grignard reagents. However, these zinc and boron compounds must be prepared from the corresponding aryl halides prior to the catalytic reactions. Thus, the direct carboxylation of the parent aryl halides is most desirable as this is a more straightforward transformation.

The catalytic carboxylation of aryl halides employing CO<sub>2</sub> was first developed as *electrochemical* reactions in the presence of <sup>25</sup> nickel<sup>21a-c</sup> and palladium catalysts (Scheme 13a). <sup>21d,e</sup> Unfortunately, these were not efficient synthetic methods, and the scope of possible substrates was quite restricted. Later, the non-electrochemical carboxylation of aryl bromides and chlorides using CO<sub>2</sub> was carried out in the presence of *stoichiometric* amounts of Ni(0) complexes and 2,2'-bipyridine (bpy) (Scheme 13b). <sup>22a</sup> Recently, the carboxylation of aryl bromides employing CO<sub>2</sub> was performed *catalytically* using a palladium complex as the catalyst. <sup>22b</sup>

**Scheme 13** Carboxylation of aryl halides with CO<sub>2</sub>

Scheme 14 Stoichiometric reactions relevant to the catalytic cycle

#### 3.6 Carboxylation of arenes having acidic C-H bonds

The Kolbe-Schmitt salicylic acid synthesis from sodium phenolate and CO<sub>2</sub> is an early example of carboxylation of C-H bonds. Such direct carboxylation of C-H bonds must be more 5 sustainable transformations. Recently, Nolan et al. reported that (IPr)AuOH can perform the activation of C-H bonds of electrondeficient arenes. 23a They found that the same Au(I) complex is very active for carboxylation of aromatic acidic C-H bonds (pKa values < 30) with CO<sub>2</sub> (Table 7). <sup>23b</sup> The possible catalytic cycle 10 for the present C-H carboxylation (Scheme 14) is similar to the mechanism postulated for Cu(I) catalyzed carboxylation of organoboronic esters (Scheme 5). The authors confirmed the mechanism by stoichiometric reactions (Scheme Protonolysis of (IPr)AuOH by 1 equiv of oxazole gave the 15 intermediate 4, which was isolated in 93% yield (step 1). Reaction of 4 by saturating the solution with CO2 at -100 °C afforded carboxylate complex 5 in 86% isolated yield (step 2). Finally, (IPr)AuOH was regenerated upon metathesis of 5 with 1 equiv of KOH with precipitation of potassium oxazole-2-20 carboxylate (step 3). Hou et al. 23c and Nolan and coworkers 23d independently reported that Cu catalysts are active as catalysts in the carboxylation reaction of the acidic C-H bonds (Table 8). 23c Hou isolated catalytic intermediates (arylcopper and copper carboxylate) in a stoichiometric reaction and determined the 25 structures by X-ray crystallographic analysis. 23c However, these C-H carboxylations could proceed with only a base (Cs2CO3 is most suitable) at 125 °C. 23e Iwasawa et al. found the first C-H carboxylation with CO<sub>2</sub> via chelation-assisted C-H activation. 2-arylpyridines and 1-arylpyrazoles 30 carboxylated products in the presence of a Rh catalyst and a stoichiometric amount of AlMe<sub>2</sub>(OMe) (Table 9).

Table 8 Cu-catalyzed carboxylation of aromatic C-H bonds

$$\begin{array}{c} \text{Ar-H} + \text{CO}_2 \\ \hline & \text{1 atm} \end{array} \begin{array}{c} \text{Cu(IPr)CI (5 mol\%)} \\ \text{KOfBu (1.1 equiv)} \end{array} \begin{array}{c} \text{C}_6 \text{H}_{13} \\ \hline \text{DMF, 80 °C} \end{array} \begin{array}{c} \text{Ar-CO}_2 \text{C}_6 \text{H}_{13} \\ \hline \text{yield} \end{array} \\ \\ \text{Me} \\ \hline & \text{87\%} \end{array} \begin{array}{c} \text{N} \\ \text{85\%} \end{array} \begin{array}{c} \text{C}_6 \text{H}_{13} \\ \hline \text{DMF, 80 °C} \end{array} \begin{array}{c} \text{Ar-CO}_2 \text{C}_6 \text{H}_{13} \\ \hline \text{yield} \end{array} \\ \\ \text{Br} \\ \hline & \text{CO}_2 \text{C}_6 \text{H}_{13} \\ \hline & \text{62\%} \end{array} \begin{array}{c} \text{N} \\ \text{CO}_2 \text{C}_6 \text{H}_{13} \\ \hline & \text{50\%} \end{array} \\ \\ \text{NC} \\ \hline \\ \text{NC} \\ \hline \\ \text{86\%} \end{array} \begin{array}{c} \text{N} \\ \text{CO}_2 \text{C}_6 \text{H}_{13} \\ \hline \\ \text{38\%} \end{array}$$

**Table 9** Rh-catalyzed direct carboxylation of arenes via chelation-assisted C–H bond activation

CO<sub>2</sub> (1 atm)

#### 35 Conclusions

Carbon dioxide ( $CO_2$ ) can be utilized in carbon–carbon bond forming reactions with the aid of transition-metal catalysts. Organozine compounds usually do not react with  $CO_2$  without a transition-metal catalyst. and Ni and Pd catalysts are active to

afford the corresponding carboxylic acids. Organoboronic esters are also useful substrates which are reacted with CO<sub>2</sub> to provide the carboxylic acids in the presence of Rh and Cu catalysts. In the Cu-catalyzed reaction, aryl-Cu or ArCOOCu intermediates were isolated and fully characterized. Dienes, alkenes, and alkynes were efficiently reacted with CO<sub>2</sub> in the presence of Ni, Pd, and Cu catalysts. Usually, highly reactive and pyrophoric reducing reagents such as Et<sub>2</sub>Zn and Et<sub>3</sub>Al are indispensable in the reactions. Recently, easy-to-handle hydrosilanes such as HSi(OEt)<sub>3</sub> and polymethylhydrosiloxane (PMHS) can be employed with a Cu catalyst. Carboxylation of aryl halides and arenes having acidic C–H bonds are realized.

#### Notes and references

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