Carboxylic acid derivatives via catalytic carboxylation of unsaturated hydrocarbons: whether the nature of a reductant may determine the mechanism of CO2 incorporation?

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Abstract. Application of CO\textsubscript{2} as a renewable feedstock and C\textsubscript{1} building block for production of commodity and fine chemicals is a highly challenging but obvious industry-relevant task. Of particular interest is the catalytic coupling of CO\textsubscript{2} with inexpensive unsaturated hydrocarbons (olefins, dienes, styrene, alkynes), providing direct access to carboxylic acids and their derivatives. Although not brand new for the scientific community, it is still a complete challenge, as no truly effective catalytic system has been reported to date. In this Perspective, we discuss the available experimental, theoretical and mechanistic data for such homogeneously catalyzed carboxylation processes. A special focus is placed on the understanding of the key elementary steps and of some thermodynamic and kinetic constraints.

1. Introduction

In a global context aiming at a more sustainable development, catalytic activation of CO\textsubscript{2} mediated by transition metal complexes with complete integration of the whole CO\textsubscript{2} molecules into...
the final products has become a highly topical research area. Many international leading research
groups have been working on such catalytic CO₂ conversion into valuable chemicals, or on general
methods for fixation of CO₂. Besides two very recent major contributions,¹ many reviews have
been released on topics relevant to catalytic CO₂ transformation and its utilization as C₁ building
block.² Reactions of particular interest include catalytic reduction (hydrogenation, hydrosilylation)
of CO₂,³ synthesis of carboxylic acids,⁴ of acrylic acid derivatives from olefins/alkynes,⁵ and of
polycarbonates from epoxides;⁶ also, much work has been devoted to computational aspects of
some catalytic reactions implies CO₂,⁷ the stoichiometric chemistry of CO₂,⁸ and of other
processes.⁹

Among the above processes, formation of C–C bonds by carboxylation of readily available
unsaturated hydrocarbons (alkenes, dienes, styrenics, alkynes) offers a direct access to valuable
saturated and unsaturated carboxylic acids and their derivatives.¹⁰ Modern industrial processes for
production of carboxylic acids do not involve reactions with CO₂. Only the Monsanto/Cativa
processes implicate indirectly CO₂ as a component for the water-gas shift reaction. Otherwise,
carboxylic acids and their derivatives are traditionally synthesized by multi-step oxidation protocols
starting from petrochemicals (propylene, butadiene, toluene, xylenes, naphthalene) or by
carbonylation of alkenes with CO/H₂O-alcohols (e.g. Koch acids, Reppe chemistry,
hydroesterifications (Lucite process)).¹¹ The direct catalytic carboxylation of unsaturated
hydrocarbons bestows, in principle, obvious benefits in terms of energy and raw materials with
respect to regular multistep oxidation protocols. This is fundamentally different from other
activation processes implying severe reduction of CO₂ (sometimes complete when hydrocarbons-
type products are targeted), which are highly energy-intensive (reduction of C=O bonds) and
inherently poorly atom-efficient.

This Perspective article discusses a few existing strategies for the direct coupling of
unsaturated hydrocarbons (alkenes, alkynes) with CO₂ towards carboxylic acids.; a special attention
is drawn to potential catalytic processes enabling the atom-economic reductive carboxylation of
alkenes (Scheme 1). The article is focused on unraveling the essential problems of catalytic coupling of CO₂ with unsaturated hydrocarbons and it analyzes the available information about the nature of key intermediates and side-products of the operative (actual and putative) catalytic mechanisms.

Scheme 1. Catalytic reductive carboxylation of unsaturated hydrocarbons into carboxylic acids and esters.

2. State-of-the-Art and Main Features of Carboxylation of Unsaturated Hydrocarbons

2.1. Some Thermodynamic Considerations

The high thermodynamic stability of CO₂ makes unfavorable its conversion to other molecules,⁷ᵃ,¹² therefore, the corresponding processes require significant energy input in a form of heat,¹³ or, more common, as highly energetic co-reagents. For instance, some non-extensive thermochemical calculations on relevant reactions involving CO₂ (Table 1) demonstrated that formation of the corresponding acids from CO₂ and alkane/alkene is generally an endothermic process (reactions (2) and (3), respectively). Yet, although a significant thermodynamic constraint is also characteristic for the reduction of CO₂ with H₂ into formic acid (reaction (1)), the related reaction conducted in the presence of ethylene, to deliver eventually propionic acid, is calculated to be quite thermodynamically favorable (reaction (7)). Because a saturated acid forms in the latter coupling reaction, the corresponding energy is virtually driven by the exergonic ethylene hydrogenation reaction (4). Similarly, highly negative enthalpies and Gibbs energies are estimated.
for the reactions implying strong reductants such as hydrosilanes or hydroboranes (reactions (8)–(11)), which provide the corresponding carboxylic acids esters.

**Table 1.** Calculated Enthalpies and Free Energies (in kcal·mol⁻¹) for some basic reactions implying CO₂ and ethylene.ᵃ

<table>
<thead>
<tr>
<th>Reaction</th>
<th>ΔH</th>
<th>ΔG</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. CO₂ + H₂ → HCOOH</td>
<td>+7.4</td>
<td>(+5.1) b</td>
</tr>
<tr>
<td>2. CO₂ + CH₃CH₃ → CH₃CH₂COOH</td>
<td>+9.7</td>
<td></td>
</tr>
<tr>
<td>3. CO₂ + CH₂=CH₂ → CH₂=CHCOOH</td>
<td>+5.8</td>
<td>(+5.7) b</td>
</tr>
<tr>
<td>4. CH₂=CH₂ + H₂ → CH₃CH₃</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. CH₂=CH₂ + Et₃SiH → Et₄Si</td>
<td>−32.6</td>
<td>−15.3</td>
</tr>
<tr>
<td>6. CH₂=CH₂ + HBpin → EtBpin</td>
<td>−35.6</td>
<td>−19.3</td>
</tr>
<tr>
<td>7. CO₂ + CH₂=CH₂ + H₂ → CH₃CH₂COOH</td>
<td>−26.7</td>
<td>−7.5</td>
</tr>
<tr>
<td>8. CO₂ + Et₃SiH → HCOOSiEt₃</td>
<td>−16.0</td>
<td>−11.7</td>
</tr>
<tr>
<td>9. CO₂ + HBpin → HCOOBpin</td>
<td>−14.8</td>
<td>−5.0</td>
</tr>
<tr>
<td>10. CO₂ + CH₂=CH₂ + Et₃SiH → CH₃CH₂COOSiEt₃</td>
<td>−49.9</td>
<td>−23.4</td>
</tr>
<tr>
<td>11. CO₂ + CH₂=CH₂ + HBpin → CH₃CH₂COOBpin</td>
<td>−48.3</td>
<td>−21.7</td>
</tr>
</tbody>
</table>

ᵃ M06/6-31(d,p) in gas phase. b taken from ref. 12b (G3B3 in gas phase).

The above thermochemical estimations unequivocally illustrate the feasibility of alkene carboxylation reactions towards the corresponding carboxylic acids or some of their derivatives, provided they are carried out in the presence of an appropriate reducing agent (H₂ or hydrosilane/hydroborane, respectively). Thus, a quest for elaboration of new systems competent for catalyzing these processes has been ongoing.

### 2.2. Direct carboxylation of unsaturated hydrocarbons (alkenes, alkynes, dienes) to carboxylic acids. Role of the (reverse) water-gas shift reaction ((r)WGSR).

Straightforward production of carboxylic acids from parent alkenes can be realized via two principal catalytic ways – hydrocarboxylation and hydroxycarbonylation –, both of them using end-
products of the water-gas shift reaction (WGSR), namely, CO/H$_2$O and CO$_2$/H$_2$ combinations, respectively.

It has been previously disclosed that carboxylic acids (as mixtures of linear and branched products) can be obtained via Pd-catalyzed hydroxycarboxylation of $\alpha$-olefins with CO/H$_2$O. The so-called hydride mechanism, elucidated by Van Leeuwen and coworkers (Scheme 2), presumed that the Pd-acyl species, formed via insertion of CO into the Pd-alkyl intermediate, reacts with H$_2$O eliminating the corresponding carboxylic acid and, thus, regenerating the catalyst. In spite of the obvious thermodynamic preference of the WGSR ($\Delta H = -9.8$ kcal·mol$^{-1}$), this step (even if operational) seemed not to play any crucial role in the process.

![Scheme 2](image.png)

**Scheme 2.** Proposed mechanism for the Pd catalyzed hydroxycarbonylation of olefins with CO/H$_2$O (only formation of the linear product is illustrated).

On the other hand, there are only a few reports disclosing the straightforward synthesis of carboxylic acids from CO$_2$. In seminal contributions, it has been demonstrated that the “direct” carboxylation of ethylene with CO$_2$, mediated by homogeneous catalytic systems ((Ph$_3$P)$_2$PdCl$_2$/HCl or (Ph$_3$P)$_3$RhCl/HBr), required very harsh conditions (120–180 °C, 150–700
atm) and resulted in the formation of propionic acid in 38% yield, with ethanol and ethyl propionate as the major byproducts.¹⁸

Very recently, another catalytic protocol for hydrocarboxylation of both linear and cyclic alkenes has been disclosed by Leitner et al.;¹⁹ this one uses [RhCl(CO)₂]₂/PPh₃/p-TsOH·H₂O as precatalysts coupled with CH₃I as promoter and H₂ as hydrogen source/reductant. In this case, the proposed catalytic cycle incorporates two interlinked loops (Scheme 3): the first one sets up an equilibrium between CO₂/H₂ and CO/H₂O through a reverse water-gas shift reaction (rWGSR), while the second one leads to formation of a Rh-alkyl intermediate, followed by a carbonylation reaction sequence. The intrinsic feature of this mechanism is the absence of carboxylate intermediates, as well as that the principal hydroxycarbonylation cycle parallels the one of the mechanism of hydroxycarboxylation using CO/H₂O.¹⁶ The overall reaction occurs in acetic acid under, however, still drastic conditions (60 bar CO₂, 10 bar H₂, 180 °C).

Scheme 3. Proposed mechanism for the Rh-catalyzed hydroxycarbonylation–“hydrocarboxylation” of olefins.¹⁹

Interestingly, under virtually similar conditions (CO₂/H₂, 30–60 bar, 130–140 °C, NMP as solvent and Li salts as additive) upon using H₄Ru(CO)₁₂²⁰ and Ru₃(CO)₁₂/phosphine²¹ as catalysts,
domino hydroformylation/reduction reactions of alkenes have been observed, which resulted in isolation of the corresponding alcohols (mixtures of linear and branched products). The possible intermediacy of the carboxylic acid formation step for these processes has not been established.

Yet, electrochemical coupling of CO$_2$ with $\alpha$-olefins (propylene and 1-butene) mediated by Ni(II) complexes has been reported to afford mixtures of two major products, namely, a substituted succinic acid and the corresponding unsaturated monocarboxylic acid (Scheme 4).$^{22}$ Under the same conditions, butadiene has been transformed into a mixture of mono- and dicarboxylic acids: predominantly, 3-pentenoic, 2-ethenyl-succinic and cis-/trans-3-hexene-1,6-dioic acids. The mechanism of formation of these products remains unclear. By a similar electroreductive coupling of both terminal and internal alkynes and CO$_2$, catalyzed by Ni complexes, unsaturated carboxylic acids have been prepared.$^{23}$

![Scheme 4. Ni-catalyzed electrocarboxylation of propylene and 1-butene.]

Benzonitrile-promoted electrocarboxylation of styrene into 3-phenylpropionic acid has been also documented to operate presumably via a radical mechanism.$^{24}$

**2.3. Acrylates from alkenes and CO$_2$: from stoichiometric studies towards catalysis**

The catalytic coupling of alkenes with CO$_2$ to produce acrylic acids$^{25}$ has long been of interest from both fundamental and industrial points of view. In pioneering studies aimed at
identifying potential catalytic pathways for the straightforward production of acrylic acid from ethylene and CO$_2$, several Ni and Pd complexes have been recognized as effective precatalysts. The key step of such catalytic processes is presumably the oxidative coupling reaction of the alkene with CO$_2$ onto low valent metal centers in the catalytically active species, which affords metallacyclic product 1-M (Scheme 5).$^{26,27}$ Such intermediacy has been experimentally established for Ni(0),$^{28}$ Fe(0)$^{29}$ and Pd(0)$^{30}$ (with dienes) precursors in the pioneering studies of Hoberg and coworkers and others.$^{31,32,33}$

![Scheme 5. Hypothetic mechanism for the synthesis of acrylic acids from alkenes and CO$_2$ via formation of metallalactone 1-M (M = Ni, Fe).$^{28,29}$](image)

While it had been proposed that the metallalactone derivative 1-M can follow β-H and reductive elimination reactions giving rise to the corresponding acrylic acid (Scheme 4), it has eventually appeared that the metallacycles are essentially robust; in fact, cleavage of the M–O and M–C bonds does not occur without decomposition of these complexes. Only in one case, transformation of several nickelalactone complexes into hydrido-acrylate and then into acrylate derivatives, however without further reductive elimination step, induced by addition of dppe ligand, has been documented.$^{34}$
It has been experimentally demonstrated that some additional components can provoke release of the carboxylic fragment (Scheme 6). For instance, acidic hydrolysis yielded the corresponding carboxylic acid (Scheme 6, eq (1)), while treatment of nickelalactone 1-Ni with CH$_3$OH under acidic conditions provided the corresponding ester (eq (2)). In particular, stoichiometric reactions of 1-Ni with ethylene, styrenes and other functionalized alkenes (eq (3)) followed by hydrolysis resulted in isolation of mainly linear carboxylic acids. In this case, it was proposed that the insertion of the substrate C=C double bond into the Ni–C bond of 1-Ni resulted in the ring expansion of the five-membered metallacycle, the derived thereof seven-membered intermediate being more prone for β-H elimination reaction. The reductive elimination reaction towards the corresponding carboxylic acid and regeneration of the {L}$_n$Ni(0) catalytic species appeared, however, to be unfavorable for the above systems, which hampered catalytic turnover; nonetheless, this step was found operational for the Pd- and Ni-catalyzed catalytic coupling of CO$_2$ with dienes (functionalization/telomerization) or with diynes.

Scheme 6. Stoichiometric reactions of nickelalactones 1-Ni.
As demonstrated by DFT studies, the direct β-H elimination reaction from metallalactone 1-Ni (L = DBU), which would have resulted in formation of a metal-hydrido-acrylate intermediate (Scheme 5), is not only kinetically problematic (computed $\Delta G_{298}^{\neq} = \text{ca. } 40 \text{ kcal} \cdot \text{mol}^{-1}$) but also thermodynamically unfavorable (computed $\Delta G_{298} = \text{ca. } +10 \text{ kcal} \cdot \text{mol}^{-1}$). Moreover, the subsequent reductive elimination step involving coupling of the metal-hydride and acrylate groups was also found to feature a high activation barrier, which, depending on the mechanism of elimination, can vary between 20 and 40 kcal·mol$^{-1}$. It has been predicted for the given system that preliminary cleavage of the Ni–O bond and decoordination of the carboxylate group can release the ring strain and stabilize the transition state leading to β-H elimination. This estimate has been further corroborated experimentally by Rieger and coworkers and others on the reactions between unsubstituted nickelalactones 1-Ni bearing different ligands L with an excess of CH$_3$I as the metallalactone cleaving agent (Scheme 6, eq (4)). However, the low yields (2–56% vs Ni) of methyl acrylate product over 3–48 h and significant concomitant decomposition of the Ni species make these reactions catalytically impractical so far. Additional insights in the nature of possible intermediates in this process and hypothetic decomposition pathways have been recently gained from combined computational and experimental studies. For example, for the reactions involving CH$_3$I as co-reagent, the corresponding activation barrier $\Delta G_{298}^{\neq}$ for the β-H elimination of methyl acrylate was computed in the reasonable range 15–25 kcal·mol$^{-1}$ (depending on the theoretical method and model). On the other hand, a high endothermicity ($\Delta G_{298} = +5–25$ kcal·mol$^{-1}$) was computed for the HI reductive elimination step (which can in principle be surmounted by an introduction of a base), required to regenerate a $\{L\}_n$Ni(0) catalyst; this set up another obstacle for catalytic turnover.

By analogy, addition of stoichiometric amounts of different reagents (B(C$_6$F$_5$)$_3$)$_3$, Na$^+$$[B(m-(CF$_3$)$_2$Ph)$_4$]$^-$, $X_2$ (X = Cl, Br, I), Ph$_2$Zn$^+$) to 1-Ni complexes has been shown to eventually promote cleavage of the nickelalactone cycle and elimination of the acrylate fragment.
Yet, the carboxylation reactions of Mo(0)- and W(0)-ethylene complexes, affording the corresponding acrylates 2 and 3, originally reported by Carmona and coworkers, constitute another model for coupling of CO₂ with olefins (Scheme 7). As proposed from DFT calculations, the formation of 2-W and 3-W is intermediated by formation of a metallacyclic species analogous to 1-M. Thus, the corresponding barrier for the β-H elimination reaction yielding 2-W and 3-W was estimated to be significantly lower (computed $\Delta E^{298} = \text{ca. } 10 \text{ kcal mol}^{-1}$) than those typically found in Ni-based systems.

![Scheme 7](image)

**Scheme 7.** Reaction between Mo(0)- and W(0)-ethylene complexes and CO₂.

Of particular interest is the reactivity of hydride complex (Triphos)MoH₄(PPh₃) (Triphos = Ar₂PCH₂CH₂)₂PPh; Ar = Ph) disclosed by Bernskoetter and coworkers (Scheme 8). Several products, formate, acrylate and propionate complexes can be derived from the corresponding precursor, the selectivity of the reaction depending on the ratio of the C₂H₄/CO₂ feed.
Scheme 8. Reactivity of (Triphos)MoH₄(PPh₃) with ethylene and CO₂.⁵⁰

Selective formation of similar acrylate products in reactions of zerovalent Mo-Triphos complexes (Triphos)Mo(C₂H₄)(N₂)₂ has been found to take place with different rates depending on the nature of aryl substituents on the ligand (the following rates order has been established: Ph > m-Me₂-Ph ≥ p-CF₃-Ph > p-Me-Ph ≈ p-F-Ph > p-MeO-Ph > m-tBu₂-Ph).⁵¹ Yet, the above acyclic metal-hydrido-carboxylate products of Mo and W did not reveal any propensity to reductively eliminate the corresponding carboxylic acids. Alike 1-Ni, the monomeric tungsten acrylate ((MeO)₃P)₅Mo(H)(OC(O)CH=CH₂) upon treatment with CH₃I slowly released methyl acrylate again with low yield.⁵²

For a completely different system based on a cationic Pd-methoxycarbonyl complex (Scheme 9), it has been established that methyl acrylate is readily released upon exposure to an ethylene atmosphere under mild conditions (room temperature; 1 bar pressure).⁵³ Nature of the solvent influenced the yield of the reaction; significantly better yields than those having been obtained in CH₂Cl₂/MeCN mixture or in neat CH₂Cl₂ were achieved in DMF. According to the DFT model, the essential role of DMF is the stabilization of intermediates and transition states via formation of relatively strong metal-solvent bonds that grants more favorable pathway in terms of free energy. The rate-determining step for this process was suggested to be the C–C bond coupling.
reaction, that is, the insertion of ethylene into the Pd–C bond of the methoxycarbonyl complex (computed $\Delta G^\circ_{298} = 26$ kcal·mol$^{-1}$), while the $\beta$-H scission reaction from the open form of the alkyl-Pd intermediate featured a lower barrier (computed $\Delta G^\circ_{298} = 17$ kcal·mol$^{-1}$).

Scheme 9. Computed mechanism for methyl acrylate formation from the reaction of complex $[\{dppe\}_2Pd(COOMe)]^+[SO_3CF_3]^{-}$ with ethylene in DMF.$^{53}$

Under somewhat more drastic conditions (30 bar ethylene, 30 bar CO$_2$) complex $[\{dppe\}_2Pd(COOMe)]^+[SO_3CF_3]^{-}$, along with stoichiometric production of methyl acrylate, has been found to catalyze the formation of ethyl acrylate.$^{53}$ A putative mechanism involving the intermediacy of Pd-hydrido species was proposed (Scheme 10). However, the results of both stoichiometric and theoretical studies have ruled out the proposed mechanism.$^{53}$ No alternative mechanism for this catalytic reaction matching the experimental observations has been suggested so far.
Scheme 10. Disclaimed putative hydride mechanism for Pd-catalyzed synthesis of ethyl acrylate from ethylene and CO$_2$.$^{53}$

2.4. Catalytic coupling of CO$_2$ with alkenes/alkynes/dienes in the presence of organic/organometallic substrates as reductants

The lack of catalytic activity systematically observed in the above studies about alkene carboxylation into acrylic acid derivatives, and also fundamental thermodynamic calculations,$^{39}$ suggest unfavorable overall thermodynamics of this type of reaction. That is, the thermodynamic driving force of the coupling reaction of alkene with CO$_2$ is directly related to the formation of a species with a strong M–O bond (a metallalactone or a carboxylate), but release of a “free” carboxylic acid and regeneration of a reduced metal species, which is mandatory to enable the catalytic turnover, are disfavored.

A workaround has been identified, which consists in implementing in the reaction (super)stoichiometric amounts of “energetic” reactants (organometallics or related organoelementals) that compensate the initially unfavorable thermodynamics of the reaction by the more favored formation of the corresponding metal/organoelemental carboxylate derivatives, and simultaneously regenerate the low valent metal active species.
A breakthrough along these lines has been achieved with alkoxy/phenoxy, amido and borate salts of Na, which enabled the synthesis of sodium acrylates with a Ni-based catalytic system (Scheme 11); so far, the maximal TON reached ca. 100 (10 bar CO₂, 5 bar ethylene, THF, 100 °C; 300 equiv o-F-C₆H₄-ONa vs. Ni). As it has been corroborated by theoretical investigations, the key factor in this system is the presence of a base with high proton affinity (typically, RO⁻), which modifies a transition state leading to the β-H elimination reaction in the nickelalactone and reduces the barrier of this rate determining step (the corresponding calculated barrier ΔG°₂⁹₈ drops down to 12–16 kcal·mol⁻¹). Also, as long as the corresponding carboxylic acid is produced in a form of a sodium salt, this allows maintaining favorable thermodynamics of the global process (computed ΔG°₂⁹₈ = ca. −15 kcal·mol⁻¹).

Scheme 11. Proposed mechanism for the Ni-catalyzed formation of acrylic acid derivatives from alkenes and CO₂ in the presence of inorganic or organometallic salts M⁺X⁻ (and an additional base).

The first catalytic reductive hydrocarboxylation of styrenes under mild conditions (1 bar CO₂; room temperature; THF), using an excess of Et₂Zn as nucleophilic interceptor, has been
successfully achieved by Rovis and coworkers. The NiR₂/additive/Et₂Zn (R = 1,5-COD, acac; additive = Cs₂CO₃, DBU, pyridine) catalytic system has been surmised to operate via a completely different mechanism, which excludes formation of nickelalactone intermediates and involves distinct hydrometallation and carboxylation steps leading regioselectively to saturated α-substituted carboxylic acid derivatives (Scheme 12). Several limitations have been identified for the Ni-based system: (1) it is poorly active in carboxylation of both linear olefins, cyclic dienes and β-substituted styrenes; (2) the activity of the catalytic system strongly depends on the nature of the additive; 3) also, the nature of the reducing agent plays one of the key roles, as the system is not efficient if Et₂Zn is replaced with iPrOH, Ph₃SiH or H₂, while Me₂Zn and Ph₂Zn afforded different products (no alkylated product and benzoic acid, respectively). Very recently, the mechanism has been computationally revisited, asserting transient Ni-hydrides as catalytically active species, whereas the other possible pathway involving formation of nickelalactone has been found to be kinetically disfavored.

Scheme 12. Proposed mechanism for the Ni(COD)₂/Cs₂CO₃/Et₂Zn-catalyzed hydrocarboxylation of styrenes.
FeCl₂/bis(iminopyridine)-catalyzed reductive hydrocarboxylation of styrenes has been also reported to proceed under mild conditions (1 bar CO₂; room temperature; THF) using excess EtMgBr as reductant.⁵⁹ The proposed mechanism (Scheme 13) involves an organomagnesiation step, followed by carboxylation of the resulting secondary Grignard reagent to yield mostly the corresponding α-substituted carboxylic acid derivatives. This Fe-based system is only active with ethyl Grignard co-reagents; many other traditional hydride sources (NaBH₄, LiAlH₄, NaHB₃Et₃, nBuLi, Et₂Zn, Et₃Al, iBu₂AlH) were found completely inefficient.

![Scheme 13. Proposed mechanism for FeCl₂/bis(iminopyridine)/EtMgBr-catalyzed carboxylation of styrenes.⁵⁹](image)

There are also a few reports on reductive carboxylation of functionalized alkenes: terminal alkene hydroboration followed by Cu-catalyzed carboxylation in the presence of KOrBu,⁶⁰ and reductive carboxylation of allyl esters and halides on Ni catalysts carried out using metallic Zn and Mn powders as stoichiometric reducing agents.⁶¹

Dienes and alkynes are substantially more reactive substrates that can be converted to the corresponding unsaturated carboxylic acids derivatives. Thus, the ring-closing carboxylation of
bis(1,3-dienes)\textsuperscript{62} and hydrocarboxylation of 1,3-dienes\textsuperscript{63} and allenes,\textsuperscript{64} catalyzed by Ni and Pd complexes have been reported to proceed in the presence of superstoichiometric amounts of organometallic reagents (R\textsubscript{2}Zn (R = Et, Ph) and Et\textsubscript{3}Al, respectively). Similarly, both terminal and internal alkynes and some functional derivatives have been used in Ni- and Cu-catalyzed reactions with CO\textsubscript{2} in the presence of different reagents (R\textsubscript{2}Zn and RZnX (R = Me, Et, Bu, Bn, Ph; X = Cl, Br, I), M\textsubscript{2}CO\textsubscript{3} (M = K, Cs)),\textsuperscript{65,66} and in related bora-\textsuperscript{57} and silacarboxylation\textsuperscript{68} reactions using B\textsubscript{2}(pin)\textsubscript{2}/LiO\textsubscript{t}Bu and Et\textsubscript{3}SiB(pin)/NaO\textsubscript{t}Bu co-reagents. The synthesis of propiolic acids by carboxylation of terminal alkynes in the presence of bases M\textsubscript{2}CO\textsubscript{3} (M = K, Cs) catalyzed by Cu(I) and Ag(I) complexes\textsuperscript{69} and heterogeneous Ag-doped MOFs\textsuperscript{70} has been also developed.

The coupling of styrene derivatives with CO\textsubscript{2} in MeOH into the corresponding methyl esters catalyzed by Ni hydrides has been described by Garcia and coworkers.\textsuperscript{71} The proposed hydroesterification mechanism (Scheme 14) parallels that of hydroxycarbonylation (Scheme 2);\textsuperscript{16} it also incorporates a step of formation of an acyl-metal intermediate from the corresponding Ni-alkyl species and CO generated from CO\textsubscript{2}.\textsuperscript{72} The CO\textsubscript{2} origin of the carbonyl group in the final product has been corroborated by \textsuperscript{13}C labelling experiments.

\textbf{Scheme 14.} Proposed mechanism for the Ni-catalyzed hydroesterification of styrenes.\textsuperscript{71}
A similar mechanistic pathway has been proposed by Beller and coworkers for the Ru-catalyzed hydroesterification of many industrially important olefins; the reduction of CO₂ to CO in MeOH (solvent) was also suggested.

There are only a few examples describing the use of hydrosilanes as reductants in carboxylation of unsaturated substrates (typically, alkynes). For example, the hydrocarboxylation of disubstituted alkynes catalyzed by (NHC)CuF (NHC = N-heterocyclic carbene) has been reported to proceed under mild conditions (1 atm CO₂, 65–100 °C), giving the corresponding silyl esters of α,β-unsaturated carboxylic acids (Scheme 15). Stoichiometric studies revealed that the rate-determining step of the process is the insertion of CO₂ into the Cu–C bond of the alkenyl intermediate. Remarkably, the reaction of the Cu(I)-carboxylate intermediate with hydrosilane occurred readily, already at room temperature; different hydrosilanes have been tested as reducing agents, exhibiting the following order of reactivity: (EtO)₃SiH ≥ PMHS (= polymethylhydrosiloxane) > (iPrO)₃SiH > Ph₂SiH₂.

Scheme 15. Proposed mechanism for the (NHC)CuF/R₃SiH-catalyzed hydrocarboxylation of alkynes.

3. Perspectives
Depending on the nature of the catalytic system used for the carboxylation of alkenes, three distinct types of mechanisms can be operational: (1) a rWGSR-mediated hydroxycarbonylation that affords straightforwardly saturated carboxylic acids; (2) the formation of acrylates via oxidative addition of CO$_2$ and alkene onto a low-valent metal center, followed by $\beta$- and reductive eliminations steps; (3) the production of metallic salts/organoelemental derivatives of the corresponding saturated acids via an hydrocarboxylation mediated by metal-hydrido or -alkyl species.

For the first process, the key and indispensible step is the rWGSR utilizing CO$_2$/H$_2$ mixtures as CO surrogates. Since dihydrogen is widely preferred as reductant in organic synthesis and catalysis$^{76}$ as an inexpensive, clean and sustainable reagent,$^{77}$ it obviously grants atom-economy for the given reaction. The use of dihydrogen as reducing agent in this process admittedly limits its applicability to homogeneous catalytic systems that are capable of maintaining efficiently the rWGSR step in the putative catalytic cycle (Scheme 3). While many heterogeneous, both single- and multicomponent systems are known to catalyze this reversible process,$^{78}$ only for a few homogeneous systems (typically based on Rh and Ru) the rWGSR has been proposed to operate as a part of global carbonylation processes utilizing CO$_2$/H$_2$ mixtures as CO surrogates.$^{2a,j,19,20,21}$ In this regard, the identification of other homogeneous catalytic systems, based preferably on more abundant, less expensive metals (Fe, Co, Ni etc) competent for catalyzing the rWGSR under mild conditions, would also be of high industrial relevance. Unfortunately, to date, the complete deficiency of experimental (kinetic) and computational data for this process does not allow a rational optimization/development; identification of the optimal conditions remains still serendipitous.

For the second and third processes, efficient high-energy reductants are mandatory in order to compensate the unfavorable thermodynamics of the carboxylation reactions: (1) for cleavage of the metallacycle in the divalent Ni$^{(II)}/$Pd$^{(II)}$ intermediates formed in the oxidative coupling step of unsaturated hydrocarbon with CO$_2$; and/or (2) for regeneration of the low valent transition metal
species from the corresponding carboxylate intermediates; and (3) for conversion via transmetallation of metal-carboxylate intermediates to the catalytically active metal-alkyl or -hydrido species. These reactions thus result in release of the carboxylate products as metallic (Na\(^+\), Zn\(^{2+}\) or Mg\(^{2+}\), respectively) salts, which have to be treated acidically in order to recover the corresponding pure carboxylic acids. Therefore, this way of achieving the carboxylation process is plagued by the consumption of a stoichiometric amount of an expensive, difficult-to-handle and waste-generating organometallic reagent.

On the other hand, use of dihydrogen as reductant is hardly compatible with both mechanisms (Schemes 11–13). Indeed, the pivotal step of the reductive carboxylation process, namely, the reduction of metal-carboxylate intermediates with dihydrogen to give the corresponding carboxylic acid and a metal-hydrido species, is thermodynamically difficult (BDE\(_{298}(\text{H-H bond}) = 104\) kcal\(\cdot\)mol\(^{-1}\), BDE\(_{298}(\text{Si-H bond}) = 80\) kcal\(\cdot\)mol\(^{-1}\)).\(^{79}\)

A modified scenario can be envisaged, by implementing reduction with a molecular hydrosilane or the more economically-viable polymethylhydrosiloxane (PMHS); this would provide direct access to valuable carboxylic acid silyl esters. Readily available, easy-to-handle and relatively inexpensive (although admittedly incompatible with the production of commodities) hydrosilanes are widely used as reducing organic agents for many functional groups.\(^{80}\) Given the high propensity of hydrosilanes to induce formation of M–H bonds in metal complexes via the simultaneous, thermodynamically favored formation of Si–O bonds (BDE\(_{298}(\text{Si-O bond}) = 190\) kcal\(\cdot\)mol\(^{-1}\))\(^{79}\) and their highly tunable reactivity by the substituents at the silicon atom, these are eminently competitive reducing agents to organometallic substrates; they are also a frequent alternative to dihydrogen in many metal-catalyzed processes. PMHS, a low molecular weight (\(M_n = 1,700–3,200\) g\(\cdot\)mol\(^{-1}\)) polymeric byproduct of the silicone industry, can also be used as a very cheap and environmentally friendly reducing agent. PMHS is more air- and moisture-stable than other hydrosilanes and can be stored for long periods of time without loss of activity. The mechanism of action and reactivity pattern of PMHS are the same as those featured by usual
hydrosilanes; application of PMHS for reduction of C=O groups in aldehydes and ketones has been reported. The envisioned mechanism for reductive carboxylation of unsaturated hydrocarbons with hydrosilanes (Scheme 16, Cycle A) is composed of a series of elementary chemical reactions which, individually, are all well-documented: (i) alkene insertion into the M–H bond of active metal-hydrido species 1 to afford metal-alkyl 2; (ii) insertion of CO$_2$ into the M–alkyl bond of 2 yielding metal-carboxylate 3; (iii) a transelementation reaction of 3 with a hydrosilane delivering the corresponding alkyl silyl ester 4. A competing mechanism may be envisioned (Scheme 16, Cycle B), which involves rapid insertion of CO$_2$ into the M–H bond of 1 to give a metal-formate intermediate of the type 5, which upon reaction with the hydrosilane would release the corresponding silyl formate 6 as a by-product besides the desired 4. The latter process, that is hydrosilylation of CO$_2$, is well documented in the literature and it is known to be catalyzed by complexes and salts of non-transition elements.

Scheme 16. Putative catalytic cycle (A) for the formation of silyl esters by reductive carboxylation of alkenes, and CO$_2$ hydrosilylation (B) as anticipated side reaction.
In both cases (Scheme 16, Cycles A and B), hydrolysis of silyl esters 4 and 6 under mild conditions in a distinct, final workup step shall yield the corresponding alkyl-carboxylic and formic acids, respectively, and nontoxic (poly)siloxane byproducts (use of PMHS). However, silyl esters primarily resulting from the above global process are also much valuable precursors and synthetic intermediates. For instance, they are employed as intermediates and cross-linking reagents in the synthesis of poly(silyl esters), some biodegradable polymers widely used for medical applications (drug delivery systems, surgical devices, recyclable materials). Synthesis of silyl esters generally relies on condensation of carboxylic acids with chlorosilanes with concomitant elimination of HCl. An alternative synthetic protocol exists, which is based on dehydrogenative coupling of carboxylic acids with hydrosilanes catalyzed by transition metal complexes (Ru-Pd, Pt, Cu).

4. Conclusions

In this Perspective, we have highlighted advances and recent breakthroughs in the “direct” catalytic synthesis of carboxylic acids and their derivatives starting from unsaturated hydrocarbons (alkenes, alkynes, dienes) and CO₂. All of the existing systems are plagued either by a poor catalytic productivity and/or the use of expensive and waste-generating stoichiometric reductants. Efforts must be paid to optimize the cost, safety and global sustainability of such processes. These objectives could be achieved by two interlinked approaches: (1) implementation of catalytic systems that are competent for carboxylation of unsaturated hydrocarbons (alkenes) using H₂ as reducing agent under mild conditions (temperature, pressure); and/or (2) implementation of relatively inexpensive and efficient reducing agents in the catalytic systems (e.g., molecular hydrosilanes or the polymeric analogue, polymethylhydrosiloxane (PMHS)). Understanding of the nature and role of key intermediates and mechanisms of CO₂ activation from experimental approaches and multi-scale theoretical simulations is the key for elaborating processes that enable efficient transformation of CO₂ into value-added chemicals. This is also of high interest for future renewable energy technologies and addressing the global issue of energy transition.
REFERENCES


10 Ethylene is the largest volume organic chemical produced in the world (140 MT/y), followed by propylene (87 MT/y), see: E. B. Stechel, J. E. Miller, Journal of CO₂ Utilization, 2013, 1, 28-36.


Acrylic acid is produced by a multi-step oxidation of propylene. The market of acrylic acid derivatives is around 2.5 MT/year with annual expansion rate of 4%, see: ref. 2c.


A revisited mechanism has been described recently: P. N. Plessow, A. Schafer, M. Limbach, P. Hofmann, *Organometallics*, 2014, 33, 3657-3668.


A thermally stable metallalactone complex of Ti(IV) has been reported to be the product of the reaction of the parent Ti(II) precursor Cp₂^*Ti(η²-C₂H₄) and CO₂: S. A. Cohen, J. E. Bercaw, *Organometallics*, 1985, 4, 1006-1014.


82 This reaction constitutes the principal step of the mechanism of polymerization of alkenes using Ziegler-Natta, metallocene and post-metallocene catalysts: J. A. Gladysz, Special issue: Frontiers in Metal-Catalyzed Polymerization, Chem. Rev. 2000, 100, 1167.

83 To our knowledge, this reaction has been intentionally substantiated only in a few cases: (a) hydrosilane-mediated reduction of Ir-carboxylate complexes into the corresponding hydrides, M. A. Esteruelas, O. Nurnberg, M. Olivan, L. A. Oro, H. Werner, Organometallics, 1993, 12, 3264-3272; (b) quantitative formation of silyl esters from the parent U(III)-carboxylates by Me3SiX halides (X = Cl, I): E. M. Matson, W. P. Forrest, P. Fanwick, S. C. Bart, J. Am. Chem. Soc., 2011, 133, 4948-4954; (c) alkoxide and alkylperoxy group transfer reactions from \{L\}_nZn–OR and \{L\}_nZn–OOR complexes to organosilanes have been reported to yield
