Heterogeneous Phase Microwave-Assisted Reactions under CO₂ or CO Pressure

Emanuela Calcio Gaudino 1, Laura Rinaldi 1, Laura Rotolo 1, Diego Carnaroglio 1,2, Camillo Pirola 2 and Giancarlo Cravotto 1,*

1 Dipartimento di Scienza e Tecnologia del Farmaco and NIS—Centre for Nanostructured Interfaces and Surfaces, University of Turin, Via P. Giuria 9, Turin 10125, Italy; emanuela.calcio@unito.it (E.C.G.); laura.rinaldi@unito.it (L.R.); laura.rotolo@unito.it (L.R.); diego.carnaroglio@unito.it (D.C.)
2 Milestone srl, Via Fatebenefratelli, 1-5, Sorisole 24010, Italy; c.pirola@milestonesrl.com
* Correspondence: giancarlo.cravotto@unito.it; Tel./Fax: +39-011-670-7687

Abstract: The present review deals with the recent achievements and impressive potential applications of microwave (MW) heating to promote heterogeneous reactions under gas pressure. The high versatility of the latest generation of professional reactors combines extreme reaction conditions with safer and more efficient protocols. The double aims of this survey are to provide a panoramic snapshot of MW-assisted organic reactions with gaseous reagents, in particular CO and CO₂, and outline future applications. Stubborn and time-consuming carbonylation-like heterogeneous reactions, which have not yet been studied under dielectric heating, may well find an outstanding ally in the present protocol.

Keywords: microwave-assisted organic reactions; heterogeneous phase; gas pressure; CO carbonylation; CO₂ insertion

1. Introduction

The development of environmentally benign and efficient synthetic protocols is still a central goal of current research in chemistry [1,2]. In this regard, heterogeneous multiphase reactions with gaseous reagents in closed microwave (MW) reactors stand out as key alternatives to conventional protocols and provide “greener” chemical conversions. While synthetic chemistry still depends strongly on fossil fuels as an energy source and source of raw materials, the search for renewable raw feedstocks is a prerequisite for sustainable chemical processes [3]. The ambitious aim of decreasing the use of depleting fossil fuel deposits has stimulated an intense research effort in recent years that has been focused on exploiting carbon monoxide (CO) and carbon dioxide (CO₂) as renewable and inexpensive C₁ building blocks [4].

Despite their huge potential, CO and CO₂ are seldom used as reagents in industrial organic synthesis because of the general reluctance to use these gases as reagents and their need for specific pressure-resistant equipment [5]. In particular, CO is an extremely useful carbonyl synthon in organic synthesis. Carbonylation reactions, such as formylation [6], hydroformylation [7], alkoxy-carbonylation [8], aminocarbonylation [9], carbonylative Heck [10], Suzuki Miyaura [11], and Sonogashira reactions [12], are industry’s core strategies for converting various bulk chemicals into an extensive set of useful products, such as alcohols, aldehydes and carboxylic acid derivatives. CO works as a high affinity ligand for transition metal catalysts (especially palladium), because of its dual ability to act as a σ-donor and π-acceptor, making “CO chemistry” an easy and practical route for the introduction of a carbonyl group into an organic substrate. Despite the above-mentioned advantages, CO’s reputation as “the silent killer”...
that leads to asphyxiation (as an oxygen competitor in haemoglobin binding), has generated major concern and warnings concerning its use. A number of protocols that see CO released in situ have been developed to circumvent the safety issues surrounding this invisible, odourless, tasteless and highly toxic gas [13–19]. These methods include the use of CO-equivalents (alkyl formates [20], formic acid, formic anhydride, formamide [21–23], N-substituted formamide [24], carbamoylstannanes [25], carbamoylsilane [26]), and CO-releasing reagents (metal carbonyls [27]). Unfortunately, these reagents are relatively expensive and sometimes require harsh reaction conditions for CO release (strong base combined with high temperatures and long reaction times).

Only 1% of the total amount of CO2 on Earth is currently being used by chemical industries, because of its chemical inertness and the high cost of capture and storage [28]. The number of scientific documents that deal with CO2 use has grown exponentially over the last decade, with the notable exception of a sudden decrease in 2012, when the Fukushima disaster in Japan also affected the scientific production of that country (Figure 1). The large-scale transformation of CO2 into basic bulk chemicals remains a long-standing challenge and fascinating early dream. The insertion of CO2 into metal-element bonds is already a common procedure, mainly exploited for the synthesis of fine chemical and a few materials, rather than for a broader use on ton scale [29,30].

![Scientific papers with “CO₂” in the title](image)

**Figure 1.** Scopus® (Elsevier) scientific document analysis.

In particular, CO2 gas plays an important role in the 100% atom economy synthesis of organic carbonates (including cyclic carbonates and polycarbonates) [31,32], urea derivatives [23], oxazolidinones [33], formic acid [34], solar fuels [35] and so forth [36]. However, it should be noted that the carbon atom in CO2 is present in its most oxidized state, which results in higher stability and lower reactivity than CO [37]. High energy input is necessary for the sustained use of CO2 as a C1 environmentally friendly building block and the use of alternative energy source or technologies is thus recommended.

In synthetic organic chemistry, one of the most direct ways to pursue green chemistry is the use of enabling technologies that include heterogeneous phase synthesis as well as relatively new techniques such as MW-assisted synthesis, continuous flow reactors, etc. that have been developed to speed up synthetic transformations and achieve process intensification [38,39]. Since the first report in 1986 [40,41], MW-assisted chemistry has fulfilled its promise of being a fast synthetic technique and it has now been recognized as one of the most powerful and sustainable tools in organic chemistry [42]. MW irradiation provides efficient, in-core volumetric reaction mixture heating via the direct coupling of electromagnetic waves with polar entities (solvents, reagents, catalysts), providing rapid energy transfer (less than nanosecond) to the reaction mixture [43]. The efficiency of MW flash heating has resulted in dramatic reductions in reaction times, improvements in product yields and purity, even in heterogeneous gas-phase [44] or in a synergistic combination with other energy sources [45]. Despite MW assisted organic synthesis (MAOS) being a flourishing topic, little attention has been paid to the possibility of combining MW technology with gaseous reagents especially at low pressure (hydrogenation [46], hydroformylation [47], and alkoxy- [48] or aminocarbonylation [49]).
This is feasible because all MW reactors have been designed to resist the pressure generated by solvents used in it and can be considered as small autoclaves that can carry out reactions with pressurized gasses. Only a few reports have dealt with the combined use of CO or CO\(_2\) in MW at higher gas pressures. The recent introduction onto the market of new pressure resistant (up to 200 bar) high power (up to 1.5 kW) MW reactors equipped with separate multiple gas inlets, which can be considered autoclaves, could pave the way for heterogeneous gas-phase MW reactions [50]. This review aims to summarize recent work in this area and evoke some potential applications, including in the field of flow techniques thanks to the recent development of commercial MW flow reactors for organic synthesis under pressure [51–54].

2. MW-Assisted Reactions under CO\(_2\) Pressure

As stated above, only a limited number of studies into the MW-assisted synthesis of fine chemicals have been reported in the literature. This is normally due to the instrumental difficulty of performing MW reactions under gas pressure. The synthesis of cyclic and linear carbonates is the most common application of MW. The synthesis of five-membered ring carbonates from epoxides and CO\(_2\) is a promising methodology from a resource utilization standpoint because of the 100% atom economy of the reaction and the wide variety of applications of cyclic carbonates, which include use as polar aprotic solvents, precursors of polymeric materials and intermediates in the syntheses of pharmaceuticals [55].

In 2004, Nüchter et al. [56] described the first examples of the MW-assisted reactions of different oxiranes with CO\(_2\) in ionic liquids (ILs). Being highly polar, ILs are rapidly heated by MW irradiation [57] affording in shorter reaction times comparable yields (Table 1). Work-up procedures have been explained in detail and help to envisage the scale-up of the process. Compared to conventional conditions, these reactions are much faster, with higher energy efficiency and precise parameter monitoring and control. MW heating has been used in one of the most active catalytic systems for CO\(_2\) coupling with epoxides; namely zinc salts in ILs [58]. Zinc phenosulfonate octahydrate and C\(_4\)NBr efficiently catalysed the solvent-free synthesis of cyclic carbonates with remarkable turnover frequency (TOF). Park and co-workers [59,60] have proposed novel silica-supported ILs for the cycloaddition of epoxides and CO\(_2\) under MW irradiation as a means to overcome the major drawback of IL catalysis, their recovery from the reaction mixtures and recycling.

Heterogeneous catalysis is generally preferred by the chemical industry as it leads to easier separation and the possible use of a fixed-bed reactor. However, supported catalysts suffer from the limited diffusion of reagents and products between the solution and the catalyst surface. MW irradiation can nevertheless strongly enhance activity via the selective overheating of catalyst particles [61]. The rapid and selective MAOS of cyclic carbonates has been established using homogeneous (Entry 3, Table 1) and heterogeneous IL (Entry 4, Table 1) catalysts. Both catalysts showed excellent activity and short reaction times (15–30 min) under relatively low CO\(_2\) pressures (9.7 bar). Unlike conventional reactions, MW technology was able to selectively provide cyclic carbonates rather than mixtures of cyclic and linear products. Even higher selectivity towards the cyclic carbonates was achieved by supporting ILs on silica. This solid catalyst was recycled three times without any significant loss in activity.

An environmentally-friendly synthesis of cyclic carbonates from CO\(_2\) and epoxides with a HCOOH/KI catalytic system that exploits the synergistic effect between KI and hydrogen bonding donors has been studied under MW irradiation (Table 1, Entry 5) [62]. The search for more efficient processes has prompted Park [63] to propose the use of pyridinium-based ILs as an efficient catalyst for the rapid, solvent-free MW-assisted cycloaddition to yield cyclic carbonates under moderate reaction conditions (Entry 6, Table 1). The cycloaddition reaction occurred over a short reaction time (30 s), resulting in a high TOF which ranged from 200 to 7000 h\(^{-1}\). Moreover, the catalyst was recovered and reused three times without losing its catalytic properties.
Table 1. MW-assisted synthesis of cyclic carbonates.

| Entry | Catalytic System | MW (Watt) | Temp. (°C) | CO₂ (bar) | Time (min) | Yield (%) | Conv. (%) | Ref. |
|-------|------------------|-----------|------------|-----------|------------|-----------|-----------|------|
| 1     | [C₄MIm]Br (0.05 mol %), ZnPO (0.043 mol %), C₄NBr (2.174 mol %) | /         | 140        | 10        | 60         | 86        | 97        | [56] |
| 2     | TABC (0.027 mol %) | 300       | 120        | 30        | 60         | 97        | /         | [58] |
| 3     | SiO₂/C₄ImBr (4 wt %) | 100       | 120–160    | 9.7       | 15         | 87.9      | /         | [59] |
| 4     | HCOOH (0.145 mol %), KI (0.016 mol %) | 200       | /          | 9.6       | 15         | 59        | 60        | [62] |
| 5     | C₄PyBr (0.054 mol %) | 200       | 0-180      | 9.6       | 0.5        | 67.2      | 70        | [63] |

The second interesting field of use for CO₂ is the synthesis of dimethyl carbonate (DMC, Scheme 1).

Scheme 1. Dimethyl carbonate synthesis.

Direct synthesis of DMC from CO₂ and methanol is a very attractive reaction, not only because the product is a well-known low polluting substitutive intermediate of toxic phosgene or dimethyl sulfate, but also because CO₂ can be consumed as a greenhouse compound in this process. In 2001, the first example of this synthesis was reported as a facile method for the quick synthesis of DMC from CO₂ and methanol under MW irradiation [64] and it was compared with conventional heating. In that work, rapid MW heating resulted in higher selectivity to DMC, while conductive heating (slow heating rate) may favour the formation of DMC. Another MW-promoted DMC synthesis reaction with methanol and CO₂ was carried out over a K₂CO₃/CH₃I catalyst in the presence of IL by Du et al. (2012) [65]. IL may affect the polarity of the intermediates and the dipole moment at the transition state, by which selective MW energy absorption and CO₂ solubility in the reaction medium were increased. IL may also enhance mass transfer by increasing molecular collisions. Computational studies confirmed the hypothesis that the kinetic and thermodynamic factors driving DMC synthesis may be affected by IL that effectively activate CO₂ molecules by altering the angle and length of the C=O bond. Such activation facilitates the base-catalyzed insertion step by means of K₂CO₃ without affecting the total reaction mechanism.

The last example of MAOS under CO₂ pressure is a new, green and efficient procedure to obtain isocyanates and urea libraries [66]. A versatile pressure-resistant reactor gave a fast reaction with poor reagent excess. An optimized protocol for a Staudinger-aza-Wittig reaction with polymer bound diphenylphosphine (PS-PPh₂) under CO₂ pressure was carried out. This study also synthesized a series of symmetric and asymmetric alkyl/aryl urea derivatives from alkyl bromides in a one-pot procedure.

3. MW-Assisted Reactions under CO Pressure

Palladium-catalyzed C-C bond formation with aryl halides is currently one of the most important organometallic reactions in synthetic organic chemistry [67]. It offers the practical preparation of a broad spectrum of organic derivatives from simple, commercially available building blocks (Scheme 2). In particular, the carbonylation of aryl halides offers a one-step route to a number of important products, including carboxylic acids, esters and amides [68].

When transferring this chemistry from conventional to dielectric heating a number of critical issues had to be considered to circumvent the difficulties of working in a multiphase gas/liquid (solid) system. Up to fifteen years ago, the lack of technological progress in the design of high pressure resistant MW reactors, resulted in a scarcity of synthetic reports on this topic. Most of the commercially available MW reactors had a pressure limit of 20–30 bar, with no possibility of a prior gas pressurization. The latest
generation of pressure-resistant MW reactors may meet all these requirements. Thus, Leadbeater and Kormos have published the first report on the MW-promoted hydroxy- and alkoxy carbonylations of aryl iodides in the presence of Pd(OAc)$_2$ as a catalyst using heavy-walled quartz reaction vessels that are pre-pressurized with CO. The authors described a MW-promoted hydroxycarbonylation of aryl iodides in water [69]. Reactions were performed at 165 °C (CO 14 bar) using Pd(Ac)$_2$ as a catalyst (0.01 mol % or 1 mol % loadings) in the absence of a phase transfer agent. This is a valid alternative to the use of solid CO sources, giving the corresponding benzoic acids in good yields (59%–80%) inside 20 min, including the ortho-substituted analogues.

![Scheme 2. Carbonylation of aryl-X compounds.](image)

Furthermore, Iannelli et al. [70], focused their attention on the use of simple ligandless palladium complexes as catalysts in carbynylation chemistry as they described the ethoxycarbonylation of iodobenzene on a 1 mol scale. Six different fast ethoxycarbonylation reactions of several aryl iodides were performed simultaneously using a near-stoichiometric loading of CO gas (1.08 stoichiometric equiv.) in a sealed multimode MW reactor. Excellent conversions (91%–99%) were achieved under MW irradiation in only 30 min at 125 °C on both 50 mmol and 1 mol reaction scales using Pd(Ac)$_2$ as a catalyst.

Besides the obvious safety concerns, a large excess of highly toxic CO gas may affect the efficiency of palladium catalysts because of the strong π-acceptor character of CO as a ligand. Furthermore, the rate of oxidative addition is greatly reduced and the clustering of palladium atoms is facile in the presence of CO, leading to non-active palladium species. To overcome these drawbacks, a palladium mediated alkoxy carbonylation procedure using near-stoichiometric quantities of CO gas was also presented by Kormos and Leadbeater [71]. They performed the alkoxy carbonylation reactions using CO in a close to stoichiometric ratio, under MW heating which, as well as the inherent advantages of rate acceleration, offers a convenient method for loading vessels with gas in a safe manner. Precise CO pressure could be monitored in real time by means of a pressure sensor directly placed on the top of the reaction vessel in the monomode reactor. They successfully performed a series of carbonylations with Pd(OAc)$_2$ in alcohols (MeOH, i-PrOH and t-BuOH) at 125 °C under CO (0.97 bar)/N$_2$ (9.38 bar) pressure in 20 min. The same authors described the alkoxy carbonylation of several aryl iodides to the corresponding esters using primary, secondary and, the less reactive, tertiary alcohols [72]. These reactions were still performed at 125 °C using 0.1 mol % of Pd(OAc)$_2$ as the catalyst and DBU as the base with good to excellent yields in 20–30 min. It is worth noting that the conditions for alkoxy carbonylation are milder than those previously reported by the same authors [62], due to the greater solubility of CO in ethanol than in water.

Heterogeneous catalysts such as Pd/C strongly absorb MW energy, and the local temperature on the catalyst surface is much higher than the bulk temperature measured in the solvent (using optical-fiber thermometers or thermocouples) particularly in the case of low tanδ solvents [73]. The hot catalyst surface may dramatically accelerate chemical reactions when an efficient mass transfer is provided. At this regards, a MW-assisted carbonylation under heterogeneous catalysis with Pd/C has been reported by Salvadori et al. [74]. Alkoxy carbonylations were performed using stoichiometric amounts of different primary and secondary alcohols in DMF in the presence of CO gas (9 bar), Pd/C as catalyst and DBU as the base. In 10–20 min at 130 °C, they achieved 75%–99% product yields. Analogously, iodobenzene, CO and amines were transformed into the corresponding amides in good yields (75%–95%) after simple filtration to remove the heterogeneous catalyst. Pd/C was recycled twice without relevant differences being noted. The versatility of Pd/C as a heterogeneous catalyst
for the carbonylation reaction is also confirmed by the possibility of continuing the MW-assisted
cyclohydrocarbonylation of o-iodoaniline with acyl chlorides, producing benzoazinones in satisfactory
yields (70%–82%) with CO gas (9 bar) (Scheme 3).

Scheme 3. MW-assisted carbonylation and cyclocarbonylation of aryl iodides under ligand-free
heterogeneous catalysis.

Liptrot et al. [75] has reported the first practical synthesis of aryl and heteroaryl N-acylureas via
the MW-assisted palladium-catalyzed carbonylation of aryl or heteroaryl halides in the presence of
urea nucleophiles (Scheme 4). The reactions tolerate a wide range of substituted ureas and substituted
aryl or heteroaryl halides while proceeding in good to excellent yields (55%–90% at 100 °C in 4 h), when
using either CO (4.5 bar) or molybdenum hexacarbonyl (Mo(CO)6) as the CO source in pre-pressurised
or standard MW vials. To illustrate the usefulness of this method, the authors also reported the
one-step synthesis of diflubenzuron (an important insecticide) in acceptable yield (45%) using PdCl2
(dppf)CH2Cl2 as catalyst.

Scheme 4. Synthesis of aryl and heteroaryl N-acylureas via MW-assisted palladium catalysed carbonylation.

There has been a recent increase in interest in MW carbonylation reactions that combine the
use of CO, as a benign carbon atom source, and new eco-friendly catalysts. In fact, Pizzetti et al.
(2011) [76] have developed the first catalytic MW-assisted aminocarbonylation of ynamides using
[Fe3(CO)12] as the catalyst precursor and TEA as the only ligand. The aminocarbonylation products
were obtained in only 20 min of irradiation (90 °C and 250 W) in good to excellent yields (67%–90%).
The authors described an atom-efficient aminocarbonylation procedure in which all reactants were
used in a stoichiometric amount under low CO pressure (1.3 bar). The same MW procedure was
easily applied to the alkoxy carbonylation of terminal alkynes to give a new class of (E)-acylamides
that was regioselectively obtained in only 20 min (44%–78% yields). Furthermore, this procedure
was easily exploited for the synthesis of acryl esters from ynamides and alkynes when using MeOH
as nucleophile (Scheme 5). MW irradiation allowed milder reaction conditions to be used; lower
temperatures and gas pressures and shorter reaction times.

Scheme 5. MW-assisted iron-catalysed monocarbonylation of ynamides and terminal alkynes with
different nucleophiles.
The safe and synergistic use of CO in a closed MW reactor has led to a technological breakthrough in aminocarbonylation reactions. To the best of our knowledge, very few reports have dealt with the use of CO gas as an aminocarbonylation reagent inside a MW oven. In this regard, Calcio Gaudino et al. [77] have described a fast MW-assisted Pd(II)-catalysed protocol for the synthesis of amides using CO gas (Scheme 6).

![Scheme 6. MW-assisted aminocarbonylation using CβCAT.](image)

Their investigation dealt with the optimization of a synthetic heterogeneous aminocarbonylation protocol which made use of CO gas and a supported palladium catalyst (CβCAT) which was based on Pd(II)-triphenylphosphine embedded in cross-linked β-cyclodextrins (β-CD). Several amide derivatives were synthesized in good yields (59%-95%) in acetonitrile using an aryl iodide/nucleophilic amine ratio of 1:2 and a CO/N₂ mixture in a MW reactor equipped with a multiposition rack for simultaneous reactions and fine pressure control. The green β-CD based catalyst showed excellent performance, including easy recyclability and no significant loss in activity aside from negligible entrapped metal catalyst leaching.

4. A Promising Future for the Use of MW-Assisted Protocols under CO₂ or CO Pressure

In this section, we report a few relevant examples of reactions that could potentially gain significant advantages from MW-assisted heterogeneous phase protocols under CO₂ or CO pressure. Available published data leads to expectations of striking increases in reaction rate in all cases.

As mentioned above, CO₂ plays a pivotal role as a reagent in many fields of modern chemistry. The use of CO₂ and waste biomass as a chemical feedstock would provide us with a sustainable basis for the future of the chemical industry, if we could balance the high energy requirements of dielectric heating.

The production of formamides using a MW-autoclave is an eloquent example of potential applications. An alternative green route to the N-formylation of amines is the use of cheap, abundant and safe CO₂ along with H₂ as a formylating reagent [78]. Zhang et al. [79], have recently proposed a highly efficient catalyst for the N-formylation of various amines with CO₂ and H₂. This catalyst is based on ruthenium-pincer-type complexes and gives the corresponding formamides with excellent yield and selectivity (Scheme 7).

![Scheme 7. Catalyzed N-formylation of amines with H₂ and CO₂.](image)

Another hot topic at the moment is the development of new materials that are able to capture CO₂. With this aim in mind, Takada et al. [80], have set up a new procedure for the synthesis of optically pure oxazolidinones from CO₂ and β-amino alcohols that are derived from natural amino acids (Scheme 8).

![Scheme 8. Synthesis of oxazolidinone from β-amino alcohols.](image)
The synthetic protocol reported here is simple, straightforward and can be easily replicated. The only limits of this reaction are the harsh conditions and long reaction time. Irradiation with MW under CO\textsubscript{2} pressure may dramatically simplify the reaction outcome.

The transition-metal-catalysed simultaneous insertion of heteroatom functional groups and CO into organic molecules is one of the most important tools for the direct synthesis of heteroatom-functionalized carbonyl compounds. Dicobalt octacarbonyl and palladium complexes [Pd(PPh\textsubscript{3})\textsubscript{2} and Pd(OAc)\textsubscript{2}] were found to exhibit excellent catalytic activity for the thiolative lactonization of internal alkynes, that bear a hydroxyl group, in the presence of CO and organic disulfides [81]. The authors successfully prepared γ-lactone derivatives (71%–98% yields) that bear exo-methylene and thio groups with excellent regio- and stereoselectivities (140 °C for 20 h) (Scheme 9). The procedure can also be applied to δ-lactone synthesis. The obtained thiolated lactone derivatives are promising synthetic intermediates, as the thio group can be displaced by a variety of nucleophiles, while exo-methylene groups make them a possible substrate for Michael additions.

\begin{equation}
\text{HO} \quad \text{C} \quad \text{H} \quad \text{R}_{1} + (\text{R}_{2}S)_{2} \quad \text{cat. } \text{Co}_{2}(\text{CO})_{8} \quad \text{or} \quad \text{cat. } \text{Pd}(\text{PPh}_{3})_{4} \quad \begin{array}{c}
\text{140 °C, 20 h, CO (20 bar)}
\end{array} \quad \begin{array}{c}
\text{O} \\
\text{O} \\
\text{S} \\
\text{S}
\end{array}
\end{equation}

\textbf{Scheme 9}. Thiolative lactonization of internal alkynes bearing a hydroxyl group with CO and organic disulfides catalyzed by transition-metal complexes.

Porous organic polymers (POPs) have recently emerged as metal catalyst supports. In particular, Pd supported on triphenylphosphine-functionalized microporous knitting aryl network polymers (Pd@KAPs(Ph-PPh\textsubscript{3})) have attracted much attention. In this context, Lei et al. [82] have reported the efficient alkoxy carbonylation of aryl iodides with alcohols and phenols using this supported palladium catalyst in the presence of low pressure CO (1 bar). The corresponding alkoxy carbonylation products were obtained in moderate to excellent yields (74%–96%) at 80 °C in 6–10 h (Scheme 1). The Pd@KAPs(Ph-PPh\textsubscript{3}) catalyst was easily separated by simple filtration and recycled up to ten times without a significant decrease in activity being observed. The salient features of this protocol are the simplicity in catalyst handling, low CO pressure, negligible palladium leaching and good catalyst recyclability.

Pd-catalyzed carbonylation reactions have already become practical tools for the modern organic chemist and offer promising options for the preparation of carbonyl containing compounds. By incorporating one or even more molecules of CO into the parent structure, a carbon chain can be easily increased and the resulting products ready for further modification, which is a factor that holds its own importance. The advantages of carbonylation reactions mean that it is even more attractive to use carbonylations in the synthesis of biologically active heterocyclic compounds. In fact, an interesting and convenient procedure for the carbonylative synthesis of isoindoloquinazolinones was first described by Chen et al. [83]. They performed the carbonylative reactions in the presence of Pd(OAc)\textsubscript{2} at 120 °C from 1,2-dibromobenzenes and 2-aminobenzyl amine, which are readily available substrates. They isolated the isoindoloquinazolinone products in good yields (48%–84% in 16 h), while two CO molecules were installed (Scheme 10).

\begin{equation}
\text{NH}_{2} \quad \text{NH}_{2} \quad \begin{array}{c}
\text{Br} \\
\text{Br}
\end{array} \quad \begin{array}{c}
\text{Pd(OAc)_{2}}
\end{array} \quad \text{CO (2 bar) \quad base, DMA}
\end{equation}

\textbf{Scheme 10}. Pd-catalysed carbonylative synthesis of isoindoloquinazolinones.

The development of new synthetic strategies that work via direct C(sp\textsuperscript{3})-H bond functionalization is a hot topic in organic chemistry. It has recently been established that Pd(Xantphos)Cl\textsubscript{2} is able
to catalyse alkyl aromatic carbonylation to afford the arylacetic acid esters via C-H activation. In this context, Xie et al. [84] have reported a new synthetic route for arylacetamides based on C-H activation via a Pd-catalysed oxidative aminocarbonylation of alkyl aromatics with different amines in the presence of CO gas via C(sp^3)-H activation (Scheme 11).

![Scheme 11. Palladium-catalysed oxidative aminocarbonylation via C-H activation.](image)

To further explore the synthetic potential of this process, a series of alkyl aromatics were subjected to the reaction under optimized reaction conditions (CO 10 bar, 120 °C for 16 h). While substrates that bear electron-rich substituents on the aryl ring furnished the corresponding products in high yields (82%–85%), those containing an electron-withdrawing group were less reactive and afforded the desired amides in lower yields (52%–74%). It is worth noting that the tolerance of the halogens on the aromatic ring in this transformation offers an opportunity for subsequent cross-coupling, which facilitates the expedient synthesis of complex arylacetamides.

Transition-metal-catalysed carbonylations with the simultaneous introduction of a heteroatom are one of the most important tools for the synthesis of carbonyl compounds that bear heteroatom functional groups. Higuchi et al. [85] have proved that cobalt carbonyl (Co₂(CO)₈) is an excellent catalyst for the carbonylative cyclization of internal alkynes with CO. When Co₂(CO)₈-catalyzed reactions of internal alkynes with organic thiols are conducted in acetonitrile under CO pressure (4 MPa), the thiolative lactonization of internal alkynes successfully takes place in 17 h and two molecules of CO are incorporated at 140 °C (Scheme 12). Moreover, this carbonylation provides a useful tool with which to prepare the corresponding α,β-unsaturated γ-thio-γ-lactones (butenolide derivatives) in good yields (50%–70%). In the case of unsymmetrical alkynes, such as 2-octyne and 6-methyl-2-heptyne, thiolative lactonization proceeds with moderate regioselectivity to give the butenolide derivatives whose carbonyl group preferentially bonds to the less hindered acetylenic carbon. The present carbonylation of internal alkynes will open up new routes to the cobalt-catalysed reactions of heteroatom compounds with internal alkynes.

![Scheme 12. A highly selective cobalt-catalysed carbonylative cyclization of internal alkynes with CO and organic thiols.](image)

Most synthetic procedures for quinazolinone derivatives rely on the use of anthranilic acid or its derivatives as the starting materials and generally suffer from low yields, multistep reactions or
relatively harsh reaction conditions. The use of Pd/C as a heterogeneous catalyst for the carbonylation of 2-iodoanilines in the presence of trimethyl orthoformate and different amines was first proposed by Natte et al. \[86\]. Excellent yields of 4(3H)-quinazolinones were reported (84\%–96\%) using this four-component reaction approach (110 °C, 20 h) in presence of CO (10 bar) (Scheme 13).

The novelty of this procedure is that it avoids the use of expensive phosphine ligands as well as having the additional advantage of catalyst recovery. Furthermore, five new quinazolinone scaffolds that contain the trifluoroethyl group were introduced using this procedure and gram scale experiments were successfully reported as well.

This short overview highlights the broad scope of the potential applications of MW-assisted protocols under CO pressure in batch mode. Even though several CO and CO\(_2\) gas-liquid transformations have been optimized in flow-through reactors under conventional heating \[87\], so far MW flow chemistry remain a rather unexplored field of investigation \[53\]. This challenging technique should provide valuable guidelines for the scale-up of greener gas-liquid synthetic reactions procedures.

5. Conclusions

The conversion of CO\(_2\), an abundant renewable carbon reagent, and CO into chemicals of academic and industrial interest is of crucial relevance to creating a higher degree of sustainability in chemical processing and production. Moreover, the intensification of chemical processes, which means improving their efficiency and cutting down on energy consumption, requires increased automation and the use of non-conventional energy sources, as well as new, efficient and scalable protocols that can be implemented in continuous-flow reactors. This review aims to indicate some solutions, such as the combined use of gas and MW activation that will have potential applications in the near future of newly-born MW flow reactors.

Acknowledgments: The University of Turin is acknowledged for its financial support (Fondi Ricerca Locale 2014).

Author Contributions: Emanuela Calcio Gaudino and Laura Rinaldi first draft preparation, Laura Rotolo literature search and scheme draw, Diego Carnaroglio and Camillo Pirola literature selection and technical comments, Giancarlo Cravotto manuscript planning and revision.

Conflicts of Interest: The authors declare no conflict of interest.

References

1. Watson, W.J.W. How do the fine chemical, pharmaceutical, and related industries approach green chemistry and sustainability? Green Chem. \textbf{2012}, \textit{14}, 251–259. [CrossRef]
2. Beach, E.S.; Cui, Z.; Anastas, P.T. Green chemistry: A design framework for sustainability. Energy Environ. Sci. \textbf{2009}, \textit{2}, 1038–1049. [CrossRef]
3. Kondratenko, E.V.; Mul, G.; Baltrusaitis, J.; Larrazabal, G.O.; Pérez–Ramirez, J. Status and perspectives of CO\(_2\) conversion into fuels and chemicals by catalytic, photocatalytic and electrocatalytic processes. Energy Environ. Sci. \textbf{2013}, \textit{6}, 3112–3135. [CrossRef]
4. Liu, Q.; Wu, L.; Jackstell, R.; Beller, M. Using carbon dioxide as a building block in organic synthesis. Nat. Comms. \textbf{2015}. [CrossRef] [PubMed]
5. Mercadante, M.A.; Leadbeater, N.E. Development of methodologies for reactions involving gases as reagents: Microwave heating and conventionally-heated continuous-flow processing as examples. Green Process. Synth. \textbf{2012}, \textit{1}, 499–507.
6. Beller, M.; Wu, X.-F. \textit{Transition Metal Catalyzed Carbonylation Reactions}; Springer: Berlin, Germany, 2013; pp. 13–52.
7. Franke, R.; Selent, D.; Borner, A. Applied hydroformylation. Chem. Rev. \textbf{2012}, \textit{112}, 5675–5732. [CrossRef] [PubMed]
8. Liu, B.; Jiang, H.-Z.; Shi, B.-F. Pd(II)-catalyzed oxidative alkoxy-carbonylation of 2-phenoxypyridine derivatives with CO and alcohols. Org. Biomol. Chem. \textbf{2014}, \textit{12}, 2538–2542. [CrossRef] [PubMed]
9. Hermange, P.; Lindhardt, A.T.; Taaning, R.H.; Bjerglund, K.; Lupp, D.; Skrydstrup, T. Ex situ generation of stoichiometric and substoichiometric $^{12}$CO and $^{13}$CO and its efficient incorporation in palladium catalyzed aminocarbonylations. *J. Am. Chem. Soc.* 2011, 133, 6061–6071. [CrossRef] [PubMed]

10. Wu, X.F.; Neumann, H.; Spannenberg, A.; Schulz, T.; Jiao, H.; Beller, M. Development of a general palladium-catalyzed carboxylative Heck reaction of aryl halides. *J. Am. Chem. Soc.* 2010, 132, 14596–14602. [CrossRef] [PubMed]

11. Cai, M.; Zheng, G.; Zha, L.; Peng, J. Carbonylative Suzuki–Miyaura coupling of arylboronic acids with aryl iodides catalyzed by the MCM-41-supported bidentate phosphane palladium(II) complex. *Eur. J. Org. Chem.* 2009, 10, 1585–1591. [CrossRef]

12. Fusano, A.; Fukuyama, T.; Nishitani, S.; Inouye, T.; Ryu, I. Synthesis of alkyl alkylnyl ketones by Pd/light-induced three-component coupling reactions of iodoalkanes, CO, and 1-alkynes. *Org. Lett.* 2010, 12, 2410–2413. [CrossRef] [PubMed]

13. Wu, L.; Liu, Q.; Jackstell, R.; Beller, M. Carboxylations of alkenes with CO surrogates. *Angew. Chem. Int. Ed.* 2014, 53, 6310–6320. [CrossRef] [PubMed]

14. Konishi, H.; Manabe, K. Formic acid derivatives as practical carbon monoxide surrogates for metal-catalyzed carbonylation reactions. *Synlett* 2014, 25, 1971–1986. [CrossRef]

15. Natte, K.; Dumrath, A.; Neumann, H.; Beller, M. Palladium-catalyzed carbonylations of aryl bromides using paraformaldehyde: Synthesis of aldehydes and esters. *Angew. Chem. Int. Ed.* 2014, 53, 10090–10094. [CrossRef] [PubMed]

16. Ueda, T.; Konishi, H.; Manabe, K. Palladium-catalyzed reductive carbonylation of aryl halides with N-formylsaccharin as a CO source. *Angew. Chem. Int. Ed.* 2013, 52, 8611–8615. [CrossRef] [PubMed]

17. Wang, H.; Cai, J.; Huang, H.; Deng, G.-J. Palladium-Catalyzed phthalazinone synthesis using paraformaldehyde as carbon source. *Org. Lett.* 2014, 16, 5324–5327. [CrossRef] [PubMed]

18. Steffen, V.F.; Ulven, H.; Ulven, T. Oxalyl chloride as a practical carbon monoxide source for carbonylation reactions. *Org. Lett.* 2015, 17, 2832–2835. [CrossRef]

19. Christensen, S.H.; Olsen, E.P.K.; Rosenbaum, J.; Madsen, R. Hydroformylation of olefins and reductive carbonylation of aryl halides with syngas formed ex situ from dehydrogenative decarbonylation of hexane-1,6-diol. *Org. Biomol. Chem.* 2015, 13, 938–945. [CrossRef] [PubMed]

20. Schareina, T.; Zapf, A.; Gotta, C.M.; Beller, M. An improved protocol for palladium-catalyzed alkoxy carbonylations of aryl chlorides with alkyl formats. *Adv. Synth. Catal.* 2010, 352, 1205–1209. [CrossRef]

21. Nouira, I.; Kostakis, I.K.; Dubouilh, C.; Chosson, E.; Iannelli, M.; Besson, T. Decomposition of formamide assisted by microwaves, a tool for synthesis of nitrogen-containing heterocycles. *Tetrahedron Lett.* 2008, 49, 7033–7036. [CrossRef]

22. Loidreau, Y.; Besson, T. Microwave-assisted thermal decomposition of formamide: A tool for coupling a pyrimidine ring with an aromatic partner. *Tetrahedron* 2011, 67, 4852–4857. [CrossRef]

23. Wan, Y.; Alterman, M.; Larhed, M.; Hallberg, A. Formamide as a combined ammonia synthon and carbon monoxide source in fast palladium-catalyzed aminocarbonylations of aryl halides. *J. Comb. Chem.* 2003, 5, 82–84. [CrossRef]

24. Sawant, D.N.; Wagh, Y.S.; Bhatte, K.D.; Bhanage, B.M. Palladium-catalyzed carbon-monoxide-free aminocarbonylation of aryl halides using N-substituted formamides as an amide source. *J. Org. Chem.* 2011, 76, 5489–5494. [CrossRef] [PubMed]

25. Lindsay, C.M.; Widdowson, D.A. Organic synthesis with carbon monoxide: The synthesis of carbamoylstannanes and aromatic carbamoylation. *J. Chem. Soc. Perkin Trans. 1* 1988, 1, 569–573. [CrossRef]

26. Cunico, R.F.; Pandey, R.K. Palladium-catalyzed conversion of benzylic and allylic halides into α-aryl and β,γ-unsaturated tertiary amides by the use of a carbamoylsilane. *J. Org. Chem.* 2005, 70, 9048–9050. [CrossRef] [PubMed]

27. Wieckowska, A.; Fransson, R.; Odell, L.R.; Larhed, M. Microwave-assisted synthesis of Weinreb and MAP aryl amides via Pd-catalyzed Heck aminocarbonylation using Mo(CO)$_3$ or W(CO)$_3$. *J. Org. Chem.* 2011, 76, 978–981. [CrossRef] [PubMed]

28. Cokoja, M.; Bruckmeier, C.; Rieger, B.; Herrmann, W.A.; Kuhn, F.E. Transformation of carbon dioxide with homogeneous transition-metal catalysts: A molecular solution to a global challenge? *Angew. Chem. Int. Ed.* 2011, 50, 8510–8537. [CrossRef] [PubMed]
29. Meng, X.L.; Nie, Y.; Sun, J.; Cheng, W.G.; Wang, J.Q.; He, H.Y.; Zhang, S.J. Functionalized dicyandiamide-formaldehyde polymers as efficient heterogeneous catalysts for conversion of CO₂ into organic carbonates. *Green Chem.* 2014, 16, 2771–2778. [CrossRef]

30. Xu, B.H.; Wang, J.Q.; Sun, J.; Huang, Y.; Zhang, J.P.; Zhang, X.P.; Zhang, S.J. Fixation of CO₂ into cyclic carbonates catalyzed by ionic liquids: A multi-scale approach. *Green Chem.* 2015, 17, 108–122. [CrossRef]

31. Cokoja, M.; Wilhelm, M.E.; Anthofer, M.H.; Herrmann, W.A.; Kuhn, F.E. Synthesis of cyclic carbonates from epoxides and carbon dioxide by using organocatalysts. *ChemSusChem* 2015, 10, 2436–2454. [CrossRef] [PubMed]

32. North, M.; Pasquale, R.; Young, C. Synthesis of cyclic carbonates from epoxides and CO₂. *Green Chem.* 2010, 12, 1514–1539. [CrossRef]

33. Hu, J.; Ma, J.; Zhu, Q.; Zhang, Z.; Wu, C.; Han, B. Transformation of atmospheric CO₂ catalyzed by protic ionic liquids: Efficient synthesis of 2-oxazolidinones. *Angew. Chem. Int. Ed.* 2015, 54, 5399–5403. [CrossRef] [PubMed]

34. Wang, W.H.; Himeda, Y.; Muckerman, J.T.; Manbeck, G.F.; Fujita, E. CO₂ Hydrogenation to formate and methanol as an alternative to photo- and electrochemical CO₂ reduction. *Chem. Rev.* 2015, 115, 12936–12973. [CrossRef] [PubMed]

35. Kortlever, R.; Shen, J.; Schouten, K.J.P.; Calle-Vallejo, F.; Koper, M.T.M. Catalysts and reaction pathways for the electrochemical reduction of carbon dioxide. *J. Phys. Chem. Lett.* 2015, 6, 4073–4082. [CrossRef] [PubMed]

36. Bai, D.; Wang, Q.; Song, Y.; Li, B.; Jing, H. Synthesis of cyclic carbonate via epoxide and CO₂ catalyzed by magnetic nanoparticle-supported porphyrin. *Catal. Commun.* 2011, 12, 684–688. [CrossRef]

37. Mikkelsen, M.; Jørgensen, M.; Krebs, F.C. The tetron challenge. A review of fixation and transformation of carbon dioxide. *Energy Environ. Sci.* 2010, 3, 43–81. [CrossRef]

38. Kirschning, A.; Solodenko, W.; Mennecke, K. Combining enabling techniques in organic synthesis: Continuous flow processes with heterogenized catalysts. *Chem. Eur. J.* 2006, 12, 5972–5990. [CrossRef] [PubMed]

39. Cravotto, G.; Bonrath, W.; Tagliapietra, S.; Speranza, C.; Calcio Gaudino, E.; Barge, A. Intensification of organic reactions with hybrid flow-reactors. *Chem. Eng. Process. Process Intensif.* 2010, 49, 930–935. [CrossRef]

40. Gedye, R.; Smith, F.; Westaway, K.; Ali, H.; Baldisera, L.; Laberge, L.; Roussell, J. The use of microwave ovens for rapid organic synthesis. *Tetrahedron Lett.* 1986, 27, 279–282. [CrossRef]

41. Giguere, R.J.; Bray, T.L.; Duncan, S.M.; Majetich, G. Application of commercial microwave ovens to organic synthesis. *Tetrahedron Lett.* 1986, 27, 4945–4948. [CrossRef]

42. De La Hoz, A.; Loupy, A. *Microwaves in Organic Synthesis*, 3rd ed.; Springer: New York, NY, USA, 2012.

43. Moseley, J.D.; Kappe, C.O. A critical assessment of the greenness and energy efficiency of microwave-assisted organic synthesis. *Green Chem.* 2011, 13, 794–806. [CrossRef]

44. Durka, T.; Gerven, T.V.; Stankiewicz, A. Microwaves in heterogeneous gas-phase catalysis: Experimental and numerical approaches. *Chem. Eng. Technol.* 2009, 32, 1301–1312. [CrossRef]

45. Cravotto, G.; Cintas, P. The combined use of microwaves and ultrasound: Improved tools in process chemistry and organic synthesis. *Chem. Eur. J.* 2007, 13, 1902–1909. [CrossRef] [PubMed]

46. Taktak, F.; Bulduk, I. Microwave-assisted hydrogenation of codeine in aqueous media. *ISRN Org. Chem.* 2012. [CrossRef] [PubMed]

47. Petricci, E.; Mann, A.; Schoenfelder, A.; Rota, A.; Taddei, M. Microwaves make hydroformylation a rapid and easy process. *Org. Lett.* 2006, 8, 3725–3727. [CrossRef] [PubMed]

48. Watson, D.A.; Fan, X.; Buchwald, S.L. Carbonylation of aryl chlorides with oxygen nucleophiles at atmospheric pressure. Preparation of phenyl esters as acyl transfer agents and the direct preparation of alkyl esters and carboxylic acids. *J. Org. Chem.* 2008, 73, 7096–7101. [CrossRef] [PubMed]

49. Friis, S.D.; Skyrdstrup, T.; Buchwald, S.L. Mild Pd-catalyzed aminocarbonylation of (hetero)aryl bromides with a palladacycle precatalyst. *Org. Lett.* 2014, 16, 4296–4299. [CrossRef] [PubMed]

50. Caporaso, M.; Cravotto, G.; Georgakopoulos, S.; Heropoulos, G.; Martina, K.; Tagliapietra, S. Pd/C-catalyzed aerobic oxidative esterification of alcohols and aldehydes: A highly efficient microwave-assisted green protocol. *Beilstein J. Org. Chem.* 2014, 10, 1454–1461. [CrossRef] [PubMed]

51. Bergamelli, F.; Iannelli, M.; Marafie, J.A.; Moseley, D. A commercial continuous flow microwave reactor evaluated for scale-up. *Org. Process Res. Dev.* 2010, 14, 926–930. [CrossRef]
52. Rinaldi, L.; Carnaroglio, D.; Rotolo, L.; Cravotto, G. A microwave-based chemical factory in the lab: From milligram to multigram preparations. J. Chem. 2015. [CrossRef]

53. Choedkiatsakul, I.; Ngaosuwan, K.; Assabumrungrat, S.; Mantegna, S.; Cravotto, G. Biodiesel production in a novel continuous flow microwave reactor. Renew. Energy 2015, 83, 25–29. [CrossRef] [PubMed]

54. Matsuzawa, M.; Togashi, S. Microwaves in Catalysis: Methodology and Applications; Wiley-VCH: Weinheim, Germany, 2015; pp. 141–154.

55. Schaffner, B.; Schaffner, V.; Verevkin, S.P.; Borner, A. Organic carbonates as solvents in synthesis and catalysis. Chem. Rev. 2010, 110, 4554–4811. [CrossRef] [PubMed]

56. Nüchter, M.; Ondruschka, B.; Hoffmann, J.; Tied, A. Microwave-assisted reactions of oxiranes with carbon dioxide in ionic liquids. Stud. Surf. Sci. Catal. 2004, 153, 131–136.

57. Leadbeater, N.E.; Torenius, H.M.; Tye, H. Microwave–promoted organic synthesis using ionic liquids: A mini review. Comb. Chem. High Throughput Screen. 2004, 7, 511–528. [CrossRef]

58. Ono, F.; Oiao, K.; Tomida, D.; Yokoyama, C.J. Rapid synthesis of cyclic carbonates from CO$_2$ and epoxides under microwave irradiation with controlled temperature and pressure. Mol. Catal. A Chem. 2007, 263, 223–226. [CrossRef]

59. Dharman, M.M.; Hye-Ji Choi, H.J.; Park, S.W.; Park, D.W. Microwave assisted synthesis of cyclic carbonate using homogeneous and heterogeneous ionic liquid catalysts. Top. Catal. 2010, 53, 462–469.

60. Dharman, M.M.; Choi, H.J.; Kim, D.W.; Park, D.W. Synthesis of cyclic carbonate through microwave irradiation using silica-supported ionic liquids: Effect of variation in the silica support. Catal. Today 2011, 164, 544–547. [CrossRef]

61. Palmisano, G.; Bonrath, W.; Boffa, L.; Garella, D.; Barge, A.; Cravotto, G. Heck Reactions with Very Low Ligandless Catalyst Loads Accelerated by Microwaves or Simultaneous Microwaves/Ultrasound Irradiation. Adv. Synth. Catal. 2007, 349, 2338–2344. [CrossRef]

62. Tharun, J.; Mathai, G.; Kathalikkattil, A.C.; Roshan, R.; Kwaka, J.Y.; Park, D.W. Microwave-assisted synthesis of cyclic carbonate using a formic acid/KI catalytic system. Green Chem. 2013, 15, 1673–1677. [CrossRef]

63. Tharun, J.; Kathalikkattil, A.C.; Roshan, R.; Kang, D.H.; Woo, H.C.; Park, D.W. Microwave-assisted, rapid cycloaddition of allyl glycidyl ether and CO$_2$ by employing pyridinium-based ionic liquid catalysts. Catal. Commun. 2014, 54, 31–34. [CrossRef]

64. Chun, Y.; He, Y.G.; Zhu, J.H. Microwave-assisted synthesis of dimethyl carbonate. React. Kinet. Catal. Lett. 2001, 74, 23–27. [CrossRef]

65. Du, J.; Shi, J.; Li, Z.; Liu, Z.; Fan, X.; Tao, C. Ionic liquid mediated CO$_2$ activation for DMC synthesis. J. Nat. Gas Chem. 2012, 21, 476–479. [CrossRef]

66. Carnaroglio, D.; Martina, K.; Palmisano, G.; Penoni, A.; Domini, C.; Cravotto, G. One-pot sequential synthesis of isocyanates and urea derivatives via a microwave-assisted Staudinger-aza-Wittig reaction. Beilstein J. Org. Chem. 2013, 9, 2378–2386. [CrossRef] [PubMed]

67. Mägerlein, W.; Indolese, A.F.; Beller, M. Alkoxycarbonylation of aryl iodides using gaseous carbon monoxide and pre-pressurized reaction vessels. Angew. Chem. Int. Ed. 2007, 46, 4114–4133. [CrossRef] [PubMed]

68. Kormos, C.M.; Leadbeater, N.E. Microwave-promoted hydroxy-carbonylation in water using gaseous carbon monoxide and pre-pressurized reaction vessels. Synlett 2006, 11, 1663–1666. [CrossRef]

69. Iannelli, M.; Bergamelli, F.; Kormos, C.M.; Paravisi, S.; Leadbeater, N.E. Application of a batch microwave unit for scale-up of alkoxy-carbonylation reactions using a near-stoichiometric loading of carbon monoxide. Org. Process Res. Dev. 2009, 13, 634–637. [CrossRef]

70. Kormos, C.M.; Leadbeater, N.E. Alkoxy-carbonylation reaction performed using near stoichiometric quantities of CO. Synlett 2007, 13, 2006–2010. [CrossRef]

71. Kormos, C.M.; Leadbeater, N.E. Alkoxy-carbonylation of aryl iodides using gaseous carbon monoxide and pre-pressurized reaction vessels in conjunction with microwave heating. Org. Biomol. Chem. 2007, 5, 65–68. [CrossRef] [PubMed]

72. Kappe, C.O.; Stadler, A.; Dallinger, D. Microwaves in Organic and Medicinal Chemistry, 2nd ed.; Wiley-VCH: Weinheim, Germany, 2013; pp. 41–81, ISBN: 3527647848.
74. Salvadori, J.; Balducci, E.; Zaza, S.; Petricci, E.; Taddei, M. Microwave-assisted carbylation and cyclocarbonylation of aryl iodides under ligand free heterogeneous catalysis. J. Org. Chem. 2010, 75, 1841–1847. [CrossRef] [PubMed]

75. Liptrot, D.; Alcaraz, L.; Roberts, B. New synthesis of aryl and heteroaryl N-acylureas via microwave-assisted palladium-catalysed carbonylation. Adv. Synth. Catal. 2010, 352, 2183–2188. [CrossRef]

76. Pizzetti, M.; Russo, A.; Petricci, E. Microwave-assisted aminocarbonylation of ynamides by using catalytic [Fe₃(CO)₁₂] at low pressures of carbon monoxide. Chem. Eur. J. 2011, 17, 4523–4528. [CrossRef] [PubMed]

77. Calcio Gaudino, E.; Carnaroglio, D.; Martina, K.; Palmisano, G.; Penoni, A.; Cravotto, G. Highly efficient microwave-assisted CO aminocarbonylation with a recyclable Pd(II)/TPP-β-cyclodextrin cross-linked catalyst. Org. Process Res. Dev. 2015, 19, 499–505. [CrossRef]

78. Tlili, A.; Blondiaux, E.; Frogneux, X.; Cantat, T. Reductive functionalization of CO₂ with amines: An entry to formamide, formamidine and methylamine derivatives. Green Chem. 2015, 17, 157–168. [CrossRef]

79. Zhang, L.; Han, Z.; Zhao, X.; Wang, Z.; Ding, K. Highly Efficient Ruthenium-Catalyzed N-Formylation of Amines with H₂ and CO₂. Angew. Chem. Int. Ed. 2015, 54, 6186–6189. [CrossRef] [PubMed]

80. Takada, Y.; Foo, S.W.; Yamazaki, Y.; Saito, S. Catalytic fluoride triggers dehydrative oxazolidinone synthesis from CO₂. RSC Adv. 2014, 4, 50851–50857. [CrossRef]

81. Higashimae, S.; Tamai, T.; Nomoto, A.; Ogawa, A. Selective Thiolative Lactonization of Internal Alkynes Bearing a Hydroxyl Group with Carbon Monoxide and Organic Disulfides Catalyzed by Transition-Metal Complexes. J. Org. Chem. 2015, 80, 7126–7133. [CrossRef] [PubMed]

82. Lei, Y.; Wu, L.; Zhang, X.; Mei, H.; Gu, Y.; Li, G. Palladium supported on triphenylphosphine functionalized porous organic polymer: A highly active and recyclable catalyst for alkoxycarbonylation of aryl iodides. J. Mol. Catal. A Chem. 2015, 398, 164–169. [CrossRef]

83. Chen, J.; Neumann, H.; Beller, M.; Wu, X.-F. Palladium-catalyzed synthesis of isoindoloquinazolinones via dicarbonylation of 1,2-dibromoarenes. Org. Biomol. Chem. 2014, 12, 5835–5838. [CrossRef] [PubMed]

84. Xie, P.; Xia, C.; Huang, H. Palladium-catalyzed oxidative aminocarbonylation: A new entry to amides via C-H activation. Org. Lett. 2013, 15, 3370–3373. [CrossRef] [PubMed]

85. Higashimae, S.; Higashimae, S.; Tamai, T.; Ogawa, A. A highly selective cobalt-catalyzed carbonylative cyclization of internal alkynes with carbon monoxide and organic thiols. Tetrahedron 2013, 69, 11197–11202. [CrossRef]

86. Natte, K.; Neumann, H.; Wu, X.-F. Pd/C as an efficient heterogeneous catalyst for carbonylative four-component synthesis of 4(3H)-quinazolinones. Catal. Sci. Technol. 2015, 5, 4474–4480. [CrossRef]

87. Mallia, C.J.; Baxendale, I.R. The use of gases in flow synthesis. Org. Process Res. Dev. 2015. [CrossRef]