Impact of Microwaves on Organic Synthesis and Strategies toward Flow Processes and Scaling Up

Katia Martina, Giancarlo Cravotto,* and Rajender S. Varma*

ABSTRACT: Microwave-assisted organic synthesis has been widely studied and deliberated, opening up some controversial issues as well. Nowadays, microwave chemistry is a mature technology that has been well demonstrated in many cases with numerous advantages in terms of the reaction rate and yield. The strategies toward scaling up find an ally in continuous-flow reactor technology comparing dielectric and conductive heating.

In the more than 30 years of microwave (MW) chemistry, many green protocols in organic synthesis have been reported, showing significant advantages compared to the conventional heating. Selective, volumetric dielectric heating has resulted in time and energy savings, enabled the deployment of safer solvents or solvent-free processes, selective catalysis in fewer steps, and generally attaining higher selectivity and yields.1 Although in the beginning, an improper temperature measurement sometimes overestimated the positive effect on the reaction rate, a fast and efficient energy transfer was demonstrated; however, for years, MW applications in organic synthesis were limited to the laboratories of a limited number of experts. The paucity of data available pertaining to the dielectric properties of materials as a function of temperature and frequency generated an empirical approach. In the meantime, the design of MW reactors was mainly focused on applications in analytical chemistry because of the rapid diffusion in sample mineralization. The lack of dedicated equipment often led to scarcely reproducible results and varying kinetics, which limited scaling-up studies for the industrial applications.

Despite the evident production efficiency under dielectric heating, the interest in industrial processing was delayed by the demand for safer equipment and plants.2 The candidates for further development were the most efficient and reproducible protocols.

However, continuous-flow synthetic processes were shown to be less flexible when working with different reactions. After the optimization of the residence time, mixing, and flow rate, the reaction could be easily scaled up with parallel reactors or in bigger size. In all cases, the main goal was to preserve the reaction kinetics.3 Aiming to improve heat and mass transfer, a progressive transition from batch to continuous processing is expected, which is a mandatory condition for exploiting the MW energy in industrial production. MW-assisted continuous-flow organic synthesis (MACOS) has been successfully investigated in a plethora of organic reactions (e.g., heterocycle synthesis and metal-catalyzed chemistry) and, in particular, for the semi-industrial preparation of aromatic compounds.4 Numerous studies on MW-assisted synthesis have nowadays revealed the possibility of transferring the methodology from the MW batch mode to a conventionally heated or MW-heated flow mode.

Because this field of research has been previously reviewed and a number of publications have reported the advances in MW-assisted organic synthesis (MAOS) in batch and in continuous flow mode,5−10 this Perspective has focused on recent studies with an emphasis on the advantages and limitations of MW heating. In this regard, synthetic protocols aimed at scaling up of the process and the comparison between different approaches directed to large-scale synthesis have been selected to provide a comprehensive picture of the state of the art. Recent advancements in stereoselective MAOS are included as well.

Special Issue: Enabling Techniques for Organic Synthesis

Received: April 14, 2021
Published: June 14, 2021
**RECENT ADVANCES IN MICROWAVE TECHNOLOGY**

The use of alternative energy sources to improve mass or heat transfer has tremendous potential for chemical reactions, and dielectric heating may strongly enhance the reaction rate.

Scrutinizing the chemical engineering advancements to intensify synthetic protocols under MW irradiation, special attention is being paid to the design of the MW cavities that provides a homogeneous field and uniform heating.11

Single-mode MW instruments have higher energy efficiency when compared to multimode counterparts, as the reactor vessel geometry and position enable a well-defined field. A monomodal MW instrument is well suited for the mechanism study of the chemical reaction, but only relatively small vial diameters may receive/adsorb significant power.12

A recent study reported a fixed bed of 200 mg of NaY zeolite in a monomode rectangular resonant cavity wherein a temperature gradient was correlated to the nonuniform electric-field distribution.13 As presented in the Figure 1, the region at the higher temperature corresponds to the higher electric field region, and almost 60% of the sample has a temperature from 160 to 197 °C.

The attenuation of the electromagnetic wave, measured at a small penetration depth of the most adsorbing solvent, is another limitation of MW irradiation when large-scale synthesis is studied. Different approaches have been pursued to overcome this limitation, and the combination of MW irradiation with acoustic waves (ultrasound) may help generate an increase in the penetration depth beyond intensifying the mixing and mass transfer.14,15 The mechanical stirring would yield volumetric heating, and an example of a large-scale MW batch reactor devoted to synthesis has been described.16

Understanding how the MW deployment can enhance the reaction rates in organic synthesis requires a good background with knowledge of MW interactions with matter and other fundamentals. Moreover, a strict control of the applied energy with suitable hardware and software and reliable tools for temperature and pressure measurement is necessary. A rational balance of the pros and cons of MW energy deployment is relevant in deciding when and where MW irradiation can replace other energy input sources, and for this purpose, a study by Mase et al. reported a method that combines experimental design and 3D surface approximation for the rapid optimization of reaction conditions.17 The principles behind and the factors determining the successful scale-up of MW-assisted technology are the frequency, power, penetration depth, and energy consumption; thus studies in this field are collected in this comprehensive review focused on instrument design and reactor configuration.18–20

Modern MW technology can finely tune the electromagnetic radiation at high field strength and frequency, leading to fast and safe heating. MW heating can, a priori, be independent of the macroscopic temperature; MW penetrability is dependent on the frequency. National and international institutions allow only a few frequencies for laboratory, industry, and medical (ISM) applications. The main frequency bands for commercially
available equipment are 2.45 GHz (most common), 915 MHz (USA), and 896 MHz (U.K.). Although equipment at 5.8 GHz is presently being created, an industrial setup requires further developmental work. MW irradiation at 5.8 GHz displays a shorter penetration depth but a better energy conversion. The selected frequency impacts the dielectric constants and the dissipation factors ($\tan \delta$). Apart from MW reactors working at 2.45 GHz, all other devices require ad hoc designs because they are not available from commercial suppliers.21

Relevant advances in the “kilolab” continuous-flow MW mode have been reported by Morschhäuser et al. using a transmission-line short-circuited waveguide unit, combining the features of mono- and multimode systems.22,23

The new generation of MW reactors have come a long way in terms of efficiency and compatibility with the continuous-flow systems; they can promote chemical reactions at varying temperatures up to 300 °C and pressures up to 200 bar with safety as a priority and excellent parameter controls.24 The combination of several modes in the direction of the reaction flow allows a homogeneous field distribution in resonator cavities, therefore, the continuous reactions have advantages related to MW irradiation and flow chemistry (Figure 2).

Two examples of the evaluation of MW applicators for flow synthesis are represented in Figures 3 and 4. A resonator-type MW reactor for continuous flow mode is depicted in Figure 3. The system comprises a rectangular MW cavity equipped with a helical tubular borosilicate glass reactor with an internal volume of 1.0 mL.25 In Figure 4, another example of an MW tubular reactor heated by means of an Al–Cu helical antenna is represented and, in particular, the IR camera image that represents the temperature is shown.26 The temperature is rather homogeneous, and lower temperatures are registered at the top and bottom of the reactor, which correspond to the entrance and exit of the flow.

Large laboratory MW units have been produced to achieve the kilogram scale in loop, in stopped flow or continuous flow, and with parallel vessels; nevertheless, these systems have a few inherent limitations.27

The industry is adopting strict strategies to minimize risks and to avoid batch failures, thus moving toward the continuous-flow synthetic processes. Proper modeling and simulation tools are fundamental for the design of safe and efficient reactors. This would help to get rid of bottlenecks that currently prevent scaling up. In addition to the reaction behavior during the reaction course, modern MW reactors should strictly monitor all of the parameters. Several variables affect the output of MW reactor and its scale, reflecting that MW technology cannot be implemented in a straightforward manner like classic conductive heating.

Industrial MW installations have been developed by specialized companies on the basis of the chemical conversion and the productivity required. Equipment manufacturers, who offer on-demand specifications (reactor geometry, frequency, power, etc.), can then make recommendations and offers.

■ IMPACT OF HIGH TEMPERATURE AND PRESSURE ON HOMOGENEOUS ORGANIC SYNTHESIS

Recent examples of batch and continuous MW organic synthesis take advantage of modern instruments in which the temperature and pressure are controlled. Esterification, aromatic nucleophilic substitution, and Claisen rearrangement, among others, have been optimized under high pressure and consequently at higher temperature.
been successfully exploited. Comparing different studies also underlined the importance of the MW reactor design, and the depth of the MW irradiation has sometimes been measured. The typical limitations due to MW absorbance and its conversion have been largely centered on continuous operations with a thinner reaction cell, tube, or tubular coil.

In 2017, the esterification of leucine with PTSA (p-toluene sulfonic acid) and butanol was efficiently performed in an MW reactor (Scheme 1A, compound 1). The authors compared the Fischer–Speier esterifications performed in a round-bottomed flask or in two types of tubular reactors of different inner diameters and lengths. By means of the in situ measurement of the dielectric properties of the reaction mixture in methanol, the penetration depth was calculated (~13 mm), and the reaction was therefore performed in MW batch mode and continuous-flow mode with an MW oven equipped with tubular reactors made of glass or Teflon with varying dimensions (Scheme 1A). In the case of a large-scale reaction with a Teflon tube deployment, the authors observed that the highest impact on the reaction conversion was correlated to the inner diameter of the reactor. With a 40 mm internal diameter, the beneficial effect of flow was apparent, but the reaction was not complete. As depicted in Scheme 1A, the full conversion was attained when the MW batch reaction was performed at reflux with a thinner tubular reactor heated to 120–140 °C. Esterification that was performed in a conventional batch mode gave only 64% conversion. Furthermore, this study demonstrated that the MW-assisted flow reaction could be successfully scaled up to a tubular system of a ~600 mL inner volume with a flow of 20 mL min⁻¹ at an elevated temperature of 140 °C, but an optimal reactor tube was necessary to secure the highest yield.

The esterification of benzoic acid under MW irradiation has been proven to be fast and efficient, and a few studies reported this reaction in the presence of acid catalysts. The most recent protocol of the continuous MW-assisted esterification of benzoic acid in the presence of PTSA was optimized at 140 °C on the basis of an MW-assisted protocol studied in a batch mode. On the basis of the order of reactivity of alcohols (nBuOH, iBuOH > nPrOH, nPentOH, iOctOH > EtOH > iPrOH), the authors accomplished the reaction in flow mode with a variable residence time of 20 to 30 min (Scheme 1B). The reaction was performed in a CEM Discover focused MW synthesizer equipped with a CEM 10 mL flow-cell accessory (irradiated volume 7 mL), wherein 35 mmol of benzoic acid was converted to its ester form in >95% yield. Another example of the esterification of levulinic acid using sulfuric acid or PTSA as the catalyst in MW batch mode and in conventional flow mode performed above the atmospheric boiling point of the solvent has been described. MW heating provided complete conversion at 120 °C in 5 min with catalyst loading of 2.5 mol % and a levulinic acid–ethanol ratio of 1:10 (Scheme 1C). To mimic the batch reaction conditions in conventional flow mode, two solutions of acid and of levulinic acid in ethanol were pumped in a 10 mL perfluorooalkoxy alkane (PFA) coil that could operate at up to 150 °C. When the reaction was performed with 2 mL/min flow, a conversion of 78% was observed; therefore,
reducing the flow rate to 1 mL/min was found to be necessary to attain complete conversion in one pass.

Large interest in organophosphorus transformations has garnered the attention of the MW-assisted protocol, and the esterification of phenyl-H-phosphinic acid took advantage of the high temperature, namely, in the alcoholysis of dimethyl phosphonate. Both MW reactions were performed and scaled up in MW/flow in a flow cell housed in a CEM MW reactor. The esterification was promoted by the addition of [bmim][PF6], which exhibited activity as an enhancer of MW irradiation, whereas transesterification generally did not require catalyst addition (Scheme 2A). Butanol reacted at 175–200 °C in 30 min in an MW flow cell of 10 mL, and the authors showed that roughly the same conversion was obtained in both the batch and flow modes (Scheme 2B).

The investigation of the transcarbamylation/transesterification of sulfonyl carbamate evidenced that the MW flow procedures shortened the reaction time when compared to the MW procedure performed at the boiling point of the solvent. The reaction was shortened from 20 min to 40 s when butanol was heated from 120 °C in MW batch mode to 180 °C in MW flow mode (Scheme 3).

The SNAr reaction has been extensively studied by means of MW irradiation in a preliminary attempt to scale up the aromatic nucleophilic substitution of chloroarenes. The MW batch procedure could be adapted to a stop-flow continuous MW protocol. The reaction was performed between 1,2-dichloro 4-nitrobenzene and 4-methoxy phenol in an 80 mL reaction vessel (50 mL of operating volume) in the presence of an organic base (DBU) in DMA, and the vessel was filled and emptied along different lines (Scheme 4A, reaction conditions above). The automated repetition of the small-batch MW protocol ensured the large-scale production of product 7. In 2011, the Watts group translated the methodology to a continuous-flow microreactor, enabling a reduction in the reaction time from 10 min to 60 s by means of a glass microreactor of 10 μL volume (Scheme 4A, reaction conditions below). Other advantages of the microreactor have been exploited because under pressure, a high-boiling solvent such as DMA could be replaced with MeCN. This could speed up the reaction outcome and the product isolation.

To extend the scope of MW continuous-flow SNAr, a two-step procedure via an epoxide opening/SNAr cyclization was reported by Organ et al. for the synthesis of benzo fused sultams (Scheme 4B). The optimal protocol in batch mode was performed under MW irradiation in DMSO at 180 °C in 1 min, and DBU and tBuOK were selected as the optimal bases. Scaling up in continuous-flow mode by means of a single-mode Biotage Smith Creator synthesizer equipped with a 1700 μm (int Ø) borosilicate capillary allowed the production of 12 derivatives on a 2 to 3.5 g scale in a 55% average yield. To avoid clogging, the reactions were performed in the presence of DBU.

Claisen rearrangement is accelerated under high-temperature and high-pressure conditions, and the efficacy of MW irradiation to enhance the reaction conversion has been exploited in continuous MW synthesis. Horikoshi et al. studied the influence of MW irradiation on the conversion of 1-allyloxy-4-methoxybenzene (Scheme 5A, reaction conditions above), as they observed that no significant improvement could be discerned when the reaction was performed under conventional conditions or under MW heating in DMSO. Neat conditions enabled the increase in the yield from 2.5 to 36%, and a two-fold yield improvement was observed compared to the reaction heated in an oil bath. A similar study was reported by Larhed et al. to access the applicability of a nonresonant MW system for the continuous MW/flow chemistry. As depicted in the Scheme 5A (reaction conditions below), the reaction has been performed in N-methyl-2-pyrrolidone (NMP) and neat at 270 °C, proving the efficacy of higher temperature conditions. In 5 to 15 min, the 2-allyl phenol was obtained in 78 to 85% yield,
respectively, with a calculated throughput of 13.6 and 15.0 mmol/h.

Another study focused on a resonator-type MW reactor in which a helical tubular borosilicate glass reactor was placed within a rectangular resonant cavity where the electromagnetic wave intensity was maintained at the maximum. The Claisen reaction of alkyloxy naphthalene was tested as a model reaction, and a detailed evaluation of the effects of the MW power, concentration, and flow rate were described by Koyama et al. An extremely efficient protocol was described in cyclopentyl methyl ether (CPME) with the substrate dissolved at an optimal concentration of 2.0 M in the presence of a back pressure of 2.5 MPa. The solution was heated to 232 °C and flowed at 1 mL/min with a residence time of 1 min to obtain the desired product in >95% yield (Scheme 5B). The scaling up of the reaction was validated with 11 g of starting material that reacted in 30 min, and a productivity of 20.26 g/h was attained.

The Johnson–Claisen rearrangement has been demonstrated to be a suitable option for the MW continuous-flow protocol. Allyl alcohol and triethyl orthoacetate can be converted to ethyl pent-4-enoate with an excellent productivity of 89.5 g/h (Scheme 5C). The desired product was collected in high purity with a residence time of ~1.5 min. 10 to 25 mol % acetic acid was necessary to obtain the product in good yield while the reaction temperature was maintained at 224 °C.

Many examples of the Diels–Alder reaction have been reported in MW irradiation and in flow mode. The beneficial effect of MW irradiation was exploited to optimize the protocol and then was applied to the conventional flow mode. In 2012, Abele et al. reported the Diels–Alder reaction of (cyclohexa-1,5-dien-1-yloxy)trimethylsilane with acrylonitrile and observed that under MW irradiation the side reaction of acrylonitrile polymerization was highly reduced when compared to the conventional oil bath (Scheme 6A). Because the polymerization of acrylonitrile leads to microreactor clogging, the optimization of MW irradiation was performed at different temperatures up to 175 °C, and the authors observed that in 2 h, at 175 °C, acrylonitrile was completely consumed, but 6–8 mol/mol % of dienophile was still present. Therefore, they decided to translate the procedure in flow mode in the presence of excess flow. Working with a stainless-steel tube of 4.5 mL, the reaction was scaled up in flow under solvent-free conditions. The reaction temperature was increased to 30 °C, and the reaction time was decreased to 2 min. When 2 equiv of dienophile was used, in 1.25 min, the reaction was complete, and no precipitation was observed. The conventional continuous flow was performed at 215 °C with a residence time of 60 s to generate 164 g of compound 12 in 45 min in 95% recovery.

The Diels–Alder cycloaddition between furan and diethyl acetylenedicarboxylate has been reported as a model reaction to assess the performance of two different continuous MW equipment setups. Two recent studies reported the reaction performed on mesoscopic scale. A 10 mL MW vessel tapered to a thread-inlet was used as a reactor in one case, whereas the other case exploited a helical tubular borosilicate glass reactor of 5.5 to 6.0 mL placed in the resonant cavity (Scheme 6B). In the first case, the reaction was performed at 150 °C for 10 min under neat conditions, which revealed that the MW/flow reaction yield was lower when compared with the conventional batch reaction. The 10 mL flow cell was therefore packed with a strong MW adsorber (10% w/w Fe3O4/sand) to increase the sample temperature, and an increase in the reaction yield from 60 to 70% was observed, but it was still lower than the reaction conventionally performed batch mode in the absence of the matrix (Scheme 6B, reaction conditions above). The second study reported the optimization of an MW-assisted flow Diels–Alder reaction in a resonant cavity. The optimal reaction conditions were obtained by varying the flow rate, the MW power, and the concentration of reagent. The highest yield of 85% was obtained with a short residence time (1.2 min) at the highest concentration of 4 M in n-PrOH at 200–210 °C measured as the exiting temperature (Scheme 6B, reaction conditions below). The authors observed that the advantages of continuous flow in this reaction were correlated to the rapid heating with a residence time shorter than 1.5 min as well as the rapid cooling to reduce the impact of the retro Diels–Alder reaction, which affects the reaction at an elevated temperature.

Furthermore, the Diels–Alder reaction has been successfully studied in the chemical derivatization of graphene and fullerenes, and MW irradiation has been shown to improve its efficiency (Scheme 6C). A recent example includes the synthesis of an indene-C60 monoadduct and an indene-C60 bis-adduct. Because of their high open cell voltages, these products are used in polymer solar cells, and the scaling up of their synthesis is highly desirable. To achieve maximum productivity, the authors optimized the reaction at a temperature of up to 270 °C with the highest concentration of indene in o-xylene. The flow rate was maintained at 11.6 mL/min (residence time of 0.4 min), and the productivity was found to be 1.07 g/h.

With the aim to secure arylbutanamides on a large scale, a pharmacologically relevant substructure, the C-alkylation of dimethylacetamide with styrene has been studied using MW irradiation and scaled up in MW continuous flow mode (Scheme 7). Interestingly, an optimized protocol allowed the use of t-BuOK instead of a stronger base such as BuLi with superb results in the synthesis of a large array of derivatives. When the reaction was scaled up in flow mode in a resonant MW cavity, excellent results were obtained, and the reaction could be scaled up to reach a productivity of almost 65 g/h. A high selectivity of mono- versus dialkyalted product was obtained (10:1 mono/dialkylated).

### APPLIED HOMOGENEOUS AND HETEROGENEOUS CATALYSIS

MW-assisted synthesis has been largely applied to organometallic reactions including a review in 2019 on the continuous-flow Suzuki–Miayaura and Mizoroki–Heck reactions describing the efficacy of MW heating in scaling up of the reaction protocols in continuous mode. In addition to palladium-catalyzed synthesis, a few examples of homogeneous organometallic catalysis with other metals have been scaled up under MW continuous flow mode, and several examples have documented the attempts to optimize the scalable processes in MW batch synthesis.

Ru-catalyzed metathesis has been reported under MW irradiation, and an example of the scalable procedure was proposed by Grela et al. with the aim to contribute to the scaling up of the synthesis of Apremilast, an orally active drug for the treatment of psoriatic arthritis and psoriasis; nevertheless, the reaction has been reported on only a small scale.

An example of the nonconventional scaling up of ring-closing metathesis (RCM) was proposed by Organ et al. The generation-II Grubbs catalyst was tested, and the authors compared the conventional conditions, MW batch irradiation, and continuous flow irradiation with a capillary-based flow system in the synthesis of compound 16. As depicted in Scheme...
8A, interestingly, the reaction was performed in DCM, an MW transparent solvent (\(\delta_{0.040}\)); nevertheless, the study showed that MW irradiation showed higher conversion when compared with the conventional conditions, suggesting a direct

**Scheme 8. Ruthenium-Catalyzed Ring-Closing Metathesis**

| Ru Grub's II gen 1 mol %, DCM | 35 °C, 16 h  
|-----------------------------|-----------------  
| Ru Grub's II, 1 mol %, DCM | 200 W, 30 min  
| MW 100 W, 40 \(\mu\)L min\(^{-1}\) | 3 min 47 sec  
| capillary tube (200 \(\mu\)m int) |

**Scheme 9. Macrocyclization**

**Scheme 10. Large-Scale Cu(0) Nanoparticle-Catalyzed Reduction of Nitrobenzene in Conventional and in MW Batch Mode**

**Figure 5.** Density of temperature recorded by an IR camera in an MW-irradiated reduction of nitrobenzene Cu(0) nanoparticles cat. Reproduced with permission from ref 70. Copyright 2020 The Authors.
effect on the reagent and catalyst. As already demonstrated, the expectations based on the temperature can be exceeded when MW irradiation is exploited with polar substrates in non-adsorbing solvents.56 MW energy can thus be considered as an important variable for selectivity attainment in organic synthesis. 

Recently, another example of ruthenium-catalyzed metathesis for the ring-closing of linalool and of citronellene was studied by Leadbeater et al. (Scheme 8B).57 The reaction was performed in MW batch irradiation and then scaled up to mesoscale conventional flow irradiation in the presence of a modified second-generation Grubbs Ru catalyst. The successful reaction ensued in the presence of 0.5 mol % of catalyst at 80 °C in 10 min for the ring closing of linalool, whereas 20 min was necessary for citronellene. The results obtained in the MW batch irradiation could be perfectly transferred to conventional flow irradiation.

An unsuccessful study on the applicability of MW batch and flow irradiation in macrocycle (C-16) RCM revealed that the higher temperature is responsible for the decomposition of the Ru catalyst and that flow techniques trapped the ethylene produced during the reduction, thus generating a number of side reactions (Scheme 9A).58 Similar results were obtained in the synthesis of a C-16 macrocycle having a strong musky odor at 70 °C in toluene under MW irradiation and flowing catalyst and reagent at 1 mL/min (residence time of 1 min). When the reaction was performed conventionally in DCM at room temperature, the reaction rate was quite slow, but higher yields were obtained.

Another example focused on the synthesis of a macrocycle under MW irradiation and under flow conditions was described by Collins in 2017.59 The proposed new approach aimed to synthesize Vanepir, a protease inhibitor for hepatitis C treatment (Scheme 9B). The study is based on a catalytic macrocyclization by means of a phase separation strategy wherein poly(ethylene)glycol (PEG) lipophilic aggregates are formed, and the slow diffusion of the precursor toward the polar methanol solvent preserves the reaction outcome, avoiding the side reactions. The macrocyle was obtained by a copper-catalyzed Glaser−Hay oxidative coupling in the presence of a nickel cocatalyst, and the optimization of the protocol under MW irradiation allowed shortening of the reaction time for scale up in conventional flow mode. As depicted in Scheme 9B, the reaction yield under nonconventional conditions increased from 50 to 64% with a reaction time of 4 h; when compared with the conventional conditions, the reaction was shortened by 20 h while maintaining a comparable yield.

The application of continuous MW protocols in the preparation of metallic nanoparticles of small regular-size nanocatalysts has been the object of large interest, as summarized in recent reviews.8,60−62 The beneficial effect of MW irradiation on solid-supported catalysts is well-recognized,63−66 as is the impact of flow chemistry on process intensification.9

Because of their large industrial interest, innovative protocols in catalytic oxidation or reduction reactions represent one of the most appealing exploitations of MW chemistry in batch mode and flow mode.67,68

MW-assisted hydrogenation reactions cannot be easily performed on a large scale because of the limitations due to gaseous hydrogen. By means of a specially designed MW oven that allows a single reaction chamber to perform single or multiple reactions at temperatures up to 300 °C and pressures up to 199 bar with a gas inlet that enables the operation in a modified atmosphere (Synthwave Milestone), hydrogenation
reactions can be performed safely under MW irradiation and H2 pressure (1 bar). A recent example of Au-catalyzed levulinic acid hydrogenation was performed at 50 bar of H2 at 200 °C to produce 1,4-pentadiol in high yield with high selectivity.69 Large batch hydrogenation under MW irradiation has not been reported in the literature, however.

Reduction by transfer hydrogenation has addressed this limitation, and recently, an example of the copper-catalyzed (Cu(0) nanoparticles) reduction of nitrobenzene in glycerol was described in MW large batch mode (Scheme 10).70 Because glycerol is a biobased and eco-friendly solvent with a high MW adsorption capacity and a low vapor pressure, and because the reaction demanded a high temperature, a large-scale synthesis of the MW batch mode was studied. The reaction was performed on an almost 1 L scale in a pilot MW reactor in 95% yield, and the study demonstrated that the sample temperature in the reaction mixture was between 130 and 115 °C, with a high absorbance of delivered MW power when the power was maintained constant (Figure 5).

An original setup of the MW-assisted semihydrogenation of alkyne in continuous-flow (gas/liquid) mode is shown in Figure 6.71 The reaction of the reduction of butyndiol was performed in MW batch mode, wherein a continuous and a controlled flow of 7.5 mL of H2 was maintained to achieve not only high conversion but also high selectivity toward the cis-2-butenediol (Scheme 11). The selected catalyst was based on the solid supported Pd nanoparticle on an alumina sphere, which displayed good stability, and gradual deactivation was observed after 20 L of flowing reaction mixture. The productivity of the scaled-up procedure reached 70 mol g M−1 h−1.

The synthesis of aryl bromides via oxidative bromination with DMSO and hydrogen bromide is an efficient and inexpensive reaction that can be performed on a kilo scale.72 An attempt to perform the scaling up by MW irradiation in batch mode and in continuous flow mode illustrated the beneficial influence of high temperature (Scheme 12).73 When the reaction was performed in flow mode, the desired 4-bromo-3-methylanisole 22 could be collected in a short retention time of 1.3 min, and the synthesis was performed on an almost half a kilo scale. In 5 h, 0.32 kg of product could be obtained in 79% yield.

The use of MW irradiation in the conversion of biobased products has gained interest in the last several years.24 MW irradiation has been successfully applied in glycosylation reactions,25 and a continuous two-step process was studied in 2011 for the preparation of adenosine by glycosylation and deprotection.76 Several appliances under process intensification conditions have been reported as nonconventional protocols for the catalytic oxidation of 5-hydroxy furfural (HMF) produced from cellulose.27 Recently, aqueous H2O2 and air were deployed as oxidants in a commercial Ru/C catalyzed oxidation of HMF (Scheme 13) by means of different nonconventional technologies such as MW irradiation and continuous-flow MW irradiation.78 The reaction displayed excellent results in MW batch mode under optimized conditions, but a lower yield in continuous flow mode was obtained due to the instability of the catalyst. Not only the leaching of Ru but also the disruption of the porous structure of the catalyst was discerned by the performed analysis after usage.

### SYNTHESIS OF HETEROCYCLES AND MULTICOMPONENT REACTION

A green and sustainable approach to the synthesis of heterocycles is well documented.79,80 Because of the broad interest in medicinal chemistry and in polymer synthesis and material science, the synthesis of heterocycles has been explored, taking into account not only the use of technologies to save time and energy but also the deployment of greener solvents and even catalyst-free protocols.

The synthesis of indole has been intensely investigated because of its importance in tryptamine-based pharmaceuticals. In addition, diverse methods for its preparation via Fischer synthesis have immensely benefitted from the use of MW irradiation. Synthetic protocols for indole synthesis were proposed in MW batch mode with a homogeneous or heterogeneous acidic catalyst or a heterogeneous acid catalyst and scaled up in MW continuous mode. Because of its high efficiency and purity, MW-assisted Fischer synthesis has been practiced in the undergraduate laboratory starting from dehydroepiandrosterone acetate and phenyl hydrazine hydrochloride salt in glacial acetic acid (Scheme 14).24

The synthesis of methyl benzimidazole under MW irradiation from o-phenylenediamine in acetic acid is a simple condensation procedure that has been studied by different research groups (Scheme 15). Because a higher temperature favors attaining complete conversion in a shorter time and because acetic acid has a high loss tangent of 0.174, MW irradiation could accomplish the reduction of reaction time. Scaling up of this reaction under solvent-free conditions with a moderate to high scale of production has been documented. In 10 min, the synthesis was performed in parallel in a multimode MW
instrument with multivessel rotor systems and a total reaction volume up to 960 mL (16 Teflon vessels of 60 mL each) by Kappe et al.\textsuperscript{85} A few years later, the same group in collaboration with Clariant Produkte Deutschland reported the kilo scale production of methyl benzimidazole in continuous mode by means of an MW system operating at high temperature/high pressure processing 20 L/h.\textsuperscript{22} In 2014, Organ et al. presented another example of this reaction performed in a continuous-flow MW reactor in which the back-pressure regulators guaranteed no pressure fluctuations during the reaction and the system was equipped with a SiC reactor tube to shorten the reaction time to 6 s at 313 °C.\textsuperscript{86,87} In all of the cases operating with a high-temperature/high-pressure process, the yield was maintained at >90%.

The beneficial effect of MW irradiation on the heterogeneously catalyzed synthesis of heterocycles has been of great interest in view of the large applicability of the heterocycle in pharmaceutical chemistry. MW irradiation facilitates a reduction in reaction time, and by deploying multicomponent reactions, high atom efficiency can be reached. Additionally, a number of successful protocols have appeared in the literature, and it is not always that the reaction yield by MW irradiation is higher than that under conventional conditions. As a proof of this concept, in 2014, Török et al. studied six different reactions catalyzed by semisynthetic montmorillonite K-10, a good MW absorber, and measured the energy consumption (kWh/% yield). The electrophilic annulation of pyrrole, the synthesis of substituted pyrazole by the domino condensation−cyclization−aromatization reaction, the domino Picte−Spengler cyclization−
dehydrogenation reactions, and the multicomponent domino cyclization–aromatization reactions were compared (Scheme 16).88 Pyrazole synthesis from chalcone (product 27) and the synthesis of quinoline 28 and of carbazole 29 reached a high $E_{\text{oil/bath}}/E_{\text{MW}}$ ratio, with a value of 5.6, 9.1, and 5.7, respectively. When the indole was obtained by electrophilic condensation, the reaction represented an exception, and the conventional heating revealed a lower energy consumption with respect to MW irradiation. The quinoline synthesis was scaled up by 40 times to 17 g; interestingly, the MW irradiation produced the final product in 90% yield, whereas the conventional heating afforded a 70% yield, and the $E_{\text{oil/bath}}/E_{\text{MW}}$ ratio was 7.6, underlining the higher energy efficiency of the MW-assisted reaction. On the basis of the reported results, the conventional heating generally consumed more energy when compared with MW irradiation in heterogeneously catalyzed heterocycle synthesis; nevertheless, the reaction should be studied on a case-by-case basis.

The use of MW irradiation in multicomponent organic synthesis has been well exploited.89 Because of their potential to limit the number of synthetic steps while increasing the molecular complexity and diversity, multicomponent reactions have been largely used in pharmaceutical chemistry.

The Ugi reaction has been introduced for the preparation of multifunctional peptides and heterocyclic compounds, as summarized in a recent review.90 MW heating contributes to the efficacy of the Ugi reaction because of the high polarity of the intermediates. Furthermore, the MW irradiation power as well as the solvent selection have been demonstrated to influence product selectivity.91 Because of the long reaction time, MW irradiation also shows good advantages in increasing the reaction rate.

In 2015, Salvador et al. reported an MW-promoted sequential Ugi reaction/Cu-catalyzed alkyne azide cyclization to produce cyclopeptoids that was scaled up in conventional flow mode.92 In 2019, the same group described another scalable procedure for the multicomponent isocyanide-based Passerini reaction optimized in the MW system (Scheme 17).93 From the optimization studies in batch mode, it was observed that in toluene and CHCl$_3$, the conversion was very low or near-zero because of their low MW dielectric heating properties. In acetonitrile, the reaction efficiently delivered the desired product. When performed in the conventional flow setup, the $\alpha$-acyloxy ketone derivative was obtained in comparable yield with a productivity of 0.312 g/min.

Because of the well-known pharmacological activity of dihydropyridines for the treatment of thrombosis, atherosclerosis, and cardiovascular disease, the four-component Hantzsch synthesis has been extensively studied under conventional and unconventional conditions.94-95

The first example of scaling up in the continuous MW irradiation of 4-aryl-1,4-dihydropyridines was reported in 2001, deploying a domestic MW oven in which a round glass reactor of 65 mL was placed along the circumference of a glass plate, and the synthesis was performed in an aqueous hydrotrioxide solution (50% sodium $p$-toluene sulfonate) to maintain a homogeneous reaction medium.96 More recently, the limits of Hantzsch multicomponent synthesis in continuous mode were underscored in two publications. In the first study, the reaction was performed with MW irradiation on a 100 mmol scale of the three components, and the reaction conversion was compared with that of the conventional flow mode (Scheme 18A).97 In addition to the fact that the conversion was lower in the flow mode when compared to the MW batch mode, the low solubility of the of 4-phenyl-1,4-dihydropyridines in ethanol imposed the authors to add ethyl acetate to the reaction mixture once out from the heated aluminum block. The energy consumption in Hantzsch synthesis was measured, and comparing 100 mmol reaction scale heated for 15 min, the energy efficiency of the MW batch mode measured in Wh was more than two times that in the flow mode.

Another study confirmed the same trend by performing the reaction under MW irradiation at 140 °C for 10 min and in conventional flow mode.98 Formaldehyde was used instead of benzaldehyde, and an attempt to reduce the molar ratio was also pursued. The yield in conventional flow mode was lower than that in MW batch mode, ranging from 82% in MW batch mode to 68% in flow mode. Better results were observed when the synthesis of enamine in situ was avoided and acetooacetate was replaced with ethyl $\beta$-aminocrotonate and phenylpropargyl aldehyde instead of formaldehyde (Scheme 18B). When the reactions in batch mode and in continuous MW irradiation mode were compared, higher yields were attained, and the flow synthesis reached a higher yield. The better performance of the multicomponent Hantzsch reaction in continuous mode was achieved in the presence of a heterogeneous catalyst ($\gamma$-Fe$_2$O$_3$ nanoparticles) packed in a polytetrafluoroethylene (PTFE) capillary column.99 Because the magnetic nanoparticles catalyzed the multicomponent reaction, a 98.7% yield was obtained at 100 °C in 3 min residence time, but surprisingly, a large excess of catalyst was necessary to obtain full conversion (Scheme 18A).

An interesting example of three-component spiro-oxindole synthesis was proposed by Larhed et al.100 As depicted in Scheme 19 the influence of MW irradiation in combination with continuous flow in a silicon carbide tubular reactor shortened the reaction time and increased the yield.87
STEREOSELECTIVITY IN MICROWAVE-ASSISTED SYNTHESIS

High-temperature and high-pressure chemical transformations may suffer from low stereochemical control; nevertheless, nowadays, several studies have reported the stereoselective synthesis under MW irradiation conditions. Because selectivity is a crucial objective in organic synthesis, a careful evaluation of the reaction conditions is always warranted, and in addition to solvent selection, an appropriate study of temperature and time allows kinetic versus thermodynamic control. The possibility of controlling the regio/stereoselectivity by choosing the appropriate conventional MW heating is an attractive field of research.

As depicted in Scheme 20A, vinylallenes can be obtained enantioselectively from the alkyne in the presence of a chiral auxiliary under MW irradiation. Optically active propargylamine, in which \((S)-\)prolinol was the stereogenic group, reacted in the presence of \(\text{AgNO}_3\) in acetonitrile at \(70^\circ\text{C}\) under MW irradiation (50 W) in 20 min. The reaction proceeded to generate the desired allene derivatives in 70% average yield, exemplified by a series of 11 cases with high stereoselectivity (79 → 99% ee).

Scheme 20. Stereoselective Synthesis

Another recent example reported the palladium-catalyzed decarboxylation/alkylation of propargyl cyclopentane carboxylate (Scheme 20B) to produce the final product \(35\) in 64% yield with 89:11 er in presence of \((R,R)-\text{N-PINAB}\).

Unforeseeable results were recently obtained by Repo et al. when the condensation of \(L\)-phenylalanine and benzaldehyde allows kinetic versus thermodynamic control. The possibility of controlling the regio/stereoselectivity by choosing the appropriate conventional MW heating is an attractive field of research.

As depicted in Scheme 20A, vinylallenes can be obtained enantioselectively from the alkyne in the presence of a chiral auxiliary under MW irradiation. Optically active propargylamine, in which \((S)-\)prolinol was the stereogenic group, reacted in the presence of \(\text{AgNO}_3\) in acetonitrile at \(70^\circ\text{C}\) under MW irradiation (50 W) in 20 min. The reaction proceeded to generate the desired allene derivatives in 70% average yield, exemplified by a series of 11 cases with high stereoselectivity (79 → 99% ee).

Another recent example reported the palladium-catalyzed decarboxylation/alkylation of propargyl cyclopentane carboxylate (Scheme 20B) to produce the final product \(35\) in 64% yield with 89:11 er in presence of \((R,R)-\text{N-PINAB}\).

Unforeseeable results were recently obtained by Repo et al. when the condensation of \(L\)-phenylalanine and benzaldehyde

Scheme 21. Multicomponent Diastereoselective Synthesis

Scheme 22. Mechanism of Multicomponent the Diastereoselective Synthesis of Dicarboxylic Acid 36

Scheme 23. Wolff–Staudinger Reaction of \(\beta\)-Lactams
was performed in alkaline ethanol under MW irradiation (Scheme 21). A single crystalline compound with the structure of a dicarboxylic acid 36 was isolated and characterized by X-ray crystallography as a racemic mixture of RRR and SSS enantiomers. Intriguingly, both the pyruvic acid derivative and cinnamaldehyde were obtained by means of MW heating wherein the first one was formed by the condensation of an amino acid with benzaldehyde followed by isomerization and hydrolysis, whereas the second one could be synthesized via a Cannizzaro-type reaction followed by aldolic condensation (Scheme 22). The diastereoselectivity of this reaction has been driven on the basis of a [3,3]-sigmatropic rearrangement that involves cinnamaldehyde and phenylpyruvic acid. From the hydrated derivative of aldehyde II, it is assumed that the Na+ ion favors a chairlike conformation intermediate that delivers the final product by hydride transfer.

The Wolff–Staudinger reaction affords the formation of β-lactams via the [2 + 2] cycloaddition of ketenes with imines. The diastereoselectivity for the trans-substituted reaction has been studied in MW batch mode as well as under continuous MW operation. Because the protocol is considered problematic due to the release of gaseous nitrogen and in view of the high temperature, continuous MW mode is an appealing alternative to the batch synthesis. A set of 18 different compounds with yields ranging from 33 to 85% were synthesized with general selectivity toward the trans configuration. Electron-poor imines gave the lowest yield and stereoselectivity, whereas electron-rich imines were efficiently converted to the trans isomer. In general, high temperature and MW heating provided the optimal diastereoselectivity in the Staudinger reaction (Scheme 23).

Aimed to predict the stereoselectivity or to rationalize it, density functional theory (DFT) calculations were also included in the study.

CONCLUSIONS

On the basis of the recent literature, numerous advances in MW technologies have been reported. The possible applications of MAOS are extremely broad. When highly adsorbing polar solvent, reagents, or catalysts are deployed, a tremendous reaction rate increase is observable under MW irradiation conditions. Nevertheless, protocols may be affected by a number of variables related to the diverse equipment and different technologies used for the measurement of temperature. The multitude of reactions require a case-by-case study, as an automatic green label for MAOS is not warranted. However, MACOS represents the best setup option for any scaling up efforts geared toward the potential industrialization of dielectric heating. This also requires a careful measurement of the dielectric properties, namely, the permittivity, of reagents/reactants, solvents, and catalysts. The new age of MAOS is far from the empirical approach of the first enthusiastic practitioners and may address the strict rules of good manufacturing practice (GMP) production.

AUTHOR INFORMATION

Corresponding Authors

Giancarlo Cravotto — Dipartimento di Scienze e Tecnologia del Farmaco and Centre for Nanostructured Interfaces and Surfaces (NIS), University of Turin, University of Turin, 10125 Turin, Italy; orcid.org/0000-0001-9731-6228; Email: varma.rajender@epa.gov

Author

Katia Martina — Dipartimento di Scienze e Tecnologia del Farmaco and Centre for Nanostructured Interfaces and Surfaces (NIS), University of Turin, University of Turin, 10125 Turin, Italy; orcid.org/0000-0003-2256-5021

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.joc.1c00865

Notes

The authors declare no competing financial interest.

Biographies

Katia Martina was born in Italy and received a Ph.D. in Chemistry at the University of Turino in 2008. After 7 years of experience in pharmaceutical R&D, she joined the group of Prof. Cravotto in the Department of Drug Science of Technology of the University of Turin, where she is now an Associate Professor. Her research activities focus on organic synthesis under nonconventional conditions, heterogeneous catalysis, and the optimization of an innovative protocol for surface derivatization. Her research interests are documented in 75 publications, 9 patents, and 5 book chapters.

Giancarlo Cravotto is a Full Professor of Organic Chemistry at the University of Turin (Italy). His research activity in the domain of green organic synthesis and processing is documented by about 450 scientific peer-reviewed papers and 20 patents. In the period from 2007 to 2018, he was the Director of the Department of Drug Science and Technology, and currently, he is vice-Director. He collaborates with a variety of international industrial partners. He received the Scientific Research Award 2018 “Organic Chemistry for the Environment, Energy and Nanosciences” and Gold Medal “E. Paterno” 2020 by from the Italian Chemical Society.
The Journal of Organic Chemistry

Rajender Varma (H-Index 114, Highly Cited Researchers 2016, 2018–2020) was born in India (Ph.D., Delhi University 1976). He was a faculty member at Baylor College of Medicine and Sam Houston State University prior to joining the U.S. EPA in 1999 with a visiting position at Palacky University Olomouc, Czech Republic. He has over 48 years of research experience in the management of multidisciplinary technical programs ranging from natural products chemistry to the development of more environmentally friendly synthetic methods using microwaves, ultrasound, etc. Lately, he has been focused on greener approaches to the assembly of nanomaterials and sustainable applications of magnetically retrievable nanocatalysts in benign media. He is a member of the editorial advisory board of several international journals, has published over 710 papers, has been awarded 17 U.S. Patents, and has 6 books, 26 book chapters, and 3 encyclopedia contributions with 48 800 citations.

ACKNOWLEDGMENTS

The University of Turin is acknowledged for the financial support (Ricerca locale 2020).

REFERENCES

(1) Varma, R. S. Solvent-Free Organic Syntheses Using Supported Reagents and Microwave Irradiation. Green Chem. 1999, I, 43.
(2) Nikáčević, N. M.; Huesman, A. E. M.; Van den Hof, P. M. J.; Stankiewicz, A. I. Opportunities and challenges for process control in process intensification. Chem. Eng. Proc. 2012, 52, 1.
(3) Wegner, J.; Ceylan, S.; Kirschning, A. Flow Chemistry - A Key Enabling Technology for (Multistep) Organic Synthesis. Adv. Synth. Catal. 2012, 354, 17.
(4) Alcazar, J.; de Munoz, M. In Microwave-Assisted Continuous Flow Organic Synthesis (MACOS). In Microwaves in Organic Synthesis, 3rd ed.; de la Hoz, A.; Loupy, A., Eds.; Wiley-VCH Verlag GmbH: Weinheim, Germany, 2012; pp 1173–1204.
(5) Porta, R.; Benaglia, M.; Puglisi, A. Flow Chemistry: Recent Developments in the Synthesis of Pharmaceutical Products. Org. Process Res. Dev. 2016, 20, 2.
(6) Egami, H.; Hamashima, Y. Practical and Scalable Organic Reactions with Flow Microwave Apparatus. Chem. Rec. 2019, 19, 157.
(7) Monguchi, Y.; Ichikawa, T.; Yamada, T.; Savama, Y.; Sajihi, K. Continuous-Flow Suzuki-Miyaura and Mizoroki-Heck Reactions under Microwave Heating Conditions. Chem. Rec. 2019, 19, 3.
(8) Tagliapietra, S.; Calcio Gaudino, E.; Martina, K.; Barge, A.; Cravotto, G. Microwave Irradiation in Micro-Meso-fluidic Systems: Hybrid Technology has Issued the Challenge. Chem. Rec. 2019, 19, 98.
(9) De Ris, C.; Bortolini, O.; Brandolese, A.; Di Carnine, G.; Ragno, D.; Massi, A. Recent advances in continuous-flow organocatalysis for process intensification. React. Chem. Eng. 2020, 5, 1017.
(10) Horikoshi, S.; Serpone, N. Microwave Flow Chemistry as a Methodology in Organic Syntheses, Enzymatic Reactions, and Nanoparticle Syntheses. Chem. Rec. 2019, 19, 118.
(11) Estel, L.; Poux, M.; Benamara, N.; Polaert, I. Continuous flow-microwave reactor: Where are we? Chem. Eng. Process. 2017, 113, 56.
(12) Goyal, H.; Mehda, A.; Lobo, R. F.; Stefanidis, G. D.; Vlachos, D. G. Scaleup of a Single-Mode Microwave Reactor. Ind. Eng. Chem. Res. 2020, 59, 2516.
(13) Nigar, H.; Sturm, G. S. J.; Garcia-Baños, B.; Peñaranda- Foix, F. L.; Catalá-Civera, J. M.; Mallada, R.; Stankiewicz, A.; Santamaría, J. Numerical analysis of microwave heating cavity: Combining electromagnetic energy, heat transfer and fluid dynamics for a NaY zeolite fixed-bed. Appl. Therm. Eng. 2019, 155, 226.
(14) Morte, M.; Dean, J.; Kitajima, H.; Hascakir, B. Increasing the Penetration Depth of Microwave Radiation Using Acoustic Stress to Trigger Piezoelectricity. Energy Fuel 2019, 33, 6327.
(15) Martina, K.; Tagliapietra, S.; Barge, A.; Cravotto, G. Combined Microwaves/Ultrasound, a Hybrid Technology. Top. Curr. Chem. 2016, 374, 1.
(16) Buttress, A. J.; Hargreaves, G.; Ilchev, A.; Monti, T.; Sklavenonou, A.; Katrib, J.; Martin-Tanchereau, P.; Unthank, M. G.; Irvine, D. J.; Dodds, C. D. Design and optimization of a microwave reactor for kilo-scale polymer synthesis. Chem. Eng. Sci.: X 2019, 2, 100022.
(17) Vamosi, P.; Matsuo, K.; Masuda, T.; Sato, K.; Narumi, T.; Takeki, K.; Mase, N. Rapid Optimization of Reaction Conditions Based on Comprehensive Reaction Analysis Using a Continuous Flow Microwave Reactor. Chem. Rec. 2019, 19, 77.
(18) Barham, J. P.; Koyama, E.; Norikane, Y.; Ohneda, N.; Yoshimura, T. Microwave Flow: A Perspective on Reactor and Microwave Configurations and the Emergence of Tunable Single-Mode Heating Toward Large-Scale Applications. Chem. Rec. 2019, 19, 188.
(19) Dabrowska, S.; Chudoba, T.; Wojnarowicz, J.; Lojkowski, W. Current Trends in the Development of Microwave Reactors for the Synthesis of Nanomaterials in Laboratories and Industries: A Review. Crystals 2018, 8, 379.
(20) Stefanidis, G. D.; Muñoz, A. N.; Sturm, G. S. J.; Stankiewicz, A. A helicopter view of microwave application to chemical processes: reactions, separations, and equipment concepts. Rev. Chem. Eng. 2014, 30, 233.
(21) Baxendale, I.; Hayward, J.; Ley, S. Microwave Reactions Under Continuous Flow Conditions. Comb. Chem. High Throughput Screening 2007, 10, 802.
(22) Morschhäuser, R.; Krull, M.; Kayser, C.; Boberski, C.; Bierbaum, R.; Püschner, P. A.; Glasnov, T. N.; Kappé, C. O. Microwave-assisted continuous flow synthesis on industrial scale. Green Process. Synth. 2012, 1, 281.
(23) Patil, N. G.; Benaskar, F.; Rehrov, E. V.; Meuldijk, J.; Hulshof, L. A.; Hesselt, V.; Schouten, J. C. Microwave Setup Design for Continuous Fine-Chemicals Synthesis. Chem. Eng. Technol. 2014, 37, 1645.
(24) Rinaldi, L.; Carnaroglio, D.; Rotolo, L.; Cravotto, G. A Microwave-Based Chemical Factory in the Lab: From Milligram to Multigram Preparations. J. Chem. 2015, 2015, 879531.
(25) Koyama, E.; Ito, N.; Sugiyama, J.-i.; Barham, J. P.; Norikane, Y.; Azumi, R.; Ohneda, N.; Ohno, Y.; Yoshimura, T.; Odajima, H.; Okamoto, T. A continuous-flow resonator-type microwave reactor for high-efficiency organic synthesis and Claisen rearrangement as a model reaction. J. Flow Chem. 2018, 8, 147.
(26) Öhngren, P.; Fardost, A.; Russo, F.; Schanche, J.-S.; Fagrell, M.; Larhed, M. Evaluation of a Nonresonant Microwave Applicator for Continuous-Flow Chemistry Applications. Org. Process Res. Dev. 2012, 16, 1053.
(27) Moseley, J. D.; Lenden, P.; Lockwood, M.; Ruda, K.; Sherlock, J.-P.; Thomson, A. D.; Gilday, J. P. A Comparison of Commercial Microwave Reactors for Scale-Up within Process Chemistry. Org. Process Res. Dev. 2008, 12, 36.
(28) Furuta, A.; Fukuyama, T.; Ryu, I. Efficient Flow Fischer Esterification of Carboxylic Acids with Alcohols Using Sulfonic Acid-Functionalized Silica as Supported Catalyst. Bull. Chem. Soc. Jpn. 2017, 90, 607.
(29) Nagahata, R.; Nakamura, T.; Mori, Y.; Takeuchi, K. Microwave-assisted facile and rapid esterification of amino acids: esterification of
L-leucine from batch to flow processes and scale-up. Nat. Sci. 2017, 9 (4), 110.

(30) Li, X.; Xu, J. Effects of the Microwave Power on the Microwave-assisted Esterification. Curr. Microwave Chem. 2017, 4, 158.

(31) Tajii, A.; Toth, N.; Balint, E.; Keglevich, G. Esterification of benzoic acid in a continuous flow microwave reactor. J. Flow Chem. 2018, 8, 11.

(32) Negus, M. P.; Mansfield, A. C.; Leadbeater, N. E. The preparation of ethyl levulinate facilitated by flow processing: the catalyzed and uncatalyzed esterification of levulinic acid. J. Flow Chem. 2015, 5, 148.

(33) Balint, E.; Tajii, A.; Toth, N.; Keglevich, G. Continuous Flow Alcoholysis of Dialkyl H-Phosphonates with Aliphatic Alcohols. Molecules 2018, 23, 1618.

(34) Kiss, N. Z.; Henyez, R.; Keglevich, G. Continuous flow esterification of a H-phosphinic acid, and transesterification of H-phosphonates and H-phosphonates under microwave conditions. Molecules 2020, 25, 719.

(35) Balint, E.; Tajii, A.; Keglevich, G. Application of the microwave technique in continuous flow processing of organophosphorus chemical reactions. Materials 2019, 12, 788.

(36) Isaksson, R.; Kumpija, I.; Larhed, M.; Wannberg, J. Rapid and straightforward transesterification of sulfonyl carbamates. Tetrahedron Lett. 2016, 57, 1476.

(37) Kumpija, I.; Isaksson, R.; Saevmarker, J.; Wannberg, J.; Larhed, M. Microwave Promoted Transcarbamylation Reaction of Sulfonyl-carbamates under Continuous-Flow Conditions. Org. Process Res. Dev. 2016, 20, 440.

(38) Marafie, J. A.; Moseley, J. D. The application of step-flow microwave technology to scaling-out SNAr reactions using a soluble organic base. Org. Biomol. Chem. 2010, 8, 2219.

(39) Wiles, C.; Watts, P. Translation of microwave methodology to continuous flow for the efficient synthesis of diaryl ethers via a base-mediated SNAr reaction. Beilstein J. Org. Chem. 2011, 7, 1360.

(40) Organ, M. G.; Hanson, P. R.; Rolfe, A.; Samarakoon, T. B.; Ullah, F. Accessing Stereochemically Rich Sultams via Microwave-assisted, Continuous-flow Organic Synthesis (MACOS) Scale-Out. J. Flow Chem. 2012, 1, 32.

(41) Horikoshi, S.; Watanabe, T.; Kamata, M.; Suzuki, Y.; Serpone, N. Microwave-assisted organic syntheses: microwave effect on intramolecular reactions -- the Claisen rearrangement of allylphenyl ether and 1-allyloxy-4-methoxybenzene. RSC Adv. 2015, 5, 90272.

(42) Abele, S.; Hock, S.; Schmidt, G.; Funel, J.-A.; Marti, R. High-Temperature Diels-Alder Reactions: Transfer from Batch to Continuous Mode. Org. Process Res. Dev. 2012, 16, 1114.

(43) Karney, M. J.; Porter, K. A.; Barnhardt, E. K.; Vanier, G. S. Meso-scale microwave-assisted continuous flow reactions utilizing a selective heating matrix. RSC Adv. 2013, 3, 7106.

(44) Yokozawa, S.; Ohneda, N.; Muramatsu, K.; Okamoto, T.; Odajima, H.; Iwata, T.; Sugiyama, J.-I.; Fujita, M.; Sawairi, T.; Egami, H.; Hamashima, Y.; Egí, M.; Akai, S. Development of a highly efficient single-mode microwave applicator with a resonant cavity and its application to continuous flow synthesis reactions. RSC Adv. 2015, 5, 10204.

(45) Barham, J.; Baron, S.; Hock, S.; Schmidt, G.; Funel, J.-A.; Martí, R. Microwave-Assisted Green Synthesis of Silver Nanostructures. J. Organo-Metall. Chem. 2019, 5808.

(46) Kokel, A.; Schäfer, C.; Török, B. Application of microwave-assisted heterogeneous catalysis in sustainable synthesis design. Green Chem. 2017, 19, 3729.

(47) Kopylovich, M. N.; Ribeiro, A. P. C.; Alegra, E. C.B.A.; Martins, N. M. R.; Martins, L. M.D.R.S.; Pompeiro, A. J.L. Microwave Dielectric Heating. J. Org. Chem. 2017, 82, 7576.

(48) Petricci, E.; Cini, E.; Taddei, M. Catalytic reactions in continuous flow microwave reactor. J. Org. Chem. 2020, 4, 1360.

(49) Godin, E.; Bedard, A.-C.; Raymond, M.; Collins, S. K. Phase transfer separation in a complex pharmaceutical setting: Application toward the synthesis of vaniprevir. J. Org. Chem. 2017, 82, 1360.

(50) Alexander, K. A.; Paulhus, E. A.; Lazarus, G. M. L.; Leadbeater, N. E. Exploring the reactivity of a ruthenium complex in the metathesis of biorenewable feedstocks to generate value-added chemicals. J. Organomet. Chem. 2016, 812, 74.

(51) Morin, E.; Sosso, J.; Raymond, M.; Amorelli, B.; Boden, R. M.; Collins, S. K. Synthesis of a Green and Crude Cyclic Musc Evaluation of Batch, Microwave, and Continuous Flow Strategies. Org. Process Res. Dev. 2019, 23, 283.

(52) Varma, R. S. Greener routes to organics and nanomaterials: sustainable applications of nanocatalysts. Pure Appl. Chem. 2013, 85, 1703.

(53) Li, H.; Zhang, C.; Pang, C.; Li, X.; Gao, X. The Advances in the Special Microwave Effects of the Heterogeneous Catalytic Reactions. Front. Chem. 2020, 8, 355.

(54) Gawande, M. B.; Shelve, S. N.; Zboril, R.; Varma, R. S. Microwave-Assisted Chemistry: Synthetic Applications for Rapid Assembly of Nanomaterials and Organics. Acc. Chem. Res. 2014, 47, 1338.

(55) Varma, R. S. Journey on greener pathways: from the use of alternate energy inputs and benign reaction media to sustainable applications of nano-catalysts in synthesis and environmental remediation. Green Chem. 2014, 16, 2027.

(56) Kokel, A.; Schäfer, C.; Török, B. Application of microwave-assisted heterogeneous catalysis in sustainable synthesis design. Green Chem. 2017, 19, 3729.

(57) Cini, E.; Petricci, E.; Taddei, M. Pd/C Catalysis under Microwave Heating. Catalysts 2017, 7, 89.
(69) Bucciol, F.; Tabasso, S.; Grillo, G.; Menegazzo, F.; Signoretto, M.; Manzoli, M.; Cravotto, G. Boosting levulinic acid hydrogenation to value-added 1,4-pentanediol using microwave-assisted gold catalysis. J. Catal. 2019, 380, 267.
(70) Moran, M. J.; Martina, K.; Stefanidis, G. D.; Jordens, J.; Gerven, T. V.; Goovaerts, V.; Manzoli, M.; Groffils, C.; Cravotto, G. Glycerol: An Optimal Hydrogen Source for Microwave-Promoted Cu-Catalyzed Transfer Hydrogenation of Nitrobenzene to Aniline. Front. Chem. 2020, 8, 34.
(71) Calcio Gaudino, E.; Manzoli, M.; Carnaroglio, D.; Wu, Z.; Grillo, G.; Rotolo, L.; Medlock, J.; Bonrath, W.; Cravotto, G. Sonochemical preparation of alumina-spheres loaded with Pd nanoparticles for 2-butyne-1,4-diol semi-hydrogenation in a continuous flow microwave reactor. RSC Adv. 2018, 8, 7029.
(72) Song, S.; Sun, X.; Li, X.; Yuan, Y.; Jiao, N. Efficient and Practical Oxidative Bromination and Iodination of Aromes and Heteroarenes with DMSO and Hydrogen Halide: A Mild Protocol for Late-Stage Functionalization. Org. Lett. 2015, 17, 2886.
(73) Jin, Y.; Yang, J.; Feng, X.; Li, J.; Xu, J.; Chen, X.; Wang, S.; Lv, Y.; Yu, J. Development of large-scale oxidative Bromination with HBr-DMSO by using a continuous-flow microwave system for the subsequent synthesis of 4-Methoxy-2-methylphenol. J. Flow React. 2019, 267.
(74) Bundhoo, Z. M. A. Microwave-assisted conversion of biomass and waste materials to biofuels. Renewable Sustainable Energy Rev. 2018, 82, 1149.
(75) Manvar, A.; Shah, A. Continuous Flow and Microwave-Assisted Vortrügen Glycosylations: Historical Perspective to High-Throughput Strategies. Asian J. Org. Chem. 2014, 3, 1134.
(76) Sniady, A.; Bedore, M. W.; Jamison, T. F. One-Flow, Multistep Synthesis of Nucleosides by Bromined Acid-Catalyzed Glycosylation. Angew. Chem., Int. Ed. 2011, 50, 2155.
(77) Su, T.; Zhao, D.; Wang, Y.; Lue, H.; Varma, R. S.; Len, C. Innovative Protocols in the Catalytic Oxidation of S-Hydroxymethylfurfural. ChemSusChem 2021, 14, 266.
(78) Zhao, D.; Rodriguez-Padron, D.; Triantafyllidis, K. S.; Wang, Y.; Luque, R.; Len, C. Microwave-Assisted Oxidation of Hydroxymethyl Furfural to Added-Value Compounds over a Ruthenium-Based Catalyst. ACS Sustainable Chem. Eng. 2020, 8, 3091.
(79) Martina, K.; Tagliapietra, S.; Veselov, V. V.; Cravotto, G. Green protocols in heterocycle syntheses via 1,3-dipolar cycloadditions. Front. Chem. 2019, 7, 95.
(80) Driowya, M.; Saber, A.; Marzag, H.; Demange, L.; Benhida, R.; Bourgin, K. Microwave-Assisted Synthesis of Bioactive Six-Membered Heterocycles and Their Fused Analogues. Molecules 2016, 21, 492.
(81) Gutmann, B.; Gottsmann, M.; Elsner, P.; Cantillo, D.; Roberge, D. M.; Kappe, C. O. On the Fischer Indole Synthesis of 7-Ethylthiophenol—Mechanistic and Process Intensification Studies under Continuous Flow Conditions. Org. Process Res. Dev. 2013, 17, 294.
(82) Panther, J.; Rechmann, J.; Muller, T. T. J. Fischer indole synthesis of 3-benzyl-1H-indole via conductive and dielectric heating. Chem. Heterocycl. Compd. 2016, 52, 897.
(83) Xu, J. G.; Yu, J. G.; Jin, Y.; Li, J.; Yu, Z. Q.; Lv, Y. W. A continuous flow microwave-assisted Fischer indole synthesis of 7-Ethylthiophenol. Chem. Eng. Process. 2017, 121, 144.
(84) Cirillo, P. F.; Caccavale, A.; DeLuna, A. Green Fischer Indole Synthesis Using a Steroidal Ketone in a Conductively Heated Sealed-Vessel Reactor for the Advanced Undergraduate Laboratory. J. Chem. Educ. 2021, 98, 567.
(85) Damm, M.; Glasnov, T. N.; Kappe, C. O. Translating High-Temperature Microwave Chemistry to Scalable Continuous Flow Processes. Org. Process Res. Dev. 2010, 14, 215.
(86) Sauks, J. M.; Mallik, D.; Lawryshyn, Y.; Bender, T.; Organ, M. A Continuous-Flow Microwave Reactor for Conducting High-Temperature and High-Pressure Chemical Reactions. Org. Process Res. Dev. 2014, 18, 1310.
(87) Kappe, C. O. Unraveling the Mysteries of Microwave Chemistry Using Silicon Carbide Reactor Technology. Acc. Chem. Res. 2013, 46, 1579.
(88) Cho, H.; Török, F.; Török, B. Energy efficiency of heterogeneous catalytic microwave-assisted organic reactions. Green Chem. 2014, 16, 3623.
(89) Fairoosa, J.; Saranya, S.; Radhika, S.; Anilkumar, G. Recent Advances in Microwave Assisted Multicomponent Reactions. Chem. Select 2020, 5, 5180.
(90) Fouad, M. A.; Abdel-Hamid, H.; Ayoup, M. S. Two decades of recent advances of Ugi reactions: synthetic and pharmaceutical applications. RSC Adv. 2020, 10, 42644.
(91) Barreto, A. d. F. S.; Andrade, C. K. Z. Microwave-mediated synthesis of a cyclic heptapeptoid through consecutive Ugi reactions. Tetrahedron 2018, 74, 6861.
(92) Salvador, C. E. M.; Pieber, B.; Neu, P. M.; Torvisco, A.; Kleber Z.; Andrade, C.; Kappe, C. O. A sequential Ugi multicomponent/Cu-catalyzed azide-alkyne cycloaddition approach for the continuous flow generation of cyclic peptoids. J. Org. Chem. 2015, 80, 4590.
(93) Salvador, C. E. M.; Andrade, C. K. Z. A Mild, Fast, and Scalable Synthesis of Substituted α-Acloylo Ketones via Multicomponent Reaction Using a Continuous Flow Approach. Front. Chem. 2019, 7, 531.
(94) Slobbe, P.; Ruijter, E.; Orru, R. V. A. Recent applications of multicomponent reactions in medicinal chemistry. MedChemComm 2012, 3, 1189.
(95) Abdella, A. M.; Abdelmoniem, A. M.; Abdelhamid, I. A.; Elwahy, A. H. M. Synthesis of heterocyclic compounds via Michael and Hantzsch reactions. J. Heterocycl. Chem. 2020, 57, 1476.
(96) Khadilkar, B. M.; Madyar, V. R. Scaling Up of Dihydropyridine Ester Synthesis by Using Aqueous Hydrotrope Solutions in a Continuous Microwave Reactor. Org. Process Res. Dev. 2001, 5, 452.
(97) Devine, W. G.; Leadbeater, N. E. Probing the energy efficiency of microwave heating and continuous-flow conventional heating as tools for organic chemistry. ARKIVOC 2011, 2011, 127.
(98) Bagley, M. C.; Fusillo, J.; Jenkins, R. L.; Lubini, M. C.; Mason, C. One-step synthesis of pyridines and dihydropyridines in a continuous flow microwave reactor. Beilstein J. Org. Chem. 2013, 9, 1957.
(99) He, W.; Fang, Z.; Zhang, K.; Tu, T.; Lv, N.; Qiu, C.; Guo, K. A novel micro-flow system under microwave irradiation for continuous synthesis of 1,4-dihydropyridines in the absence of solvents via Hantzsch reaction. Chem. Eng. J. 2018, 331, 161.
(100) Koukabi, N.; Kolvari, E.; Khazaei, A.; Zolfigol, M. A.; Shirmardi-Shaghasemi, B.; Khavasi, H. R. Hantzsch reaction on free nano-FeO3 catalyst: excellent reactivity combined with facile catalyst recovery and recyclability. Chem. Commun. 2011, 47, 9230.
(101) Engen, K.; Saevmarker, J.; Rosenstroem, U.; Wannberg, J.; Lundback, T.; Jemnalm-Jensen, A.; Larhed, M. Microwave Heated Flow Synthesis of Spiro-oxindole Dihydroquinazolinone Based IRAP Inhibitors. Org. Process Res. Dev. 2014, 18, 1582.
(102) Lo, V. K.-Y.; Chan, Y.-M.; Zhou, D.; Toy, P. H.; Che, C.-M. Highly Enantioselective Synthesis Using Prolinol as a Chiral Auxiliary: Silver-Mediated Synthesis of Axially Chiral Vinyllalenes and Subsequent [Hetero]-Diels−Alder Reactions. Org. Lett. 2019, 21, 7717.
(103) Eronen, A. E. K.; Mannisto, J. K.; Moslova, K.; Niegner, M.; Heliovaaera, E.; Repo, T. Synthesis of Diaryl Hydroxyl Dicarboxylic Acids from Aminos Acids. J. Org. Chem. 2020, 85, 7599.
(104) Xu, J. Stereoselectivity in the synthesis of 2-azetidinones from ketenes and imines via the staudinger reaction. ARKIVOC 2008, 2009, 21.
(105) Musio, B.; Mariani, F.; Sliwinski, E. P.; Kabeshov, M. A.; Odajima, H.; Ley, S. V. Combination of Enabling Technologies to Improve and Describe the Stereoselectivity of Wolff-Staudinger Cascade Reaction. Synthesis 2016, 48, 3515.