Formic acid: A versatile renewable reagent for green and sustainable chemical synthesis

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ABSTRACT

Formic acid is available as a major byproduct from biorefinery processing and this together with its unique properties, including non-toxicity, favorable energy density, and biodegradability, make it an economically appealing and safe reagent for energy storage and chemical synthesis. This review provides an overview of novel recent achievements in green catalytic transformations that use biogenic formic acid as an efficient and versatile reagent. The examples selected demonstrate the advantages of formic acid in addressing the key issues (minimizing the use and generation of hazardous substances while maximizing productivity under mild and benign reaction conditions) in clean chemical transformation. Special emphasis is put on the prospects of formic acid for delivering new catalytic technology to produce a plethora of tailor-made products via the flexible and selective conversion of renewable biomass resources. The potential of formic acid as a renewable C1 feedstock for both bulk and fine chemical syntheses is also outlined with examples. The role of multifunctionality in catalyst design as a key aspect in developing new catalytic concepts capable of promoting new transformations to give unprecedented selectivity and efficiency is also discussed. This article is expected to advance research on sustainable, green and affordable bio-based processes as alternatives to traditional ones with the goal to develop a fully sustainable chemical industry based on renewable resources.

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1. Introduction

The search for sustainable chemical processes has never been more important than it is today in light of current dwindling nonrenewable energy resources such as petroleum and, more importantly, their unabated negative environmental impacts. For this purpose, the exploration of new strategies to develop catalytic methods that work under ambient conditions, maximize atom utilization, and directly transform natural resources, such as renewable biomass, into useful chemical products is highly desirable [1]. A major challenge to the transition from petroleum-based feedstocks to biorenewable resources is the effective deoxygenation of bio-molecules. To this end, reactions involving biogenic carboxylic acids, in particular, formic acid (HCOOH, FA), have been identified as important reactions for the conversion of biomass feedstocks to hydrogen, chemicals, and fuel grade alkanes [1]. The most attractive features of FA, apart from its accessibility as a major byproduct from lignocellulosic biomass processing, are efficient H2 production by selective FA decomposition (Scheme 1(a)) and the
possibility of the direct formation of CO-free clean H₂ that cannot be produced from other substrates under practical and convenient conditions [2]. Given its non-toxicity and high energy density (far exceeding that of most other H₂ storage materials used today), FA is an ideal H₂ storage material with enormous potential to meet the power needs of small fuel cells for portable applications [2]. A major research endeavor in FA chemistry is finding a suitable catalytic material that can facilitate effective and controlled H₂ evolution under ambient conditions, which is possibly a critical step to establish a "carbon-neutral" H₂-energy storage system (Scheme 1(b)) based on reversible FA-CO₂ interconversion [2].

Aside from the above incentives, there is an additional interest to use FA as a bio-renewable feedstock for chemical synthesis with the aim to develop new enabling catalytic technologies based on renewable resources. In this connection, fundamental advances in the catalysis of selective FA decomposition have promoted the search for new efficient and selective reductive transformations to replace conventional catalytic hydrogenation processes [3–6]. This is largely driven by the fact that H₂ gas currently derived from non-renewable natural gas or petroleum is highly explosive, thus, posing significant safety concerns and the need for alternative safer and more effective technologies [7]. Whereas current efforts mainly involve the use of liquid FA as a convenient hydrogen source for green organic synthesis, the true potential offered by FA for efficient and selective transformations, especially toward the conversion of biogenic feedstocks, has been recognized only very recently. Scheme 2 exemplifies a diverse range of highly attractive products and building blocks that can be derived from FA [8]. Due to its reactive carbonyl and hydroxyl functionalities, FA has many kinds of reactions, including esterification, dehydration, formylation, addition to olefins, and cyclization of toluene diamines. Of further interest is that sequences of the relevant reactions such as tandem reduction-formylation, addition-amination, cascade esterification-dehydroxylation, etc., can lead to the direct "one-pot" construction of diverse, interesting, and complex structures from readily available starting materials. A key prerequisite for such advanced synthetic concepts lies in the development of new versatile catalytic approaches for flexible and controlled FA activation.

In this review, we describe recent progress in the synthetic transformations enabled by catalytic FA activation using selected examples from the literature and our own research. Although the investigation of FA as a suitable liquid medium for cleaner H₂ production and as a hydrogen storage material has been extensively reviewed [2], no review has been devoted to a discussion of the synthetic potentials offered by FA in the context of sustainable green catalysis since the classical review in 1969 covering the chemistry of FA and its simple derivatives [8]. This review will demonstrate that in many cases FA-mediated catalytic reaction systems are advantageous for better efficiency or even give previously unknown reaction routes as the basis of new sustainable chemical technologies. In
order to highlight future opportunities, emphasis is placed on understanding the key catalytic aspects in developing FA-mediated transformation strategies for the conversion of abundantly available feedstocks, with special attention to FA as a benign C1 feedstock. Emphasis is also put on the selective transformation of biomass and biomass-related model compounds by FA catalysis, which may give new attractive biorefinery strategies. The attractiveness of FA as a green solvent for organic synthesis and biomass valorization is also discussed.

2. A historical overview of the chemo-catalytic aspects of FA

In general, FA is an environmentally acceptable and highly efficient organic acid. Noticeably different from mineral acids is that FA evaporates without leaving any residue. Due to its acidity, its aldehyde nature, and its reducing properties, FA is used in a variety of fields ranging from leather processing to food preservation. In contrast to the rich chemistry already established by the organic community, efforts to explore the catalytic aspects of FA have only focused primarily on selective H2 production by FA decomposition [2]. The history behind catalytic FA decomposition dates back to the pioneering work of Sabatier et al., who identified that the decomposition of FA occur by two reaction pathways, namely, decarboxylation yielding H2 and CO2 and dehydration into CO and H2O (Scheme 3) [9]. Since then, FA has been utilized as a probe molecule for heterogeneous reactions on metals, metal oxides, metal carbides, and oxide-supported metals. Researchers employing theoretical and computational methods have also investigated FA as a molecule to probe reactivity [7]. From the standpoint of catalysis, especially interesting is that the selectivity of the two possible pathways can be controlled by the choice of catalyst. Moreover, the parallel pathways in Scheme 3 are connected by the water-gas shift (WGS) reaction, which is important in several industrial catalytic technologies such as ammonia synthesis, methanol synthesis, and Fischer-Tropsch processes [7]. The classic "volcano curve" (Fig. 1) [7] derived from the study on FA decomposition over metallic powders has been particularly valuable in shaping our current understanding of heterogeneous catalysis [7].

In the past decade, the use of FA as an alternative hydrogen or carbon source has seen renewed and increasing interest as FA has been identified as the most common and important byproduct of the biorefineries (Scheme 1) and bio-based industrial technologies [1]. The efficient and effective utilization of FA can play an important role in building resilient supply chains for a future economy as FA can serve as an energy carrier and a flexible non-petroleum feedstock that enables safe production with exceptional versatility. Another stimulus is the concept to use FA as a liquid organic hydrogen carrier to address the supply challenge associated with fuel cell technology [2]. This has prompted endeavors directed toward the selective decomposition of liquid phase FA where the production of CO-free H2 at convenient temperatures is of great importance. Along with fundamental advances on the hydrogenation of CO2 to FA at moderate temperatures and high pressures, this gives a good entry point to use cheap, non-toxic CO2 as a renewable and eco-friendly C1 source for the synthesis of value-added chemicals or fuels, and also has the promise to establish an integrated energy storage scheme for a dispatchable solar and wind powered energy system. In the last five years, reactions involving FA have experienced rapid growth. Recent applications of the homogeneous and heterogeneous catalytic behavior of metal-based compounds or related materials have also given many new challenging directions motivated by industrial demands.

3. FA-mediated selective functional group transformations

The ultimate goal in synthetic organic chemistry is to achieve the ideal synthesis in terms of activity, selectivity, as well as atom and step economy. For this objective, the developing of more sustainable catalytic processes that reduce wastes together with the use of renewable feedstocks, eco-friendly reagents and catalysts remain highly urgent. The employment of renewable, clean and relatively safe regents, such as the versatile biogenic FA, provides a potential route to this goal. In this section, we focus on the recent achievements in several types of FA-mediated selective functional group transformations with FA as a reductive agent or C1 building block. We highlight some examples of catalytic reduction of specific functional groups that use FA as an efficient hydrogen source, such as the hydrogenation of carbonyl compounds to alcohols, imines, nitriles or nitros to amines, alkynes to alkenes, and alkenes to alkanes as well as the hydrogenolysis of alcohols or epoxides. A detailed discussion of appealing tandem reactions
for the production of value added fine chemicals with the use of FA as a benign synthon is also described. These reactions include direct reductive N-formylation of quinoline derivatives, N-formylation and methylation of amines, carbonylation of olefins, and reductive hydration of alkenes. We also emphasize a novel transformation of FA, very recently demonstrated in the literature [10,11], that methanol can be efficiently and selectively produced through a catalytic disproportionation of FA.

3.1. Reduction of specific functional groups

Regarding the reduction of chemical substances using FA as a green hydrogen donor, the most studied reaction is the selective reduction of carbonyl functionalities to the corresponding alcohols of synthetic and biological interest. In the past decade, efforts have been invested in the investigation of FA-mediated asymmetric transfer hydrogenation of ketones to chiral alcohols using organometallic compounds, such as Ru [12], Rh [12], Ir [13], and Fe [14] complexes. As a consequence of the increasing demand for greener laboratory and practical industrial applications, the development of chiral catalysts covalently attached to an insoluble support has created fast growing interest. Research efforts have been devoted to the preparation of immobilized homogeneous catalysts, esp. Ru complexes, on a variety of different polymeric organic and inorganic supports for the FA-mediated chemo- and stereo-selective reduction of C=O groups, sometimes in the presence of other reducible functional groups as C=C [15–17]. For instance, the chemoselective hydrogenation of α,β-un saturated aldehydes with FA to allylic alcohols was successfully promoted by a recoverable polymer-bound Rh complex catalyst [15]. While this heterogenized catalyst has some advantages over the traditional homogenous ones in terms of ease of catalyst and product separation, its tedious preparation and catalyst recyclability remain far from satisfactory. Yu et al. [18] developed a more "user-friendly" protocol comprising catalytic Pd(0) clusters encapsulated in polyurea ([Pd(En-Cat)]) for the chemoselective hydrogenation of a wide range of aryl ketones to benzyl alcohols with FA as hydrogen source (Scheme 4). Although this polymer-based supported Pd catalyst offers simple procedures for catalyst-product separation and recovery, a catalyst supported on a solid oxide surface is more desirable from a practical point of view.

The preparation of amines, the key intermediates for the production of pharmaceuticals, dyes and ligands for transition metal-catalyzed reactions, can also use the FA-mediated catalytic reduction of imines, nitriles and nitro compounds. As a matter of fact, the chiral Ru complex systems employed for the above asymmetric saturation C=O linkages have often been applied for the asymmetric reduction of C=N group [15–17]. In 2009, Li et al. [17] developed a magnetic siliceous mesoporous foam immobilized ruthenium complex (Ru-TsDPEN, TsDPEN =N-(p-toluenesulfonyl)-1,2-diphenylethlenediamine) catalyst. This heterogeneous Ru catalyst exhibited high catalytic activity and good enantioselectivity in the asymmetric transfer hydrogenation of imine in the HCOOH-Et3N system (Scheme 5). It could be easily reused for at least nine runs with a slight ee decrease from 94% to 90%. In the case of FA-mediated nitrile hydrogenation, Beller’s group [19] demonstrated a simple catalytic reduction system for the transfer hydrogenation of (hetero)aryl nitriles to the corresponding primary amines with HCOOH-NEt3 in the presence of commercially available Pd/C (10 wt% Pd, the straightforward and selective reduction could be smoothly achieved without any additive under mild conditions (40 °C or r.t.). In the chemoselective reduction of nitro compounds with FA, various heterogeneous catalysts, such as Pd supported on carbon [20], alginate/ionic liquid highly porous monoliths [21] or zirconium phosphate [22], have been successfully prepared and used in amine production. Beller’s group [23] recently demonstrated that a cobalt oxide based nanomaterial (Co3O4-NGr@c, 2 mol% Co) prepared by the pyrolysis of cobalt-phenanthroline complexes-impregnated carbon is a highly selective catalyst for the transfer hydrogenation of nitroarenes using FA as hydrogen source (yield up to 97%, Scheme 6). This non-noble metal material performed excellently in the reduction of many structurally diverse and functionalized nitroarenes with unprecedented chemoselectivity and with halide, olefin, aldehyde, ketone, ester, amide and nitro functionalities.

Besides alcohol and amine production, the FA-mediated reduction systems can also be applied to the hydrogenation of double or triple carbon-carbon bonds. The selective catalytic hydrogenation of C≡C bonds, particularly to C=C bonds, has been the subject of numerous investigations due to its synthetic utility and also for the selectivity of metal catalysts for the semi-hydrogenation [3]. Although several heterogeneous catalytic protocols with H2, CO/H2O, trialkylammonium formate, 1,4-dihydropyridine, and organo-silanes as hydrogen sources

\[ \text{Ar} \begin{array}{c} \text{O} \\ \text{C} \end{array} \begin{array}{c} \text{R} \\ \text{HCOOH} \end{array} \begin{array}{c} \text{CO} \end{array} \begin{array}{c} \text{R} \\ \text{Pd}^\text{En-Cat} \end{array} \begin{array}{c} \text{OH} \end{array} \begin{array}{c} \text{Ar} \\ \text{FG} \end{array} \begin{array}{c} \text{HCOOH, Et3N, THF} \\ 100 \text{ °C}, 11-17 \text{ h} \end{array} \begin{array}{c} \text{NH}_2 \end{array} \begin{array}{c} \text{FG} \end{array} \begin{array}{c} \text{>50 examples} \\ \text{89%-97%} \end{array} \begin{array}{c} \text{CO}_2 \end{array} \begin{array}{c} \text{11 examples} \\ \text{90%-99%} \end{array} \]

Scheme 4. [Pd(En-Cat)]-catalyzed hydrogenation of aryl ketones to benzyl alcohols [18].

\[ \text{Ar} \begin{array}{c} \text{O} \\ \text{C} \end{array} \begin{array}{c} \text{R} \\ \text{FG} \end{array} \begin{array}{c} \text{HCOOH} \\ \text{CO}_2 \end{array} \begin{array}{c} \text{R} \\ \text{Ar} \end{array} \begin{array}{c} \text{OH} \end{array} \begin{array}{c} \text{Ar} \\ \text{FG} \end{array} \begin{array}{c} \text{HCOOH, Et3N, THF} \\ 100 \text{ °C}, 11-17 \text{ h} \end{array} \begin{array}{c} \text{NH}_2 \end{array} \begin{array}{c} \text{FG} \end{array} \begin{array}{c} \text{>50 examples} \\ \text{89%-97%} \end{array} \]

Scheme 6. Co3O4-NGr@c-catalyzed hydrogenation of nitroarenes using FA [23].
have been reported [3], those using FA as a safer and more expedient hydrogen donor are surprisingly quite limited. The first example was reported by Yu [24] in 1998, in which the mechanism of Pd/C-catalyzed transfer hydrogenation of phenylpropiolate to cis-alkene with FA was investigated. The corresponding D-labeling studies proved that palladium diformate is a key intermediate in this transformation. Wagh et al. [25] recently developed an excellent unsupported nanoporous gold-catalyzed transfer reduction strategy for chemo- and stereoselective semihydrogenation of alkynes to Z-olefins with one equivalent of FA. The system provides a facile method for the semihydrogenation of a broad range terminal/internal and aromatic/aliphatic alkynes with good functional group tolerance. In addition to the semihydrogenation of C≡C bond by FA, the FA-mediated selective reduction of the C=C bond was also achieved. In parallel to the vapor phase hydrogenation of ethylene and propylene over Pd/C with FA at relatively low temperatures (< 167 °C) [26], Li’s group [27] recently found that a Mott-Schottky type catalyst based on g-C3N4 supported Pd nanoparticles (Pd/CN) showed high activity for the hydrogenation of various olefins and bio-derived 2-methylfuran at 25 °C (yield up to 99%). All the reactions were conducted in an aqueous solution of FA under ambient atmosphere and the catalyst can be reused at least 26 times while maintaining a high 2-methyl-tetrahydrofuran yield over 99% (Fig. 2).

3.2. Hydrogenolysis and deprotection

Catalytic hydrogenolysis involving the reductive cleavage of sigma bonds, such as C-O, C-N, C-S, and C-X bonds, is a very important technology widely applied in organic synthesis, water conservation and the petroleum industry. Several recent studies have demonstrated that the hydrogenolysis of the C-O bonds in epoxides and saturated alcohols can be achieved with FA [3]. In general, epoxides are liable to hydrogenolysis under rather mild conditions because of a strong tendency to release FA [3]. In particular, the epoxide bond is especially susceptible to hydrogenolysis with FA in water solution [31]. The hydrogenolysis of epoxides is a facile reaction, and the process proceeds smoothly. Based on the significant primary kinetic aspect, FA is an efficient hydrogen donor and FA was much better than formate salts and other agents as a hydrogen donor [30]. Studies investigating the hydrogen-donating ability of the hydrogen source and the effect of the pH value indicated that the protons in the reaction medium have a promotion effect on this catalytic hydrogenolysis. Under the optimized conditions, excellent selectivity (98.4%) for isopropylbenzene was achieved at 95.2% conversion at 80 °C. Other saturated alcohols, including primary, secondary, and tertiary benzylic alcohols, have also been hydrogenolozed by the Pd/C-FA combination. Very recently, Samec’s group [31] developed a very efficient FA-mediated catalytic system using Pd/C as the catalyst for the transfer hydrogenolysis of different benzyl alcohols to the corresponding hydrocarbons in moderate to excellent yields (50%-99%). An interesting feature was that only catalytic amounts of the added base (5 equivalents relative to palladium) was required to inhibit a competing disproportionation reaction of alcohol to alkane and ketone, allowing the transfer hydrogenolysis to proceed smoothly. Based on the significant primary kinetic isotope effects identified with D-labeled FA, a mechanism was proposed which is shown in Scheme 7. Initially, a formate anion is adsorbed on the low coordinated palladium sites by inhibiting the competing disproportionation. Formate-palladium species were generated, which was followed by reversible proto-

![Fig. 2. Pd/CN-catalyzed hydrogenation of 2-methylfuran and reuse of the Pd/CN catalyst [27].](image-url)
nation and a rate determining hydride transfer to generate the active Pd with chemisorbed hydrogen. Finally, the hydrogenolysis of the alcohol was promoted by the chemisorbed hydrogen species to form the corresponding hydrocarbon.

Apart from the catalytic hydrogenolysis of C–O bond linkages discussed above, there are also a few reports dealing with the reductive cleavage of C–N bonds. On investigating the catalytic hydrogenolysis of tertiary allylic amines, Heck's group [32] reported that the Pd/C-FA-Et3N system was useful for selectively removing morpholino and piperidino groups from the tertiary amines. However, the yields for the mixture of two isomeric olefins were moderate (62%–68%) (Scheme 8). Another interesting FA-mediated hydrogenolysis is deprotection, which is widely utilized for removing a protecting moiety introduced in order to protect a reactive functional group during a series of reactions [33,34]. FA, which is known as a good solvent for most peptides, is now mainly employed for the simple cleavage of benzyl-type protecting groups from peptides as an in situ hydrogen donor. As demonstrated by Gowda [34], benzyl-type protecting groups, including Nα-benzyloxycarbonyl, Nα=2-chlorobenzyloxycarbonyl, C-terminal benzyl ester, O-benzyl ether of O-benzyllytrosine, serine or threonine, nitro of nitroarginine and Nα-benzoyloxymethyl of histidine, were successfully removed by a commercially available 10%Pd/C catalyst with FA at 25 °C. In addition, FA-mediated desulfurization and dehydrochlorination, closely connected to the wastewater treatment and fuel industry, have also been investigated with a Pd-based catalyst supported on various minerals [3].

3.3. Tandem reactions to give complex molecules

Green and sustainable chemistry necessitates the development of new compact, simple, and reliable technologies and more efficient processes to minimize environmental impacts. In this regard, tandem reactions, also known as cascade or domino reactions, involving flexible FA activation, have been considered as sustainable process intensification strategies to allow access to complex molecules with exceptionally high efficiency and atom economy. The wide application of FA as a green source for hydrogen described above has encouraged its utilization in simple reductive amination (RA) sequences for the one-step construction of higher amines from a mixture of cheaply available compounds [35]. This is particularly attractive for its step economy as compared to traditional multistep procedures. Meanwhile, FA is also known as a straightforward and convenient source for a formyl group. For example, it is used in the formylation of alcohol under solvent-free and neutral conditions catalyzed by free I2 or I2 generated in situ from Fe(NO3)3-H2O/Nal [36] as well as N-formylation of amines using Amberlite IR-120 [37], ZnO [38], CeO2 [39] and Natrolite zeolite [40] catalysts. To advance the usage of FA in more complex synthetic systems, Török's group [41] reported a two-step, one-pot domino reaction methodology using Pd/C as catalyst with FA both as a reducing and formylating agent to synthesize N-formylindolines and N-formyltetrahydroquinolines (FTHQs), an important class of intermediates for bio-active compounds, by reductive N-formylation of the corresponding indoles and quinolines. However, the pyrophoric nature of the catalyst, high catalyst loading (10 mol%Pd), and the requirement of a large excess of FA (approx. 55 equivalents) have restricted the utility of this procedure.

Very recently, we devised a novel catalytic system capable of the one-pot conversion of quinoline compounds to FTHQs via a reduction-formylation domino reaction sequence utilizing FA as a source of both hydrogen and formyl groups in the presence of supported gold NPs (Scheme 9) [42]. To test the reducing ability of FA, we initially investigated the catalytic selective transfer hydrogenation of quinolines to 1,2,3,4-tetrahydroquinolines (THQs). The gold NPs exhibited far superior catalytic activity and selectivity than those of traditional platinum-group-metal-based catalytic systems. Among the gold catalysts tested, a single phase rutile titania-supported gold (Au/TiO2-R) with a mean diameter of 2.2 nm showed the highest activity, affording the THQs in good to excellent yields (from 88% to 98%) under convenient reaction conditions within very short time of 10–50 min, e.g. in a mixed organic solvent (Et3N/DMF in a 1:8 volume ratio) and an appropriate amount of FA (15–45 equivalents) at 130 °C. The solid Au/TiO2-R catalyst can be easily separated from the reaction mixture and reused at least three times without any loss in activity. As a further demonstration of the effectiveness of this system, we found that upon switching the mixed organics to Et3N/H2O (3:10) together with 25 equivalents of FA, a Au/TiO2-R catalyst was also highly efficient for the tandem synthesis of a diversity of formyl-substituted aromatic N-heterocycles directly from quinolines, isquinoline and 7,8-benzoquinoline substituted with mono- or dimethyl, halogen, hydroxyl, and methoxy groups in good to excellent yields (from 86% to 97%) at 100 °C. This FA-involved Au/TiO2-R catalytic system was also applicable in a preparative scale reaction, with 10 mmol of isquinoline successfully formylated to afford valuable N-formyltetrahydroisoquinoline in 97% isolated yield, where the TON reached 3880. Thus, this new method provides a more cost effective, practical and scalable approach to efficiently access this type of compound.

Scheme 8. Hydrogenolysis of tertiary allylicamines with FA [32].
To fully access the valuable chemical diversity and explore the synthetic potential of FA, a general methylation of amine protocol with Pt complexes (Karstedt’s catalyst, [Pt(CH2=CHSiMe2)2O]) using FA as the C1 building block and silanes as the reducing agent was developed by Beller’s group [43]. A broad range of amines (more than 35 different amines), including aromatic and aliphatic, and primary and secondary amines, were efficiently converted to their corresponding tertiary amines in good to excellent yields (68% to 99%) under mild conditions (r.t. or 60 °C). Notably, two [N-13C]-labelled drugs, [N-13C]-4-dimethylaminoantipyrine and [N-13C]-diphenhydramine, were also obtained using H13CO2H as an inexpensive and convenient labelling reagent in 91% and 85% yield, respectively (Scheme 10). Following this pioneering study, an improved methodology was further presented by Cantat’s group [44] for the direct methylation of amines via a sequential formylation/transfer hydrogenation pathway using FA as a source of carbon and hydrogen. In contrast to the Pt catalytic system, this Ru (II)-based protocol avoided the use of hydrocarbons as extra sacrificial reductants. However, concerns still remained about the limited selectivity and relatively harsh conditions of this FA-based synthetic strategy. Another attractive example for FA utilization in cascade reactions is the hydroformylation of alkenes using FA as a C1 building block. In 2012, Porcheddu’s group [45] developed a Ru-based two-chamber system to produce a variety of alcohols from alkenes and FA via an oxo-synthesis, where FA decomposed into CO2 and H2 in one chamber and domino RWGS- hydroformylation of alkenes proceeded smoothly in the other chamber (Scheme 11). Along the same vein, FA was also found effective in the direct transformation of alkynes into alcohols or amines via a FA-promoted alkyne-to-ketone hydration coupled with FA-participated transfer hydrogenation and transfer hydrogenative RA in the presence of homogeneous Ir [46] or Ru [47] catalysts.

3.4. Disproportionation to methanol

The works presented above all feature the use of FA as a versatile reagent for fine chemical synthesis. Recently, there were also several reports on the potential of FA as an alternative C1 feedstock to generate basic commodity chemicals, specifically, the selective conversion of FA to methanol, which is an essential bulk chemical and an emerging energy resource with
many important industrial uses. Methanol is currently produced from fossil fuels, such as natural gas, coal and oil products (e.g. heavy refinery residues, naphtha). As a result of both the high demand for new reactions that utilize renewable resources and the importance of methanol, a methanol synthesis that fully or partially uses renewable resources is a highly desirable goal. In this context, the direct synthesis of methanol from FA is attractive as it will be a new renewable route to large volume organic chemicals and renewable fuels. Despite this potential, presumably due to perceived problems associated with the unproductive decomposition of FA to produce CO2/H2 or CO/H2O, developing methodologies for direct methanol synthesis from FA has long been neglected. It may be noted, however, that in an early work, Sabatier et al. [9] had already disclosed that some dimethoxymethane was produced in the thermolysis of FA over ThO2, thereby providing indirect evidence for a methanol producing pathway. In 2013, Miller et al. [10] showed for the first time that it is possible to produce methanol directly by the catalytic disproportionation of FA. A preliminary screening of various molecular iridium species led to the discovery of the well-defined [Cp*Ir-bpy(H2O)][OTf]2 (1, bpy = 2,2’-bipyridine, OTf = trifluoro-methanesulfonate) compound as an effective catalyst for the conversion of FA to MeOH at 80 °C. One salient feature of this unusual transformation, albeit it is still very limited in overall methanol yield (< 2%), is that the reaction can occur smoothly in an acidic aqueous solution (3 mol/L, pH = 1.4) without the need for an organic solvent or externally added hydrogen.

A possible hydrogen transfer reaction mechanism for the disproportionation of FA to generate methanol involving the formation of formaldehyde as a central intermediate with FA acting both as hydrogen source and substrate was proposed (Scheme 12) [10]. The first step involves the fast protonolysis of the Ir catalyst by the abstraction of the proton of FA by the basic ligand to generate cationic Ir-H complexes. The [Ir-H]+ species then undergoes dehydrogenation to give an [Ir-OH2]2− species together with the reduction of protonated FA to give the intermediate formaldehyde (step 2). Subsequent reduction of the resultant formaldehyde by the Ir-H complexes provides methanol (step 3), thereby completing the catalytic cycle. This competing concerted and sequential reaction pathway was well supported by the following experimental results: (i) 1H nuclear magnetic resonance (NMR) and electrospray ionization mass spectrometry (ESI-MS) studies revealed that dissolving precursor 1 in HCO2H/D2O (3 mol/L) caused rapid conversion at ambient temperature to a mixture of [Cp*Ir(bpy)D][OTf]2 [10] and the corresponding formate species in an approximately 9:1 ratio; (ii) heating a HCO2H/D2O solution (3 mol/L) containing 13C-enriched paraformaldehyde and 1 to 60 °C resulted in the rapid formation of 13CH3OD and 13CH2DOD, with an exceedingly high TOF of 240 h−1 for formaldehyde conversion. It is known that all the iridium complexes tested can form Ir-hydride species in FA disproportionation reactions. The higher disproportionation activity of [Cp*Ir-bpy][H2O]][OTf]2 over those other Ir complexes can be attributed to its unique activity for formaldehyde formation (Path 1→2 in Scheme 12).

To make the process more attractive in terms of the overall FA-to-MeOH efficiency, a more cost effective catalyst that can improve the selectivity to methanol while keeping the unproductive FA decomposition minimal is preferred. A Ru complex catalyst known to catalyze a wide range of FA-mediated transformations, including FA dehydrogenation and transfer hydrogenation, is one alternative to replace the Ir-based catalyst in this reaction. Ruthenium is far less expensive than iridium ($75 per oz versus $830 per oz in 2013). Very recently, Cantat’s group [11] demonstrated that using Ru (II) complexes supported by external phosphine ligands “switch off” the competitive dehydrogenation of FA to H2 and CO2. Specifically, under relatively mild conditions (150 °C) in the presence of 0.6 mol% [Ru(COD)(methylallyl)]2 + triphos and 1.5 mol% methane-sulfonylic acid (MSA), FA was fully decomposed within 1 h to give MeOH in 50.2% yield. In analogy to the above Ir-based systems, the Ru-catalyzed FA disproportionation also appeared to proceed by means of a distinct transfer hydrogenation pathway involving formaldehyde (Scheme 13). Regarding the essential factors that play a role in the desired reaction pathway, one aspect that deserves special mention is that acid promoters such as MSA significantly boosted the catalytic efficiency of [Ru(COD)(methylallyl)]2 + triphos in FA disproportionation. In line with the detailed mechanistic studies investigating the methylation of amines with CO2/H2 by Beller’s [48] and Leitner’s group [49], the crucial role of MSA can be attributed to its effectiveness in enabling a facile formation of the reactive Ru-H species. While ongoing efforts to optimize the process parameters are necessary for further enhancing MeOH productivity, one can anticipate that the successful design of a metal catalyst abundant in the Earth with superior selectivity would render the FA-based C1 conversion process a feasible option for renewable MeOH production.

**Scheme 12.** Proposed pathways for the Ir-catalyzed disproportionation of formic acid to methanol [10].

**Scheme 13.** Pathways for Ru-catalyzed disproportionation of FA to methanol [11].
4. FA-mediated selective biomass valorization

The preceding section has shown that biogenic FA is a versatile and safe reagent for many useful functional group transformations. The continuing interest in the development of new atom-efficient reactions to afford key chemicals from sustainable resources is one of the central themes in modern chemical research. This driving force has also led to efforts to develop renewable alternatives to diminish the reliance on fossil fuels. Biomass, being a globally distributed resource, can serve as a valuable source for both energy and organic carbon. Due to its renewable nature, it is the only sustainable source of functional compounds for the chemical industry. A further advantage of its chemical production from biomass is the potential to lower greenhouse gas emission because the CO2 released in the conversion process is recycled by the subsequent growth of biomass. The selective conversion of renewable biomass resources into tailor-made products, namely a "bio-refinery", is thus an important and attractive new area of research. By utilizing new green technologies, this envisaged biorefinery would provide a more energy efficient and environmentally sustainable chemical economy. The principle challenge to this advanced bio-based concepts lies in the development of new selective processes to maximize the economic and environmental benefits. A major part of recent research in the field of biorefineries is focused on the exploitation of innovative and economically-viable catalytic methods for the selective transformation of highly functionalized bio-derived feedstocks to value added chemicals and materials. Recent breakthroughs in the development of a number of FA-mediated biomass processing strategies have made this approach more attractive in terms of overall sustainability and resource utilization efficiency.

4.1. Biomass pretreatment and fractionation

Lignocellulosic biomass, the most abundant and bio-renewable resource on Earth, is considered one of the most attractive candidates to replace fossil resources for the production of fuel, fuel additives and fine chemicals. A prerequisite of lignocellulose utilization is to fractionate it into its three major components (Fig. 3) [50], namely, cellulose (35%-50%), hemicellulose (20%-35%), and lignin (10%-25%). For further downstream processing, the degradation of sugars and formation of inhibitors should be avoided or kept to a minimum. Various processes are being pursued globally for lignocellulose pretreatment, including physical, chemical, physicochemical and biological technologies. In this connection, acid pre-hydrolysis is one of the most effective pretreatment methods to disrupt the lignin structure, remove and recover hemicellulose as dissolved sugars, and reduce the crystallinity of cellulose while increasing the porosity of the biomass to facilitate the subsequent bioconversion processes like the enzymatic production of bioethanol. As opposed to the commonly used sulfuric acid, FA is a promising alternative agent to pretreat biomass that does not introduce any additional inorganic salt species, which are potential inhibitors to the downstream heterogeneous or enzymatic catalytic process. FA can be easily recycled by distillation due to its low boiling point. For example, Marzialetti et al. [51] recently reported that diluted FA (8%) can function as an efficient acidic reagent yielding a high dissolution for switchgrass (52 wt%) at an elevated temperature around 200 °C. This FA-based acid pretreatment achieved a comparable dissolution yield to H2SO4 while generating less degradation products such as furfural and HMF that cause loss of carbohydrates and inhibition to the subsequent hydrolysis and fermentation steps. Similarly, Xu et al. [52] reported the pretreatment process of corn stover based on in situ formed FA in low concentration, with an overall glucan recovery of 94% and xylan recovery of 75% at 195 °C.

Organosolv processes, on the other hand, is also an effective strategy for biomass pretreatment in which the lignocellulosic biomass is treated with a mixture of an organic solvent and water. In contrast to other chemical pretreatments that merely liberate the cellulose fraction for further processing, the main aim of the organosolv process is to fractionate lignocellulose into its individual major fractions, typically recovering relatively pure lignin as a byproduct for the papermaking industry or further valorization. As a readily available and highly biodegradable organic solvent, FA in high concentration has been successfully used in the pulping process and it shows interesting features as an agent for the organosolv fractionation of ligno-cellulosics (Fig. 4). One advantage of this organosolv processes is that a large fraction of lignin dissolves into black liquor with simultaneous hemicellulose degradation into mono- and oligosaccharides at good pulp yield, leaving solid cellulose in the residue. In general, effective delignification of the pulp requires a higher concentration (> 80%) of FA. Several approaches for the organosolv fractionation in FA have been reported under quite mild conditions (< 130 °C) that give relatively high delignification yields [53-56]. In some studies, FA was combined with catalytic amounts of H2O2, where the fractionation process relies on the action of in situ generated peroxy FA, which degrades, solubilizes and oxidizes lignin into small fragments. For example, Brosse’s group [55] reported that the FA treatment of typha grass with H2O2 exhibited much better delignification yields than an ethanol organosolv process under microwave conditions. On the other hand, Sun’s group [56] showed that the combination of FA and acetic acid provided a more convenient and cost effective way for the fractionation of native biomass such as wheat straw. Thus, by virtue of the FA and acetic acid based organosolv system, FA in
production of many useful C5-based compounds. One key derivative of LA is γ-valerolactone (GVL), which was identified as an essential component for the production of both energy and carbon-based consumer products. Given that FA is invariably co-produced in equimolar amount along with LA in the biomass dehydration process, the ideal way for making GVL is the direct use of bio-derived FA as a hydrogen source, which eliminates the need for an external H2 supply and avoids the costly purification of LA (Scheme 15) [65]. However, the implementation of this truly convenient and green approach for making GVL remains challenging, largely due to the lack of readily accessible, applicable and reusable solid catalysts that are sufficiently active and selective for H2 production from FA under the hydrothermal processing conditions. Several relevant studies on the use of homogeneous Ru-based catalysts for the direct synthesis of GVL from LA/FA feeds have been reported [62–64]. We discovered an efficient heterogeneous catalytic system for the direct conversion of bio-derived aqueous LA/FA streams into GVL catalyzed by zirconia supported small Au nanoparticles (Au/ZrO2) under mild and convenient conditions (150 °C) [65]. The high catalytic performance of the supported Au NPs for in situ generation of H2 gas by selective FA decomposition under mild aqueous conditions is essential. Recently, we further developed a noble metal-free production of GVL from FA/LA streams based on inexpensive copper-based catalysts (Cu/ZrO2) under mild aqueous conditions (160–200 °C) [66]. Although these were over 130 times less active in terms of the mass-specific reaction rates (mmol/(g·h)) based on total metal mass compared to that of the corresponding Au/ZrO2 catalytic system under identical reaction conditions, the use of earth abundant copper can be justified by its cost (less than 0.008069 $/g) and availability.

Glycerol, obtained as a high volume byproduct in the bio-

Levulinic acid (LA), a versatile platform molecule from the acid catalyzed degradation of biomass, is of interest for the

20%–30% concentration already gave the best yield of valuable hemicellulose degradation products, especially xylose.

4.2. Transformation of bio-based platform chemicals

As a flexible approach for generating tailored renewable compounds, one particularly attractive bio-based strategy is the initial generation of a set of defined platform molecules, which can be used as key intermediates to produce diverse value added chemicals. In this respect, sugars derived from lignocellulose are a very promising class of platform chemicals with versatile applications. The feasibility of a FA-catalyzed biphasic reaction system for sugar dehydration into furan derivatives (HMF and furfural) was evaluated by Hassan’s group [57]. As a result of the continuous extraction of furans from the organic methyl-isobutylketone phase, furan formation was enhanced in the aqueous fraction by inhibiting its further conversion into undesired levulinic or formic acid compounds and humins. Moreover, FA was also shown to be an excellent hydrogen donor for the subsequent reduction of HMF into 2,5-bis(hydroxymethyl)-furan (BHMF) [58] or hydrogenolysis into 1,6-hexanediol [59]. Due to that FA can be an acid catalyst, a solvent, a source of hydrogen and a deoxygenation agent, several multistep one-pot transformations using FA as the main reagent were explored. Rauchfuss’s group [60] demonstrated a facile FA-mediated synthesis of liquid fuel 2,5-dimethylfuran (DMF) from fructose, where HMF produced by FA-catalyzed fructose dehydration was first reduced to BHMF with FA in the present of Pd/C. This was followed by the generation of the targeted DMF by FA-mediated deoxygenation of the diformate ester of BHMF as a result of H2SO4-assisted BHMF esterification with FA (Scheme 14). In this case, FA provided a milder pathway for DMF synthesis and achieved an excellent yield (> 95%). The same group also described the production of hybrid fuels from sugars [61], further highlighting the potential utility of FA as both solvent and catalyst for the reactions of fructose, glucose and cellulose. 5-mesitylmethylfurfural in moderate yields of 20%–70% was obtained by a simple one-pot process comprising sugar dehydration followed by a subsequent Friedel-Crafts arylation of HMF under mild conditions.

Levulinic acid (LA), a versatile platform molecule from the acid catalyzed degradation of biomass, is of interest for the
diezel process, has also emerged as a promising platform molecule for the production of a variety of value added products. One option for future chemical utilization of glycerol is its conversion to 1,2-propanediol (1,2-PDO) by hydrogenolysis because of the huge market for 1,2-PDO as an industrial solvent, and in the antifreeze and pharmaceutical industries. Gandarias et al. [67] demonstrated that in the absence of externally fed H2 it is possible to produce 1,2-PDO by FA-mediated glycerol hydrogenolysis over a Ni-Cu/Al2O3 catalyst with 90% glycerol conversion and 82% selectivity in 24 h at 220 °C. We have recently contributed to the field of FA-mediated glycerol hydrogenolysis by discovering that Cu/ZrO2 is more effective for the conversion of glycerol into 1,2-PDO under milder conditions (94% yield after 18 h at 200 °C) [68]. Consistent with the pathway for the FA-mediated LA-to-GVL process described above, this FA-mediated process does not proceed by a simple transfer hydrogenolysis, but rather by a less straightforward transformation of glycerol with H2 in situ generated from the FA decomposition. Meanwhile, Ellman’s group [69] disclosed another FA-mediated didehydroxylation pathway to produce allyl alcohol from glycerol at 240 °C. This reaction proceeded through an orthoester-type intermediate and the further thermal transformation of the cyclic orthoesters finally generates the double bond, as described in the route depicted in Scheme 16. This unexpected route involves the FA-induced direct removal of two hydroxyl groups from glycerol with no hydride transfer process, which is clearly a departure from the conventional FA-mediated dehydration processes via a 1, 2-hydrogen shift pathway followed by reduction. A similar dehydroxylation reaction to synthesize benzoic acid from glucose-derived quinic acid was also reported by the same group [70]. This one-pot, one-reagent deoxygenation of biogenic polyhydroxyl compounds with FA will be a valuable alternative for the manufacture of reduced oxygen content products in a biorefinery.

### 4.3. Depolymerization and deoxygenation of lignin

As the most unreactive of the three components of lignocellulosic biomass, lignin has long been treated as a waste product in the pulp and paper industry, where it is burned to supply energy and recover pulping chemicals in the operation of paper mills. The production of higher value compounds from lignin is increasingly recognized as being crucial to the economic viability of integrated biorefineries. The most prominent lignin valorization strategy so far appears to be its depolymerization to generate valuable aromatic chemicals or provide a source of low molecular mass feedstocks for downstream processing. For this purpose, sustainable, selective, and catalytic procedures must be developed. Catalytic processes for both oxidative and reductive lignin depolymerization have been reported, although the use of oxidants and/or oxidizing protocols is in principle undesirable because the presence of radicals will lead to the partial re-polymerization of lignin and thus to more complex structures. Results from recent research in this field indicate a general consensus that catalytic reductive methodologies are a valuable approach to maximize the production of phenolic compounds from lignin. To this end, biogenic FA has been identified as a reagent with unique favorable properties compared to H2 gas in facilitating the efficient disassembly of lignin [71–74]. For example, switchgrass lignin can be directly converted into phenolic monomers with a yield of up to 50% by combining the depolymerization and (hydro)-deoxygenation reactions in a single step [72]. As depicted in Scheme 17, the conversion process involves the thermal treatment of lignin in ethanol or water at 300–350 °C with FA as an active hydrogen donor. Upon heating, FA decomposes completely into CO2 and active hydrogen, which combines with oxygen from the methoxy groups of lignin to form water. The presence of a typical hydrodeoxygenation catalyst, Pt/C, is essential for achieving good results. Since both depolymerization and hydrodeoxygenation occur simultaneously, this splitting reaction can result in monomers with low oxygen contents in a single step.

In the search for new energy-efficient procedures for deconstructing lignin under industrially more favorable conditions, several recent studies showed that FA-mediated catalysts may be of special importance for expedited aromatic extraction under mild and convenient conditions. Owing to the intrinsic unreactive nature of this bulky biomass, most recently developed methods are multistep or require harsh reaction conditions (high hydrogen pressures, temperatures > 350 °C). Apart from the prohibitively high capital investment, a major drawback of these procedures is that repolymerization and self-condensation of lignin and its derivatives frequently occur under the processing conditions (due to the formation of radicals and/or C–C bonds forming self-condensation reactions in acidic media). This would eventually lead to a complex pool of poorly controlled re-condensed aromatics. It is therefore very significant that Samec and co-workers [75] have found that a simple combination of FA and a heterogeneous palladium catalyst supported on activated charcoal (Pd/C) showed great promise for producing aromatic compounds from lignin under very mild conditions. The method is a mild and robust procedure in which the catalysis is performed at 80 °C in air without specialized equipment (Scheme 18). This heterogeneous palladium-catalyzed FA-mediated procedure was particular effective for the C–O bond cleavage of various lignin models con-

Scheme 16. Mechanism for FA-mediated didehydroxylation [69].

Scheme 17. Lignin depolymerization and hydrodeoxygenation in the presence of FA [72].
taining the $\beta$-O-4'-linkage, the most abundant structural motif in lignin. By slightly modifying the reaction conditions, the corresponding aryl ketones and phenols were selectively generated with > 95% yield even for models having ortho-methoxy substitution in both the 2- and 6-positions of the arylxy group. Kinetic deuterium labeling experiments supported an initial palladium-catalyzed alcohol dehydrogenation mechanism involving a palladium-formate complex as the key reaction intermediate. Degradation experiments with more challenging pine-derived native lignin, however, afforded only a moderate shift toward lower molecular weight fragments.

As a further demonstration of the possibility of achieving facile C–O cleavage for extensive lignin depolymerization, it is worthwhile to mention a recent groundbreaking discovery by the Stahl’s group [76] that FA chemistry can be designed for lignin extraction to meet the ultimate goal of producing a significant fraction of higher value mono-aromatic streams for further upgrading (Fig. 5). In contrast to previous procedures, this FA-mediated depolymerization process is notable for its mild conditions with relatively low temperatures (110 °C) and low pressure, as well as the lack of need for expensive metal catalysts. The overall yield (more than 60 wt%) of structurally identified, monomeric aromatics obtained by this FA-mediated system was the highest reported so far for lignin depolymerization. Key to this process was the subjecting of oxidized lignin to FA processing, in which the crucial formylation of the oxidized $\beta$-O-4'-linkage enabling a facile C–O cleavage involving subsequent FA elimination from the formyl intermediates was essential. Mechanistically, the beneficial effect of lignin oxidation can be attributed to the ability of the benzylic carbonyl group to polarize the C–H bond and lower the barrier for the rate-limiting E2 elimination reaction. These observations highlight the importance of developing chemical conversion technology for S-, G- and H-derived aromatics and suggest that plants containing lignin with high $\beta$-O-4 content (as much as 85% has been observed) can be feedstocks for biomass valorization. Of significant practical importance is that there is no net consumption of FA during the process of aromatic production, thus opening up a new redox-neutral pathway for lignin valorization to bulk chemicals. This feature distinguishes this approach from reported lignin conversion methods that employ FA as a source of H2 in transfer hydrogenation/hydrogenolysis reactions with heterogeneous catalysts. The high efficiency of this redox-neutral process, operating under mild ambient conditions, exemplifies once more the unique opportunities offered by FA-mediated biorefining.

### 4.4. Upgrading of bio-oil and related model compounds

Within the frame of second generation biorefinery development, bio-oil derived from fast pyrolysis has emerged as a viable choice for making liquid fuels and chemicals from lignocellulosic biomass. Unfortunately, owing to its high oxygen content and instability, the direct use of bio-oil as a transportation fuel is not feasible. A promising route to improve the properties of bio-oil is the removal of oxygenated groups by a catalytic hydrodeoxygenation (HDO). However, the use of petrochemically-derived H2 gas may not be economically suitable and it also poses safety concerns. As a result, a method that can reduce and even eliminate the external H2 consumption is of great fundamental and practical interest. Several efforts have been directed to refine bio-oil by in situ hydrogen generated in aqueous phase reforming (APR), while most of these previously reported methods suffer from problems like low product quality and/or the necessity of a prolonged operation. Thus, the development of refining methods with more convenient reagents, simpler procedures and higher HDO efficiency is highly desired. As with the case with many of the conversions of various forms of bio-derived feedstocks, one can also envision that the FA-mediated reductive strategy can afford a green, efficient and cost effective protocol for bio-oil upgrading under HDO conditions, since FA is increasingly accepted as a carbon neutral, renewable and readily available hydrogen source. Xiong and co-workers [77] conceptually demonstrated that for the processing of crude bio-oil in methanol with liquid FA, the properties of the liquid product can be significantly improved (Fig. 6). Among the catalysts investigated, Ru was more effective than Pd and Ni based catalysts. Furthermore, the reaction routes of the main components showed that the partial hydrogenation was achieved without coke or tar formation. There was no obvious deep reduction
that took place, yet the compounds containing alkenyl or aldehyde groups were almost completely reduced.

Preliminary efforts are also devoted to FA-mediated upgrading of bio-oil model compounds, such as phenolic monomers and furans, with the aim to understand the requirements of this type of biorefining and the mechanisms. It should also be mentioned that, although many Group VIII and IB metals, notably palladium and gold, have been shown to facilitate FA dehydrogenation, the high cost and scarcity of these metals make the relevant systems difficult for practical implementation. Huge challenges but also opportunities are present for developing less expensive metal catalysts that are stable and active for FA-mediated biorefining. In a recently published study focusing on the HDO of phenol and furfural by in situ H2 generation for FA-mediated biorefining. I n a r e c e n t l y p u b l i s h e d s t u d y f o r m o r e b e n i g n  c a t a l y t i c  s y n t h e s i s . D e s p i t e  t h e  r e l e v a n t  s y s t e m s  d i f f i c u l t  f o r  p r a c t i c a l  i m p l e m e n t a t i o n .

...resources. Although much progress has been made in the past few years, more research needs to be conducted in the area of FA-mediated catalysis. Given the high demands of the modern industry for environment, economy and sustainable development, green and affordable FA-mediated processes have great potential and significance for developing sustainable solutions, especially when the following aspects can be realized: (1) select and exploit the appropriate catalytic material containing a suitable active metal species for useful catalytic reactions; (2) controllably activate FA in the presence of other reactants and reagents; (3) understand the mechanism of the complex reactions that take place, and (4) stabilize the catalyst during the catalytic reaction process. It is expected that FA-based chemistry, particularly FA-mediated catalytic transformations, will be of increasing scientific and engineering interest in the coming years.

5. Conclusions and outlook

This review described recent efforts on green chemical reactions and processes based on the use of biogenic FA as a reagent. Many chemical transformations based on renewable feedstocks have been developed in response to the demand for more benign chemical syntheses. This development promoted the use of FA because of its unique properties as described in this review. As illustrated by a variety of transformations, FA is a powerful and versatile reagent building block for sustainable organic synthesis. Of specific interest to modern industry is the carbon-neutral production of many bio-renewable chemicals achieved by the development of new FA-mediated catalytic process concepts for the integrated valorization of biomass resources.

References

Despite its structural simplicity, formic acid is a versatile bio-renewable feedstock for opening up new chemical space to lead to the discovery of new sustainable reactions of unprecedented selectivity.
利用多功能、多用途的可再生甲酸实现化学品的绿色与可持续合成

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摘要：近年来，随着化石资源日趋短缺以及由此带来的人类生存环境日益恶化，生物质等可再生资源的高效可持续利用已成为世界各国科学家关注的热点。甲酸的多酸性及良好的溶剂性可应用于多种官能团的选择转化过程。甲酸化学将得到产业界与学术界越来越多的关注和研究。在生物质催化转化方面，甲酸作为一种环境友好可再生的多功能试剂可应用于包括喹啉衍生物的还原甲酰化、胺类化合物甲酰化、生物基甲酸在绿色有机合成和生物质转化等方面表现出巨大潜力。甲酸的酸性质及良好溶剂特性可应用于生物质资源预处理过程，相较依赖对无机酸预处理体系具有沸点低、易分离、不引入无机离子、对下游反应兼容性强等优点；而作为高效氢源，甲酸也被广泛研究应用于生物质平台化合物选择催化转化制高附加值化学品、木质素降解制芳烃化合物和生物油加氢脱氧精制处理等过程。鉴于甲酸亦可用作烷基化原料，甲酸相较传统氢气具有操作简便可控、条件温和、具有良好化学选择性等优点。该领域目前已取得了一定成果并得到了快速发展，但将其转化为可利用的资源形式仍然面临挑战。综上所述，生物基甲酸在绿色有机合成和生物质转化等方面表现出巨大潜力，而其多功能性和多用途性对于实现原料的高效利用及目标产物的高选择性至关重要。该领域目前取得了显著成果，其在相关过程的挑

关键词：甲酸；新型催化；可再生氢能源；环境友好型试剂；化学合成；生物精炼

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