

Solid Acid Catalysts for the Selective Conversion of Biomass to Levulinic Acid

Mohamed Faiz Farook R. Mohamed Aboobucker¹, Indra Neel Pulidindi^{2,*}

Abstract

Levulinic acid (LA) has emerged as a versatile platform chemical with significant potential for producing renewable fuels like gamma-valerolactone (GVL), biodegradable polymers, and fine chemicals from biomass (both terrestrial and marine which are rich in carbohydrate). The selective conversion of biomass-derived carbohydrates to LA requires efficient catalytic systems that can overcome the recalcitrance of the biomass, namely the stiff-necked structural integrity of cellulose and the kind of strong interactions between the cellulose fraction and the lignin component. This often makes pretreatment a necessary as well as a costly incurrent step in the conversion of biomass to fuels and chemicals in general and to LA in particular. Solid acid catalysts have gained considerable attention due to their environmental benefits, ease of separation, and potential for reuse compared to conventional mineral acids apart from being highly active as well as selective. An insight into the recent advances in the development of the solid acid catalysts for the conversion of biomass to LA is provided. Apart from highlighting the current understanding on the mechanistic pathways of biomass conversion, a detailed account of various classes of solid acid catalysts including sulfonated activated carbon materials, simple metal oxides, zeolites (aluminosilicates), heteropolyacids (polyoxometallates or complex metal oxides), ion-exchange resins, metal organic frameworks (MOFs), covalent organic frameworks (COFs), aerogels and evaluate their catalytic performance. The effects of reaction parameters, catalyst structure-activity relationships, and challenges related to catalyst stability and biomass preprocessing are critically analyzed. Finally, the emerging trends and future research directions towards the commercial viable solid acid-catalyzed levulinic acid production from renewable biomass feedstock is exemplified.

Keywords: Levulinic acid, Biomass, Solid acid catalyst, Carbon based catalysts; Zeolites; GaHPMo; Ga modified zeolite;

*Author for Correspondence

Indra Neel Pulidindi

E-mail: rababmahdi49@gmail.com

¹Student, Saveetha Medical College (SMC) and Saveetha Institute of Medical and Technical Sciences (SIMATS), Saveetha Nagar, Thandalam, India

²Assistant Professor, Department of Ear, Nose and Throat, Saveetha Medical College (SMC) and Saveetha Institute of Medical and Technical Sciences (SIMATS), Saveetha Nagar, Thandalam, India

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INTRODUCTION

The transition from a fossil fuel-based economy to a sustainable bio-based economy necessitates the development of efficient processes for converting renewable biomass into valuable chemicals and fuels. Lignocellulosic biomass, comprising cellulose, hemicellulose, and lignin, represents the most abundant renewable carbon resource on Earth and offers tremendous potential as a feedstock for producing platform chemicals. Among these platform chemicals, levulinic acid (4-oxopentanoic acid) has been identified by the U.S. Department of Energy as one of the top value-added chemicals derivable from biomass.

Levulinic acid serves as a versatile building block for synthesizing numerous high-value products

including γ -valerolactone, alkyl levulinates, diphenolic acid, and various polymers. Additionally, levulinic acid can be converted to fuel additives, solvents, and pharmaceutical intermediates, making it an attractive target for biomass valorization. The global market for levulinic acid has been expanding rapidly, driven by increasing demand for bio-based chemicals and the push toward sustainable industrial processes [1–31].

Conventionally, levulinic acid production from biomass involves the acid-catalyzed hydrolysis of cellulose to glucose, followed by dehydration to 5-hydroxymethylfurfural (5-HMF), and subsequent rehydration to levulinic acid and formic acid. While mineral acids such as sulfuric acid and hydrochloric acid have been traditionally employed for this transformation, they present significant drawbacks including equipment corrosion, difficulties in catalyst separation and recovery, generation of waste salts during neutralization, and environmental concerns. These limitations have stimulated extensive research into developing heterogeneous solid acid catalysts as environmentally benign and economically viable alternatives. Solid acid catalysts offer several advantages over homogeneous mineral acids: they eliminate corrosion issues, simplify product separation and catalyst recovery, reduce waste generation, and enable catalyst reuse across multiple reaction cycles. Moreover, the tunable acidity, porosity, and surface properties of solid acid catalysts provide opportunities to enhance selectivity toward levulinic acid while suppressing undesired side reactions such as humin formation and excessive degradation.

Despite these advantages, significant challenges remain in developing solid acid catalysts that can match or exceed the performance of mineral acids. Key obstacles include achieving sufficient catalytic activity, maintaining stability under harsh hydrothermal reaction conditions, preventing catalyst deactivation from coking and leaching, and ensuring good accessibility of biomass-derived substrates to active sites. This review comprehensively examines the state-of-the-art in solid acid catalyst development for biomass conversion to levulinic acid, discussing mechanistic insights, catalyst design strategies, performance evaluation, and future prospects for advancing this technology toward commercial implementation.

REACTION MECHANISM AND PATHWAYS FOR LEVULINIC ACID FORMATION

The conversion of lignocellulosic biomass to levulinic acid involves a complex series of reactions that depend on the nature of the biomass feedstock, reaction conditions, and catalyst properties. Understanding these reaction pathways is essential for designing effective solid acid catalysts and optimizing process conditions. The overall transformation can be divided into several key steps. First, the cellulosic fraction of biomass must undergo hydrolysis to release glucose monomers. This depolymerization step requires breaking the β -1,4-glycosidic bonds that link glucose units in the cellulose polymer. The efficiency of this step depends on both the crystallinity of cellulose and the accessibility of acid sites to the glycosidic bonds. Amorphous cellulose hydrolyzes more readily than crystalline cellulose, and pretreatment strategies are often employed to disrupt the crystalline structure and enhance reactivity.

Once glucose is liberated, it undergoes a series of dehydration reactions to form 5-hydroxymethylfurfural. This transformation proceeds through multiple pathways involving various intermediates. In acidic aqueous media, glucose can isomerize to fructose, which dehydrates more readily than glucose to form 5-HMF. The fructose dehydration pathway is generally favored under acid catalysis and proceeds through cyclic intermediates with stepwise elimination of three water molecules. Alternatively, glucose can directly dehydrate through open-chain intermediates, though this pathway is typically slower. The subsequent conversion of 5-HMF to levulinic acid involves acid-catalyzed rehydration. Water molecules add to the furan ring of 5-HMF, leading to ring opening and formation of levulinic acid along with formic acid as a co-product. This step is relatively facile under acidic conditions and generally proceeds with high selectivity when 5-HMF is the starting material.

However, the selectivity toward levulinic acid is often compromised by competing side reactions. Glucose and fructose can undergo reverse aldol condensation, leading to formation of smaller sugar fragments. 5-HMF is prone to polymerization reactions, particularly under acidic conditions, forming insoluble humins that reduce yield and deactivate catalysts through pore blockage. Additionally, levulinic acid itself can undergo further degradation or participate in condensation reactions at elevated temperatures. The reaction mechanism is significantly influenced by the acidity type and strength of the catalyst. Brønsted acid sites facilitate proton transfer steps involved in hydrolysis, isomerization, and dehydration reactions. The acid strength must be carefully balanced—too weak and the catalyst shows insufficient activity, too strong and excessive degradation or humin formation occurs. Lewis acid sites can also contribute to certain steps, particularly glucose-fructose isomerization, and bifunctional catalysts containing both Brønsted and Lewis sites have shown promise for enhancing overall conversion efficiency.

Solvent effects play a crucial role in determining reaction pathways and product distribution. Water is necessary for hydrolysis steps but also promotes undesired side reactions and can leach certain solid acid catalysts. Biphasic systems employing both aqueous and organic phases have been investigated to continuously extract 5-HMF or levulinic acid from the reactive aqueous phase, thereby minimizing degradation. The choice of solvent system must be optimized alongside catalyst selection to achieve maximum levulinic acid yields.

CLASSES OF SOLID ACID CATALYSTS

Sulfonated Carbon-Based Catalysts

Sulfonated carbon materials have emerged as highly promising solid acid catalysts for biomass conversion to levulinic acid due to their strong acidity, thermal stability, and resistance to poisoning. These materials feature sulfonic acid groups ($-\text{SO}_3\text{H}$) grafted onto carbon frameworks, providing Brønsted acidity comparable to concentrated sulfuric acid while maintaining the advantages of heterogeneous catalysts.

The synthesis of sulfonated carbon catalysts typically involves carbonization of carbohydrate precursors or biomass followed by sulfonation using concentrated sulfuric acid or other sulfonating agents. Incomplete carbonization yields materials with both rigid aromatic carbon frameworks and flexible polycyclic aromatic carbon sheets decorated with sulfonic acid groups. This unique structure provides accessible acid sites while maintaining structural integrity under reaction conditions.

Numerous studies have demonstrated the effectiveness of sulfonated carbon catalysts for levulinic acid production. The high concentration of sulfonic acid groups, combined with the hydrophobic nature of the carbon support, creates a favorable environment for glucose dehydration while limiting excessive water-promoted degradation. The carbon framework can also facilitate adsorption of substrate molecules, increasing their local concentration near acid sites and enhancing reaction rates.

Different carbon precursors have been investigated, including glucose, sucrose, cellulose, lignin, and various types of biomass waste. The choice of precursor influences the textural properties, surface area, and acid site distribution of the resulting catalyst. Generally, materials derived from sugar carbonization exhibit higher acid site densities but lower surface areas, while those from biomass pyrolysis show greater porosity but may have less uniform acid site distribution.

Despite their advantages, sulfonated carbon catalysts face challenges related to acid site leaching in hot water, particularly at the elevated temperatures required for effective biomass conversion. Research has focused on developing strategies to enhance catalyst stability, including increasing the degree of carbonization, optimizing sulfonation conditions, and incorporating additional structural elements to anchor sulfonic acid groups more firmly to the carbon framework.

Acidic Ion-Exchange Resins

Ion-exchange resins, particularly sulfonated polystyrene-divinylbenzene copolymers such as Amberlyst and Nafion resins, represent commercially available solid acid catalysts that have been extensively studied for biomass conversion processes. These materials contain high concentrations of sulfonic acid groups attached to polymer backbones and are widely used in industrial applications.

Amberlyst-15 and similar resins have been investigated for cellulose and glucose conversion to levulinic acid. These catalysts exhibit good activity at moderate temperatures and can achieve reasonable levulinic acid yields from simple sugars. However, their performance is limited by relatively low thermal stability, with significant loss of activity occurring above 120–140°C. This temperature constraint reduces reaction rates and necessitates longer reaction times to achieve high conversions.

The polymer matrix of ion-exchange resins can also undergo degradation under hydrothermal conditions, leading to loss of sulfonic acid groups and reduced catalytic activity over multiple cycles. Swelling of the polymer in aqueous media affects pore accessibility and can alter the local environment around acid sites. Despite these limitations, the commercial availability, well-characterized properties, and ease of handling make ion-exchange resins useful benchmark catalysts for comparative studies.

Perfluorosulfonic acid resins like Nafion offer improved thermal and chemical stability compared to conventional polystyrene-based resins due to their perfluorinated backbone. The super acidic nature of Nafion provides high catalytic activity, though the high cost of these materials limits their practical application for large-scale biomass processing. Research continues on developing more robust and cost-effective resin-type catalysts through modifications to polymer structure and crosslinking density.

Zeolites and Mesoporous Silica-Based Catalysts

Zeolites are crystalline aluminosilicates with well-defined pore structures and tunable acidity, making them attractive candidates for solid acid catalysis. Various zeolite frameworks including H-ZSM-5, H-Y, H-Beta, and H-Mordenite have been evaluated for biomass conversion to levulinic acid. The Brønsted acidity in zeolites arises from bridging hydroxyl groups associated with framework aluminum atoms, and the acid strength can be adjusted by varying the Si/Al ratio.

However, conventional zeolites face significant limitations for processing bulky biomass-derived molecules. The microporous nature of zeolites (pore sizes typically less than 1 nm) severely restricts diffusion of cellulose oligomers and even glucose molecules, leading to slow reaction rates and poor catalyst utilization. External surface acid sites may contribute to catalytic activity, but the majority of the acid sites within the micropores remain inaccessible to substrates.

To overcome these mass transfer limitations, hierarchical zeolites with multimodal pore structures have been developed. These materials combine the strong acidity of zeolites with mesopores or macropores that facilitate substrate diffusion. Synthesis strategies for creating hierarchical zeolites include dealumination, desilication, templating with mesoporous structure-directing agents, and construction from zeolite nanocrystals. Such modified zeolites show enhanced activity for biomass conversion compared to conventional microporous zeolites.

Mesoporous silica materials such as MCM-41, SBA-15, and related structures offer large, uniform pore sizes that accommodate biomass-derived molecules. However, pure silica mesoporous materials lack catalytic activity and require functionalization with acid groups. Various approaches have been employed including incorporation of framework aluminum to generate Brønsted acid sites, grafting of sulfonic acid groups onto pore walls, and immobilization of acidic species such as heteropoly acids. The challenge lies in achieving high acid site density while maintaining structural integrity and preventing leaching during reaction.

Sulfonated mesoporous silica catalysts combine the advantages of controlled pore architecture with strong Brønsted acidity from sulfonic acid groups. These materials can be synthesized through co-condensation of silica precursors with organosilanes bearing thiol or sulfonic acid groups, or through post-synthetic grafting. The ordered mesostructure provides good accessibility to active sites, though care must be taken to prevent hydrolysis of Si-C bonds under hydrothermal conditions.

Heteropoly Acids and Supported Heteropoly Acid Catalysts

Heteropolyacids (HPAs), particularly those with Keggin structure such as $\text{H}_3\text{PW}_{12}\text{O}_{40}$ (phosphotungstic acid) and $\text{H}_3\text{PMo}_{12}\text{O}_{40}$ (phosphomolybdic acid), are strong Brønsted acids with catalytic activity exceeding that of conventional mineral acids. These discrete polyoxometalate anions possess well-defined structures and can function as both homogeneous and heterogeneous catalysts depending on their physical state and solubility.

In their bulk form, HPAs exhibit high catalytic activity for glucose and cellulose conversion to levulinic acid. The strong acidity and ability to form pseudoliquid phases that accommodate polar reactants make HPAs effective for biomass processing. However, bulk HPAs suffer from very low surface areas and high solubility in water, leading to practical difficulties in catalyst recovery and reuse. Leaching of the HPA into the aqueous reaction medium results in loss of heterogeneity and the same separation challenges associated with homogeneous acid catalysts.

To address these limitations, researchers have developed supported HPA catalysts by dispersing HPAs on various solid supports including activated carbon, silica, zirconia, and titania. The support provides high surface area for HPA dispersion while potentially reducing water solubility through interactions between the HPA and support surface. Activated carbon supports are particularly attractive due to their hydrophobic character, which may help retain HPAs under reaction conditions.

The stability of supported HPAs remains a critical challenge. Even with supports, significant leaching can occur during aqueous-phase reactions, especially at elevated temperatures. Strategies to improve stability include using supports with stronger HPA-support interactions, incorporating HPAs into the support framework, creating Cs or NH_4 salts of HPAs with reduced solubility, and encapsulating HPAs within mesoporous materials. Despite progress, developing truly stable supported HPA catalysts for multiple-cycle biomass conversion remains an active area of research.

Metal Oxide-Based Solid Acid Catalysts

Metal oxides and mixed metal oxides can exhibit both Brønsted and Lewis acidity depending on their composition and surface properties. Materials such as niobium oxide (Nb_2O_5), zirconium oxide (ZrO_2), titanium oxide (TiO_2), and tungsten oxide (WO_3) have been investigated as solid acid catalysts for biomass conversion, either in their pure forms or as mixed oxides and composites.

Niobium oxide has received particular attention due to its relatively strong acidity, thermal stability, and water tolerance. Hydrated niobic acid ($\text{Nb}_2\text{O}_5 \cdot n\text{H}_2\text{O}$) possesses both Brønsted and Lewis acid sites arising from surface hydroxyl groups and coordinatively unsaturated Nb sites. Studies have shown that niobic acid can catalyze glucose conversion to levulinic acid, though typically with moderate activity compared to stronger solid acids like sulfonated carbons. The advantage of niobic acid lies in its excellent stability under hydrothermal conditions with minimal leaching.

Sulfated metal oxides, particularly sulfated zirconia ($\text{SO}_4^{2-}/\text{ZrO}_2$), exhibit super acidic properties arising from the electron-withdrawing effect of sulfate groups on adjacent metal centers. These materials show high activity for various acid-catalyzed reactions and have been studied for biomass conversion. However, sulfated zirconia can suffer from sulfate leaching in hot water, leading to gradual deactivation. Modifications such as incorporation of tungsten or other elements have been pursued to enhance stability.

Phosphated metal oxides represent another class of solid acids with improved hydrothermal stability compared to sulfated analogs. Phosphate groups bound to metal oxide surfaces provide Brønsted acidity while forming more stable M-O-P linkages that resist hydrolysis. Phosphated titania, zirconia, and niobia have been investigated, with performance depending strongly on phosphate loading and calcination temperature.

Mixed metal oxides combining different elements can offer synergistic effects and tunable properties. For example, tungstated zirconia, niobium-tungsten mixed oxides, and various other combinations have been explored. The ability to adjust acid strength, hydrophobicity, and structural stability through compositional control makes mixed metal oxides versatile platforms for catalyst design, though achieving optimal performance for levulinic acid production remains challenging.

CATALYST PERFORMANCE AND STRUCTURE-ACTIVITY RELATIONSHIPS

The catalytic performance of solid acid catalysts for biomass conversion to levulinic acid depends on multiple interrelated factors including acid site type and strength, surface area and porosity, hydrophobic/hydrophilic balance, and stability under reaction conditions. Understanding structure-activity relationships is essential for rational catalyst design and optimization.

Acidity characteristics represent the most fundamental catalyst property influencing performance. Both the number and strength of acid sites affect reaction rates and selectivity. Generally, higher acid site concentrations correlate with increased catalytic activity, though excessively strong acidity can promote undesired degradation reactions and humin formation. An optimal acid strength exists that balances sufficient activity for cellulose hydrolysis and glucose dehydration while minimizing over-degradation of products and intermediates.

The type of acid sites—Brønsted versus Lewis—also influences reaction pathways. Brønsted acid sites directly donate protons and are essential for hydrolysis of glycosidic bonds and dehydration reactions. Lewis acid sites can facilitate glucose-fructose isomerization and may assist in certain dehydration steps through coordination with substrate hydroxyl groups. Some studies suggest that bifunctional catalysts containing both Brønsted and Lewis sites may offer advantages by enabling complementary reaction pathways.

Textural properties significantly impact catalyst performance by controlling substrate accessibility to active sites. High surface area generally improves catalyst utilization, but surface area alone does not determine activity. Pore size distribution is equally important—micropores may provide high site density but restrict diffusion of bulky cellulose fragments, while large mesopores and macropores facilitate mass transfer but may yield lower volumetric activity. Hierarchical pore structures combining different pore size regimes often show optimal performance.

The hydrophobic/hydrophilic character of the catalyst surface influences substrate adsorption and product desorption. Hydrophobic surfaces can selectively adsorb glucose while repelling water, potentially enhancing dehydration reactions and suppressing hydrolysis of desired products. Carbon-based catalysts inherently possess hydrophobic character, which may contribute to their effectiveness. Conversely, some degree of hydrophilicity is necessary to maintain good dispersion in the aqueous reaction medium and prevent catalyst agglomeration.

Catalyst stability under hydrothermal conditions represents a critical performance criterion. Leaching of active species into solution compromises heterogeneity and reduces activity over multiple reaction cycles. Different catalyst types exhibit varying degrees of stability—sulfonated carbons and metal oxides generally show better retention of activity than supported heteropoly acids or certain ion-exchange resins. Evaluating stability requires testing over multiple reaction cycles rather than relying solely on single-batch results.

Comparative studies have attempted to rank different solid acid catalysts based on performance metrics such as levulinic acid yield, substrate conversion, and selectivity. While sulfonated carbon-based catalysts frequently emerge as top performers due to their combination of strong acidity, appropriate surface properties, and good stability, no single catalyst has been universally superior across all reaction conditions and substrate types. The optimal catalyst choice depends on the specific biomass feedstock, reaction system, and process requirements.

INFLUENCE OF REACTION PARAMETERS

Beyond catalyst properties, various reaction parameters significantly influence the conversion of biomass to levulinic acid and must be optimized to achieve maximum yields and productivity.

Temperature exerts a dominant effect on reaction kinetics and product distribution. Higher temperatures accelerate all reaction steps including cellulose hydrolysis, glucose dehydration, and 5-HMF rehydration. However, elevated temperatures also promote undesired side reactions such as glucose degradation, humin formation, and levulinic acid decomposition. Most studies report optimal temperatures in the range of 150–200°C, balancing reaction rate against selectivity loss. The specific optimal temperature depends on the catalyst type, substrate concentration, and reaction time. Reaction time determines the extent of conversion and must be balanced against selectivity considerations. Short reaction times may yield incomplete substrate conversion, while excessively long times lead to over-degradation of levulinic acid. The optimal reaction time depends strongly on temperature, catalyst loading, and substrate concentration. Kinetic studies tracking concentration profiles of reactants, intermediates, and products over time provide valuable insights for identifying conditions that maximize levulinic acid yield. Catalyst loading affects both the reaction rate and economic viability of the process. Higher catalyst concentrations generally increase conversion rates, allowing shorter reaction times or lower temperatures. However, excessive catalyst loading leads to diminishing returns and increases process costs. Typical catalyst loadings range from 2–20 wt % relative to substrate, with optimal values depending on catalyst activity and reaction conditions.

Substrate concentration influences reaction kinetics and also impacts process economics. Higher substrate concentrations improve productivity and reduce processing volumes, but may lead to increased viscosity, mass transfer limitations, and formation of insoluble humins that foul catalysts. Most laboratory studies employ relatively dilute solutions (1–10 wt% substrate), while commercial viability likely requires processing more concentrated feeds. The choice of solvent or solvent system can dramatically affect levulinic acid yields. Pure water is the most environmentally benign solvent but also promotes degradation reactions. Organic solvents or water-organic biphasic systems have been investigated to extract 5-HMF or levulinic acid from the reactive aqueous phase, thereby preventing their further degradation. Common organic phases include γ -valerolactone, tetrahydrofuran, methyl isobutyl ketone, and various alcohols. Salt addition to the aqueous phase can enhance product partitioning into the organic phase. While biphasic systems often improve yields, they add complexity and separation costs. Mechanical agitation and mass transfer considerations become important when scaling up beyond laboratory studies. Proper mixing ensures uniform temperature and concentration distributions, facilitates substrate-catalyst contact, and minimizes external mass transfer limitations. The physical form of the catalyst (powder versus granules) affects suspension behavior and influences the degree of mass transfer control.

BIOMASS FEEDSTOCK CONSIDERATIONS AND PRETREATMENT

The nature of the biomass feedstock significantly influences the efficiency of conversion to levulinic acid and dictates requirements for catalyst design and reaction conditions. Real biomass is compositionally and structurally complex, presenting challenges beyond those encountered when processing model substrates like pure glucose or cellulose.

Lignocellulosic biomass consists of cellulose (40–50%), hemicellulose (20–30%), and lignin (15–30%), with composition varying among different biomass sources such as agricultural residues,

forestry waste, energy crops, and municipal solid waste. Only the cellulose and hemicellulose fractions are readily convertible to levulinic acid, while lignin is largely recalcitrant to acid-catalyzed conversion under typical conditions. The presence of lignin can complicate processing by reducing substrate accessibility, contributing to catalyst deactivation, and forming undesired products. Cellulose crystallinity represents a major barrier to efficient conversion. Crystalline cellulose regions resist hydrolysis due to extensive hydrogen bonding networks and limited accessibility to catalyst active sites. The crystallinity index of cellulose varies among biomass types and decreases with pretreatment. Even with strong solid acid catalysts, conversion of highly crystalline cellulose to levulinic acid may remain incomplete without extended reaction times or harsh conditions that promote degradation.

Hemicellulose, being less crystalline and more easily hydrolyzed than cellulose, can contribute to levulinic acid production through conversion of hexose sugars (glucose, mannose, galactose) released upon hydrolysis. However, pentose sugars (xylose, arabinose) derived from hemicellulose do not follow the same pathway to levulinic acid and instead form furfural. The fate of pentoses depends on reaction conditions and can influence overall product distribution and byproduct formation. Biomass pretreatment is often necessary to enhance reactivity and improve levulinic acid yields from raw biomass. Common pretreatment methods include: Physical pretreatments such as milling and grinding reduce particle size and increase surface area, improving catalyst-substrate contact. However, mechanical pretreatment alone typically provides limited enhancement of cellulose accessibility due to persistent crystallinity and lignin shielding.

Chemical pretreatments including dilute acid treatment, alkaline treatment, and organosolv processes can partially remove lignin and hemicellulose while disrupting cellulose crystallinity. These pretreatments significantly improve subsequent conversion but add process complexity and costs. The choice of pretreatment must be optimized in conjunction with the catalytic conversion step. Hydrothermal pretreatment using hot compressed water or steam explosion can effectively reduce biomass recalcitrance without requiring additional chemicals. These methods partially hydrolyze hemicellulose and increase cellulose accessibility, though care must be taken to avoid excessive degradation during pretreatment. An emerging approach involves one-pot conversion processes where pretreatment and catalytic conversion occur in a single reactor. This simplification can reduce capital costs and energy consumption but requires catalysts capable of maintaining activity in the presence of biomass impurities and degradation products. Few solid acid catalysts have been successfully demonstrated for direct conversion of raw biomass to levulinic acid with acceptable yields. Impurities present in real biomass can adversely affect catalyst performance. Inorganic elements such as alkali and alkaline earth metals can neutralize acid sites, while proteinaceous materials and other nitrogen-containing compounds may adsorb strongly to active sites. Developing robust catalysts that tolerate these impurities or implementing effective washing procedures represents an important practical consideration.

CHALLENGES AND FUTURE RESEARCH DIRECTIONS

The diversity of the biomass feedstock is a challenge for the conversion of the cellulose component to levulinic acid. The suitability of the catalyst should be demonstrated over a wide range of biomass feedstock [26–28]. Major agricultural residues that can be considered for the industrial scale production of levulinic acid, include, rice straw, corn stover, sweet sorghum bagasse, miscanthus, *Pinus radiata*, *Cicer arietinum*, cotton, and sugar cane bagasse (Figure. 1).

Despite significant progress in developing solid acid catalysts for biomass conversion to levulinic acid, like, fine tuning of the Bronsted and Lewis acid sites, as in the case of heteropoly acid modified zeolites, several challenges must be addressed to enable commercial implementation of this technology. The interplay of the suitable combination of Lewis acid sites for the isomerization of glucose to fructose and the subsequent dehydration and rehydration reactions (Figure 2) catalyzed by the Bronsted acid sites is well exploited in the design of the solid acid catalysts (Figure 3).



Figure 1. Suitable biomass for the industrial scale production of levulinic acid [Adapted with permission from 26, 27].

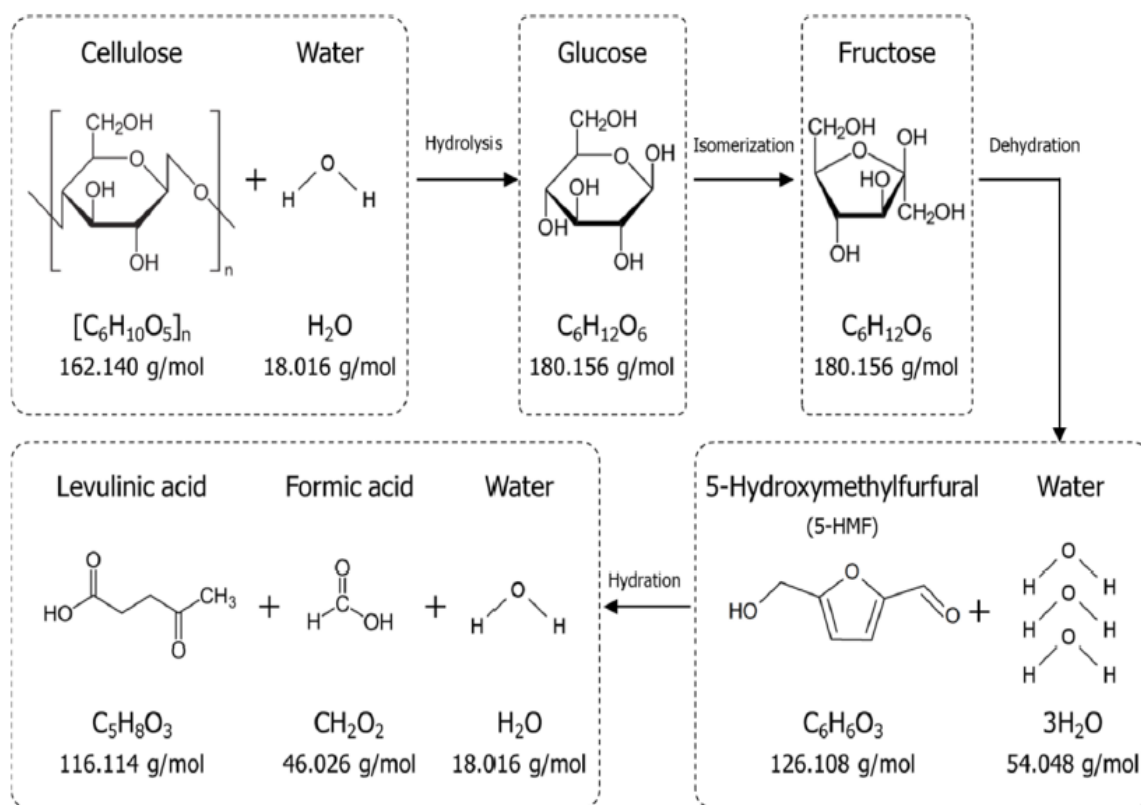


Figure 2. Molecular level mechanistic steps for the conversion of cellulose to levulinic acid [Adapted with permission from 26].

The SEM image with the EDAX images of the Ga salt of molybdophosphoric acid with the Lewis acid sites (Ga^{3+}) and the Bronsted acid sides of the molybdophosphate anion $[PMo_{12}O_{40}]^{3-}$ were shown in Figure 3A. Likewise, the Lewis acid sites (Ga^{3+}) uniformly dispersed over the aluminosilicate (mordenite zeolite support) were shown in Figure 3B. Such fine interplay of the Lewis and Bronsted acid sites was found play a crucial role in the performance of the solid acid catalyst for the conversion of glucose to levulinic acid [29, 30].

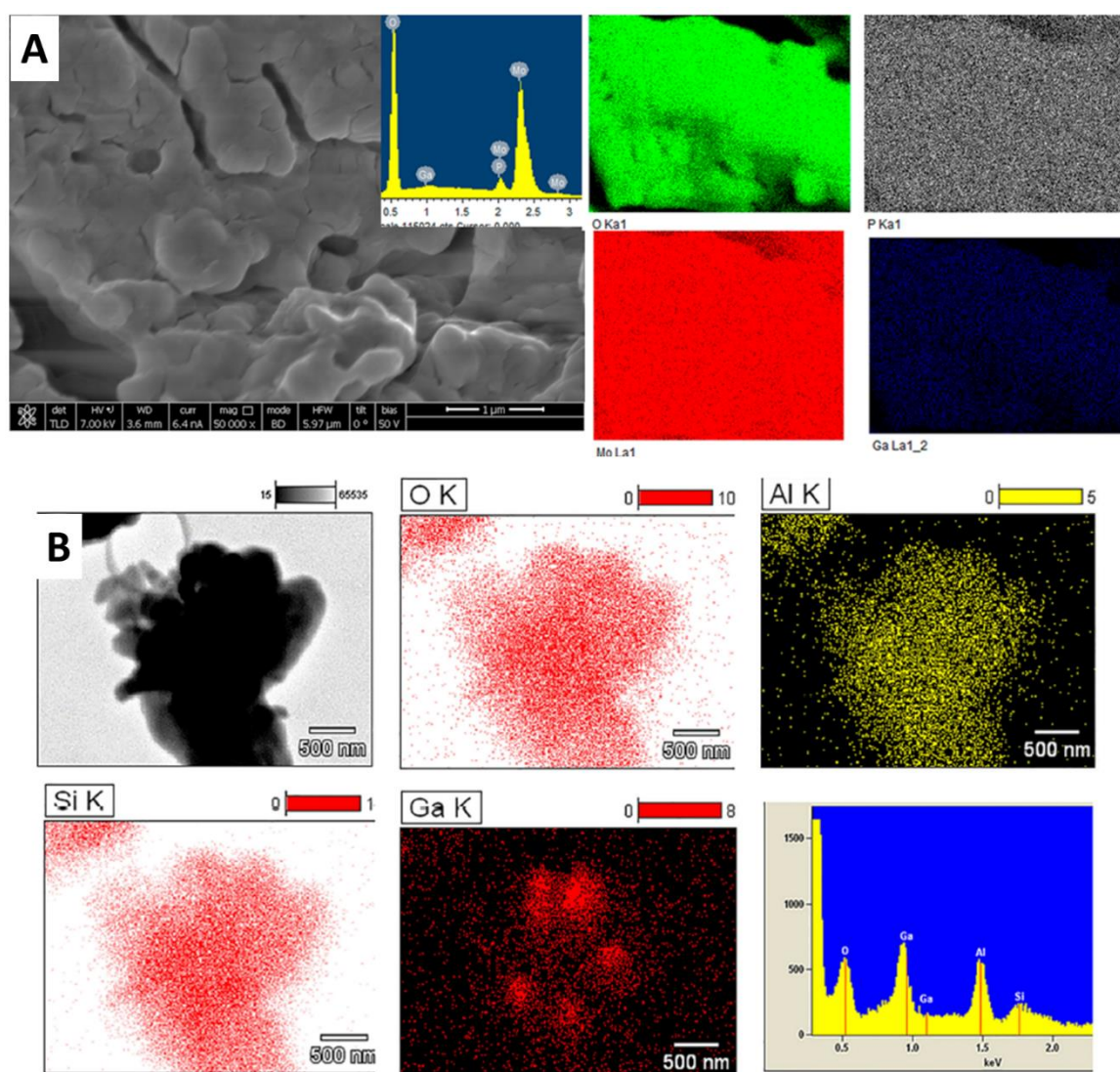


Figure 3. Interplay of Lewis acid and Bronsted acid sites for the selective conversion of glucose to levulinic acid; A. Ga salt of molybdophosphoric acid and B. [Adapted with permission from 29, 30].

Catalyst deactivation during long-term operation remains a primary concern. Mechanisms of deactivation include leaching of active species, pore blockage by humins and other carbonaceous deposits, and structural degradation under hydrothermal conditions. While short-term recyclability tests (typically 3–5 cycles) often show acceptable performance retention, extended operation over hundreds or thousands of cycles as required for industrial application has been rarely demonstrated. Development of robust catalysts with proven long-term stability is essential. Achieving high levulinic acid yields from real biomass feedstock continues to be challenging. While many studies report excellent yields from pure glucose or cellulose, performance typically degrades when processing actual lignocellulosic materials due to compositional complexity, structural recalcitrance, and presence of impurities. Bridging this gap requires either development of more effective catalysts or integration with practical pretreatment strategies. Process intensification and optimization present opportunities to improve economic viability. Most current research employs batch reactors with relatively long reaction times and dilute substrate concentrations. Continuous flow reactors could enable better temperature control, more efficient heat integration, and higher productivity. Microwave and ultrasound-assisted processing have shown promise for accelerating reactions and improving selectivity [31]. Integration with in-situ product separation or extraction could shift equilibria favorably and reduce degradation. Understanding reaction mechanisms at a molecular level would facilitate rational catalyst design. While the general

pathway from biomass to levulinic acid is established, detailed mechanistic insights including transition state structures, rate-determining steps, and origins of selectivity loss remain incompletely understood. Advanced characterization techniques and computational modeling can provide mechanistic clarity to guide development of improved catalysts. Sustainability and life cycle analysis should inform catalyst development priorities. An ideal catalyst should be synthesizable from abundant, low-cost materials using environmentally benign methods. The carbon footprint, energy requirements, and waste generation associated with catalyst production and use should be evaluated alongside performance metrics. Some highly effective catalysts may prove impractical if their environmental or economic costs outweigh benefits. Valorization of byproducts and co-products represents an important aspect of process economics. Formic acid is co-produced stoichiometrically with levulinic acid and could provide additional revenue if recovered efficiently. Humins, while problematic as catalyst foulants, might be converted to valuable products such as carbon materials or fuel additives. Lignin fractions could potentially be separated and utilized for the production of aromatic chemicals or materials. Integration of catalytic biomass conversion with biorefinery concepts offers a pathway to economic feasibility. Rather than focusing solely on maximizing levulinic acid yield, a biorefinery approach would co-produce multiple value-added chemicals from different biomass fractions, improving overall economics and sustainability. Solid acid catalysts capable of selective conversion of specific biomass components while leaving others intact for alternative processing could enable such integrated schemes. Techno-economic analysis and process modeling are needed to identify key performance targets and guide research priorities. Such analyses can reveal which catalyst properties (activity, selectivity, stability, cost) most strongly influence process economics and should therefore receive primary research attention. Optimal operating conditions and reactor configurations that may differ from those used in laboratory studies may be identified.

CONCLUSIONS

The conversion of biomass to levulinic acid using solid acid catalysts represents a promising approach for sustainable production of this valuable platform chemical. Significant advances have been made in developing diverse classes of heterogeneous acid catalysts including sulfonated carbons, modified zeolites, heteropolyacids, ion-exchange resins, and metal oxides, each offering distinct advantages and facing specific challenges. Sulfonated carbon-based catalysts have emerged as particularly effective due to their strong Brønsted acidity, favorable surface properties, and reasonable stability under reaction conditions. However, challenges related to acid site leaching and catalyst deactivation from coking still require solutions. Zeolite-based catalysts benefit from well-defined structures and tunable acidity but must overcome mass transfer limitations through hierarchical pore engineering. Heteropoly acids exhibit exceptional activity but suffer from water solubility issues that complicate catalyst recovery. Metal oxide-based catalysts offer excellent stability but often show moderate activity compared to stronger solid acids. Understanding the complex reaction network from biomass to levulinic acid, including competing pathways and deactivation mechanisms, has enabled more rational catalyst design. The importance of balancing acid strength, optimizing pore architecture for substrate accessibility, and ensuring stability under harsh hydrothermal conditions has become increasingly clear. However, translating fundamental insights into practical catalysts that match or exceed the performance of traditional mineral acids remains challenging. Process optimization through careful control of reaction temperature, time, catalyst loading, and solvent systems can significantly improve levulinic acid yields. The use of biphasic solvent systems and in-situ product removal strategies shows promise for minimizing degradation reactions. Nevertheless, most studies still rely on model substrates like pure glucose or cellulose, and performance with real biomass feedstock typically falls short of theoretical potential due to compositional complexity and structural recalcitrance. Looking forward, several research directions appear particularly important: developing catalysts with demonstrated long-term stability across hundreds of cycles; achieving high yields from real lignocellulosic biomass without extensive pretreatment; integrating catalytic conversion with effective pretreatment and product separation; understanding mechanisms at a molecular level to guide rational design; and performing techno-economic analyses to identify performance targets most critical for

commercial viability. The vision of economically viable, environmentally sustainable production of levulinic acid from biomass using solid acid catalysts remains compelling. While significant technical hurdles persist, the progress achieved thus far provides reason for optimism that continued research will ultimately enable commercial realization of this important biorefinery pathway.

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