Volume 23 Number 1 7 January 2021 Pages 1-614

Green Chemistry

Cutting-edge research for a greener sustainable future

rsc.li/greenchem



ISSN 1463-9262



CRITICAL REVIEW Javier Pérez-Ramírez, Ning Yan *et al.* Biomass valorisation over polyoxometalate-based catalysts



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CRITICAL REVIEW

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Cite this: Green Chem., 2021, 23, 18

Biomass valorisation over polyoxometalate-based catalysts

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The efficient utilization of biomass, the only globally available, renewable and abundant carbon-neutral source, is of high significance in green and sustainable chemistry. Polyoxometalates (POMs) and POMbased composites have been widely applied in green catalytic reactions, due to their tunable Brønsted/ Lewis-acidity and redox properties enabling high reactivity in a wide range of chemical transformations. This review covers recent advances in the chemocatalytic conversion of biomass into chemicals and fuels over POMs and POM-metal composites. For biomass valorisation over POMs, the advances of acid catalysis including hydrolysis, dehydration, etherification, alcoholysis, transesterification and esterification are summarised. Furthermore, applications in chemical oxidation for the synthesis of organic acids and furan chemicals are discussed. For biomass valorisation over metal-POM composites, an overview of tandem reactions (e.g. hydrolysis-hydrogenation, hydrolysis-oxidation, and hydrogenolysis-hydrodeoxygenation) is highlighted. The future prospects of biomass valorisation over POM-based catalysts are finally presented.

Received 21st September 2020, Accepted 2nd November 2020 DOI: 10.1039/d0gc03190a

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Introduction 1

Since the industrial revolution, the ever-increasing consumption of fossil carbon resources including coal, crude oil and natural gas for energy and chemicals has resulted in enormous emissions of greenhouse gases (GHGs) into the atmosphere. The increasing demand for resources, the diminishing finite fossil reserves as well as the urgent need to reduce GHG emissions have driven the innovative development of alternative



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energy sources derived from green and sustainable feedstocks in a post-petroleum era.

Biomass, including carbohydrates, lignin, and lipids, among others, is an attractive alternative feedstock. It is the only globally available, renewable and abundant carbonneutral source. Due to the relevant debate regarding "food *versus* fuel", lignocellulosic biomass composed of non-edible cellulose (35–50%), hemicelluloses (20–35%) and lignin (10–25%) has attracted significant attention. A biorefinery that provides low-value/high-volume liquid fuels and low-volume/ high-value chemicals, in particular from lignocellulosic biomass, has been the subject of intense research efforts.^{1,2}

Polyoxometalates (POMs) are a unique class of anionic polynuclear metal-oxo clusters with structural diversity at the atomic level, with their negative charge being balanced by countercations. In general, POMs are mainly classified into isopolyoxometalates $[M_x O_y]^{n-}$, which feature only metal and oxygen atoms [M = typically high-valent (d⁰ or d¹ electronicconfiguration) group V and VI transition metals (e.g., Mo, W, V)], and heteropolyoxometalates $[X_x M_y O_z]^{n-}$, containing additionally a heteroelement X (e.g., Si and P).^{3,4} Due to the tunable Brønsted/Lewis-acidity and redox properties, POMs and POM-based composites have long been applied in heterogeneous catalysis.^{3–5} In the field of catalytic biomass valorisation, POMs have been employed as environmentally benign acid or oxidation catalysts^{3,4} in the synthesis of renewable chemicals and fuels including platform chemicals,6-8 organic acids, furans,9-11 and biodiesels.12-14 Similarly, metal-POM composites have also been used for biomass conversion into chemicals and fuels.6,15,16

Many excellent reviews on biosourced chemicals and fuels have been published, devoted to specific feedstock such as carbohydrates,^{1,9,17} cellulose,^{8,18,19} lignin,²⁰ C₅–C₆ sugars,^{21,22} and glycerol,^{23,24} focusing on specific reactions such as hydrolysis,^{7,8,25–27} dehydration,^{21,23} esterification, transesterification,^{12–14} oxidation,^{9,10} hydrogenolysis,^{24,28} and hydrodeoxygenation,^{29,30} or dealing with specific catalytic



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systems such as solid acids,^{8,14,25,31} metal catalysts,¹⁵ functionalised heterogeneous catalysts (e.g., carbon-based catalysts, metal-organic frameworks (MOFs), solid phase ionic liquid based catalysts, and magnetic iron oxide based catalysts).³² An updated and in-depth review on the theme of biomass valorisation over POM-based catalysts, to our knowledge, is not available. The aim of this review is to provide a detailed analysis integrating the state-of-the-art of the chemocatalytic conversion of biomass over POMs and metal-POM composites. While the focus is to show the wide applicability of POM-based catalysts in biorefinery, discussions on the relationship between the catalytic performance and the chemical/electronic/structural properties of the materials, as well as the reaction mechanisms, are also provided. For biomass valorisation over POMs, firstly the advances of acid catalysis including hydrolysis, dehydration, etherification, alcoholysis, transesterification and esterification are examined. Furthermore, applications in chemical oxidation for the synthesis of organic acids and furans are analysed. For biomass valorisation over metal-POM composites, an overview of tandem reactions (e.g. hydrolysis-hydrogenation, hydrolysis-oxidation, and hydrogenolysis-hydrodeoxygenation) is provided.

2 Properties and preparation of POM-based catalysts

Currently, most studies are focused on the Keggin and Wells-Dawson type of POM. The Keggin anion family is based on the $[XM_{12}O_{40}]^{n-}$ anion (X = Si, P, etc.; M = Mo, W, etc.), while the Wells-Dawson anion family bears the chemical formula of $[X_2M_{18}O_{62}]^{n-4}$ POMs are versatile catalysts due to the multiple active sites, including protons as Brønsted acids, substituted metals with strong Lewis acidity, or metals with redox properties, and thus they are usually applied in acid-catalysed and oxidation reactions.3 POMs with protons as the only countercausually tions are called heteropolyacids (HPAs). Phosphotungstic acid (PTA, H₃PW₁₂O₄₀, HPW), silicotungstic acid (STA, H₄SiW₁₂O₄₀, HSiW), phosphomolybdic acid (PMA, H₃PMo₁₂O₄₀, HPMo) and silicomolybdic acid (SMA, H₄SiMo₁₂O₄₀, HSiMo) with strong Brønsted acidity are widely applied in acid catalysis. As for oxidation catalysis, the reported POMs $[X_x M_y O_z]^{n-}$ mainly contain P as the heteroelement X, and a combination of Mo and/or V as metal M.^{4,9,10}

POMs are normally soluble in both water and polar organic solvents; thus inorganic cations (*e.g.* Cs^+ , Ag^+ , *etc.*) are usually employed as counter-cations to enhance the insolubility of POMs. In addition to inorganic cations, organic compounds with designable and flexible functional groups (*e.g.*, organic amines, quaternary ammonium ions, organic surfactants, and ionic liquid cations) have also been adopted for the solidification of POMs.^{3,33} Meanwhile, various heterogeneous porous supports including transition metal oxides, clays, carbons, zeolites, mesoporous silicas, polymers, and MOFs, as well as magnetic nanoparticles (NPs) have been adopted for the immobilization of POMs.^{3,33} A variety of methods such as impreg-

Critical Review

nation, co-precipitation, deposition precipitation, equilibrium adsorption and ion exchange have been developed.^{3,5,12,33} Due to the increased thermal/hydrolytic/oxidative stability, these heterogeneous solidified/immobilised POMs have been widely applied as solid catalysts.^{3–5}

Metal–POM composites, containing both the metal sites (Ru, Au, Pd, Pt, *etc.*) and POM sites, can be prepared by impregnation, encapsulation, physical mixing and other methods.^{3,15,16} These hybrid materials have also been applied in a variety of one-pot conversions of biomass feedstock in a tandem reaction sequence including the hydrolysis–hydrogenation, hydrolysis–oxidation, and hydrogenolysis–hydrodeoxy-genation combinations, owing to the facilitated migration of the substrates/intermediates between the metal sites and POM sites. Since the preparation and characterization of POM-based catalysts have been covered in other excellent reviews, ^{3,5,12,33} in this review, the emphasis is on the functionality of POM-based catalysts in biomass transformations.

3 Biomass valorisation into renewable chemicals and fuels over POMs

3.1 Acid catalysis

3.1.1 Hydrolysis, dehydration and related reactions

3.1.1.1 Monosaccharide and derivative synthesis. As the most abundant and valuable component of lignocellulosic biomass, cellulose is a linear biopolymer consisting of glucose units connected through β -1,4-glycosidic bonds. The hydrolysis of cellulose or raw lignocellulosic biomass by acid catalysts leads to the generation of oligosaccharides or monosaccharides such as glucose.^{7,8,26} Further dehydration results in the formation of hydroxymethylfurfural (HMF).^{21,22,34–37} HMF has been widely accepted as an important platform chemical included in the "Top 10 + 4" list revised on the "Top 10" list described by the US Department of Energy (DOE).³⁸ The "Top 10 + 4" list also includes other critical platform chemicals such as furfural (FAL), glycerol, levulinic acid (LA), lactic acid, sorbitol, which will be discussed in this review.

So far, multiple POM-based catalysts have been reported for glucose synthesis.^{7,8,26} The representative POMs for fructose production are summarised in Table 1. Mizuno *et al.* found that highly negatively charged HPAs (*e.g.*, $H_5BW_{12}O_{40}$, $H_5AlW_{12}O_{40}$, and $H_5GaW_{12}O_{40}$), in particular $H_5BW_{12}O_{40}$, displayed higher activity than other typical HPAs (*e.g.*, HPW and HSiW) for the saccharification of polysaccharides, due to the high acidity and hydrogen-bonding accepting ability to decrease the crystallinity of polysaccharides (Table 1, entry 1).³⁹ Wang *et al.* reported that a micellar HPA catalyst [$C_{16}H_{33}N(CH_3)_3$] $H_2PW_{12}O_{40}$ displayed a remarkable performance in the hydrolysis of polysaccharides (*e.g.*, cellulose and starch), which is attributed to the facilitated access to catalytic sites by polysaccharide molecules, and the enhanced reaction rate and improved mass transport in the micellar catalyst

Table 1

Entry

Ref (%) Yield (Sel.^b (%) $Conv.^{a}$ (%) Product Glucose Time (h) ſemp. (°C) CH_3OH : toluene (1:2) $MIBK:H_2O\;(10:1)$ MIBK : $H_2O(10:1)$ $MIBK : H_2O(10:1)$ $MIBK : H_2O (3:1)$ sec-Butanol EMIM]CI Solvent DMSO OSMC H₂0 H₂0 chcl H_2^- O H_2O H_2O H,0 H₂0 H₂0 catalysts $Cr[Cr_{3}O(OOCCH_{2}CN)_{6}(H_{2}O)_{3}]_{3}[PW_{12}O_{40}]_{2} \cdot 69H_{2}O_{40}]_{3}$ Hydrolysis of polysaccharides/monosaccharides over POM $SS[Cr_3O(OOCC_2H_5)(H_2O)_3]_2[SW_{12}O_{40}]$ $Cr[(OSO_3C_{12}H_{25})H_2PW_{12}O_{40}]$ $T[(OSO_3C_{12}H_{25})H_2PW_{12}O_{40}]$ $(C_6H_{15}O_2N_2)_{3-x}H_xPW_{12}O_{40}$ $ChH_4PW_{11}TiO_{40}$ $Tr[(OSO_3C_{12}H_{25})H_2PW_{12}O_4$ HOCH2CH2N(CH3)3)H2PW $C_{16}H_{33}(CH_3)_3|H_2PW_{12}O_4|$ $C_{16}H_{33}(CH_3)_3|H_2PW_{12}O_4|$ H₃PW₁₂O₄₀/MCM-41 H₃PW₁₂O₄₀/MIL-101 MIM-PS]₃PW₁₂O₄₀ FePW₁₂O₄₀ Ds_{2.5}H_{0.5}PW₁₂O₄₀ ${
m H_5BW_{12}O_{40}} \\ {
m Cs_1H_2PW_{12}O_{40}} \\$ HSiW/graphene ChH₄AlW₁₂O₄₀ Ag4SiW12O40 g4SiW12O40 Catalyst Glycogen Fructose Fructose Sucrose Cellulose Corn stover Husk Cellulose Cellulose Cellulose Substrate Fructose Fructose Cellulose Cellulose Cellulose Fructose Fructose Fructose Glucose Starch

Green Chemistry

(Table 1, entries 3 and 4).⁴⁰ Gedanken *et al.* prepared HSiW/ graphene *via* a sonochemical method for glucose synthesis from glycogen hydrolysis. The hydrophobic cavities on the graphene surface facilitated the anchoring of glycogen and promoted the attack of glycosidic bonds *via* protons, which resulted in selective and efficient hydrolysis (Table 1, entry 5).⁴¹

A variety of POMs have been evaluated for HMF generation.^{21,22,34-37} In addition to monophasic systems such as water, organic solvents and ionic liquids, innovative biphasic aqueous/organic systems have also been employed, which efficiently remove the produced HMF from the reactive aqueous phase to the organic phase, thus minimising side reactions.^{21,34,37} The representative POMs for the hydrolysis of polysaccharides/monosaccharides in monophasic and biphasic systems are summarised in Table 1. Wang and co-workers reported a micellar Brønsted-Lewis-surfactant-combined HPA catalyst Cr[(DS)H2PW12O40]3 (DS represents dodecyl sulfate, $OSO_3C_{12}H_{25}$) showing notable activity and stability for HMF synthesis from unpretreated lignocellulosic biomass (Table 1, entries 10-12). The enhanced performance is mainly attributed to the Brønsted and Lewis acidities, as well as the micellar structure with hydrophobic groups.⁴² The same authors prea series of $(HOCH_2CH_2N(CH_3)_3)_xH_{3-x}PW_{12}O_{40}$ pared $(Ch_xH_{3-x}PW_{12}O_{40}, x = 1, 2 \text{ and } 3)$ via choline chloride and HPW, and the highest HMF yield of 75% was achieved over ChH₂PW₁₂O₄₀ after 8 h at 140 °C (Table 1, entry 15).⁴³ The hydrophilic head of $Ch_xH_{3-x}PW_{12}O_{40}$ concentrates the cellulose substrate, while the hydrophobic tail inhibits further hydration of HMF to by-product LA, thus benefitting HMF production. POMs encapsulated in MOFs (e.g. MIL-10144) or immobilised on porous supports (e.g. MCM-41⁴⁵) have also been adopted for HMF generation. For instance, Hensen et al. reported PTA/MIL-101-PTA encapsulated in MIL-101 as a composite catalyst to convert sugars. The system achieved the same HMF yield of 63% from fructose in [EMIM]Cl at 80 °C after 1 h (Table 1, entry 17), and in DMSO at 130 °C after 30 min.44

Furthermore, various POMs have been evaluated for the production of LA or levulinate ester from carbohydrates in consecutive reactions.⁴⁶⁻⁶¹ Novel temperature-responsive HPA nanohybrids $[(CH_3)_3NCH_2CH_2OH]_nH_{5-n}AlW_{12}O_{40}$ (Ch_nH_{5-n}AlW₁₂O₄₀)⁴⁶ and ChH₄PW₁₁TiO₄₀,⁵² HPA-based ionic liquid catalysts including [PyPS]₃PW₁₂O₄₀ [PyPS = 1-(3-sulfopropyl)pyridinium]⁴⁷ and $[C_4H_6N_2(CH_2)_3SO_3H]_{3-n}H_nPW_{12}O_{40}$,⁵⁵ HPAs with Brønsted and Lewis acidic sites $H_n PW_{11}MO_{39}$ (M = Cu^{II}, Zn^{II}, Cr^{III}, Fe^{II}I, Sn^{IV}, Ti^{IV}, and Zr^{IV}; for Ti and Zr),^{48,49,52}, combination of Zr-MCM-41 and HPW catalysts with tuneable Lewis and Brønsted acidity,⁵⁰ as well as acid-base bifunctional HPA nanospheres $(C_6H_{15}O_2N_2)_{3-x}H_xPW_{12}O_{40}^{51}$ were developed. Different reaction media including supercritical methanol, water-methanol mixtures,53 and water-methyl isobutyl ketone (MIBK)^{52,55} were also explored. For example, Wang and co-workers developed novel temperature-responsive $Ch_nH_{5-n}AlW_{12}O_{40}$ catalysts, which dissolve at high temperature and precipitate from the reaction medium at room tempera-

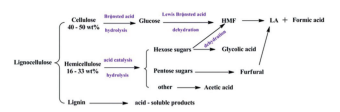


Fig. 1 Conversion pathways of cellulose into levulinic acid.⁴⁶ Reproduced from ref. 46 with permission from The Royal Society of Chemistry, copyright 2016.

ture. Among the tested catalysts, $ChH_4AlW_{12}O_{40}$ exhibited the highest LA yield of 74.8% with a cellulose conversion of 98.9% in one pot, due to the synergistic effects of the temperaturestimulus, and dual Lewis and Brønsted acidity (Table 1, entry 19).⁴⁶ As shown in Fig. 1, the Brønsted acid sites are proposed to facilitate hydrolysis, isomerization and dehydration/rehydration pathways, while Lewis acid sites are beneficial for isomerization of glucose to fructose.

3.1.1.2 Acrolein synthesis. The significant surge in biodiesel production via transesterification of triglycerides with methanol is accompanied by a substantial increase of glycerol generation as a by-product (ca. 10 wt%). Acrolein synthesis via the dehydration of glycerol is a promising method for glycerol valorisation, as acrolein is an important intermediate for valued-added chemicals such as acrylic acid and amino-acid DL-methionine.^{23,69–72} POMs supported on oxides (e.g. titania,⁷³ niobia,^{74,75} alumina,^{76,77} silica, alumosilicate,⁷⁸ and zirconium dioxide^{77,79}) and molecular sieves (MCM-41,⁸⁰ amino siloxane-functionalised MCM-41,80,81 Zr-MCM-41,82,83 SBA-15,⁸⁴ and ZSM-5⁸⁵) have been reported for acrolein synthesis from glycerol. For example, Yun et al. prepared the Keggin type H₃PW₁₂O₄₀/MCM-41 and Wells-Dawson type H₆P₂W₁₈O₆₂/MCM-41 for the dehydration of glycerol to acrolein. The selectivity to acrolein increased with the increasing ratio of Brønsted to Lewis acids, while the lower ratio of Brønsted/Lewis acids favoured the production of hydroxyacetone as undesired side products.80

3.1.2 Etherification alcoholysis. 5and Ethoxymethylfurfural (EMF) is a potential biofuel alternative due to its high energy density (30.3 MJ L^{-1}) similar to gasoline (31.9 MJ L^{-1}) and diesel (33.6 MJ L^{-1}), as well as lower emission of soot, NO_x and SO_x. Thus, EMF production from HMF or monosaccharides has attracted much attention.86 The representative POM catalysts for EMF synthesis via etherification, and dehydrative etherification in monophasic and biphasic systems are summarised in Table 2. Chen et al. prepared MOFbased POMs [Cu-BTC][HPM] (NENU-5) (BTC = benzene-1,3,5tricarboxylate, HPM = phosphomolybdic acid hydrate) for the conversion of HMF to EMF and ethyl levulinate (Fig. 2).87 [Cu-BTC][HPM] consisted of guest molecules of HPM in the pores of the insoluble Cu-BTC host and displayed remarkable catalytic activity and stability. Maximum EMF and ethyl levulinate (EL) yields of 68.4% and 20.2% were obtained at 140 °C after 12 h, respectively (Table 2).

 Table 2
 Etherification and dehydrative etherification for 5-ethoxymethylfurfural synthesis over POM catalysts

Substrate	Substrate Catalyst Solvent		Temp. ^{<i>a</i>} (°C)	Time (h)	$\operatorname{Conv.}^{b}(\%)$	Sel. ^c (%)	Yield (%)	Ref
HMF	H ₄ SiW ₁₂ O ₄₀ /MCM-41	Ethanol	90	2	92	84	81	88
HMF	[MIMBS] ₃ PW ₁₂ O ₄₀	Ethanol	70	24	98	93	91	89
HMF	Fe ₃ O ₄ @SiO ₂ -HPW	Ethanol	100	11	98	86	84	90
HMF	K-10 clay-HPW	Ethanol	100	10	98	93	92	91
HMF	[Cu-BTC][HPM] (NENU-5)	Ethanol	140	12	N/A	N/A	68	87
Fructose	$H_{3}PW_{12}O_{40}$	Ethanol : THF(5:3)	130	0.5	98	77	76	92
Fructose	[MIMBS] ₃ PW ₁₂ O ₄₀	Ethanol	90	24	N/A	N/A	91	89
Fructose	Fe ₃ O ₄ @SiO ₂ -HPW	Ethanol	100	24	N/A	N/A	55	90
Fructose	K-10 clay-HPW	Ethanol	100	24	100	62	62	91
Fructose	$H_{3}PW_{12}O_{40}$	Ethanol: DMSO(7:3)	140	2	99	64	64	93

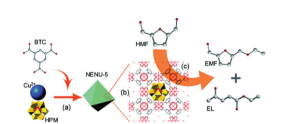


Fig. 2 Schematic illustration of the synthesis and catalysis procedure. (a) Synthesis of the catalyst. (b) The crystal structure of the catalyst. (c) The procedure of conversion of HMF into EMF and EL.⁸⁷ Reproduced from ref. 87 with permission from The Royal Society of Chemistry, copyright 2016.

3.1.3 Esterification and transesterification: biodiesel and additive synthesis

3.1.3.1 Biodiesel synthesis. Biodiesel is a non-toxic, biodegradable, environmentally friendly fuel alternative or additive of the petroleum-derived diesel. As a mixture of $C_{12}-C_{22}$ fatty acid monoalkyl esters (FAMEs), biodiesel is obtained by transesterification of the triacylglyceride (TAG) component and/or esterification of free fatty acids (FFAs) with C_1-C_2 alcohols in plant oils, animal fat or algae (Fig. 3).^{12–14,35,94} In contrast to "first-generation" biodiesel derived from edible feedstock (*e.g.*, soybean oil), increasing research interest is focused on the "second-generation" biodiesel from inedible feedstock (*e.g.* jatropha oil and horn oil) or waste cooking/frying oils.^{13,35} The fatty acid chain is mainly classified into saturated and

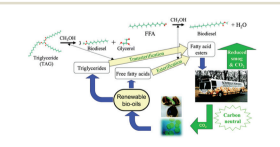


Fig. 3 Biodiesel production from renewable bio-oils *via* transesterification and esterification.¹³ Reproduced from ref. 13 with permission from The Royal Society of Chemistry, copyright 2014.

unsaturated chains with chain lengths of 16 and 18 carbons, including stearic, oleic, linolenic and palmitic acids.¹² For biodiesel production, parameters such as acid strength, structural porosity and surface hydrophobicity exert substantial impact on the catalytic performance.^{12,14} The representative inorganic cation/organo-solidified POMs and POMs supported on metal oxides, clays, zeolites, mesoporous silica, magnetic materials, and MOFs for biodiesel production are summarised in Table 3.

Guo and co-workers prepared novel organic-inorganic hybrid catalysts, HPA and ZrO₂ bifunctionalised organosilica hollow nanospheres (H₂PW₁₂O₄₀/ZrO₂-Et-HNS).⁹⁵ H₃PW₁₂O₄₀/ ZrO₂-Et-HNS: due to the strong Brønsted and Lewis acidity, unique hollow nanospherical morphology and hydrophobic surface, this material displayed encouraging activity and recyclability for transesterification of yellow horn oil with methanol to biodiesel, as well as esterification of levulinic acid (LA) with methanol to methyl levulinate (discussed in 3.1.3.2). The hollow nanospherical morphology enhanced the accessibility of guest molecules to the acid sites, shortened the diffusion pathway and enhanced the diffusion efficiency. The hydrophobic surface of the hybrid catalysts induced by the incorporation of bridging ethyl groups facilitated the enrichment of the hydrophobic reactant and expulsion of the hydrophilic products, thus accelerating the reaction (Fig. 4).

Li *et al.* prepared core-shell structured HPW-PGMA-MNPs composed of iron oxide magnetic NPs (MNP) as the core, poly (glycidyl methacrylate) (PGMA) as the shell, and HPW as the surface acid group. HPW-PGMA-MNPs exhibited remarkable activity and good recyclability for the one-pot transformation of waste grease to biodiesel *via* simultaneous esterification of FFAs and transesterification of TAG with methanol, as a result of the firm attachment of HPW on MNPs *via* covalent binding, the stable PGMA shell, and the superparamagnetic properties of the MNPs (Fig. 5).⁹⁶

Shul *et al.* prepared bifunctional core–shell HPA-functionalised zeolitic imidazolate framework-8 (ZIF-8) NPs *via* imidazolium medium. The strong O–N bonding indicated the strong interaction between the Keggin structure in HPA and imidazole units in ZIF-8 NPs. The optimised ZIF-8/HPA with both acidic and basic sites, and large surface areas displayed

Table 3 Biodiesel production over inorganic cation/organo-solidified POMs and POMs supported on metal oxides, clays, zeolites, mesoporous silica, magnetic materials, and MOFs

Support	Catalyst	Feedstock	Cat. (wt%)	Alcohol/ feedstock mole ratio	Temp. ^{<i>a</i>} (°C)	Time (h)	Conv. ^b / yield ^c (%)	Ref.
Inorganic cation	$Zn_{1,2}H_{0,6}PW_{12}O_{40}$	Waste cooking oil	2.5	28	65	12	97 ^b	98
	Cs _{2.5} H _{0.5} PW ₁₂ O ₄₀	Yellow horn oil ^e	1	12	60	0.2	96 ^c	99
	$Cs_{1.5}H_{1.5}PW_{12}O_{40}$	Jatropha oil	3	25	65	0.6	91 ^c	100
Organo-solidified	$(C_6H_{15}O_2N_2)_2HPW_{12}O_{40}$	Eruca sativa gars (ESG) oil	9	9	65	12	91^b	101
U U	(NH ₂ CH ₂ COOH)H ₂ PW ₁₂ O ₄₀	Palmitic acid	6	12	90	3	93 ^c	102
Ionic liquid	MIM-PSH H ₂ PW ₁₂ O ₄₀	Palmitic acid	7	13	80	5	92 ^c	103
Oxide	Ta_2O_5/SiO_2 -[H ₃ PW ₁₂ O ₄₀ /R] (R = Me or Ph)	Soybean oil (20% myristic acid)	2	90	65	24	90 ^c	104-106
	20%H ₃ PW ₁₂ O ₄₀ /ZrO ₂	Oleic acid	10	6	100	4	88^b	107
	25%H ₃ PW ₁₂ O ₄₀ /Nb ₂ O ₅	Used cooking oil	3	18	200	20	92 ^c	108
	15%TPA/SnO ₂	Palmitic acid	1	14	65	4	81^b	109
	25%TPA-Al ₂ O ₃	Jatropha oil ^d	4	19	65	0.8	84^c	110
	H ₃ PW ₁₂ O ₄₀ -ZrO ₂ -Si(Et)Si-60	Eruca sativa gars oil	5	90	65	24	75–99 ^c	111
	H ₃ PW ₁₂ O ₄₀ /ZrO ₂ -Et-HNS-2.0	Yellow horn oil	5	90	65	24	63–76 ^c	95
Clay	20%HPW/metakaolin flint	Oleic acid	5	30	130	2	97 ^c	112
-	20%H ₃ PW ₁₂ O ₄₀ /K10	Oleic acid	5	8	165	5	100^{b}	113
Carbon	H ₃ PW ₁₂ O ₄₀ /ACF	Palmitic acid	1.4	97	60	6	88^b	114
	20%TPA-AC	Jatropha oil ^d	4	20	65	0.66	87 ^c	115
Zeolite	$30\%H_{3}PW_{12}O_{40}/H\beta$	Oleic acid	0.1	20	60	6	84^b	116
	30% SiW ₁₂ /Hβ	Oleic acid	0.1	20	60	10	86^b	117
Mesoporous silica	23%H ₃ PW ₁₂ O ₄₀ /SBA-15	Oleic acid	0.3	40	40	4	90^{b}	118
	TSA ₃ /SBA-15	Waste cooking oil	0.3	8	65	8	86^b	119
	45%TPA/SBA-15	Canola oil	3	26	200	6	97 ^c	120
	TPA ₃ /MCM-41	Palmitic acid	0.1	40	60	6	100^{b}	121
	30%H ₃ PW ₁₂ O ₄₀ /MCM-41	Lauric acid		2	90	3	95 ^c	122
	30%SiW11/MCM-41	Oleic acid	0.1	40	65	16	81^b	123
Magnetic NPs	HPW-PGMA-MNPs	Grease	4	33	122	24	98 ^c	96
MOF	ZIF-8/HPA	Rapeseed oil	4	10	200	2	98^b	97
	AILs/HPW/UiO-66-2COOH	Soybean oil	10	35	110	6	96^b	124

^a Temp. = temperature. ^b Conv. = conversion. ^c Yield. ^d Assisted by ultrasonic energy. ^e Microwave.

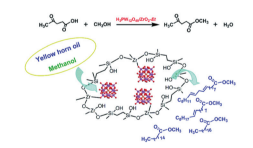


Fig. 4 The shell composition of $H_3PW_{12}O_{40}/ZrO_2$ -Et hollow nanospheres and the processes of levulinic acid esterification and yellow horn oil transesterification catalysed over $H_3PW_{12}O_{40}/ZrO_2$ -Et hollow nanosphere hybrid catalysts.⁹⁵ Reproduced from ref. 95 with permission from The Royal Society of Chemistry, copyright 2014.

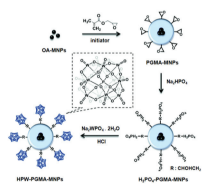


Fig. 5 Synthesis of phosphotungstic acid-functionalised iron oxide particles (HPW-PGMA-MNPs) as a magnetic nano-size solid acid catalyst.⁹⁶ Reproduced from ref. 96 with permission from The Royal Society of Chemistry, copyright 2014.

notable activity and recyclability for the transesterification of rapeseed oil with methanol to biodiesel (Fig. 6).⁹⁷

3.1.3.2 Synthesis of levulinate esters. Levulinate esters have potential applications in the fragrance and flavouring industry, as blending agents in biodiesel, *etc.* In particular, ethyl levulinate (EL) is a novel diesel miscible biofuel. Levulinate esters are generally obtained *via* esterification of levulinic acid (LA), one of the most important platform molecules in the "Top 10 + 4" list mentioned earlier. The typical POM-based catalysts for

esterification of levulinic acid to alkyl levulinate are summarised in Table 4.

Guo and co-workers prepared ZrO₂-based organic–inorganic hybrid catalysts functionalised by Keggin-type HPA and hydrophobic alkyl groups (*i.e.*, benzene-terminally bonded and ethane-/benzene-bridged organosilica moieties), $H_3PW_{12}O_{40}/$ ZrO₂-Si(Et/Ph)Si and $H_3PW_{12}O_{40}/$ ZrO₂-Si(Ph).¹²⁵ $H_3PW_{12}O_{40}/$

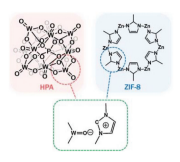


Fig. 6 Illustration of the plausible chemical bonding structure for the hybrid ZIF-8/HPA bifunctional catalyst for the transesterification of rapeseed oil.⁹⁷ Reproduced from ref. 97 with permission from Elsevier, copyright 2019.

ZrO₂-Si(Et/Ph)Si and H₃PW₁₂O₄₀/ZrO₂-Si(Ph) exhibited excellent catalytic activity and stability for the esterification of LA. The authors attributed the superior catalytic performance to the strong Brønsted and Lewis acidity, unique textural properties and the hydrophobic surface that inhibit the strong adsorption of the hydrophilic by-products. In particular, the ordered 2D hexagonal mesostructured H₃PW₁₂O₄₀/ZrO₂-Si(Ph) Si exhibited higher activity than 3D wormhole-like H₃PW₁₂O₄₀/ ZrO₂-Si(Et)Si and H₃PW₁₂O₄₀/ZrO₂-Si(Ph), due to the higher diffusion efficiency in the ordered mesoporous structure (Fig. 7). The same authors developed HPA and ZrO₂ bifunctionalised organosilica nanohybrids PW12/ZrO2-Si(Et)Si with a 1D hollow tubular nanostructure, a 2D hexagonal periodic mesostructure and a 3D interconnected mesostructure.¹²⁶ Among these, HPW and ZrO₂-bifunctionalised organosilica nanotubes PW₁₂/ZrO₂-Si(Et)Si-NTs exhibited the highest activity for alkyl levulinates by LA esterification (Fig. 8).

3.2 Oxidation

The carbohydrate feedstock has high oxygen content; thus oxidation into oxygen-containing chemicals or oxygen removal *via* dehydration, hydrogenolysis, or hydrodeoxygenation is a convenient pathway. The catalytic oxidation of carbohydrates and their derivatives provide a variety of value-added fine chemicals, including organic acids (*e.g.* formic acid, glycolic acid, gluconic acid, lactic acid, and acrylic acid),^{4,10,11} as well as furan chemicals (*e.g.* 2,5-diformylfuran).^{9,131}

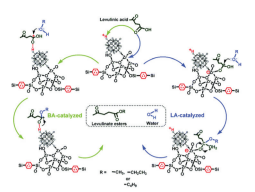


Fig. 7 The process of esterification of LA with alcohol to produce levulinate ester over $H_3PW_{12}O_{40}/ZrO_2$ -Si(Ph)Si hybrid materials. Left part: Brønsted acid site-catalysed reaction; right part: Lewis acid site-catalysed reaction.¹²⁵ Reproduced from ref. 125 with permission from The Royal Society of Chemistry, copyright 2013.

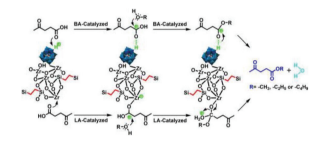


Fig. 8 Possible reaction mechanism of the synthesis of alkyl levulinates from the esterification of levulinic acid over PW_{12}/ZrO_2 -Si(Et)Si-NT nanohybrids.¹²⁶ Reproduced from ref. 126 with permission from Elsevier, copyright 2016.

3.2.1 Organic acid synthesis

3.2.1.1 Oxidation of carbohydrates to formic acid. Formic acid (FA) is a versatile chemical finding widespread applications. It is used as a H_2 donor in hydrogenation reactions, as well as a hydrogen storage compound in the context of hydrogen economy. Research studies have been focused on sustainable FA generation from oxidation of monosaccharides, oligo-saccharides, and bio-polymers such as cellulose, hemicellulose, lignin and even algae, under mild conditions with

 Table 4
 Esterification of levulinic acid to alkyl levulinate over POM-based catalysts

Catalysts	Reaction conditions ^{<i>a</i>}	Conv. ^b /yield (%)	Ref.	
HPW/desilicated-ZSM-5	0.25 wt%/6/78/4	94^b	127	
H ₄ SiW ₁₂ O ₄₀ -SiO ₂	0.1 g/50/65/6	73	128	
H ₃ PW ₁₂ O ₄₀ /ZrO ₂ -Si(Ph)Si	2 wt%/7/65/3	99.9	129	
$H_3PW_{12}O_{40}/ZrO_2-Si(Ph)Si-1.0$ 2 wt%/7/65/3		99.9	125	
$H_3PW_{12}O_{40}/ZrO_2$ -Si(Et)Si-1.0		68		
H ₃ PW ₁₂ O ₄₀ /ZrO ₂ -Si(Ph)-1.0		72		
$H_3PW_{12}O_{40}/ZrO_2$ -Et-HNS-2.0	2 wt%/7/65/1	92	95	
PW ₁₂ /ZrO ₂ -Si(Et)Si-NTs	2 wt%/7/65/1.5	99.9	126	
[Cu-BTC][HPM]	N/A/N/A/120/6	>99	130	

^{*a*} Reaction conditions: Amount of catalyst wt%/molar ratio of alcohol to substrate/reaction temperature in °C/reaction time in h. ^{*b*} Conv. = conversion.

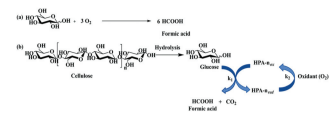


Fig. 9 (a) Overall stoichiometric reaction for oxidation of glucose into formic acid; (b) oxidative conversion of carbohydrates (*e.g.* cellulose) to formic acid over $H_{3+n}PV_nMo_{12-n}O_{40}$ (HPA) catalysts. HPA- n_{ox} : The oxidised state of HPA and HPA- n_{red} : the reduced state of HPA.⁹ Reproduced from ref. 9 with permission from The Royal Society of Chemistry, copyright 2018.

molecular oxygen or air as an oxidant.^{9–11,132,133} The oxidative conversion of cellulose includes the hydrolysis of highly polymerised cellulose into glucose over acid sites and the subsequent oxidative C–C bond cleavage of glucose into FA over redox sites, while overoxidation leads to the formation of CO_2 which is the thermodynamically favoured product. Keggin-type phosphovanadomolybdates, $H_{3+n}PV_nMo_{12-n}O_{40}$ (HPA-*n*), with n = 0-6, are typical catalysts for the oxidative conversion of biomass to FA (Fig. 9). The representative $H_{3+n}PV_nMo_{12-n}O_{40}$ (HPA-*n*) catalysts for the oxidation of carbohydrates to formic acid are summarised in Table 5.

Wasserscheid *et al.* firstly reported the oxidation of carbohydrate-based biomass to FA over $H_5PV_2Mo_{10}O_{40}$.¹³⁴ An FA yield of around 50% with nearly full conversion was obtained from monosaccharides (glucose and xylose) and disaccharides (cellobiose and sucrose) at 80 °C and 3 MPa O₂ after 26 h (Table 5, entry 1). However, water-insoluble cellulose only offered an FA yield of 1% under the same reaction conditions. Wasserscheid *et al.* further investigated the activity of $H_{3+n}PV_nMo_{12-n}O_{40}$ (n = 0-6) in selective oxidation of biomass to FA, and found that a higher degree of V-substitution enhanced the catalytic activity. The highest activity was obtained over HPA-5 (Table 5, entry 3),¹³⁵ probably due to the formation of pervanadyl (VO²⁺) cations that are known to be a strong oxidative species.

The addition of inorganic or organic acids can further increase the FA yield due to the facilitated hydrolysis of cellulose. Fu *et al.* reported an FA yield of 34% with full cellulose conversion over HPA-2 combined with HCl at 170 °C and 1 MPa O₂ after 9 h (Table 5, entry 14).¹³⁶ Wu *et al.* reported an FA yield of 61% with 100% cellulose conversion over HPA-2 and H₂SO₄ at 180 °C after 5 min (Table 5, entry 15).¹³⁷ Wasserscheid *et al.* obtained 22% FA from oxidative conversion of cellulose over HPA-2 and *p*-toluenesulfonic acid (TSA) at 90 °C and 3 MPa O₂ after 66 h.¹³⁸ The FA generation over a combination of HPA-2 and *p*-TSA can be expanded to a wide range of feedstocks including lignocellulose (*e.g.*, xylan, lignin,

Table 5 Oxidation of carbohydrates to formic acid over $H_{3+n}PV_nMo_{12-n}O_{40}$ (HPA-n)

Entry	Substrate	Feed conc. ^{<i>a</i>} (mg mL ⁻¹)	Catalyst	Temp. ^b (°C)	O ₂ (MPa)	Time (h)	Conv. ^c (%)	Yield (%)	Ref.
1	Glucose	30	$H_5PV_2Mo_{10}O_{40}$	80	3	26	>98	47	144
2	Glucose	25	$H_5PV_2Mo_{10}O_{40}$	100	5 (air)	3	100	52	136
3	Glucose	50	$H_8PV_5Mo_7O_{40}$	90	3	8	100	60	135
4	Glucose	18	$H_8PV_5Mo_7O_{40}$	90	2	48	100	85^a	141
5	Cellobiose	30	$H_5PV_2Mo_{10}O_{40}$	80	3	26	>98	47	134
6	Glucose	72	[MIMPS] ₃ HPMo ₁₁ VO ₄₀	180	1	1	100	55	140
7	Sucrose	30	$H_5PV_2Mo_{10}O_{40}$	80	3	26	>98	48	134
8	Sucrose	34	$H_8PV_5Mo_7O_{40}$	90	2	48	96	76^a	141
9	Xylan	30	$H_5PV_2Mo_{10}O_{40}$	80	3	26	N/A	33	134
10	Xylan	27	$H_5PV_2Mo_{10}O_{40} + p$ -TSA	90	3	24	97	53	138
11	Xylan	50	$H_8PV_5Mo_7O_{40} + p$ -TSA	90	3	24	100	58	135
12	Cellulose	30	$H_5PV_2Mo_{10}O_{40}$	80	3	26	N/A	1	134
13	Cellulose	10	$H_4PVMo_{11}O_{40}$	180	0.6	3	100	68	144
14	Cellulose	10	$H_5PV_2Mo_{10}O_{40} + HCl$	170	1	9	100	34	136
15	Cellulose	27	$H_5PV_2Mo_{10}O_{40} + H_2SO_4$	180	3	0.08	100	61	137
16	Cellulose	27	$H_5PV_2Mo_{10}O_{40} + p$ -TSA	90	3	66	N/A	22	139
17	Cellulose	27	$H_5PV_2Mo_{10}O_{40} + p$ -TSA	90	3	24	N/A	31	145
18	Cellulose	50	$H_8PV_5Mo_7O_{40} + p$ -TSA	90	3	24	76	28	135
19	Cellulose	67	[MIMPS] ₃ HPMo ₁₁ VO ₄₀	180	1	1	93	51	140
20	BM cellulose	10	Co _{0.6} H _{3.8} PMo ₁₀ V ₂ O ₄₀	160	$2(O_2 + N_2)$	5	N/A	66	146
21	Lignin	30	$H_5PV_2Mo_{10}O_{40}$	80	3	26	N/A	14	134
22	Lignin	27	$H_5PV_2Mo_{10}O_{40} + p$ -TSA	90	3	24	N/A	31-32	138 and 145
23	Lignin	50	$H_8PV_5Mo_7O_{40} + p$ -TSA	90	3	24	100	32	135
24	Poplar sawdust	30	$H_5PV_2Mo_{10}O_{40}$	80	3	26	N/A	19	134
25	Pomace	33	$H_5PV_2Mo_{10}O_{40} + p$ -TSA	90	3	24	N/A	55	139
26	Cane trash	33	$H_5PV_2Mo_{10}O_{40} + p-TSA$	90	3	24	N/A	49	139
27	Beech wood	27	$H_5PV_2Mo_{10}O_{40} + p$ -TSA	90	3	24	N/A	35	138
28	Beech wood	16	$H_8PV_5Mo_7O_{40} + p$ -TSA	90	2	48	N/A	61^d	141 and 145
29	Bagasse	10	$H_4PVMo_{11}O_{40}$	180	0.6	3	100	61	144
30	Hay	10	$H_4PVMo_{11}O_{40}$	180	2	3	100	55	144

^{*a*} Feed conc. = feed concentration. ^{*b*} Temp. = temperature. ^{*c*} Conv. = conversation. ^{*d*} Water : 1-hexanol (1 : 1).

Critical Review

beech wood, pomace, and cane trash), as well as the thirdgeneration biomass (*e.g.*, algae) (Table 5, entries 10, 16, 22, and 25-27).^{138,139} Similarly, high FA yields from cellulose, xylan, and lignin were obtained over HPA-5 with p-TSA as an additive (Table 5, entries 11, 18 and 23).¹³⁵

The H⁺ cation in HPA-*n* can be exchanged by organic compounds and transition metal cations. Liu *et al.* prepared a series of heteropolyanion-based ILs with $PMo_{11}VO_{40}^{4-}$ anions and $-SO_3H$ functionalised cations including $-SO_3H$ functionalised methylimidazole (MIMPS), $-SO_3H$ functionalised pyridinium (PyPS) and TEAPS.¹⁴⁰ The $-SO_3H$ functionalised cations are responsible for cellulose hydrolysis to glucose, while the $PMo_{11}VO_{40}^{4-}$ anions catalyze the glucose oxidation to FA. [MIMPS]_3HPMo_{11}VO_{40} provided the highest FA yield of 51% from cellulose at 180 °C and 1.0 MPa O₂ (Table 5, entry 19).

Novel process technologies such as *in situ* extraction of FA are adopted to further increase FA yields.¹⁴¹ Albert *et al.* reported a water–organic biphasic system to effectively boost the FA selectivity compared with monophasic aqueous media.¹⁴¹ FA yields of up to 85% and 61% were obtained from glucose and beech wood over HPA-5 in the biphasic system (Table 5 entries 4, 8 and 28). The *in situ* extraction of FA with long-chain primary alcohols such as 1-hexanol and 1-heptanol prevented the decrease of pH of the aqueous phase, and resulted in a higher FA selectivity and yield.

In addition to Keggin-type POMs, Lindqvist-type isopolyoxometalates $[V_nW_{6-n}O_{19}]^{x^-}$ have also been applied in the biomass oxidation to FA. Albert *et al.* found that $K_5V_3W_3O_{19}$ selectively oxidised hemicellulose and lignin to FA, while being inactive for cellulose conversion.^{142,143}

3.2.1.2 Oxidation of carbohydrates to glycolic acid. Glycolic acid (2-hydroxy acetic acid) is an important C2 α -hydroxy acid used as a precursor for the biodegradable polymer polyglycolic acid.¹⁴⁷ The reaction pathway of carbohydrate conversion to glycolic acid is proposed in Fig. 10. Glucose from the hydrolysis of cellulose undergoes multiple retro-aldol reactions to glycolaldehyde, which is converted to glycolic acid through oxidation. Similarly, fructose from glucose isomerization also undergoes successive retro-aldol reactions and oxidation to provide glycolic acid and FA.¹⁴⁸ Han *et al.* reported heteromolybdic acid H₃PMo₁₂O₄₀ exhibiting high activity for one-pot conversion of α -cellulose into glycolic acid. A glycolic acid yield

Fig. 10 The proposed reaction pathway for the conversion of cellulose to glycolic acid.¹⁴⁸ Reproduced from ref. 148 with permission from American Chemical Society, copyright 2012.

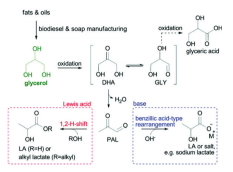


Fig. 11 Catalytic conversion of glycerol to lactic acid or alkyl lactates in water or alcohol, respectively.¹⁸ Reproduced from ref. 18 with permission from The Royal Society of Chemistry, copyright 2017.

of 49% together with 10% FA were obtained at 180 $^{\rm o}{\rm C}$ and 0.6 MPa O_2 after 1 h. 148

3.2.1.3 Oxidation of glycerol to lactic acid. Lactic acid (2-hydroxypropionic acid), as one of the US DOE's top 10 carbohydrate-derived chemicals, has widespread applications in the food, pharmaceutical and chemical industries such as building blocks of biodegradable plastic polylactic acid.¹⁸ The reaction pathway of glycerol conversion to lactic acid is shown in Fig. 11. Glycerol undergoes dehydration/oxidation to form dihydroxyacetone (DHA) in equilibrium with glyceraldehyde (GCY). The pyruvaldehyde (PAL) obtained from the dehydration of DHA further undergoes a benzylic acid-type rearrangement to provide lactic acid.¹⁴⁹ The oxidation of glycerol into lactic acid in water over representative POM-based catalysts is summarised in Table 6.

Wang and co-workers reported $H_3PMo_{12}O_{40}$ (HPMo) with optimal Brønsted acidic and redox properties exhibiting a lactic acid selectivity of 90% at 88% glycerol conversion. HPMo loaded on carbonised willow catkins (HPMo/C) achieved an even higher lactic acid selectivity of 94% at 98% glycerol conversion (Table 6).¹⁴⁹ The preparation of HPMo/C catalysts is shown in Fig. 12, including (1) surface oxidation of the carbon support; (2) covalent binding of ethanediamine to the surface carboxylic groups; and (3) functionalization with 1-bromodecane and immobilization of HPMo into a lipid-like layer. The inclusion of HPMo in a lipid-like layer prevents HPMo leaching; thus the HPMo/C exhibits high catalyst stability.

In addition, novel HPMo@lipid(n)/GO catalysts were developed with HPMo embedded in lipid-like bilayers that are covalently bonded to graphene oxide (GO). The coupling of the surface carboxylic groups on functionalised graphene oxide (GO) with diamine provided a covalently linked GO-NH₂ monolayer. Then the functionalisation of the GO-NH₂ monolayer with 1-bromodecane, followed by embedding of HPMo into the lipid-like bilayer, provided HPMo@lipid(n)/GO hybrid materials (Fig. 13).¹⁵⁰ The optimal HPMo@lipid(4)/GO catalyst featured balanced hydrophobic and hydrophilic properties, and the capillary-like microreactor configuration formed by the lipid-like bilayer structure. It achieved a high lactic acid yield of 90% with a glycerol conversion of 97% at 60 °C and 1

 Table 6
 Oxidation of glycerol to lactic acid in water over POM-based catalysts

Catalysts	Reaction conditions ^{<i>a</i>}	Yield (%)	Ref.
30%H ₃ PMo ₁₂ O ₄₀ /C	10/60/5/5	92	149
AlPMo ₁₂ O ₄₀	10/60/10/5	85	151
Ag ₃ PMo ₁₂ O ₄₀	10/60/10/5	93	152
H ₃ PMo ₁₂ O ₄₀ @lipid/GO	N/A/60/10/3.5	90	150

 a Reaction conditions: Catalyst amount (mg $\rm mL^{-1})/reaction$ temperature °C/oxygen pressure bar/reaction time h.

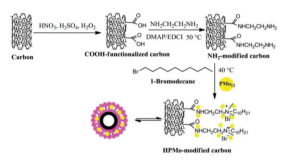


Fig. 12 Synthetic progress of HPMo-modified carbon catalysts.¹⁴⁹ Reproduced from ref. 149 with permission from WILEY, copyright 2015.

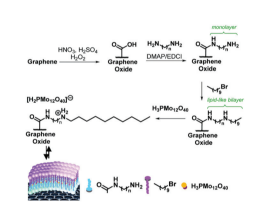


Fig. 13 Synthetic strategy for the assembly of HPMo@lipid(*n*)/GO hybrid catalyst materials, where *n* represents the length of the diamine carbon chain (n = 2, 4, 6, 8, and 10).¹⁵⁰ Reproduced from ref. 150 with permission from The Royal Society of Chemistry, copyright 2017.

MPa O_2 after 3.5 h (Table 6). The capillary reactor of HPMo@lipid(*n*)/GO enhanced the adsorption of glycerol and oxygen molecules, and concentrated them around HPMo, thus improving glycerol conversion. The oxidative ability of HPMo enhanced by GO and optimised by the length of the alkyl chain, combined with the balanced hydrophobicity of the lipid bilayer and the hydrophilicity of HPMo, led to high selectivity to lactic acid.

3.2.1.4 Dehydration-oxidation of glycerol to acrylic acid. Acrolein and acrylic acid (AA) are critical chemicals derived from glycerol. AA is generally obtained by a two-step tandem reaction including glycerol dehydration to acrolein over a solid acid and subsequent oxidation of acrolein over Mo- and

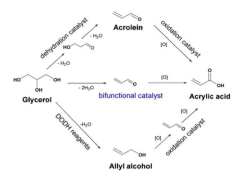


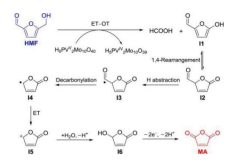
Fig. 14 Two-step and one-step conversion of glycerol to acrylic acid.¹⁵³ Reproduced from ref. 153 with permission from American Chemical Society, copyright 2016.

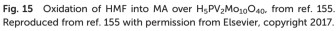
V-based catalysts (Fig. 14). AA synthesis is also realised *via* a two-step tandem reaction *via* allyl alcohol as an intermediate. In this context, the direct oxidative dehydration of glycerol to acrylic acid over bifunctional catalysts is highly desired. Zhang *et al.* reported that the incorporation of vanadium species into CsPMo and CsPW promoted the formation of acrylic acid from glycerol conversion, due to the synergism between the Keggin anions and the cesium/vanadium species at the secondary structure. The combination of Cs(VO)_{0.2}PMo and Cs(VO)_{0.2}PW as solid solution catalysts changed the oxidation ability and surface acidity, and improved selectivity for acrylic acid. Consequently, $H_{0.1}Cs_{2.5}(VO)_{0.2}(PMo_{12}O_{40})_{0.25}(PW_{12}O_{40})_{0.75}$ displayed the highest acrylic acid yield up to 60% and satisfactory resistance to coke deposition.¹⁵³

3.2.2 Synthesis of furan derivatives. Maleic anhydride (MA) is an important precursor for polyester resins, additives, pharmaceuticals, and so on. Yin *et al.* reported a vanadium-substituted HPA catalyst $(H_5PV_2Mo_{10}O_{40})$ for the aerobic oxidation of HMF to maleic anhydride (MA).¹⁵⁴ An MA yield of 41.8% and total yields of MA and maleic acid of 50.9% were achieved under optimised conditions (acetonitrile, 90 °C, 10 bar O₂, 8 h). A novel mechanism initialised by the C–C bond cleavage between the hydroxymethyl group and furan sketch of HMF *via* an ET-OT process, without the formation of 2,5-diformyl-furan (DFF), 5-formyl-2-furancarboxylic acid (FDCA), and 5-hydroxymethyl-2-furancarboxylic acid (HMFCA), was proposed for MA formation (Fig. 15).

4 Biomass valorisation into renewable chemicals and fuels over metal–POM composites

Metal NP–POM composites are referred to either metal NPs supported on heterogenised POMs or both metal NPs and POMs immobilised on an additional support. Such an intimate contact between the metal and POMs has the advantage of facilitated migration of the substrate/intermediates between the metal and acid sites at a nanometer distance. This section





describes how metal–POMs can be used in catalytic biorefinery to achieve a one-pot tandem catalytic process. The use of physical mixtures of metal catalysts and POMs as catalysts is also included to provide readers a more comprehensive overview.

4.1 Hydrolysis, hydrogenation and related reactions

4.1.1 Hydrolysis/hydrogenation of polysaccharides. The hydrolysis of polysaccharides (e.g., cellulose and inulin) and disaccharides (e.g., cellobiose) generates monosaccharides. These monosaccharides undergo hydrogenation to yield hexitols (e.g., sorbitol) as important products in the food, cosmetic and pharmaceutical industries.⁶ Since the pioneering works from Kou,¹⁵⁶ Liu,¹⁵⁷ and Fukuoka,¹⁵⁸ which proved the feasibility of one-pot reaction combining polysaccharide/disaccharide hydrolysis and monosaccharide hydrogenation, many catalytic systems have been developed. In the context of using metal-POM composites as catalysts, several factors including the substrate concentration,159 hydrophobicity,160 acidity of POMs,160-163 and size of metal NPs161 seem to have great impact on the catalytic activity and product selectivity. The hydrogenation activity is sufficiently higher compared with the hydrolysis rate; thus hydrolysis is generally believed to be the rate-determining step.159,161

Sels and co-workers reported the direct combination of $H_4SiW_{12}O_{40}/H_3PW_{12}O_{40}^{159}$ or water-tolerant $Cs_{3.5}H_{0.5}SiW_{12}O_{40}$ (CsSiW)/Cs_{2.5} $H_{0.5}PW_{12}O_{40}$ (CsPW)¹⁶⁰ with Ru/C to convert cellulose to hexitols. Palkovits *et al.* adopted physical mixtures of $H_4SiW_{12}O_{40}/H_3PW_{12}O_{40}$ and Ru/C for the transformation of cellulose and biomass feedstock spruce to sugar alcohols. High yields of C_4 - C_6 sugar alcohols up to 81% and 65% were obtained at 160 °C starting from cellulose and spruce, respectively.¹⁶³

Wang and co-workers reported Ru/Cs₃PW₁₂O₄₀ catalysts for the transformation of cellobiose and cellulose into sorbitol.¹⁶² Ru/Cs₃PW₁₂O₄₀ catalysts exhibited full cellobiose conversion, and a high sorbitol yield of 86% at 140 °C. It is proposed that the reversible Brønsted acid sites were generated on the Cs₃PW₁₂O₄₀ surface by spillover H species from H₂ dissociation on Ru NPs.¹⁶² However, the surface area and porosity of Cs_xH_{3-x}PW₁₂O₄₀ are relatively low, leading to inefficient diffusion of the reactant and product.

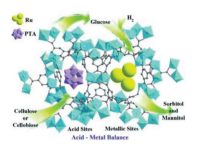


Fig. 16 Metal-acid bifunctional catalyst Ru-PTA/MIL-100(Cr) for the conversion of cellulose and cellobiose into sorbitol.¹⁶⁴ Reproduced from ref. 164 with permission from WILEY, copyright 2013.

Chen *et al.* encapsulated PTA $H_3PW_{12}O_{40}$ inside the mesoporous cavities of MIL-100(Cr), and developed a Ru-PTA/MIL-100(Cr) catalyst to convert cellulose and cellobiose into sorbitol (Fig. 16).¹⁶⁴ The ratio of the acid site density to the number of Ru surface atoms (n_A/n_{Ru}) in the catalytic performance was analysed in detail. The optimum n_A/n_{Ru} values between 8.84 and 12.90 led to the maximum conversion of cellulose and cellobiose into sorbitol. The Ru-PTA/MIL-100(Cr) catalysts with optimal acid/metal ratios exhibited high sorbitol yields of 97.1% and 57.9% at full cellobiose and cellulose conversion, respectively. However, the Ru-PTA/MIL-100(Cr) catalysts suffered from poor reusability, which might have resulted from the adsorption of insoluble substances such as oligomeric products on catalysts.

Isosorbide, a precursor of various biopolymers, pharmaceuticals and other chemicals, is obtained from the combined hydrolytic hydrogenation of cellulose to sorbitol and further dehydration of sorbitol.¹⁸ Sels and co-workers reported such an approach using a physical mixture of $H_4SiW_{12}O_{40}$ and Ru/C. 52% isosorbide was obtained from purified microcrystalline cellulose, while 63% isosorbide was generated from crude wheat straw pulps obtained by organosolv fractionation.¹⁶⁵

The hydrolysis of polysaccharides and subsequent hydrogenolysis with C-C and C-O cleavage can provide high value diols such as ethylene glycol (EG) and propylene glycol (PG).¹⁸ For instance, García-Bosch et al. reported a Ru-STA/AC catalyst to achieve a 1,2-PDO yield of 40% and selectivity up to 50% from fructose at 140 °C.166 Zhang and co-workers reported the combination of RANEY® nickel and W species for EG synthesis. The EG yield increased in the following order: $H_4SiW_{12}O_{40} < H_3PW_{12}O_{40} < WO_3 < H_2WO_4$ (Fig. 17).¹⁶⁷ The W species catalysed the cleavage of the C-C bonds in cellulose by a retro-aldol reaction pathway to generate glycolaldehyde, which undergoes further hydrogenation to form EG.167-170 Similarly, Guerrero-Ruiz et al. adopted Ru NPs supported on an HPA-carbon material [activated carbon (AC), high surface area graphite (HSAG)] for the hydrolytic hydrogenolysis of cellulose to alkanediols.¹⁷¹ Among the various POMs tested, Ru/ AC-HPA and Ru/HSAG-HPA catalysts based on tungsten-based HPA (PTA and STA) exhibited better performance in terms of EG selectivity.

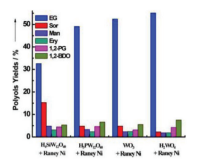


Fig. 17 Catalytic performance of different binary catalysts composed of tungsten-based compounds with RANEY® Ni. The reactions were performed with cellulose (0.5 g), water (50 mL), RANEY® Ni (40–50 mg), and a W compound (50 mg) at 245 °C and 6 MPa H₂ for 0.5 h.¹⁶⁷ Reproduced from ref. 167 with permission from WILEY, copyright 2013.

4.1.2 Hydrolysis/hydrogenation of 5-hydroxymethylfurfural/furfural. HMF is obtained from acid-catalysed dehydration of carbohydrates. A variety of fuels and commodity chemicals such as FDCA,¹⁸ DFF, 2,5-dimethylfuran (2,5-DMF) and alkyl levulinate are further produced from HMF.³⁴ For instance, selective hydrogenation-hydrogenolysis of HMF provided 2,5-DMF (Fig. 18), which is a potential substitute for ethanol and other fossil energy sources. Yadav et al. reported a Pd-Cs_{2.5}H_{0.5}PW₁₂O₄₀/K-10 clay that exhibited 98% conversion of HMF with 81% selectivity to 2,5-DMF under mild reaction conditions (90 °C, 1 MPa H₂).¹⁷² H₂ was dissociatively adsorbed on Pd sites, and the aldehyde group of HMF was hydrogenated into the hydroxyl group, thus generating bis(hydroxymethyl) furan (BHMF) as an intermediate. Further hydrogenolysis of the hydroxyl groups of BHMF to 2,5-DMF was facilitated by the acid sites of the Cs_{2.5}H_{0.5}PW₁₂O₄₀/K-10 clay.

While HMF is a key platform chemical from cellulose, furfural is an important platform molecule derived from hemicellulose for the synthesis of value-added products including furfuryl alcohol (FAL) and alkyl levulinate (AL). Furfural hydrogenation into intermediate FAL and subsequent acid-catalysed alcoholysis of FAL provide AL as an important bio-based fuel additive and a precursor for γ -valerolactone (Fig. 19). The Au-HSiW/ZrO₂ catalyst achieved a high AL yield of 80.2% with full furfural conversion at 100 °C.¹⁷³ The Au surface sites catalysed the transfer hydrogenation reaction of furfural to FAL using 2-propanol as a H-donor, while the acidic HSiW sites catalysed

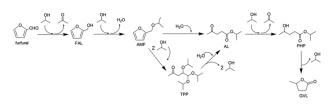


Fig. 19 Conversion of furfural to alkyl levulinate with 2-propanol as the H-donor.¹⁷³ Reproduced from ref. 173 with permission from The Royal Society of Chemistry, copyright 2016.

the alcoholysis of FAL to yield AL. Among HSiW/HPW/HPMo impregnated, ZrO₂ supported Au catalysts, a linear relationship between the acidity of Au-HPA/ZrO₂ catalysts and AL yield was established.

4.1.3 Hydrogenolysis/dehydration of glycerol to diol/acrolein. The biodiesel production *via* transesterification of plant oils and animal fats with short chain alcohols generates a large amount of glycerol as a by-product. Thus, the transformation of bioglycerol into value-added chemicals via hydrogenolysis, dehydration, and other reactions is of significance. In particular, the selective hydrogenolysis of glycerol to valueadded C3 chemicals including 1,2-propanediol (1,2-PDO),174 1,3-propanediol (1,3-PDO),^{175,176} and propanols¹⁷⁷⁻¹⁷⁹ is promising to valorize the surplus glycerol.^{23,24,180} The dehydration of glycerol to 3-hydroxypropaldehyde (3-HPA) on Brønsted acid sites followed by subsequent hydrogenation on the metal center lead to the formation of 1,3-PDO.^{175,176} The dehydration of glycerol into acetol over either metal sites or Lewis acid sites and consecutive hydrogenation over metal sites (e.g. Ru and Cu) result in the formation of 1,2-PDO.²⁴ 1-Propanol is usually formed via 1,2-PDO as an intermediate (Fig. 20).¹⁷⁷ 1,3-PDO has the highest economic value among the three, and functions as a monomer to make polypropylene terephthalate as polyester fibers.

A series of alkaline metals, including Li, K, Rb and Cs, were used to modify the Pt-H₄SiW₁₂O₄₀/ZrO₂ catalyst. Li⁺ exhibited the best promotional effect in glycerol hydrogenolysis, providing 53.6% 1,3-PDO selectivity and 43.5% glycerol conversion at 180 °C, 5 MPa (Table 7).¹⁷⁵ A linear relationship between 1,3-PDO yield and concentration of Brønsted acid sites was observed, indicating that the hydrogenolysis of glycerol to 1,3-PDO proceeded *via* dehydration of glycerol to 3-HPA on

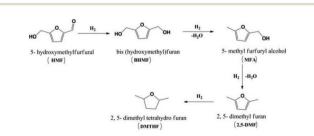


Fig. 18 Selective hydrogenation of 5-hydroxymethylfurfural to 2,5dimethyl furan.¹⁷² Reproduced from ref. 172 with permission from American Chemical Society, copyright 2016.

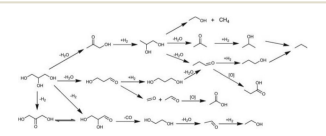


Table 7	Hydrogenolysis and dehydration of	of aqueous glycerol over metal NP-POM bifunctional catalysts	
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Catalysts	H ₂ (MPa)	Temp. ^{<i>a</i>} (°C)	Time (h)	$\begin{array}{c} WHSV \\ \left(h^{-1} \right) \end{array}$	$\begin{array}{c} \text{Conv.}^{b} \\ (\%) \end{array}$	Sel. ^{<i>c</i>} (1,2-PD)	Sel. ^c (1,3-PD)	Sel. ^c (1-PrOH)	Sel. ^c (acrolein)	Ref.
$5 \text{ wt}\% \text{Ru/C}(3 \text{ wt}\%) + 15 \text{ wt}\% \text{PTA/ZrO}_2 (6 \text{ wt}\%)$	6	180	8	_	44	64.3	_	_	_	174
1 wt%Pt-20 wt%Li ₂ H ₂ SiW/ZrO ₂	5	180	_	0.09	43.5	14.2	53.6	24.1	_	175
2 wt%Pt-15 wt%H ₄ SiW ₁₂ O ₄₀ /ZrO ₂	5	180	_	0.09	24.1	16.5	48.1	21.8	_	177
2 wt%Pt-15 wt%H ₄ SiW ₁₂ O ₄₀ /ZrO ₂	5	200	_	0.05	99.7	5.1	0.9	80.0	_	177
2 wt%Pt-15 wt%H ₃ PW ₁₂ O ₄₀ /ZrO ₂	0.1	250	_	1.02	98.6			97	_	178
0.5 wt%Pd/Cs _{2.5} H _{0.5} PW ₁₂ O ₄₀	0.1	275	5	2.80	79			_	96	182
1 wt%Pd-30 wt%H ₃ PW ₁₂ O ₄₀ /C	0.1	260	7	0.08	90			_	70	182
2 wt%Pd-30 wt%H ₃ PW ₁₂ O ₄₀ /Zr-MCM-41	0.1	320	5	0.35	90	—	—	—	83	83

^a Temp. = temperature. ^b Conv. = conversion. ^c Sel. = selectivity.

Brønsted acid sites followed by hydrogenation of the dehydration product 3-HPA on nearby metal sites. Furthermore, Pt-HSiW/ZrO₂ exhibited superior performance in glycerol hydrogenolysis to 1,3-propanediol among HSiW, HPW and HPMo modified Pt/ZrO₂ catalysts, due to the higher concentration of Brønsted acid sites.¹⁷⁵ In parallel, a close correlation between the 1,2-PDO yield and the concentration of Lewis acid sites was obtained. Thus, the Brønsted acid sites are proposed to facilitate the selective generation of 1,3-PDO, while the Lewis acid sites favor the generation of 1,2-PDO.

Zhu and co-workers reported that the Pt-H₄SiW₁₂O₄₀/ZrO₂ catalyst afforded a high 1-propanol yield of 80% and long-term stability of 160 h at 200 °C and 5 MPa H₂ (Table 7).¹⁷⁷ The Pt-H₄SiW₁₂O₄₀/ZrO₂ catalyst exhibited higher activity and propanol selectivity than its counterparts with Pd, Cu, and Ni as metal sites. ZrO₂ interacts strongly with Pt sites, and prevents the crystallisation of Pt NPs in calcination at high temperatures. The small Pt particle size with high exposure of the Pt active species on the surface resulted in superior activity of Pt-H₄SiW₁₂O₄₀/ZrO₂. The Pt-H₄SiW₁₂O₄₀/ZrO₂ catalysts also displayed high resistance to the impurities in crude glycerol.

Acrolein, an important precursor for the production of adhesives, polymers, and detergents, can be obtained *via* the dehydration of glycerol over POMs/HPAs. H₂ co-feeding and incorporation of platinum-group metals prevent coking and enhance the catalyst stability.^{73,181} Pd incorporation showed a larger stability enhancement effect than the incorporation of its Pt and Ru counterparts.^{83,182} For instance, Pd-H₃PW₁₂O₄₀/ Zr-MCM-41 exhibited high catalytic activity and stability, showing a slight decrease of glycerol conversion from 97% to 87% after 50 h.⁸³

4.2 Oxidation of carbohydrates and derivatives

The hydrolysis of cellulose and cellobiose and further oxidation lead to the generation of gluconic acid, an important food additive and intermediate for the synthesis of fine chemicals and pharmaceuticals.⁶ For metal NP–POM nanocomposites, the acidic sites facilitate the hydrolysis of cellulose and cellobiose to glucose, while the metal (usually Au) NPs facilitate the oxidation of glucose into gluconic acid (Fig. 21). Han *et al.* reported a Au/Cs₂HPW₁₂O₄₀ acidic/oxidative bifunctional catalyst, achieving a high gluconic acid yield of 96.4%

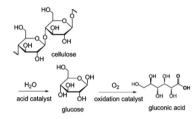


Fig. 21 Direct conversion of cellulose into gluconic acid by using hydrolysis and oxidation reactions.¹⁸⁴ Reproduced from ref. 184 with permission from WILEY, copyright 2012.

with nearly complete cellobiose conversion.¹⁸³ Cs₂HPW₁₂O₄₀ was inert to the deep oxidation of gluconic acid, and passivated the oxidative activity of Au NPs; thus Au/Cs₂HPW₁₂O₄₀ nanocomposites effectively prevented over-oxidation of gluconic acid. XPS analysis and DFT calculations demonstrated that the strong metal-support interfacial interaction effectively modulates the electronic structures of Au NPs, and subsequently the reactivity of the active oxygen species on the Au surface, thus resulting in the selective oxidation to gluconic acid. Several Au/Cs_xH_{3-x}PW₁₂O₄₀ catalysts were developed for the same reaction. 97% gluconic acid was attained over Au/ Cs_{1,2}H_{1,8}PW₁₂O₄₀, which bears stronger acidity and smaller Au NP size compared to other samples, at 145 °C for 3 h.¹⁸⁴ Indeed, both the acidity of POMs and mean-size of the Au NPs are crucial parameters in the conversion of cellobiose into gluconic acid. Cs_xH_{3-x}PW₁₂O₄₀ with stronger acidity not only facilitated the conversion of cellobiose, but also accelerated the desorption of gluconic acid and inhibited the consecutive degradation. The Au NPs with smaller size accelerated the oxidation of glucose, leading to high selectivity for gluconic acid.

4.3 Hydrodeoxygenation/deoxygenation of biomass-derived oxygenates

Due to the high oxygen content in biomass feedstock, hydrodeoxygenation (HDO) is an effective method to remove (part of) oxygen to convert biomass-derived oxygenates into drop-in bio-fuels and chemicals.^{20,29,30} The HDO process generally involves a combination of different reactions including hydrolysis, hydrogenation, hydrogenolysis, dehydration, decarboxylation and decarbonylation.²⁹ Based on the structures of substrates, the HDO process can be mainly classified into the HDO of ketones,^{185,186} ethers,¹⁸⁸ carboxylic acids,¹⁸⁹ esters,¹⁸⁸ angelica lactones,¹⁹⁰ furanic compounds,¹⁹¹ carbohydrates,¹⁹² and aromatic compounds.^{186,188}

For the HDO of aliphatic and aromatic ketones, the reaction pathway involves the hydrogenation of ketones to a secondary alcohol on metal sites (mostly Pt) followed by dehydration of the alcohol to alkene on acid sites, and finally hydrogenation of the alkene to alkane on metal sites.^{185–187} Kozhevnikov *et al.* investigated the gas-phase deoxygenation of aromatic ether anisole, aliphatic diisopropyl ether (DPE), and aliphatic ester ethyl propanoate (EP) over bifunctional catalysts comprising Pt, Ru, Ni, and Cu as the metal components and $Cs_{2.5}H_{0.5}PW_{12}O_{40}$ as the acid component (Fig. 22).¹⁸⁸ For the HDO of anisole, the model compound of lignin, consecutive hydrogenation of the aromatic ring, elimination of methanol

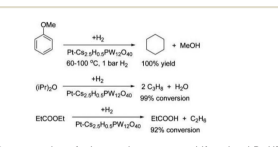


Fig. 22 Deoxygenation of ethers and esters over a bifunctional Pt-HPA catalyst in the gas phase.¹⁸⁸ Reproduced from ref. 188 with permission from American Chemical Society, copyright 2016.

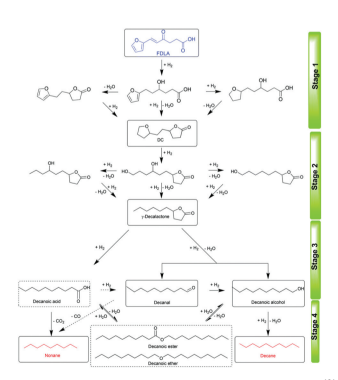


Fig. 23 Proposed reaction pathway for the HDO of FDLA.¹⁹¹ Reproduced from ref. 191 with permission from The Royal Society of Chemistry, copyright 2020.

from methoxylcyclohexane, and further hydrogenation to cyclohexane can be facilitated by bifunctional catalysts. 0.5% Pt/Cs_{2.5}H_{0.5}PW₁₂O₄₀ achieved a high cyclohexane selectivity of 89% with an anisole conversion of 87% under mild conditions (100 °C and 1 bar of H₂), and the physical mixture of Pt/C and Cs_{2.5}H_{0.5}PW₁₂O₄₀ (0.35% Pt content) provided 100% yield of cyclohexane. The Pt/C + Cs_{2.5}H_{0.5}PW₁₂O₄₀ physical mixture displayed higher activity and resistance to deactivation than Pt/ Cs_{2.5}H_{0.5}PW₁₂O₄₀. For the deoxygenation of DPE, the combined Pt/C + Cs_{2.5}H_{0.5}PW₁₂O₄₀ catalysts achieved an increased DPE conversion of 99% and a propane selectivity of 93%. Similarly, for the deoxygenation of EP, the Pt/Cs_{2.5}H_{0.5}PW₁₂O₄₀ catalyst under H₂ showed much better stability compared with Cs_{2.5}H_{0.5}PW₁₂O₄₀, due to the decreased catalyst coking in the presence of Pt and H₂.

Recently, interest in the one-pot HDO of furan derivatives and carbohydrates to alkanes has considerably increased. Zhang *et al.* reported the first one-pot HDO of high carbon furans (HCFs) and δ -furfurylidenelevulinic acid (FDLA) to long-chain alkanes over Pd/C combined with HPW.¹⁹¹ The total yield of alkanes as high as 93.2% including an 89.5% yield of decane was obtained under relatively mild conditions (3 MPa H₂, 170 °C, 4 h). The consecutive pathways include (a)

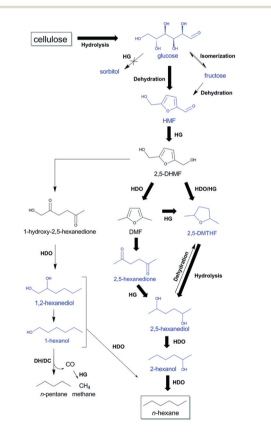


Fig. 24 Proposed reaction pathways from cellulose to *n*-hexane and *n*-pentane through HMF with TSA and htTSA-Ru/C. The most selective reaction pathway from cellulose to *n*-hexane is indicated with bold arrows. HDO, hydrodeoxygenation; HG, hydrogenation; and DH/DC, dehydrogenation/decarbonylation. Reproduced from ref. 192 with permission from The Royal Society of Chemistry, copyright 2015.

8

Critical Review

the hydrogenation of unsaturated alkenyls, keto-carbonyls and furans of FDLA over metal sites, (b) the formation of a 5-(2-(tetrahydrofuran-2-yl)ethyl)-dihydrofuran-2(3*H*)-one (DC) intermediate *via* the intramolecular dehydration reaction, (c) the hydrogenolysis and HDO of furans and lactones over metal-acid sites, and (d) the generation of alkanes *via* HDO of alde-hyde/alcohol *via* hydrogenation–dehydration–hydrogenation steps (Fig. 23).

One-pot conversion of cellulose to n-hexane has also been studied. Sels and co-workers reported the combination of tungstosilicic acid (TSA) and hydrothermally TSA-treated Ru/C (ht-STA-Ru/C) catalysts to convert microcrystalline cellulose into 52% n-hexane in a biphasic decane/water system.¹⁹² The direct conversion of cellulose to straight-chain alkanes mainly proceeded via HMF rather than sorbitol as an intermediate. This is in contrast to the widely accepted pathway, in which hydrolytic hydrogenation of cellulose to sorbitol and consecutive HDO to n-hexane is proposed. The kinetic study indicated that the hydrothermal treatment of Ru/C with TSA effectively enhanced the selectivity towards HMF hydrogenation and suppressed the glucose hydrogenation ability. The cellulose hydrolysis to glucose, dehydration of glucose to HMF, ring opening hydrolysis of 2,5-dimethylfuran (DMF) to 2,5-hexanedione, as well as ring-opening hydrolysis of 2,5-dimethyltetrahydrofuran (2,5-DMTHF) to 2,5-hexanediol were favoured over TSA in the aqueous phase, while the hydrogenation of the intermediates to the straight-chain alkanes was facilitated over ht-STA-Ru/C (Fig. 24).

5 Conclusions and outlook

Recent progress in the chemocatalytic conversion of biomass into chemicals and fuels over POMs and POM-metal composites are summarised. For biomass valorisation over POMs, the advances of acid catalysis and chemical oxidation are discussed. For biomass valorisation over metal-POM composites, an overview of the types of tandem reactions enabled by a metal-POM combination is presented. Despite these encouraging developments, there are still considerable challenges/ opportunities ahead for biomass valorisation using POMbased catalysts.

As shown in this review, a variety of methods have been adopted to transform POMs into insoluble solids or immobilize POMs on porous supports so that they can be used as recyclable heterogeneous catalysts. However, leaching still seems to be a common issue, thus compromising the catalyst stability. Future efforts should be directed to the detailed study of the POM–support interaction, and the kinetics and thermodynamics of POM adsorption/desorption on various support materials. Furthermore, new combinations of POMs anchored on emerging porous materials, such as nitrides, carbides, covalent organic frameworks (COFs), and porous organic frameworks (POFs), may be studied.

In addition, most of the reported POMs for biomass utilisation are focused on the Keggin and Wells-Dawson-type POMs, and there is still unexplored potential for application of POMs with other structures (*e.g.*, Lindqvist-type). Due to the tunable Brønsted/Lewis-acidity and redox properties, the POMs are generally applied in acid catalysis and chemical oxidation for biomass valorisation, while POM-metal composites are widely applied in the related tandem reactions. Beyond these, attention can be paid to explore other types of reactions in biomass valorisation over POM-based catalysts. For instance, it is envisaged that tungsten-containing POMs are active for retro-aldol condensation to break C–C linkages in carbohydrates. Metal–POM composites should also be effective in making organonitrogen chemicals, which appears to be a new direction in biomass conversion.^{193–195}

POMs have been proven as effective supports to stabilize a range of single atom catalysts (SACs),^{196,197} exhibiting excellent performance in CO oxidation,¹⁹⁸ selective hydrogenation,^{199,200} and Suzuki–Miyaura coupling reaction.²⁰¹ Compared to their NP counterparts, SAC-POMs have maximal atom utilization, together with strong and tuneable interactions between single metal atoms and POMs. We put forward SAC-POMs to be potentially efficient catalytic systems to enable both oxidative (*e.g.*, conversion of the –OH group into the –COOH group) and reductive (*e.g.*, C=O hydrogenation and hydrodeoxygenation) transformations of biomass to value-added fine chemicals under mild conditions.

Finally, a deep understanding of the relationship between the catalytic performance and the chemical/electronic/structural properties of the POM-based catalysts will be of significance for biomass utilisation. It is also of high interest to develop a new generation POM-based catalytic system for biomass valorisation in the future. We hope that this review has highlighted the usefulness of POMs in catalytic biorefinery, and will encourage more researchers to delve into this exciting field.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank the National University of Singapore Flagship Green Energy Program (R-279-000-553-646 and R-279-000-553-731) for financial support.

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