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Heterogeneous catalysts for advanced bio-fuel production through catalytic biomass pyrolysis vapor upgrading: a review

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Nowadays concerns regarding fossil fuel resources depletion as well as environmental issues attributed to $CO₂$ accumulation in the atmosphere force communities toward utilizing biomass as a substitute fuel source which is environmentally secure and renewable. Pyrolysis bio-oil from biomass comprises varieties of undesirable oxygenate and heavy compounds and has to be treated before feeding to biorefineries. Catalytic biomass pyrolysis vapor upgrading presently seems to be a techno-economical process toward production of fuel-like components. However, selection of stable and productive catalyst(s) to yield desirable chemicals with low coke formation is a great challenge. The three most important classes of catalysts comprising microporous zeolites, mesoporous catalysts and metal based catalysts are utilized for vapor phase bio-oil upgrading. This study offers a comprehensive review on catalytic biomass pyrolysis vapor upgrading by emphasizing particularly on catalyst types and properties, coke formation over catalysts and catalytic process conditions. **Published on 18 and 2019.**
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1. Introduction

Sustainable developments of societies in recent decades lead them to the high consumption of natural fossil fuel resources. Biomass is considered as the only available sustainable energy

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source of organic carbon which can appropriately substitute petroleum to yield carbon based materials, chemicals and fuels.¹

Pyrolysis process can be utilized to convert lignocellulosic biomass to liquid fuel.^{2,3} Fast pyrolysis process, which is distinguished by a high heating rate of particles at a short time,⁴ has recently attracted the broad attentions and can be considered as one of the most capable technologies which are

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exploited for the conversion of renewable biomass resources to bio-oil.⁵⁻⁸ The bio-oil derived from depolymerization of cellulose, hemicelluloses and lignin, three main building block of lignocellulosic biomass, is a complex mixture of different oxygenated compounds. A typical bio-oil with broad molecular weight range from 18 to 5000 g mol^{-1} or even more can contain more than 400 different compounds which most of them are oxygenated. Most of bio-oil deficiencies comprising its low heating value, corrosiveness and instability under long storage time and transportation conditions caused by these oxygenated compounds.⁹⁻¹³

Different approaches were employed targeting bio-oil's quality enhancement consisting: reduced pressure distillation,¹⁴ pyrolysis under reactive atmosphere,¹⁵⁻¹⁷ high pressure thermal treatment,¹⁸ hydro-treatment at high pressure,¹⁹ pyrolytic lignin removal,²⁰ pyrolysis vapor upgrading at low pressure,²¹ and conversion of bio-oil's acidic compounds to esters and ketons over acidic²² and basic²³ catalysts, respectively.

Bio-oil upgrading through conventional hydro-treating (HDT) at high pressure could accomplish oxygen removal by high hydrogen consumption, but it will fail to minimize carbon loss. Non-condensable undesirable C_2-C_3 gases instead liquid C_6-C_{14} hydrocarbons (appropriate for fuel applications) will be resulted from the small molecules during HDT process.²⁴

Bio-oils and the model compounds upgrading investigations showed a considerable decrease in product yield as a result of catalyst deactivation and severe tar and char formation during catalytic upgrading.^{25,26} Park et al.²⁷ carried out investigation on the catalytic upgrading of biomass pyrolysis vapor over HY and HZSM-5 zeolite catalysts in a fixed bed reactor. Their investigation outcomes, which were compared with the data from Vitolo et al.,²⁸ showed that employing biomass as feedstock instead of bio-oil increased upgraded bio-oil yield by 10 wt%.

The catalytic upgrading of the biomass fast pyrolysis vapor is considered as one of the most promising process to produce upgraded bio-oil. Deoxygenation of the produced bio-oil can be achieved in the presence of selected catalysts to enhance bio-oil properties. Investigations are being conducted towards the

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Fig. 1 Bio-oils (derived from lignocellulosic biomass) chemical composition.³⁰

design of selective catalysts to achieve production of high added value chemicals (e.g. phenol) or minimizing of the formation of undesirable bio-oil components such as acids and carbonyls.²⁹

Three main important oxygenated compounds families available in bio-oil can be characterized as: (1) aldehyde, ketones and acids (like acetone, acetic acid, acetol, etc.); (2) sugar derived compounds (like levoglucson and furfural); and (3) lignin-derived phenolics.²⁴ Different available components in the bio-oil are illustrated in Fig. 1.³⁰

Deoxygenation of these components suggests a great challenge. Accordingly, it is important to investigate the role of different catalysts play in the conversion of oxygenated compounds to fuel-like hydrocarbons. In this regard, development of highly durable and selective catalysts will be crucial and can be considered as key to the success for bio-oil upgrading processes at atmospheric condition and in the absence of hydrogen feeding.⁹

Two important targets in the biomass to bio-fuel conversion researches can be; increase the bio-fuel potential to replace petroleum and its cost competitiveness improvement. These two goals could be attained by minimizing hydrogen consumption and carbon loss. Furthermore, improvement of product quality can assist incorporation of upgraded bio-fuel in petroleum refineries or blending operations.

This review summarizes the recent researches and trends in the bio-oil catalytic vapor cracking/upgrading followed by deoxygenation focusing on catalysts properties and reaction conditions to selectively direct reactions toward production of fuel-like components and valuable chemicals.

2. Catalytic biomass pyrolysis vapor upgrading

The produced bio-oil from fast pyrolysis contains various oxygenated compounds that provide shortcomings to be used as transportation fuel. Although, it can be utilized directly for the purpose of heat and electricity generation. The high oxygen content of bio-oils has the negative effect on the energy density (16-19 MJ kg⁻¹ versus 46 MJ kg⁻¹ for conventional gasoline),

and it is caused poor stability as well as low volatility of the liquid bio-oil.^{1,31} Further, the bio-oils high viscosity and corrosiveness discourage their consumption in internal combustion engines.

One of the known solutions to stabilize the bio-oil and decrease its oxygen contents is to blend it with the hydrotreating process feed, even though bio-oil transportation and storage before its blending makes significant problems.^{31,32} Hydro-treatment, which is the bio-oil treatment at high hydrogen pressure (30–140 bar) and moderate temperature, is likely the most common route to the bio-oil compounds deoxygenation (HDO).³³⁻³⁵ In this method, bio-oil is completely deoxygenated and oxygen is removed in the form of water.

HDO is typically carried out in the presence of NiMo and CoMo catalysts.³³ It is worthwhile to mention that Pt and Ru metals exhibit higher hydrogenation activity, although they show lower tolerance of sulfur impurities.^{36,37}

The high hydrogen consumption in the bio-oil HDO process is the main drawback of this technology. Further, high pressure process which leads to high operational cost could be considered as the other disadvantage. One of the main challenges of HDO process is to hydrogenate the aliphatic compounds, whilst avoiding reduction of aromatics. This type of hydrogenation process control is difficult to achieve at high hydrogen pressure required for HDO.

Pyrolysis vapor upgrading can alternatively be carried out before vapor being condensate at atmospheric pressure and 350–500 °C, when vapors are passed through catalyst(s). The pyrolysis vapors need to pass certain stabilizing and oxygen removal processes without external hydrogen supply. At these conditions, vapors components undergo a series of reactions comprising, cracking, aromatization and dehydration. Through these reactions, oxygen is removed in the form of water, $CO₂$ and CO. Consequently, bio-oil is converted into a mixture of aromatic and aliphatic

hydrocarbons, although a large fraction of organic carbon reacts to create solid carbonaceous deposits over catalyst named coke.6,11,38–⁴⁰

Fig. 2 shows schematically this type of process. Following to upgrading process, liquid bio-oil intermediate can be fed to refinery, bio-refinery or petrochemical plants for further treatment toward fuels and/or chemicals production.

Zeolites are the most known catalysts used for the bio-oil pyrolysis vapor upgrading. Generally, coke formation over zeolite catalysts is one of the main problems for upgrading process. Investigation showed about 30% (maximum) of carbon in the feed can deposit as coke on the zeolite. It is due to low effective hydrogen available in the bio-oil. There are various oxygenated compounds in the bio-oil and therefore highly oxidized feed need to be converted to hydrocarbons. During this conversion, the excess carbon subsequently deposits as coke on the catalyst (s) .⁴¹

Chen's effective ratio (H/C_{eff}), which is defined as (H –2O)/C (H, O and C are hydrogen, oxygen and carbon moles, respectively), indicates the feedstock effective hydrogen. Generally, a low H/Ceff. ratio of feed is conducted to more coke formation than those having higher ratios.^{41,42}

Considering pyrolysis vapor upgrading approach, due to the similar operating pressure at which pyrolysis and upgrading are carried out, these two processes can be integrated. It is contrary to high pressure HDO process which cannot be easily integrated simultaneously with low pressure pyrolysis. Despite all the advantages associated with this type of upgrading process, there are some drawbacks. The yields of hydrocarbons are somehow modest. Another disadvantage of this technology is irreversible deactivation of catalysts, attributed to the partial dealumination of zeolite structures in the presence of water (usually found in bio-oils). Researches still undergo toward development of acidic catalysts with better resistance against water.⁴³ **ESC Areness**

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Fig. 2 Schematic of pyrolyis process and upgrading – highlighting pyrolysis vapor upgrading

By employing high reactive oxygenated compounds (carbonyl, hydroxyl, carboxyl, and ketonic groups) in bio-oils, reactions of C–C bonds formations such as aromatization, aldol condensation and ketonization can be carried out. It means oxygen functionalities potentials can be used to yield high carbon content deoxygenated fuel components instead of their elimination too early.²⁴ Through ketonization, two carboxylic acids are condensed into a larger ketone with the release of stoichiometric amounts of water and $CO₂$. Usually inorganic oxide like Al_2O_3 , TiO₂, ZrO₂ and CeO₂, operating at atmospheric pressure and moderate temperature (300–425 $^{\circ}$ C), are used as catalyst for these types of reactions.⁴⁴–⁴⁸

It is worthwhile to note that ketonization through carboxylic acids consumption can lead to oxygen removal in the form of carbon dioxide and water. Acids almost make about 30 wt% (maximum) of bio-oils and their conversion can improve biooils properties and mitigate their corrosiveness and chemical instability.¹ As a result, ketonization uses the oxygen functionality of acid groups to produce molecules with high heating value and consequently hydrogen consumption is reduced. Furthermore, through ketonization typical bio-oil components like esters can be condensed. Unlike zeolite catalysts, which their activity is sensitive to water presence, this type of upgrading process can be performed under moderate amounts of water.^{46,49,50}

Upon selective catalytic upgrading and deoxygenation of pyrolysis vapor, depending on the catalyst type, biomass composition and process conditions, different products with improved chemical and physical properties can be yielded. Nowadays, various researches are being conducted toward the design and selection of appropriate solid catalysts for production of high added value chemicals (e.g. phenolic compounds) or molecules with enhanced properties to be used as bio-fuel component. Recent catalytic pyrolysis of the lignocellulosic biomass for the phenolic compounds production has employed different catalysts, comprising alkaline catalysts, K_3PO_4 and activated carbon.⁵¹

Table 1 illustrates what can be anticipated for the characteristics and the compositions between raw pyrolysis oil, hydrodeoxygenated oil (HDO), zeolite cracking oil, and benchmarked crude oil.⁵²

Three categories of catalysts including microporous zeolites, mesoporous catalysts and metal based catalysts have recently attracted the considerations of researchers for the biomass pyrolysis vapor upgrading.

2.1. Microporous zeolite catalysts

Many zeolites have multi-dimensional microporous structure. This micro-porous system permits small molecules of reactants to diffuse in to the zeolite structure, therefore provide molecules access to internal acid sites. The microporous nature provides another essential feature to the zeolites, called shape-selectivity. The micro-pore channels size-restraints can somehow control the formation of unwanted products.⁴¹

The pores are frequently required to produce sufficiently high surface areas necessary for catalyst high activity. According to the IUPAC definition, porous materials are classified in three main groups; microporous (pore size \leq 2 nm), mesoporous (2–50 nm), and macroporous (>50 nm) materials.⁴³ Wide varieties of reactions could be catalyzed by zeolites attributed to their shape selectivity. The different zeolites pore size varying from 5 \AA to 12 Å affects molecules mass transfer.⁵³

Various types of shape selectivity can be identified depending upon if pore size restricts the reacting molecules entrance, or the product molecules departure, or the creation of certain transition conditions. Selectivity of the reactants achieved while among all the reactant molecules only whom which are small enough can diffuse through the catalyst pores. When parts of the products inside the pores are too large to diffuse out, product selectivity occurs. They are either transformed to smaller molecules or in the worst case block the pores and deactivate the catalyst. Restricted transition state selectivity takes place when particular reactions are avoided because the relevant transition state would need bigger space than available in the cavities. Different pore systems may employ to control the molecular traffic. Reactant molecules may favorably diffuse in the catalyst through one pore, while products leave through the other. So, counter diffusion is minimized.⁵⁴ Review Workers (may employing high reactive oxygenated compounds to the IPAC definition, poross materials are dassified in the corrections of c-C bomb formations and are computed in the corrections of the C bomb formations

One of the most important applications of zeolite catalysts is in fluid catalytic cracking (FCC) process. It provides about 45% of the global gasoline pool through the large hydrocarbon cracking into the gasoline range molecules.⁵⁵ Zeolites are appropriate catalysts for the biomass pyrolysis vapor/bio-oil upgrading due to containing Lewis and Brønsted acid sites. Reaction selectivity toward desired product can be controlled using acid site's strength and density distribution. Among all zeolites applications, catalytic conversion of oxygenates to hydrocarbons particularly has drawn the attentions. Most known is the methanol conversion to gasoline (MTG) over HZSM-5 catalyst.⁵⁶

Table 1 Comparison of characteristics of bio-oil, catalytically upgraded bio-oil, and benchmarked crude oil⁵²

Table 2 Summary of most recent researches of vapor phase bio-oil upgrading over zeolite catalysts

Table 2 (Contd.) Table 2 (Contd.)

naphthalenes

Biomass in the past decade has been considered as an important renewable resource of transportation fuels and its catalytic conversion over zeolites has been widely employed. Through several researches, a broad range of zeolites including HZSM-5, Y zeolite and beta zeolite have been investigated using bio-oils or biomass as feedstock. These studies indicated that the zeolite addition into the pyrolysis reactor could signicantly increase the formation of aromatics. $CO₂$, $CO₃$, water, tar and coke were also formed during this process.⁵⁷ The majority of these investigations resulted that HZSM-5 catalyst gave the highest aromatics yield.⁵⁸ In Huber et al ⁵⁹ patented investigations, glucose pyrolysis in the presence of HZSM-5 catalyst, maximum aromatics was yielded at catalyst Si/Al ratio of 60 and 600 °C. Agblevor^{60,61} patented fractional pyrolytic process, wherein the biomass materials were selectively converted into desired products in the presence of HZSM-5 catalyst, eliminating potential secondary and post-pyrolysis processing steps. He showed that the biomass lignin fraction could be converted to phenolic components with low char production when pryolysis and catalytic processes were carried out simultaneously. Due to the considerable demethoxylation and demethylation, the molecular mass distribution of the fractional catalytic process product was about half of the conventional pyrolysis without catalyst. **ESC Areness** in the past decade has been considered as an \sim Zeolite catalysis could be modified by incorporation decade has been considered on the modified or National Core National Core interaction and the considered

Zeolite catalysts could be modified by incorporation of metals. Incorporation of Co or Ni transition metals (1–10 wt%) into HZSM-5 catalyst indicated significant effect on the performance of the parent HZSM-5 catalyst. Compared to the $Co₃O₄$, NiO modified catalysts showed more reactivity towards increasing the gaseous products and decreasing the organic phase. All the metal-modified catalysts showed remarkable reactivity towards production of phenols and aromatics, although exhibited limited reactivity toward water production. These are attributed to different hydrocarbon conversion reactions, comprising dehydrogenation, cracking and aromatization/cyclization reactions, which are catalyzed by Brønsted acid sites of the zeolite. In addition, water production enhancement was due to increased decarboxylation/ dehydration of the oxygenated compounds on the zeolite acid sites.^{62,63} Catalytic conversion of particle board biomass over microporous zeolite catalyst exhibited that impregnation of 1 wt% Ga on HZSM-5 through incipient-wetness technique enhanced catalyst selectivity toward aromatic production. It is attributed to the dehydrocyclization of bio-oil intermediate products. Ga incorporation to the zeolite caused reduction of acid sites numbers. Although the selectivity towards aromatics was improved, but it caused lower degree of bio-oil deoxygenation (lower water yield).⁶⁴

Fig. 3 Reaction pathways for pyrolysis and catalytic pyrolysis vapor upgrading of lignocellulosic biomass over HZSM-5 catalyst. Adapted from Wang et al.⁷⁰

2.1.1. Summary of the fast pyrolysis vapor upgrading studies on microporous zeolites. Table 2 summarizes the most recent researches performed on fast pyrolysis vapor upgrading over zeolite catalysts. In this regard, some of the key aspects are as follows:

 HZSM-5 zeolite catalysts showed very good performance. It yielded bio-oil with low oxygen contents, less acidic, less viscous and stable with high energy content.^{21,58,65-67}

 HZSM-5 zeolite catalysts led to water increase in the bio-oil via dehydration reactions, and enhancement of organics, aromatic hydrocarbons and gaseous products caused by decarbonylation, dealkylation, decarboxylation, cracking, and aromatization reactions. Coke formation over catalysts was also increased during catalytic upgrading.^{21,58,66,67}

 Pore size and Si/Al ratio played important role on HZSM-5 catalyst performance, product distribution and selectivity. Metal substituted HZMS-5 enhanced bio-oil yield and properties.^{21,65-68}

• β-Zeolite, Y-zeolite and ferrierite zeolite showed very good performance in the bio-oil deoxygenation and aromatic compounds production. b-Zeolite showed high activity in the de-oxygenation reactions followed by Y and ferrierite zeolites. Ca–Y-zeolite deactivated less quickly and offered a superior ability to produce aromatics, compared to β -zeolite.^{58,60,69}

2.1.2. Reaction pathway for biomass pyrolysis vapor upgrading over HZSM-5 catalyst. In general, during pyrolysis and upgrading processes, lignocellulosic biomass pyrolysis vapor passes through a series of pyrolysis reactions followed by catalytic conversion of oxygenated compounds available in pyrolysis vapors.^{32,70} Recently Wang et al.⁷⁰ revealed deoxygenation pathway over HZSM-5 catalyst for cellulose, hemicellulose, and lignin (three most important building blocks of lignocellulosic biomasses). According to their investigation, the proposed reactions networks for the biomass catalytic pyrolysis vapor upgrading is shown in Fig. 3. It was assumed that there is negligible interaction effects among three biomass components during both thermal pyrolysis and catalytic conversion of pyrolysis vapors.

The biomass oxygenated organic compounds over zeolite catalysts at 350 \degree C to 500 \degree C passed through decarboxylation, cracking, alkylation, polymerization, condensation and aromatization reactions. When acidic zeolite catalysts like HZSM-5 was employed, dehydration was the dominant mechanism. Under this condition, the yielded products was a mixture of low molecular weight olefins and aromatic hydrocarbons.⁷¹

During biomass pyrolysis and catalytic upgrading, the major product from cellulose pyrolysis was levoglucsoan, which could produce smaller furanic compounds through decarbonylation, decarboxylation or dehydration reactions.⁷² These furans then could diffuse into the acidic zeolite pores to produce olefins and aromatics through oligomerization and decarbonylation reactions. On the other hand, double hydrated xylose, the predominant product from hemicellulose pyrolysis could diffuse together with other low molecular weight molecules like acetic acid, furaldehyde, acetol and formic acid into zeolite pores without any further reaction. Lignin pyrolysis initially

yielded monomeric phenolic components, which showed very low reactivity over HZSM-5 catalyst. Phenols acid-dehydration conducted to the formation of large amounts of cokes, whereas phenols cracking generated aromatics. Alkyl-phenols cracking to produce olefins might be another intermediate to yield aromatic compounds. Their investigation also showed that the aromatics yield of three main building block of lignocellulosic biomasses increased in the following order: lignin \ll hemicellulose < cellulose. Moderately higher temperature indicated lower coke generation and higher aromatics yield for three components of biomasses. It was attributed to the higher desorption of the coke precursors and generation of lower molecular weight oxygenated components during pyrolysis and upgrading. Lignin, among the three biomass components, had the most complicated structure and phenolic molecules produced from its thermal degradation were prone to the coke and char formations, which could decrease the carbon efficiency for the biomass pyrolysis and catalytic upgrading. Therefore, product distribution of the biomass pyrolysis and catalytic upgrading was highly depending on the biomass composition. Positive Warkers on Internactions, which showed by the final monomeric phenolic components, which showed stations and the monomerical monomerical monomeric schemes and the monomerical monomerical monomerical monomerical mo

2.2. Mesoporous catalysts

To eliminate the possibility of secondary reactions, which enhance the coke formation and consequently catalyst deactivation caused by a slow mass transport to and away from the catalytic center, suitable catalyst should have all advantages of microporous zeolite while provide additional diffusion pathways for larger molecules as shown in Fig. 4.⁷³

Since the pyrolysis vapor comprised various components with different sizes and molecular weight, porosity can play an important role toward production of desired products. Macroporous and mesoporous materials can be selected as the first choice for catalytic process in the presence of large molecules.⁷⁴ However, while size selectivity is desired, pores need to have a defined structure and be narrow enough to provide reagents; products and/or transition state selectivity. Macroporous materials, due to the high exposure of active site to substrates, restrict the reaction pathway toward selective reaction. To overcome this type of drawback, recently mesoporous materials with highly ordered structures have attracted the attentions.⁷⁵

diffusion of large molecules within microporous zeolites. These mesopores can be created as intercrystalline pores in nanozeolite aggregates (right) or may be formed as intracrystalline voids within zeolite single crystals (left). Adapted from Moller et al.⁷³

Biomass catalytic pyrolysis by the use of different acid catalysts has been employed to improve the bio-oil quality through deoxygenation reactions. Catalysts deactivation through coke formation is one of the main problems during deoxygenation. Coking is mostly caused by the phenolic compounds condensation. Further, the bio-oil components having large molecular volume cannot diffuse to the active sites placed inside the zeolite pores. Therefore, deoxygenation process of the bio-oil components is obstructed. To enhance the molecular transport and to prevent the pore blockage by coke generation, mesoporosity formation into the zeolite catalysts seems to be promising approach. Mesopores presence in the zeolite crystalline framework would be equivalent to external surface enhancement. It makes a large number of pore openings accessible to the large molecules. Shortened diffusion path length and enlarged external surface area would ease the coke precursors mass transfer from the micropores to the external surface of zeolite catalyst and consequently prevent its quick deactivation.⁷⁶ Therefore, catalytic performance of catalyst is enhanced.77,78 PSC Advances

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Isomerization and aromatization of 1-hexene over micro-/ mesoporous HZSM-5 catalyst (alkali-treated) indicated similar phenomena. The mechanism for catalytic stability improvement of alkali-treated HZSM-5 catalyst is illustrated in Fig. 5. As can be seen, due to the mesopores and micropores interconnection, the diffusion path in the micropores is considerably shortened. So, the isomerization and aromatization products or even precursors of coke (here naphthalene as the representative), which are created in the micropores, can diffuse out of the pores before deposition. It leads to the coke deposition in the mesopores of HZSM-5 catalyst and prevents micropores blockage. Consequently, the improvement of the stability of the catalyst in isomerization and aromatization can be attained attributed to the reduced diffusion path and coke formation in the mesopore structure.^{76,79}

2.2.1. Mesoporosity creation in the zeolites during synthesis. During the last decade various investigations

performed to synthesize zeolites with additional mesoporosity.76,80–⁸² Generally, different synthesize strategies can be used to generate mesoporosity in the zeolites structure as following:73,81 (a) dual templating method, in which the secondary template is used along with the common zeolites directing agent for mesostructuring the crystals of zeolites, (b) use a single but multifunctional templating route, having structure directing agents for the meso- and microscale in the same component, (c) adjustment of the synthesis reaction conditions, in which the secondary templates are unnecessary.

The dual templating method (a) uses the same basis that was already proven to be a very successful route in the synthesis of the microporous zeolites. In this method, a sacrificial scaffold is used to create mesoporosity during crystallization and can be eliminated from the zeolite framework without loss of its final structural characteristics. Based on the physiochemical nature of the secondary templates, they can be divided to hard and so templates. Soft templates also can be categorized to amphiphilic surfactants derivatives, macromolecular polymers and silylating agents. Applying multifunctional templates (b) produce micro- and mesopore structure in the zeolites at the same time through a single templating molecule. The third method (c) simplify the zeolite synthetic requirements and save additional cost of the production by directing the process toward synthesis of nanozeolite aggregates and mesoporous network.

2.2.2. Mesoporosity creation in the premade zeolite through leaching. Apart from the zeolites synthesis methods for mesoporosity creation (explained in Section 2.2.1), it is also possible to generate mesoporosity in the zeolites through a secondary reaction.⁸¹ This is generally performed after the micropore zeolite synthesis and calcination, when it is free from micropore templates. Different desilication or dealumination leaching methods may be employed to generate amorphous regions in the zeolite framework. Extraction of these amorphous debris can create mesoporosity in the zeolites. Generally, leaching is a destructive process, in which part of the micron-

Fig. 5 Mechanism for catalytic stability enhancement of the alkalitreated HZSM-5 zeolite with micro-mesopore porosity. Adapted from Li et al. 79

sized zeolite structure is sacrificed for the generation of larger external surfaces in the form of mesopores.^{73,81}

2.2.2.1. Mesoporosity generation through desilication. Postsynthetic desilication of pre-synthesis zeolite catalyst can be used to induce intracrystalline mesopores. Local dissolution of zeolite frameworks in a basic solution (like NaOH) is a known strategy for mesoporosity creation.⁸³ Fig. 6 depicts schematically the effect of Al content on the desilication treatment of MFI zeolites in alkali solution.⁸⁴

The desilication process is more effective for zeolites with high silica (Si/Al > 20) than high Al because SiO-removal, which is bonded directly to Al, is very difficult.⁷⁸ The zeolites desilication can be easily carried out at low concentration of alkali metal hydroxide. Mesoporosity generation highly depends on the Al distribution and concentration within zeolite crystals. Alrich textures almost remain unchanged, while silica-rich textures are easily leached out to produce large mesopores. Investigations showed that the Si/Al molar ratios in the range of 25–50 were most favorable for the uniform mesoporosity development and keeping the HZSM-5 crystal morphology.⁸⁴ HZSM-5 with Si/Al molar ratio less than 20 was very difficult to desilicate. Under mild basic conditions its framework was insoluble, while strong basic condition totally destroyed its zeolite framework. Alternatively, zeolites with high Si/Al molar ratio (>50) exhibited unselective and excessive dissolution generating too large pores.⁷⁸ Highly uniform mesopores could be generated within the zeolite frameworks by the addition of cetyltrimethylammonium bromide (CTAB) surfactant to the desilication medium.⁸⁵ This surfactant could contribute to the local desilication process, making micelles joint with base in the partly desilicated zeolite. The modified zeolite catalyst through this process showed that the acidic properties of the resultant zeolite changed slightly during desilication process. Review Workers and the secure of the generation of larger unaffected under similar situations. Furthermore, beta contract contract the form of mesopheric on the form of mesopheric contribution control of the particle of th

2.2.2.2. Mesoporosity generation through dealumination. During the decades selective dealumination has been utilized because it was understood that production of zeolite with higher ratio of Si/Al could create stable zeolites with higher strength acid sites. Generally, during calcination some parts of alumina species are removed from the zeolite structure, when it is less stable. Hydrolysis of the Si–O–Al bonds creates defect sites, therefore extra-framework alumina species can be eliminated. Using extra steam, which is commonly used for zeolite Y dealumination, increases the hydrolysis severity. Then, ultrastable Y zeolite with higher Si/Al ratio (USY), which is used as cracking catalyst in FCC (fluidized catalytic cracking) process, can be produced.⁸⁰ Amorphous alumina residues extraction is then performed by diluted nitric acid or oxalic acid. Hence, cavities and pores with broad sizes between 2 and 50 nm are generated.73,76

Apart from Y zeolite, dealumination can be applied to ferrierite, mordenite and beta zeolites mostly by direct leaching with more concentrated acids. Depends on the nature of zeolites, different acids such as oxalic, acetic, tartaric, nitric, hydrochloric and sulfuric were utilized with various concentration (even 6.0 M HCl). For example, in a comparative investigation of three different structures, dealumination of beta zeolite was easier than mordenite, whereas HZSM-5 was almost

unaffected under similar situations. Furthermore, beta zeolite dealumination conducted to higher loss of crystallinity, whereas mordenite indicated considerable mesopore volume.⁸⁶

Principally, aluminum extraction from the zeolite structure inevitably conducted to a change in Si/Al ratio, and consequently the acidity, while mesopores are created at the same time. In this conditions, understanding the effects of mesoporosity on changes of zeolite catalytic activity seems to be difficult. This type of complication may be one the main reason which mesopore generation through leaching process has been recently carried out by desilication process instead.^{73,80}

2.2.3. Summary of the fast pyrolysis vapor upgrading studies on mesoporous catalysts. Varieties of mesoporous catalysts consisting MCM-41, Al-MCM-41, metal-Al-MCM-41, MCM-48, Al-MCM-48, meso-MFI, Pd/SBA-15, MSU-S and SBA-15 were investigated for bio-oil/biomass pyrolysis vapor upgrading.⁸⁷–⁹⁴ Among different mesoporous materials, MCM-41 and meso-MFI based catalysts were extensively used for bio-oil upgrading. These mesoporous catalysts, alternatively, could resolve microporous zeolites drawbacks where it was difficult for large molecules to diffuse the catalyst pores.

The recent catalytic biomass to bio-fuel conversion investigations, conducted under different conditions over mesoporous catalysts, are summarized in Table 3. In this regard, the following key features could be concluded:

 Compare to nano Al-MCM-48, meso-MFI showed higher catalytic activity and higher yield of aromatic, phenolic and gaseous components thanks to its strong acidic sites and high porosity, which accelerated cracking reactions. High acidity caused decreasing of organic fraction. Incorporation of Ga to meso-MFI led to less cracking, increasing of aromatic components and coke formation diminishing.^{90,92}

 Mesoporous Al-MCM-48 and Al-MCM-41 catalysts showed high selectivity toward phenolic compounds, while meso-MFI (which possesses strong acid sites) indicated high selectivity toward aromatic components production. Pt incorporation to meso-MFI catalyst promoted dehydrogenation and cracking then conducted to enhanced aromatization and deoxygenation. Enlargement of MCM-41 pore size and loading of transition metals to it reduced acetic acid and water yield among the pyrolysis products.87,93

 Al-MCM-41 catalyst led to decarbonylation, decarboxylation, dealkylation, cracking and aromatization reactions. Higher coke formation, compared to zeolite catalysts, could be strong evidence of mentioned reactions. Higher Al content or in the other world lower Si/Al ratio caused an increase in the yield of the aromatic components. Incorporation of metals (like Fe and Cu) to Al-MCM-41 enhanced the phenols yield and decreased the level of both hydrocarbons and poly-aromatic hydrocarbons (PAHs).^{88,95}

2.3. Metal based catalysts

As explained within the context, there are several methods for the bio-oil upgrading comprising catalytic upgrading, steam reforming and hydrogenation. Among all these, the latter is the most widely used commercial process for the bio-oil upgrading

Table 3 Summary of most recent researches of vapor phase bio-oil upgrading over mesoporous catalysts

high acidity, promoted cracking and dehydrogenation and resulted enhanced deoxygenation and aromatization

Table 3 (Contd.) Table 3 (Contd.)

levels of both hydrocarbons and PAHs were low. (3) Fe–Al-MCM-41 and Cu–Al-MCM-41 together with the parent material with the lower Si/Al ratio were the optimum for the production of and conducting hydro-deoxygenation reactions. Through these reactions, some components like aldehydes can be converted into stable chemicals like alcohols and hydrocarbons with high heating value and low oxygen contents.^{89,96,97}

Mostly noble metal catalysts (e.g., Pt, Ru and Pd) at high temperature and pressure are employed to carry out hydrodeoxygenation reactions. These types of reactions often suffer from catalyst deactivation and clogging of the reactor at high temperatures. To overcome these problem, an approach for the bio-oil deoxygenation into high yield commodity products was employed. In this approach, hydro-processing of the bio-oil performed over supported metal catalysts (Ru/C and Pt/C) followed by conversion over zeolite catalyst (HZSM-5). Using this strategy, drawbacks associated with the conventional hydrogenation processes were overcome by operating the process at moderate temperature (\leq 250 °C), at which no reactor plugging or catalyst coking was observed. Alternatively, the bio-oil upgrading at atmospheric pressure is the other promising strategy to overcome the aforementioned problems.⁹⁸⁻¹⁰⁰

Some metal based catalysts like Fe, Zn, Al, and Mg can participate in organic reactions as strong reductants. For instance, Zn and Fe are commonly used for reducing nitro compounds to amines. Further, Zn is a key metal catalyst in the conversion of carbonyl groups (e.g. aldehyde and ketones) into methylene groups. These reactions are usually carried out with a high selectivity and yield at ambient temperature and pressure in acidic conditions.¹⁰¹

Bio-oil is a mixture of many oxygenated compounds like aldehydes, ketones and acids, which conduct bio-oil toward instability and corrosiveness. Therefore, the use of mentioned metal based catalysts can effectively enhance the bio-oil quality. Contrary to conventional hydrogenation process, low pressure pyrolysis vapor upgrading process over mentioned catalysts can be conducted without the presence of other catalysts and additional hydrogen gas.¹⁰¹

Catalytic cascade approach for the biomass pyrolysis vapor upgrading has recently attracted the attentions of researchers. The idea is to maximize carbon efficiency during bio-oil quality enhancement. In this regard, instead of oxygen functionalities (carbonyl, carboxylic, ketonic, and hydroxyl groups) elimination too early, their high reactivity is utilized to conduct C–C bond formation reactions, including aldol condensation and ketonization. Metal oxide catalysts are mostly efficient in catalyzing carboxylic acids ketonization, but reducible oxides like ceria can even catalyze the small aldehyde ketonization.²⁴ Fig. 7 depicts the proposed reaction mechanism of small aldehyde (propanal) conversion over $Ce_{0.5}Zr_{0.5}O_2$ catalyst. The contribution of two major reactions comprising ketonization and aldol

Fig. 8 Schematic of the chemical looping deoxygenation (over metal oxide catalysts) concept $(T_3 > T_1 > T_2)$.¹⁰³

Fig. 7 Proposed reaction mechanism for propanal conversion over $Ce_{0.5}Z_{0.5}O_2$ – adapted from Gangadharan et al.¹⁰²

Table 4 Summary of most recent researches of vapor phase bio-oil upgrading over metal base catalysts

proportion of heavy fraction in the bio-oil (from 22% to 15%)

with a limited decrease in its yield

Table 4 (Contd.) Table 4 (Contd.)

condensation could be observed in the network, involving several condensation steps. Furthermore, there were also different side reactions that could take place in parallel. It is well understood that aldol condensation can happen on both basic and acid sites. In the mixed oxide catalysts, the oxygen anions can behave as either Brønsted or Lewis base sites, while the exposed cations are Lewis acid sites.^{56,102}

Zeolites are also effective catalysts for C–C bond formation, but the selectivity is toward aromatics formation. For instance, propanal can be selectively converted to C_7-C_9 aromatics through aldol condensation over HZSM-5 catalyst.²⁴

Pacific Northwest National Laboratory (PNNL) in USA has recently focused on the pyrolysis vapor upgrading with the objectives of maximizing carbon efficiency and minimizing hydrogen consumption. They employed a new concept based on chemical looping and utilizing metal oxide catalysts to selectively eliminate oxygen from the pyrolysis vapors without hydrogen feeding.¹⁰³ The concept is shown in Fig. 8.

The pyrolysis vapors react with the partially reduced metal oxide (MeOx-1) while they pass over the deoxygenation catalyst. The metal oxide is oxidized (MeOx) while the pyrolysis vapors are reduced (deoxygenated). To reduce the catalyst (MeOx-1) having the ability to be recycled back to the reactor, the metal oxide is heated under N_2 stream at a higher temperature. Model compound experiments and theoretical calculations identified some promising metal oxide catalysts for such type of vapor phase deoxygenation.¹⁰³ A similar investigation was patented by Lissianski et al.¹⁰⁴ where pyrolyzing the biomass was performed in the presence of a transition metal, using microwave energy.

In an earlier study, Sanna and Andrersen¹⁰⁵ suggested new catalysts for the biomass (wheat spent grains) conversion into deoxygenated bio-oil in a fluidized bed reactor. They used two Mg-rich activated olivine (ACOL) and activated serpentine (ACSE), and alumina (ALU) as catalyst. A considerable reduction of oxygen content in the bio-oil was observed in following order: ACOL > ACSE > ALU. Particularly, compared to ACOL which was able to remove about 40 wt% of the original oxygen from the bio-oil, ACSE and ALU decreased it to less than 20–30 wt%. The oxygenated compounds of the bio-oil interacted in the catalyst's active sites with the metallic species and produced C_5-C_6 components through decarboxylation.

The catalytic vapor upgrading, which is an attractive process with lots of advantages, has been widely investigated employing acidic zeolites. Nevertheless, zeolite catalysts suffer from fast coke deposition and PAHs formation during upgrading. Furthermore, MCM-41 based mesoporous catalysts exhibited crucial disadvantages; high production cost and poor hydrothermal stability.¹⁰⁶ Due to the advantages associated with metal based catalysts and in order to likely resolve zeolites and mesoporous catalysts problems, several researches recently have been performed on varieties of metal based catalysts, which some of them are summarized in Table 4.

Some of the key features of metal oxide catalysts used for biooil vapor phase upgrading can be noted as follows:

 Several metal oxides including MgO, NiO, alumina, zirconia/titania, zirconia and titania were used as catalyst. Deoxygenation occurred up to some extents in compare to noncatalytic pyrolysis, but zirconia/titania showed the most interesting deoxygenation and highest yield of aromatic compounds. Alumina showed that bio-oil yield and composition depended on pyrolysis atmosphere and the catalysis/feed ratio. The bio-oil produced over $\text{Na}_2\text{CO}_3/\gamma$ -Al₂O₃ had comparable properties to those of fuel oil.^{15,66,107-110}

 \bullet TiO₂ and ZrO₂ investigations indicated that they both had very good heat stability compare to mesoporous catalysts. TiO₂ + $ZrO₂$ mixed oxides resulted high hydrocarbon yield and significantly decreasing of acid contents.¹¹¹

 Reaction over CaO yielded bio-oil with low acid, carbonyl and phenols contents while the molality of hydrocarbons increased. CaO/feed ratio increasing enhanced dehydration reactions.112,113

 CuO exhibited very interesting results to yield semi-volatile compounds and high bio-oil yield. Boric oxide promoted hydroxyl group removal with generation of alkyl groups which consequently reduced oxygen contents.^{114,115}

2.4. Catalyst deactivation

The catalyst deactivation is one of the challenging issues in catalytic biomass pyrolysis vapor upgrading. It is not only caused by coke formation, but also the strong adsorption of the oxygenate components on the surface of catalyst support. Generally two types of cokes are formed over catalyst during biomass catalytic vapor upgrading. One, which has thermal origin, is called thermal coke and the other, which has catalytic origin, is called catalytic coke. Thermal coke, which is often formed over outside of catalyst's particle, is due to phenolic compounds polymerization. Catalytic coke, which is mostly deposited inside the catalyst's channels, is caused by aromatization, oligomerization condensation and cyclization of oxygenate components.¹¹⁶–¹²⁰ Catalyst characteristics and biomass feedstock properties can influence catalyst deactivation and coke formation. Review Workers condensation could be observed in the network, involving eat
alytic probabis, but arizonizational showed the most interest of the
differential distribution steps. Furthermore, there were also of production
a

> 2.4.1. Effects of catalyst characteristics. The catalyst deactivation will be more pronounced on the aluminosilicate type of

Fig. 9 Ammonia temperature programmed desorption (TPD) for the fresh (solid line) and spent (dotted line) catalyst.¹²¹

catalysts, which contain acid sites. Carlson et al .¹²¹ investigated the zeolite catalyst deactivation caused by acid sites lost during biomass (pine wood sawdust) catalytic pyrolysis. They used temperature programmed desorption (TPD) to measure the total number of acid sites. The related TPD curves for the spent and fresh catalyst are illustrated in Fig. 9. As can be seen, there were two peaks with centers at \sim 275 °C and \sim 475 °C. The high temperature peak was related to the more strongly bound ammonia on Brønsted acid sites, while the low temperature peak corresponded to weakly bound ammonia on Lewis acid sites.¹²¹–¹²⁴ From the TPD curves it was concluded that the zeolite catalyst's acidity loss was attributed to a Lewis acid sites deactivation as the high temperature peak did not change much.

In zeolites, coke and tar deposits, which block the catalyst pores and cover its acid sites, significantly are influential on the catalyst activity and selectivity reduction.¹¹⁸ Dealumination of the zeolite catalysts (like HZSM-5) in the presence of steam was reported by several researchers. It can be conducted to catalyst acidity lost and irreversible deactivation.¹²¹

Physical characteristics of the zeolite catalyst comprising pore shape, pore size and crystallite size can highly affect the coke formation.¹²⁵ Catalytic upgrading of pine wood in a fluidized bed reactor using HBeta-25, HY-12, HZSM-5 and HMOR-20 (Mordenite) catalysts showed that coke formation was fairly low for both Mordenite and HZSM-5. Spent Y zeolite (HY-12) exhibited the highest coke content. This was possibly due to the highest initial surface area and large cavities in the structure of Y zeolite, which allowing bigger molecules to diffuse to the inner part of zeolite.¹²⁶

Catalytic pyrolysis, using different zeolite catalysts having small (ZK-5, SAPO-34), medium (Ferrierite, ZSM-23, MCM-22, SSZ-20, ZSM-11, ZSM-5, IM-5, TNU-9) and large (SSZ-55, Beta zeolite, Y zeolite) pore size, were studied by Jae et $al.53$ Compared to zeolites with large pore size, small pore size produced less coke. The least coke formation was resulted from medium pore size with moderate internal pore space catalysts.

In addition, coke formation over catalyst can be influenced by zeolite crystallite size. Small crystallites showed much slower deactivation and less coke formation compared to large crystallites. This was due to the shorter diffusion path and quicker removal of products from the catalyst's channels. So, products did not have sufficient time to be converted to coke precursors and coke.¹²⁷

2.4.2. Effects of feedstock properties. The availability of oxygenated compounds, such as guaiacol or phenol in bio-oil, contributes to the coke formation. Part of this coke blocks the pores thanks to the bulky oxygenated molecules (which are adsorbed on the outer zeolite crystal surface) diffusional constraints.¹²⁸ The different roles of the bio-oil components in the formation of coke have been investigated by Gayubo et $al.^{129}$ They identified the phenols and aldehydes as the main precursors of cokes.¹³⁰

Increasing H/Ceff. mole ratio in oxygenated bio-oil favored the formation of olefins and aromatics and attenuates coke formation.¹³¹ Investigations showed that feedstocks with hydrogen to carbon effective ratio $(H/C_{\text{eff.}})$ less than 1.0 were

difficult to upgrade over a HZSM-5 catalyst due to quick deactivation (coke formation) of the catalyst.¹³² The H/C_{eff} ratio of the petroleum based feedstocks varies from 1 to 2, whereas that of the biomass feeds are only from 0 to 0.3. Therefore, the biomass contained hydrogen deficient molecules, and approaches for the biomass and its derived feedstocks transformation must consider their H/C_{eff.} ratio. Co-feeding of alcohol (like methanol) and biomass is one of the possible ways to enhance H/C_{eff.} ratio.²⁹ Zhang et al.¹³³ showed that methanol cofeeding with biomass (pine wood) at $H/C_{eff.}$ ratio of 1.25 increased the aromatics yield and decreased coke formation over HZSM-5 catalyst. Fig. 10 depicts a kinetic scheme suggested for the bio-oil/biomass pyrolysis vapor and methanol mixture conversion into coke and hydrocarbons over HZSM-5 catalyst.

The quantity and composition of the deposited coke on the HZSM-5 catalyst showed the significance of catalyst acidity for the formation of catalytic and thermal coke fractions. The major fraction of the produced coke was possibly due to the polymerization of the products derived from the biomass components pyrolysis (mostly lignin). Mostly, two fractions of coke were formed on the catalyst. The fraction of coke which was burned at low temperature was formed by condensation– degradation of lignin based oxygenated compounds. This type of coke was deposited on macro- and mesoporous structure of the zeolite catalyst matrix. The other one, which was burned at higher temperature and being deposited on the catalyst's micropores, was formed by condensation reactions activated by the acid sites. Formation of this type of coke was considerable in pure methanol catalytic conversion. Methanol addition to the pyrolysis vapor decreased the coke formation attributed to the attenuation of the phenolic compounds (lignin originated) polymerization and their deposition on the catalyst. According to the literature, pure methanol catalytic conversion on HZSM-5 catalyst formed non-oxygenated aromatics and aliphatic hydrocarbons as major components.^{119,120} **PSC Areness**

eatalysis, which contain aid sits. Carlson et al.¹⁰ investigated difficult to upgrade over a 1125M-5 catalys disc equilibre three contains the contains the contains of the contains of the contains of the c

> Lignin derived phenolic compounds like anisole and guaiacol and mostly those with multiple oxygen functionalities $(-OH, -OCH₃, C=O)$ are the major deactivating components.

Fig. 10 Kinetic scheme suggested for the bio-oil/biomass and methanol mixture conversion into hydrocarbons and coke on HZSM-5 catalysts. Adapted from Valle et al.¹¹⁵

Table 5 Summary of most recent researches of vapor phase bio-oil upgrading catalyst deactivation

Table 5 (Contd.)

The presence of alkalis, as well as N- and likely S-containing compounds in the biomass, improve catalyst deactivation.134,135

2.4.3. Summary of researches on catalyst deactivation. Due to the importance of catalyst deactivation in the biomass to biofuel catalytic conversion, the present study tried to have a survey on the related investigations. Its outcome is summarized in Table 5.

Some key aspects in this regard are as following:

 Feedstock (biomass and lignin) pyrolysis product deoxygenation over HZSM-5 catalyst indicated that higher content of H-lignin, which could produce higher concentration of phenols, could cause quick catalyst deactivation. An irreversible poisoning was observed after some regeneration cycles due to the Brønsted acid sites deactivation. Strong acidic HZSM-5 zeolite catalyst led to coke formation due to dealkylation, decarboxylation, decarbonylation, aromatization and cracking reactions. The methanol co-feeding indicated a significant e ffect on coke deposition reduction on Ni-HZSM-5 catalyst during bio-oil upgrading.^{21,28,135,136}

 Cracking of the high moisture pyrolysis vapors resulted slower catalyst deactivation and coke formation on β - and
V-zeolites Incorporation of Ca to V-zeolite caused more overant Y-zeolites. Incorporation of Ca to Y-zeolite caused more oxygen removal and slower catalyst deactivation. It is due to conservation of slightly more hydrogen in pyrolysis products.^{69,137}

 MSU-S catalyst high selectivity toward aromatics, PAHs and coke formation (catalyst deactivation) were attributed to its stronger acid sites in comparison to Al-MCM-41 catalyst. Compared to HZSM-5 zeolite, Al-MCM-41 showed higher tendency toward coke formation and catalyst deactivation.^{21,94}

3. Conclusion and outlook

Catalytic biomass pyrolysis vapor upgrading process to enhance the bio-oil quality indicates immense potential to convert renewable biomass to bio-fuel. Fast pyrolysis, which is known as a promising process to convert pretreated biomass to bio-oil, is affected by the biomass types and reaction conditions. Catalytic vapor phase upgrading is aiming to treat the fast pyrolysis vapor before condensation. It recently has attracted the attentions of bio-fuel researchers due to the prominent techno-economical characteristics of this type of upgrading in comparison with conventional hydro-deoxygenation (HDO) process. Despite HDO process which consumes high hydrogen quantity and requires complicated equipment working at high pressure, this upgrading approach is carried out at atmospheric condition without hydrogen feeding. The produced bio-oils yields and qualities are strongly dependant on catalysts types and properties (e.g. structure, acidity and pore size), reaction conditions and feed type.

Three most important classes of catalysts including microporous zeolites, mesoporous catalysts, and metal based catalysts are used for vapor phase bio-oil upgrading. Among zeolite catalysts, HZSM-5 (which possesses a three dimensional pore structure, high acidity and shape selectivity) indicates the superior performance in deoxygenation, aromatic compounds production and resistance to coke formation. However, to keep the catalyst activity and selectivity for long time, deactivation through dealumination as well as coke deposition need to be minimized. The

product distribution and coke formation amount over catalyst are strong function of catalyst shape selectivity and acidity. Shape selectivity of the zeolite catalyst are mostly influenced by pore size and shape as well as crystallite size. Mesoporous catalysts having pore diameter larger than 2 nm can resolve large molecules mass transfer limitation, associated with microporous zeolites. Contrary to mesoporous MCM-41 based catalysts, which have lower thermal stability and acidity compared to zeolite based catalysts, mesoporous MFI catalysts simultaneously possess high acidity as well as large pore diameter to overcome mass transfer limitation (which is micro-zeolites drawback). On the other hand, metal based catalysts, exhibit high acidity and outstanding resistance to coke deposition; therefore they can be reputable catalysts for bio-oil upgrading.

Efforts to transform lignocellulosic biomass to intermediate and base chemicals for the biofuels production have been fruitful to a considerable extent in recent years. In the next decades, it will be expected to find more techno-economical processes which can employ advanced catalytic processes to convert biomasses from various resources into fine chemicals, base chemicals and fuels. We will be approaching to more sustainable and renewable economy, although further efforts will be required. Biofuel upgrading technologies still need development to create costcompetitive products with acceptable productivity and selectivity. Promising improvement on heterogeneously catalyzed transformation of lignocellulosic biomasses to fuel like and value added chemicals with low coke formation over catalysts has attracted intensive attention in the past few years and breakthroughs have been attained up to some extent. It might be proper to mention that conversion of biomasses to desired chemicals with low coke formation, high selectivity and yield remains in its infancy until there are considerable developments in heterogeneous catalysts. Biomasses catalytic pyrolysis vapor upgrading through a cascade system of different catalysts (micropore zeolites, mesopore and metal based catalysts), that includes several consecutive steps for various bio-oil fractions upgrading, seems to be a promising thermochemical conversion/ upgrading technology. Each individual mentioned catalysts or their employment in a cascade system indicated high potential for industrialization, although bio-oil upgrading through a cascade systems of catalysts most probably in the near future would attract the researchers' attentions. Review Works charge the computer on the computer of the compu

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