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Selective hydrodeoxygenation of 5-hydroxymethylfurfural to 2,5-dimethylfuran on Ru–MoO_x/C catalysts†

Yue Yang,^a Qiying Liu,^{*b} Dan Li,^b Jin Tan,^b Qi Zhang,^b Chenguang Wang^b and Longlong Ma^{*b}

Selective hydrogenation of 5-hydroxymethylfurfural (HMF) has potential application in high quality biofuels. Herein, the catalytic hydrodeoxygenation (HDO) of HMF to 2,5-dimethylfuran (DMF) was investigated using bi-functional Ru–MoO_x/C catalyst prepared by initial wetness impregnation. The high dispersion and electronic transfer between Ru and MoO_x were demonstrated by a series of characterization techniques. During this HDO process, the synergy effect between metallic Ru and acidic MoO_x species in the Ru–MoO_x/C catalyst plays an essential role in obtaining maximized target product DMF (79.4%) via effective aldehyde group hydrogenation by Ru followed by dehydration over MoO_x. This work also elucidated that DMF production proceeded through two distinct pathways: the 2,5-hydroxymethyl furan intermediate was preferable by the aldehyde group hydrogenation of HMF over the Ru–MoO_x/C catalyst. Over MoO_x/C catalyst, comparatively, 5-methyl furfural was the key intermediate by direct hydrogenolysis of the hydroxyl group in HMF. This kind of catalyst is stable for the first two runs by maintaining the target product yield. After the third run, the catalyst showed deactivation gradually but could be almost completely recovered after regeneration by H₂ reduction.

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1. Introduction

With the decrease in fossil resources, the demand for exploring new sustainable energy to produce chemicals and fuel is urgent for the future.^{1–3} Renewable and sustainable lignocellulose is composed of cellulose (40–50%), hemicellulose (25–35%) and lignin (15–20%), and can be used as feedstock for value-added chemicals/fuels production, which has attracted great attention during the last decade.^{4,5} However, the typical biomass-derived compounds usually have high oxygen-content and could not be used for fuel directly.^{6,7} Therefore, an additional hydrogenation step is necessary to complete or selectively remove the oxygen atoms in the compounds.^{8,9}

5-Hydroxymethylfurfural (HMF), one of the important platform compounds from cellulose,^{10–12} can be further converted into a series of valuable high-quality chemicals/fuels, such as γ -valerolactone, levulinic acid, oxidative 2,5-furandicarboxylic, 2,5-dimethylfuran and so on.^{13–17} Among those chemicals/fuels, 2,5-dimethylfuran (DMF)^{19–21} by HMF hydrogenolysis has attracted

much attention, due to the oxygen-containing DMF being an ideal renewable oxygen liquid fuel according to its higher energy density, higher octane number and higher boiling point compared to the traditional cellulose based ethanol.^{22,23}

Catalysts played the key role in producing DMF with high yields from HMF, lignocellulosic biomass and the derivatives. In 2007, Dumesic *et al.*¹ (Table 1, entry 1) firstly reported a catalytic strategy using a biphasic system to extract HMF, and a separated process using 6.8 bar of H₂ over CuRu/C catalyst at 220 °C to produce 71% yield of DMF. Under the same reaction conditions, Binder and Raines²⁵ (Table 1, entry 8) achieved 49% DMF yield by hydrogenolysis of crude HMF which obtained from corn stover. Chidambaram and Bell² (Table 1, entry 7) added acetonitrile as auxiliaries and obtained 32% DMF yield over Pd/C catalyst in ionic liquids. For improving the yield of DMF, Rauchfuss *et al.*²⁷ (Table 1, entry 6) utilized HCOOH as the hydrogen donor and obtained 95% DMF yield. However, HCOOH is corrosive and environment unfriendly, which leads to apply restrictedly.⁷ Recently, researchers reported to employ adding transitional metal additives which can form Lewis acid to obtain high DMF yield by adsorption and activation the hydroxyl and aldehyde group in HMF. And the metal sites were active for hydrogenation while the acidic sites for promoting dehydration of CH₂–OH group.^{28,29} Zu *et al.*²⁴ (Table 1, entry 4) used Ru/Co₃O₄ to produce 93.4% DMF yield at 130 °C after 24 h and Huang *et al.*²⁶ (Table 1, entry 5) reached up to 96% DMF yield over Ni–W₂C/AC at 180 °C after 3 h in THF. Judging from

^aDepartment of Chemistry, University of Science and Technology of China, Hefei 230026, P. R. China

^bKey Laboratory of Renewable Energy, Guangzhou Institute of Energy Conversion, Chinese Academy of Sciences, Guangzhou 510640, P. R. China. E-mail: mall@ms.giec.ac.cn; liuqy@ms.giec.ac.cn; Tel: +86-20-3702-9721, +86-20-87057673

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Table 1 Results of direct conversion of biomass into DMF

Order	Substrate	Medium	Catalyst	Condition	Yield%	References
1	HMF	1-Butanol	Cu/Ru/C	220 °C, 10 h	71	1
2	HMF	<i>n</i> -Butyl alcohol	Ru/C	260 °C, 1.5 h	60.3	18
3	HMF	THF	Zn/Pd/C	150 °C, 8 h	85	22
4	HMF	THF	Ru/Co ₃ O ₄	130 °C, 24 h	93.4	24
5	HMF	THF	Ni-W ₂ C/AC	180 °C, 3 h	96	26
6	HMF	THF + HCOOH	H ₂ SO ₄ + Pd/C	70 °C, 15 h	95	27
7	Glucose	EMIMCl	12-MPA + Pd/C	120 °C, 3 h	32	2
8	Cellulose	DMA-LiCl/ EMIMCl	CrCl ₂ + HCl + Cu/Ru/C	220 °C, 10 h	49	25

the results, the bi-functional catalyst was an ideal pattern for HMF hydrogenolysis, but these process involved either long reaction time or high H₂ pressure and the interaction between metal and acid sites was not clearly. Therefore, there is a need to develop a new-type metal-acid catalyst to conversion of HMF to DMF in mild condition and explore the synergy effect between metal and acid sites.

On the basis of the above-mentioned research, we designed the metal-acid Ru-MoO_x/C catalyst by step wetness impregnation for HMF hydrogenolysis to DMF in *n*-butyl alcohol. Due to the different functional groups in the HMF, such as C=O, CH₂-OH, furan ring, hydrogenation of HMF to DMF needs C=O hydrogenation to the corresponding hydroxyl groups and subsequent C-OH groups hydrogenolysis. Under the optimum condition (180 °C, 1.5 MPa, 1 h), the selectivity of DMF was 79.8%. In addition, the characterization of catalyst was conducted to explain the interaction and electronic transfer between Ru and MoO_x, and the cause of catalyst deactivation. In addition, various reaction parameters was modulated to investigate the pathway for HMF hydrogenolysis and the synergistic effect of the metal and acid sites.

2. Experimental

2.1. Materials

HMF (98%) and 5-MF (99%) was supplied by J&K chemical Co. Ltd. DMF (99%) and RuCl₃·3H₂O was purchased from Shanghai Aladdin Reagent Co. Ltd. 2,5-Hexanediol (98%) and 2,5-hexanedione (98%) was sourced by Macklin Reagent Co. Ltd. 2,5-dihydroxymethylfuran (95%), (5-methyl-2-furyl)methanol (97%), (*cis*-tetrahydrofuran-2,5-diyl)di-methanol (95%) and (NH₄)₆Mo₇O₂₄·4H₂O, 2,5-dimethyltetrahydrofuran (98%) was obtained from Maya Reagent, Energy chemical, Shu ya Reagent, Tianjin Kaida chemical and TCI Co. Ltd respectively. All the reagents was used without further purification.

2.2. Catalyst preparation

The catalyst was prepared by step-wise impregnation method. In a typical synthesis, active carbon (40–60 mesh) was chose as support materials. Firstly it was pretreated by nitric acid (0.6 mol l⁻¹) at 110 °C for 5 h, then filtered until cooled to room temperature, and dried at 100 °C overnight. RuCl₃·3H₂O was dissolved in deionized water and a certain amount of active carbon was added to a desired Ru loading by weight, and stirred

at 100 °C to remove excess water. The solid was dried at 100 °C overnight and calcined under a N₂ atmosphere at 400 °C for 4 h. After the preparation of Ru/C catalyst, a second impregnation with (NH₄)₆Mo₇O₂₄ solution by the similar procedure mentioned above. Prior to the experiment, the dried sample was reduced in flowing H₂ (40 ml min⁻¹) at 350 °C for 4 h.

2.3. Catalyst performance

Hydrogenolysis of HMF was carried out in a stainless steel reactor with a Teflon inner. In a typical experiment, a mixture of 0.5 g HMF dissolved in 40 ml *n*-butyl alcohol was loaded to the auto-clave, followed by adding 0.1 g of freshly reduced 5% Ru-10% MoO_x/C catalyst (reduced at 350 °C for 4 h). The reaction system was sealed, flushed by N₂ to remove air residue, pressurized the H₂ pressure to 1.5 MPa, and then heated to a certain temperature for a certain time. After the reaction, the reactor was cooled to room temperature immediately in a water bath. The liquid products was collected and qualitatively analyzed by a GC-MS instrument with an Agilent 7890A-5975C and quantitatively determined by GC with SHIMADZU GC-2014C gas chromatograph equipped with a FID and a HP-INNOWAX column.

The product yields were calculated based on internal standard with using cyclohexanone as the internal standard. The conversion (conv.) of HMF and the selectivity of product *i* (S_{*i*}) were defined as:

$$\text{Conv. (HMF)} = \frac{C_0 - C_t}{C_0} \times 100\%$$

$$S_i = \frac{C_i \times 6}{(C_0 - C_t) \times a_i} \times 100\%$$

here, C₀ refers to the initial concentration of HMF and C_{*t*} means the mole concentration of final HMF after reaction; C_{*i*} indicates the concentration of product *i*, and a_{*i*} represents the carbon numbers of product *i*.

2.4. Catalyst characterization

X-ray photoelectron spectroscopy (XPS) was conducted on a Thermo Fisher Scientific Escalab 250 Xi photoelectron spectrometer, the values were revised by C 1s (284.8 eV). The specific surface area was measured by N₂ isothermal (77 K) adsorption on Mesoporous ASAP-2010 automated system. The tested samples was outgassed at 200 °C for 8 h prior to analysis. The



surface area was calculated by BET method. The morphology and energy spectrum analysis of catalysts were investigated by transmission electron microscopy (TEM, JEM-2100F) and scanning electron microscopy (SEM, SU-70). Temperature programmed reduction (TPR) equipped with a TCD detector was used to measure the reduction behavior under H_2 . For H_2 -TPR experiments, 100 mg sample was laid into a quartz tube and pretreated by Ar at 300 °C for 1 h, then cooling to 50 °C, the catalysts were heated to 600 °C at the rate of 5 °C min^{-1} . Temperature programmed desorption of NH_3 (NH_3 -TPD) experiments were conducted on AutoChem II 2920 to analyze the acid content. The catalysts were purged by He at 673 K prior to adsorption, then cooling to 373 K to saturate with NH_3 and sweep by He to remove the physical adsorbed NH_3 for 30 min, whereafter, the samples were heated to 673 K at the speed of 10 K min^{-1} . All of the catalysts which needed to reduce were pre-treated under H_2 stream for 4 h at 350 °C and cooled to room temperature, then passivated by 0.6% O_2/N_2 gas for 4 h.

3. Results and discussion

3.1. Catalyst characterization

Hydrogen temperature-programmed reduction (H_2 -TPR) was conducted to investigate the reduction properties of the current

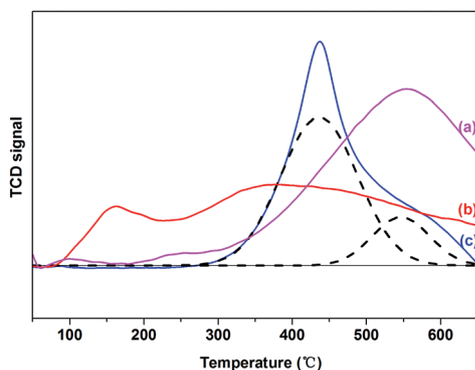


Fig. 1 H_2 -TPR profiles of (a) 5% Ru/C, (b) 5% Ru–10% MoO_x/C and (c) 10% MoO_x/C catalysts.

Ru– MoO_x/C catalysts, and the results were shown in Fig. 1. For 5% Ru/C catalyst, two reduction peaks at 100 °C and 230 °C were observed, these peaks were the reduction of RuO_x species due to the different interactions with support. The first peak was assigned to the weak interaction of Ru with support, which could be easily reduce. While, the second peak was assigned to the reduction of RuO_x species which was strongly interaction with support.^{30–32} the Ru catalytic activity can promote the break of C–C bond on the surface of active carbon, which led to the uphill of TCD signal at 500 °C.^{32,33} The profile of 10% MoO_x/C catalyst presented a main peak and a shoulder peak centered at 450 °C and 550 °C, respectively. It was attributed to the reduction of Mo^{6+} with octahedral coordination to Mo^{4+} at 450 °C, while the H_2 consumption peaks located at 550 °C belonged to the reduction of tetrahedrally coordinated Mo species to lower oxidation state species.^{34–37} Over the 5% Ru–10% MoO_x/C catalyst, the small reduction peak at 180 °C was attributed to RuO_x to Ru^0 .³⁸ However, the main Mo reduction peak located at 350 °C and it was lower than the 10% MoO_x/C catalyst (450 °C). These results suggested that Ru facilitated to the reduction of Mo species,³⁹ in which H_2 was adsorbed on Ru and dissociated to highly reactive hydrogen atoms.⁴⁰ When Ru and MoO_x species was close on the support, the H_2 spillover facilitated the reduction of MoO_x .^{39,41–43} This behavior increased a possible electronic transfer between Ru and MoO_x .³⁹

To verify the interaction between Ru and MoO_x , the surface chemical state of 5% Ru/C, 10% MoO_x/C and 5% Ru–10% MoO_x/C were conducted by X-ray photoelectron spectroscopy (XPS) characterization. Because of 5% Ru/C and 5% Ru–10% MoO_x/C catalyst were liable to oxidation, they were reduced by *in situ* technology to detect the chemical state of elements. Due to the photoemission line of Ru 3d was overlapped with C 1s,⁴⁴ Ru 3p_{3/2} line was used to analyze the surface species. As shown in Fig. 2a, for the fresh 5% Ru–10% MoO_x/C catalyst, the binding energy of 463.2 eV was ascribed to RuO_2 ,⁴⁵ and shifted to relatively low binding energy after reduction by H_2 (reduced 5% Ru–10% MoO_x/C catalyst), assigning to a reduction of RuO_2 to metal Ru. In addition, over the reduced 5% Ru/C catalyst, the binding energy of metal Ru shifted to higher binding energy compared to the reduced 5% Ru–10% MoO_x/C catalyst, which

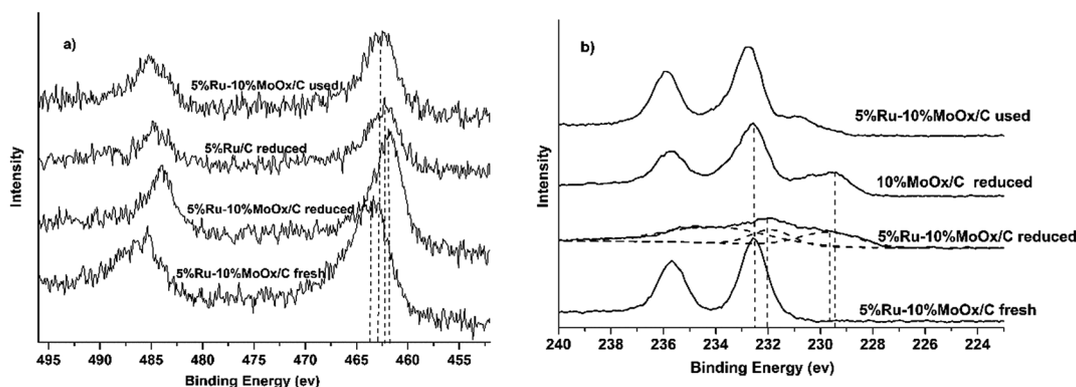


Fig. 2 XPS spectra of Ru– MoO_x/C catalysts: (a) Ru 3p and (b) Mo 3d. 5% Ru–10% MoO_x/C fresh: the catalyst had not been reduced by H_2 ; 5% Ru–10% MoO_x/C , 10% MoO_x/C and 5% Ru/C reduced: the catalysts were reduced by H_2 under 350 °C; 5% Ru–10% MoO_x/C used: the catalyst was reduced by H_2 under 350 °C and then used at the reaction condition.



indicated that there have some electrons was transferred from MoO_x to Ru.³⁹ On the other hand, the chemical status of Mo 3d was measured and the results were shown in Fig. 2b. The fresh 5% Ru–10% MoO_x/C sample presented only one doublet located at 232.6 eV which was assigned to Mo^{6+} species (MoO_3).³⁵ The reduced 5% Ru–10% MoO_x/C catalyst revealed three doublets, and the values at 232.0 eV and 229.6 eV were contributed to the Mo^{5+} and Mo^{4+} formation during H_2 reduction.^{46,47} The binding energy of Mo^{4+} over 5% Ru–10% MoO_x/C reduced catalyst was shifted to left compared to 10% MoO_x/C catalyst, which was attributed to the fact that some electrons were transferred from MoO_x to Ru. The low chemical valence of MoO_x ($x < 3$) could form surface defects, which promotes hydrogenolysis of hydroxyl groups as Lewis acid.⁴⁶ In addition, from Fig. 2a, the value of the used 5% Ru–10% MoO_x/C catalyst reflected this pattern existed two chemical state, namely Ru and RuO_2 .

Fig. 3 showed the TEM images of specified catalysts and the diameter of particles was calculated by the software Nano Measure. All of the particles was included to calculate the average particle size in the visible range. For the 10% MoO_x/C catalyst, the MoO_x particles (Fig. 3e) presented the average diameter of about 7.3 nm. The supported 5% Ru/C (Fig. 3d) catalyst showed a better dispersion by observing the smaller average diameter of about 2.6 nm. Compared to the previous two monometallic catalysts, however, the average particle size of 5% Ru–10% MoO_x/C catalyst (Fig. 3a) was about 1.8 nm which was highly dispersed on the support of activation carbon, owing to the interaction between Ru and MoO_x nanoparticles and facilitates the dispersion.^{48,49} The high dispersion was benefit to hydrogenolysis.^{48,50} While, the few larger particles in Fig. 3a were MoO_x species (distinguished by the HRTEM on the corner), which had not been combined with Ru, and their particle size was about 17 nm, it must be pointed out that this size can not reflect the actual size of MoO_x because of few statistical particles. The average size of 5% Ru–5% MoO_x/C (Fig. 3b) and 5% Ru–20% MoO_x/C (Fig. 3c) were 1.1 nm and 2.5 nm respectively, and the 5% Ru–20% MoO_x/C was showed poor dispersion, which revealed the MoO_x content could impact the size of particle and dispersion. Meanwhile, the high-resolution electron microscopy (HRTEM) image showed the morphology of the two species (Fig. 3f) of 5% Ru–10% MoO_x/C

catalyst. The MoO_x species was close contact with the Ru moiety in this two metallic catalyst, which induces the electron transferring from MoO_x to Ru. This result is well consistent with the XPS measurement.³⁹

Surface acidity and acidic strength distribution of Ru– MoO_x/C with different MoO_x loading was evaluated by NH_3 -TPD (Fig. S1†). The NH_3 -TPD profiles of 5% Ru/C, 5% Ru–5% MoO_x/C , 5% Ru–10% MoO_x/C , and 5% Ru–20% MoO_x/C consisted a wider shoulder peak in the temperature range of 390–480 K, and these peaks could be attributed to NH_3 adsorb on the weak acid site of the catalyst surface.⁵¹ For the Ru/C catalyst, almost no peak was observed (Table S1,† entry 4), with the increase of MoO_x loading, the peak area augmentation and the strength intensified which indicated that the total acid was enhanced (Table S1†).

3.2. HMF hydrogenolysis to DMF

The catalyst was tested in HMF hydrogenolysis to produce DMF. During the reaction process, 2,5-dihydroxymethylfuran (DHMF) and 5-methylfurfural (MF) were the main intermediates along with different reaction pathways. Meanwhile, the byproducts mainly contained the O-containing intermediates (e.g. 5-methyl-2-furanmethanol (MFA), 2,5-dihydroxymethyl tetrahydrofuran (DHMTHF)) and the over-hydrogenated products such as 2,5-dimethyl tetrahydrofuran (DMTHF), 2,5-hexanedione and 2,5-hexanediol by furan ring opening. The carbon balance of all detected products was about 65–86% maybe due to the formation of undetected molecules^{7,24,52} and polymers and/or cokes.⁵³ In the studied condition, the *n*-butyl alcohol had no reactivity over catalysts.

3.2.1 Ru– MoO_x/C catalysts. For all the tested Ru– MoO_x/C catalysts, the HMF conversion was 100% while the DMF selectivity significantly varied based on Ru/Mo ratios. When the 2% Ru–10% MoO_x/C was used, the target DMF selectivity was 60.8%, simultaneously accompanied with the over-hydrogenated products DMTHF (3.6%), DHMTHF (0.2%) and the furan ring opening byproducts 2,5-hexanedione (1.4%) and 2,5-hexanediol (0.7%) by trace amounts (Table 2, entry 1). Increasing the Ru loading to 5 wt%, the 5% Ru–10% MoO_x/C catalyst showed a better performance with 79.8% selectivity of DMF owing to the activity of hydrogenation increased simultaneously (Table 2, entry 2). Keep the Ru loading as a constant, when MoO_x was introduced by different loadings, the DMF selectivity gradually increased with increasing the MoO_x content to 10 wt%. When the Mo content exceeded 10 wt%, however, the DMF selectivity decreased and the furan ring opening byproducts (2,5-hexanedione and 2,5-hexanediol) increased gradually, indicating that excessive acidic site, which had been verified by NH_3 -TPD (Table S1†), was benefit to open the furan ring (Table 2, entry 2–4).

In order to understand the role of Ru and MoO_x in this hydrogenolysis reaction, 5% Ru/C and 10% MoO_x/C were prepared and tested under the same reaction condition. Over 5% Ru/C catalyst (Table 2, entry 5), the target DMF selectivity was 54.1%, in addition, over-hydrogenated DMTHF and DHMTHF selectivity was 6.7% and 10.1% respectively, which

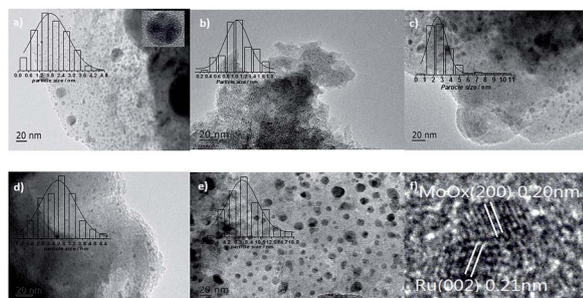
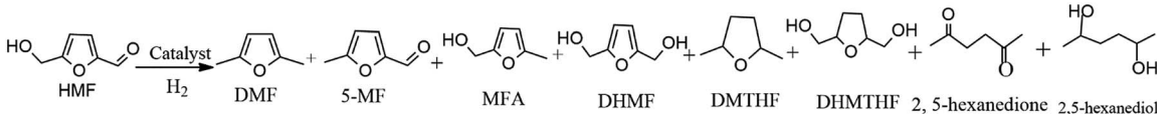


Fig. 3 TEM images of Ru– MoO_x/C catalysts: (a) reduced 5% Ru–10% MoO_x/C ; (b) reduced 5% Ru–5% MoO_x/C ; (c) reduced 5% Ru–20% MoO_x/C ; (d) reduced 5% Ru/C; (e) reduced 10% MoO_x/C and (f) HRTEM image of reduced 5% Ru–10% MoO_x/C .



Table 2 The effect of Ru/MoO_x ratio in conversion of HMF to DMF^a


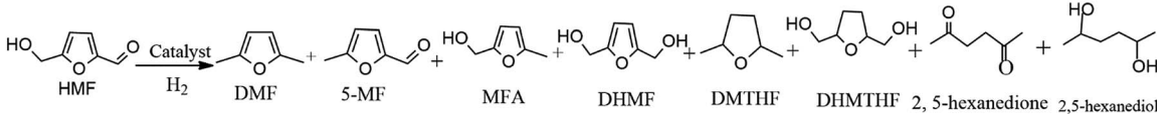
Entry	Catalyst	HMF conv. %	DMF sel. %	DMTHF sel. %	5-MF sel. %	MFA sel. %	DHMF sel. %	DHMTHF sel. %	2,5-Hexanedione sel. %	2,5-Hexanediol sel. %
1	2% Ru–10% MoO _x /C	100	60.8	3.6	—	—	—	0.2	1.4	0.7
2	5% Ru–10% MoO _x /C	100	79.8	3.4	—	—	—	0.4	0.7	1.0
3	5% Ru–5% MoO _x /C	100	71.8	2.6	—	—	—	1.1	0.4	0.9
4	5% Ru–20% MoO _x /C	100	72.6	3.1	—	—	—	0.2	1.9	0.9
5	5% Ru/C	100	54.1	6.7	—	—	—	10.1	2.2	0.5
6	10% MoO _x /C	39.9	3.9	8.5	37.5	0.4	2.6	—	6.9	11.6

^a Reaction condition: 180 °C, 1.5 MPa of H₂, 1 h, HMF 0.5 g, *n*-butyl alcohol 40 ml and 0.1 g catalyst; the Ru and Mo loading were defined by the mass fraction on support.

indicated the good hydrogenation and medium DMF selectivity for the sole Ru particle. However, if 10% MoO_x/C (Table 2, entry 6) was introduced to the reaction, the HMF conversion was only 39.8% and the DMF selectivity was 3.9%. Besides the target DMF, 37.5% of the main intermediate 5-MF was detected. Apparently, compared to Ru particle, MoO_x/C exhibited a good deoxygenation activity to crack the CH₂–OH bond in HMF, while its hydrogenation activity to activate aldehyde group (–CHO) was rather limited. Based on the above results, the synergy of Ru and MoO_x with proper ratios played a key role in obtaining high DMF yield in hydrogenolysis of HMF.

This catalytic performance could also be explained by the electronic modification between Ru and MoO_x. Compared the results of 5% Ru/C (Table 2, entry 5) with 5% Ru–10% MoO_x/C (Table 2, entry 2), the selectivity of DMTHF and DHMTHF were

6.7% and 10.1% over the 5% Ru/C catalyst, while the selectivities were decreased to 3.4% and 0.4% over 5% Ru–10% MoO_x/C catalyst. These results reflected that the hydrogenolysis ability on Ru particle was restrained when added MoO_x, which led to obtain more target DMF over Ru–MoO_x/C catalyst. The relatively weak hydrogenolysis for furan ring was attributed to the electron enrichment of Ru, which was similar to the result of hydrogenolysis of *n*-butane by Scott *et al.*⁵⁴ In the characterization section, we had interpreted the increase in the Ru electronic density due to the electronic transfer from partially reduced MoO_x to Ru. A higher electronic density over Ru increased the hydrogenolysis of the minimum electron density C–OH bond since Ru electron-donating properties.³⁹ On the other hand, the mutual electronic impact maybe was attributed to the synergistic activity. Considering the highest DMF yield

Table 3 The effect of different conditions in conversion of HMF to DMF^a


Entry	P _{H₂} (MPa)	T (°C)	t (h)	HMF conv. %	DMF sel. %	DMTHF sel. %	5-MF sel. %	MFA sel. %	DHMF sel. %	DHMTHF sel. %	2,5-Hexanedione sel. %	2,5-Hexanediol sel. %
1	1.5	120	1	97.1	21.7	0.2	3.6	11.2	36.3	0.2	0.5	0.4
2	1.5	140	1	99.8	47.9	0.8	1.0	3.5	5.0	0.4	1.5	0.6
3	1.5	160	1	100	74.4	2.1	0.1	0.8	0.1	0.6	0.8	0.8
4	1.5	180	1	100	79.8	3.4	—	—	—	0.4	0.7	1.0
5	1.5	200	1	100	69.0	9.5	—	—	—	0.4	0.3	2.3
6	1.5	120	0.25	93.2	17.8	0.8	2.2	5.4	29.4	0.1	0.3	0.3
7	1.5	120	0.5	95.6	17.4	0.5	3.4	11.3	46.6	0.3	0.4	0.4
8	1.5	120	2	99.6	46.4	0.6	1.5	5.3	13.2	0.4	1.3	0.6
9	1.5	120	4	99.7	56.1	0.6	0.6	4.7	6.8	0.5	0.6	0.7
10	1.5	120	6	99.8	50.9	0.9	0.9	5.4	6.9	1.0	0.3	0.8

^a Reaction condition: HMF 0.5 g, *n*-butyl alcohol 40 ml and 0.1 g 5% Ru–0% MoO_x/C catalyst; the Ru and Mo loading were defined by the mass fraction on support.



obtained, the 5–10% RuMo/C catalyst was selected for the following controlled experiments.

3.2.2 The pathway for DMF production from HMF. To gain insights into the reaction network, a special reaction was conducted by investigating the effect of various reaction temperature and the effect of various reaction time on products distribution at 120 °C. The intermediates were analyzed by GC and determined by GC-MS.

As shown in Table 3 (entry 1–5), when the temperature was 120 °C, only 21.7% of DMF selectivity was observed even though at the 97.1% of HMF conversion. The intermediates (5-MF, MFA and DHMF) and byproducts (DMTHF, DHMTHF, 2,5-hexanedione and 2,5-hexanediol) were detected with DHMF (36.3%) and MFA (11.2%) as the largest two. With increasing the reaction temperature from 120 °C to 180 °C, the HMF conversion and DMF selectivity were correspondingly increased from 97.1% to 100% and from 21.7% to 79.8%, respectively, which was attributed to that the intermediate products was converted to the target DMF gradually. Additionally, the over-hydrogenated byproducts were also risen but with the total amounts below 6%. However, when the temperature was further increased to 200 °C, the selectivity of DMF was decreased to 69.0%. Meanwhile, the byproducts (DMTHF, 2,5-hexanedione and 2,5-hexanediol) were significantly increased to 12.5%.

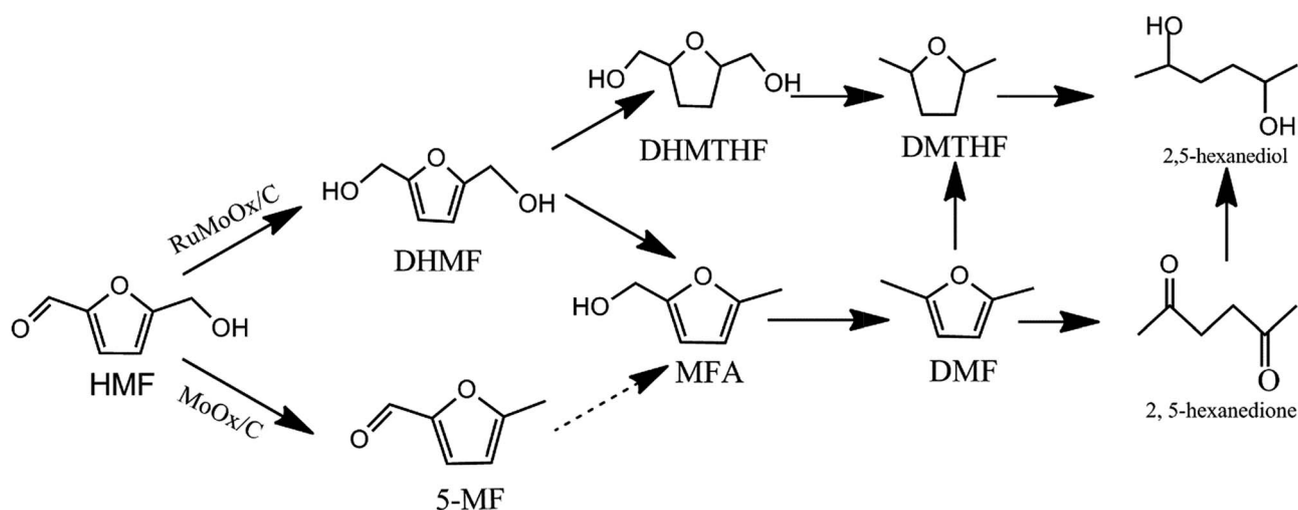
The influence of reaction time was listed in Table 3, entry 6–10. At the initial time of 0.25 h, the conversion of HMF was reached to 93.2%, nearly completely converted. However, the selectivity of the target DMF was only 17.8% and DHMF (29.4%) and MFA (5.4%) were presented as the main products. Prolonging the reaction time to 0.5 h, the conversion of HMF was increased to 95.6% and the selectivity of DMF was almost constant. While the selectivity of DHMF (46.6%) and MFA (11.3%) reached to the maximum. As the reaction time was further extended to 4 h, 56.1% of DMF was obtained while the slightly reduced MFA and DHMF intermediates were obtained. When the time was exceeded 4 h, the yield of DMF was slightly

decreased and some byproducts including DHMTHF and DMTHF were produced, which is generated by furan ring saturation *via* over-hydrogenation. The Fig. S2† showed the concentration of the products as a function of time.

Based on the above results, the aldehyde group in HMF was firstly attacked to form DHMF over the bimetallic Ru–MoO_x/C catalyst. Since DHMF could be converted to MFA and further be hydrogenated to DMF through cleavage of CH₂–OH bond, MFA was a key precursor to be converted into DMF. Interestingly, over 10% MoO_x/C catalyst (Table 2, entry 6), the result revealed that the main intermediate was 5-MF and the selectivity was 37.5%, which indicates that a different reaction pathway was observed *via* the first step of dehydroxylation in the presence of aldehyde group, comparing to the bimetallic Ru–MoO_x/C catalyst. The proposed reaction pathway was shown in Scheme 1.

According to the rules of concentration intermediates change, the catalytic step can be described as follows. Initially, Ru metal site could activated H₂ and further dissociative absorbed on the metal site. Meanwhile, the O atom of aldehyde group in HMF was activated by the MoO_x vacant orbital through Mo–O bond.⁵⁵ Firstly, activated H attacked the aldehyde group to produce DHMF, which was a key intermediate in HMF hydrolysis. Then, the MoO_x acid site accelerated dehydroxylation to generate DMF *via* MFA intermediate. After that, the DMF was desorbed and catalyst was used into the next round.¹⁹

3.2.3 Catalyst stability. In order to reflect the true activity of catalyst, 5% Ru–10% MoO_x/C was chosen to further investigate the reusability with the exception of DMF from HMF. The optimum condition (180 °C, 1.5 MPa of H₂ and 1 h) was used for the HMF hydrogenolysis to DMF. After the first run, catalyst was separated by centrifugation, washed five times with *n*-butyl alcohol, and then directly used for the next run under the same reaction condition. From Fig. 4, more than 70% of DMF selectivity and 100% of HMF conversion could be achieved after three runs, indicating that Ru–MoO_x/C catalyst was effective and stable for the HMF hydrogenolysis. After the third run, the



Scheme 1 Proposed reaction network for hydrogenolysis of HMF to DMF.



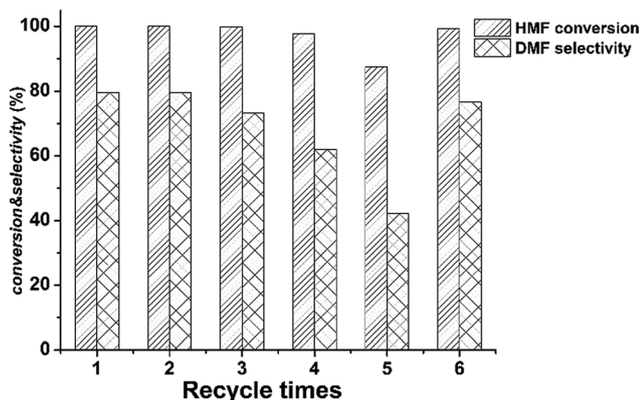


Fig. 4 Reusability of the Ru-MoO_x/C catalyst. Reaction condition: 180 °C, 1.5 MPa of H₂, 1 h, HMF 0.5 g, *n*-butyl alcohol 40 ml and 5% Ru-10% MoO_x/C catalyst 0.1 g.

selectivity of DMF decreased gradually, and for the fifth run, the selectivity of DMF decreased to 42.2%, which indicated that the catalyst deactivated during the tests. The loss of reactivity of catalyst was probably attributed to deposit of carbonaceous species like humins, which formatted during the reaction and immersed in the active sites, as notified in previous reports.^{26,56,57} To get insights into the deactivation of catalyst, the fresh and used catalysts were characterized by SEM, TEM and BET as follows. The result of SEM was presented at Fig. 5a-c. Compared to the fresh catalyst, after the first run, the size of particle had become slight larger. After five times, the diameter increased largely which was in line with the result of TEM (Table 4) and along with the agglomeration of carbon particle. From Table 4, the used sample surface area and pore volume were absorbed compounds with large molecular.

The above characterization results suggested that the deactivation of our sample maybe due to the agglomeration of carbon support and the decrease of surface area. Then the used five times catalyst was regenerated at 350 °C and the flowing H₂ (40 ml min⁻¹) for 4 h. As depicted by Fig. 4, the HMF conversion and DMF yield was regained 99.7% and 79%, indicating this catalyst could be completely recovered after regeneration. Table

Table 4 Physicochemical property of the catalysts

Catalyst	BET surface area ^a (m ² g ⁻¹)	Total pore volume ^a (cm ³ g ⁻¹)	Mean particle size diameter ^b (nm)
Fresh 5% Ru-10% MoO _x /C	447.9	0.26	1.5
Used 5% Ru-10% MoO _x /C after first run	455.6	0.29	1.7
Used 5% Ru-10% MoO _x /C after five run	194.9	0.15	2.6
Regenerated after five run	384.0	0.23	2.2

^a The BET surface area, pore volume were determined by N₂ physical adsorption. ^b Particle size was observed by TEM.

4 also revealed that the surface area and total pore volume was increased after regenerated, which was well consistent with the reaction results.

4. Conclusion

The bi-functional Ru-MoO_x/C catalysts were prepared by initial wetness impregnation and their catalytic performance was tested in DMF production from hydrogenolysis of biomass-derived HMF by finely modulating the metal-acid properties of catalysts and reaction parameters. The synergy effect between metal and acid sites plays a key role in obtaining high selectivity of DMF (79.8%). The characterizations of catalyst were interpreted the interaction and electron transfer between Ru and MoO_x and the cause of deactivation, which was attributed to the agglomeration of carbon support and the shrink of surface area of catalyst. Hydrogenation of HMF were proceeded through two pathways: for the Ru-MoO_x/C catalyst, hydrogenation of C=O group primarily occurred, followed by hydrogenolysis of two CH₂-OH bonds in DHMF to obtained the target DMF; comparatively, hydrogenolysis of CH₂-OH bonds in HMF was first taken place to 5-MF over MoO_x/C catalyst.

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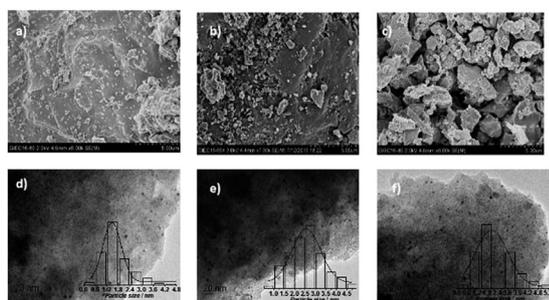


Fig. 5 SEM images of (a) the fresh 5% Ru-10% MoO_x/C catalyst, (b) the used of 5% Ru-10% MoO_x/C catalyst after first run, (c) the used of 5% Ru-10% MoO_x/C catalyst after five runs. And TEM image of (d) the used of 5% Ru-10% MoO_x/C catalyst after first run, (e) the used of 5% Ru-10% MoO_x/C catalyst after five runs and (f) the regenerated of 5% Ru-10% MoO_x/C catalyst after five runs.



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