RSC Advances



View Article Online

View Journal | View Issue

PAPER

Check for updates

Cite this: RSC Adv., 2021, 11, 48

Received 29th October 2020 Accepted 11th December 2020

DOI: 10.1039/d0ra09217g

rsc.li/rsc-advances

1. Introduction

Methyl isobutyl ketone (MIBK) is one of the most important chemical products widely used in the coatings, pharmaceutical, and petrochemical industries.¹ Methyl isobutyl methanol (MIBC), synthesized *via* MIBK hydrogenation, is an excellent medium-boiling solvent and used as a raw material for organic synthesis and mineral floaters. With the development of the global economy, the demand for MIBK and MIBC in Asia, especially China and South Korea, will increase further, due to the pull of rubber antioxidant 4020 and high-solid coatings.

At present, MIBK is mainly manufactured by a one-pot process, composed of three steps, say, acetone condensation catalyzed by acid or base to diacetone alcohol (DAA), DAA dehydration catalyzed by acid to mesityl oxide (MO), and MO hydrogenation catalyzed by Pd to MIBK.^{2,3} Normally, MIBC could also be generated in this one-pot process. To date, the only commercialized multifunctional catalyst is Pd/resin, which possess both the acid and Pd sites, which delivered an acetone conversion of 25–50% with a MIBK selectivity of 70–

^bCangzhou Dahua Group Company, Ltd, Cangzhou 061000, China

An active and stable multifunctional catalyst with defective UiO-66 as a support for Pd over the continuous catalytic conversion of acetone and hydrogen[†]

Yingjie Hu,^a Yuxin Mei,^a Baining Lin,^{*a} Xuhong Du,^a Fan Xu,^a Huasheng Xie,^b Kang Wang^b and Yonghua Zhou^b*^a

The one-pot synthesis of methyl isobutyl ketone (MIBK) and methyl isobutyl methanol (MIBC) from acetone and hydrogen is a typical cascade reaction comprised of aldol condensation-dehydration-hydrogenation. Pd loss and aggregation during long term operation are typical problems in industrial application. In this paper, an active and stable catalyst was achieved with defective UiO-66 as a support for Pd, which was synthesized with the ratio 15 : 1 of ZrOCl₂·8H₂O to ZrCl₄ as Zr-precursors. The resultant Pd catalyst remained active for at least 1000 h with a MIBK + MIBC selectivity of 84.87–93.09% and acetone conversion of 45.26–53.22% in a continuous trickle-bed reactor. Besides the increased Brønsted acid amount generated by the defect sites was favorable for the activity, the cavity confinement in the UiO-66 (R = 15 : 1) structure also efficiently prevented Pd loss and aggregation during the long term run. The contrast of the characterization of the fresh and used Pd/UiO-66 (R = 15 : 1) indicated that the deactivation of the catalyst was attributed to carbonaceous accumulation on the catalyst surface, which could be easily regenerated by calcination. This work supplied a new alternative for the design and utilization of industrial catalysts for MIBK and MIBC synthesis.

90% at 1–10 MPa and 120–150 °C.^{4–7} Nevertheless, due to the poor thermal stability of the resin, the Pd loss and aggregation during long term operation is inevitable, which leads to the deactivation of catalyst. Aiming at solving this problem, although lots of bifunctional catalysts, including Pd/metal oxide,^{8,9} Pd/molecular sieve and Pd/resin,^{10,11} Ni-based,^{12,13} Cu-based,¹⁴ and Pt-based,¹⁵ have been reported during the recent decades, the high active and stable catalyst is still desirable.

Metal-organic frameworks (MOF), in particular Zr-based MOF,¹⁶⁻¹⁸ have been extensively investigated for acid catalyzed reactions such as citronellal cyclization,19 esterification,20 transfer hydrogenation,²¹ and cross-aldol condensation.²² Besides the acid property, the cage structure of MOF also allows it to act as the excellent support for metal nanoparticles.23-25 UiO-66, as a kind of traditional Zr-based MOF, contains metal nodes composed of a zirconium oxide complex bridged by terepthalic acid ligands and possesses the high thermal stability.26 In addition, it is well recognized that the defect sites of UiO-66, mainly the missing linkers, correlates closely with the catalytic performance, since the acid-base property arises mainly from the defect sites. Therefore, to achieve the high performance of UiO-66 in catalysis, more defects are desirable to be created. In common, the defects in UiO-66 can be tailored via adjusting the synthesis conditions, including incorporation of modulators,27-29 thermal activation/dehydration,30 metal cation substitution³¹ and linker modification.³²

^aCollege of Chemistry and Chemical Engineering, Central South University, Changsha 410083, China. E-mail: baininglin@csu.edu.cn; zhouyonghua@csu.edu.cn

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/d0ra09217g

Paper

In this paper, the utilizing of the mixture Zr-precursor of $\text{ZrOCl}_2 \cdot 8\text{H}_2\text{O}$ and ZrCl_4 was firstly developed to create defect-rich UiO-66 and then Pd was loaded thereon to fabricate the multifunctional catalyst for acetone hydrogenation reaction. As expected, the catalytic performance results exhibited that a high active and stable catalyst was achieved, with the at least 1000 h stability with MIBK + MIBC selectivity of 84.87–93.07% and acetone conversion of 45.26–53.22%.

2. Experimental

2.1 Materials

All chemicals and reagents were obtained from commercial sources and used as received, unless otherwise stated. Specifically, zirconium chloride ($ZrCl_4$), zirconyl chloride octahydrate ($ZrOCl_2$ - \cdot 8H₂O), glacial acetic acid (HAc), palladium chloride (PdCl₂) and 1,4-benzenedicarboxylic acid (BDC) were purchased from Aladdin Reagent (Shanghai) Company of China, and anhydrous *N*,*N*-dimethylformamide (DMF) sodium chloride (NaCl₂) and methanol was purchased from Sinopharm Chemical Reagent Co., Ltd.

2.2 Catalyst preparation

UiO-66 was prepared following a procedure previously reported³³ with slight modifications. In a typical synthetic process, 3.28 g of ZrCl₄ and 2.32 g of BDC were first added in 160 mL of DMF. Then, 1 mL of deionized water and 24 mL of HAc were successively dropped into the mixture and stirring was initiated until all precursors were completely dissolved. The obtained mixture was transferred to a 500 mL of hydrothermal reaction vessel with Teflon liner, and kept at 120 °C for 24 h under static conditions. After cooling to room temperature, the solid MOF were collected *via* centrifugation and thoroughly washed with a mixture of methanol and DMF (v/v = 1 : 4) for three times, and methanol for two times. Finally, the obtained sample was calcined in N₂ atmosphere under 180 °C for 3 h and denoted by UiO-66.

Synthesis of UiO-66 (R = x : y). The synthesis technique of UiO-66 (R = x : y) was similar to that of UiO-66, except replacing ZrCl₄ with the mixture Zr-precursor. Typically, mix ZrOCl₂··8H₂O and ZrCl₄ in a mass ratio of 0 : 1, 1 : 1, 3 : 1, 7 : 1, 9 : 1, 15 : 1, 19 : 1 and 1 : 0 respectively. The obtained samples were denoted by UiO-66 (R = x : y), and x : y represented the mass ratio of ZrOCl₂·8H₂O to ZrCl₄.

Synthesis of Pd/UiO-66 and Pd/UiO-66 (R = x : y) catalysts. The Pd catalysts were prepared by a wet impregnation technique. At room temperature, UiO-66 or UiO-66 (R = x : y) was added into the Na₂PdCl₄ aqueous solution and the mixture was subsequently stirred for 4 h, followed by reduction by a KBH₄ solution for 30 min and washing with ethanol aqueous solution (volume ratio 1 : 1) to neutral. Eventually, vacuum drying at 50 °C was carried out. The obtained samples were denoted by Pd/UiO-66 and Pd/UiO-66 (R = x : y) catalysts, respectively.

2.3 Catalyst characterizations

Fourier transform infrared (FT-IR) spectra was measured on a Nicolet 6700 Fourier transform spectrometer (Nicolet Magana Co.). Scanning electron microscopy (SEM) images was recorded

on JSM-7610FPlus. A structural characterization was obtained by X-ray diffraction (XRD). XRD patterns of catalysts were recorded between 5° and 50° (2 θ) on a Bruker D8 Advance A25 diffractometer with monochromatic Cu K α radiation ($\lambda = 0.154$ nm). Thermogravimetric (TG) analysis was carried out in a TA Instruments TGA Q500 at a heating rate of 10 °C min⁻¹ to 800 °C in air. The measurements of the acidity and basicity of the catalysts were accomplished by Autosorb-iQ-C equipped with a thermal conductivity detector. The sample was pretreated in helium gas at 400 °C for 2 h and then cooled down to 50 °C. The surface of the materials was saturated with NH₃ (or CO₂) for 60 min. TPD profile of NH₃ (or CO₂) was recorded at the rate of 10 $^{\circ}$ C min⁻¹ from 50 °C to 800 °C by a He flow. Pd mass content was measured with a PerkinElmer Inductively Coupled Optical Emission Spectrometer (ICP), model Optima 5300DV, equipped with a peristaltic pump and across-flow nebulizer, coupled to a Ryton double pass spray chamber of the Scott type. Before analysis, Pd/UiO-66 (R =x : y) was immersed in aqua regia solution (v(HCl + HNO₃)/vH₂O = 20:30) to dissolve Pd into solution. The CO chemisorption measurements were performed in MICROMERITICS AutoChem II 2920 equipment. Before analysis, the catalysts were reduced under 10% H_2/Ar flow at 200–250 $^\circ C$ for 3 h, and subsequently under He (50 mL min⁻¹) flow at 150 °C for 1 h to blow away H_2 adsorbed on the surface of catalysts. Then, CO chemisorption analysis using double isotherm methodology was performed at 35 °C and metal dispersion (or the particle size) was calculated. Potentiometric acid-base titrations were completed with a Metrohm Titrando 905 autotitrator equipped with Dosino 800 20 and 10 mL dosing units. X-ray photoelectron spectroscopy (XPS) was performed using an ESCALAB250Xi instrument with an Al Ka Xray source (1486.6 eV), under about 2 \times 10⁻⁹ mbar at room temperature and a pass energy of 20 eV.

2.4 Catalytic testing

Acetone hydrogenation reaction was performed in a batch reactor with an inner volume of 40 mL. The reactor was equipped with a mechanically driven agitator, thermocouples, and inlet and outlet gas purge lines. Typically, 5 mL of acetone and experimentally desired amount of catalyst were added to the reactor chamber and mixed. The reactor was tightly sealed and purged three times with H_2 to remove air. Then the reactor was pressurized with H_2 at 0.75 MPa, and the reaction proceeded at various temperatures and time. After the reaction finished, cold-water quenching was applied to the reactor to stop the reaction. The liquid product was separated from the solid residue (if formed) and the catalyst by filtration.

The performance of acetone hydrogenation was also tested in a continuous trickle-bed reactor using 6 mL catalyst. The reaction conditions, reaction temperature, hydrogen pressure, liquid hour space velocity (LHSV), and feed ratio of acetone to H₂, were optimized. The long-term run of the catalyst in the trickle-bed reactor was performed under the following conditions: 0.18 mL min⁻¹ of acetone flow, 12 mL min⁻¹ of hydrogen flow, 120 °C, 2.0 MPa and LHSV of 1.8 h⁻¹. A Shimadzu GC-2010 gas chromatograph (GC) equipped with a flame ionization detector (FID) and a 30 m × 0.25 mm × 0.25 µm Insert Cap WAX capillary column was used to determine the amounts of acetone, MIBK, MIBC, MO, DAA, and others in the product. Total analysis time was 20 min. The definition of acetone conversion (x_A), selectivity to MIBK + MIBK (S_{BK+BC}) and MIBK + MIBC yield (Y_{BK+BC}) were calculated according to the following formulas.

$$x_{\rm A} = 1 - \frac{\text{moles of acetone in the product}}{\text{moles of acetone in the feed}} \times 100$$

$$S_{\rm BK+BC} = {{\rm moles \ of \ acetone \ converted \ to \ BK + BC} \over {\rm moles \ of \ acetone \ converted}} \times 100$$

$$Y_{\rm BK+BC} = x_{\rm A} \times S_{\rm BK+BC} / 100$$

3. Results and discussion

3.1 Catalytic performance

The catalytic performance of the Pd catalysts loaded on various UiO-66 (R = x:y) for acetone hydrogenation in the batch reactor are shown in Table 1. It is observed that Pd/UiO-66 delivers an acetone conversion of 25.31%, and a MIBK + MIBC selectivity of 94.00%. Comparatively, Pd/UiO-66 (R = 1:0) exhibits an

acetone conversion of 34.00%, and a MIBK + MIBC selectivity of 92.72%. This indicates that both UiO-66 and UiO-66 (R = 1 : 0) possess the acid–base property for acetone condensation. By increasing the ratio of $\text{ZrOCl}_2 \cdot 8\text{H}_2\text{O}$ to ZrCl_4 in the raw material from 1 : 1 to 19 : 1, the acetone conversion of Pd/UiO-66 (R = x : y) catalyst increases firstly and then drops gradually. In addition, the selectivity increases at the beginning and then varies little. Among them, Pd/UiO-66 (R = 15 : 1) exhibits the highest MIBK + MIBC yield of 51.95% with an acetone conversion of 54.44% and a MIBK + MIBC selectivity of 95.38%.

The catalytic performances of Pd/UiO-66 (R = 15:1) catalysts with different palladium loading are shown in Table 2. UiO-66 (R = 15:1) support itself shows the acetone conversion of 29.51%, the MIBK + MIBC selectivity of 9.72%, and MO selectivity of 68.13%. This indicates that UiO-66 (R = 15:1) support itself indeed only has acid-base function, without hydrogenation function. When Pd loading is higher than 0.3 wt%, MO is almost completely converted. With the increase of the amount of Pd, there is a climax of the MIBK + MIBC yield. It is noted that too much Pd is not beneficial to the MIBK + MIBC vield due to the decreased acetone conversion, probably due to the coverage of the acid-base sites by Pd. Therefore, 0.5 wt% Pd/UiO-66 (R =15:1) catalyst is chosen here for further investigation in the trickle-bed reactor. We compare the performance of Pd/UiO-66 (R = 15:1) with those reported in a batch liquid phase process as shown in Table S1.[†] It is seen that Pd/UiO-66 (R = 15:1)

Table 1 Effect of $ZrOCl_2 \cdot 8H_2O/ZrCl_4$ ratio on the catalytic performance in the batch reactor^a

Catalyst	Acetone conversion (%)	Selectivit						
		MIBK	MIBC	МО	DAA	IPA	Others	MIBK + MIBC yield (%)
Pd/UiO-66	25.31	92.67	1.33	0.31	2.77	1.52	1.40	23.79
Pd/UiO-66 (R = 1:1)	11.18	85.38	0.46	0.10	7.54	5.35	1.17	9.59
Pd/UiO-66 (R = 3:1)	20.46	87.54	0.89	0.81	3.08	4.80	2.88	18.09
Pd/UiO-66 (R = 7:1)	37.12	91.32	2.93	0.93	1.24	1.45	2.13	34.98
Pd/UiO-66 (R = 9:1)	39.02	91.96	2.56	1.04	1.35	1.26	1.83	36.88
Pd/UiO-66 (R = 15:1)	54.44	90.90	4.48	0.64	0.56	0.31	3.11	51.95
Pd/UiO-66 (R = 19:1)	34.64	90.88	1.96	0.93	1.88	1.50	2.85	32.16
Pd/UiO-66 ($R = 1:0$)	34.00	90.82	1.90	0.65	1.80	3.00	1.83	31.52

 a T = 120 °C, reaction time = 2 h. IPA is the abbreviation of isopropanol. Others is mainly C₉⁺. 0.5 wt% Pd loading.

Table 2 Effect of Pd loading on the catalytic performance of Pd/UiO-66 (R = 15:1) in the batch reactor^a

	Acetone conversion (%)	Selectivit						
Catalyst		MIBK	MIBC	МО	DAA	IPA	Others	MIBK + MIBC yield (%)
UiO-66 ($R = 15:1$)	29.50	9.51	0.21	68.13	2.41	0.32	19.42	2.87
0.3wt% Pd/UiO-66 (R = 15:1)	47.07	80.95	3.37	6.94	0.80	0.24	7.70	39.69
0.4wt% Pd/UiO-66 ($R = 15:1$)	53.64	87.60	4.63	2.59	0.77	0.07	4.34	46.99
0.5wt% Pd/UiO-66 (R = 15:1)	54.44	90.90	4.48	0.64	0.56	0.31	3.11	51.95
0.6wt% Pd/UiO-66 (R = 15:1)	52.67	88.60	4.86	1.07	0.67	0.28	4.52	49.22
0.7 wt% Pd/UiO-66 (R = 15:1)	52.22	90.71	2.52	1.91	0.70	0.34	3.82	48.68
0.8wt% Pd/UiO-66 (R = 15:1)	38.62	90.72	2.54	2.11	1.34	0.23	3.06	36.02
0.9wt% Pd/UiO-66 $(R = 15:1)$	31.71	90.60	2.03	2.21	1.34	1.47	2.35	29.37

^{*a*} T = 120 °C, reaction time = 2 h. IPA is the abbreviation of isopropanol. Others is mainly C₉⁺.

delivers the best performance among them under the mild reaction conditions (Table S1[†]).^{34–37}

The reaction conditions, including the reaction temperature, hydrogen pressure, LHSV and mole ratio of C₃H₆O/H₂, over 0.5 wt% Pd/UiO-66 (R = 15:1) were optimized via the orthogonal experiments with four factors and four levels. The effects of these factors on the catalytic performance are shown in Tables S2 and S3.[†] It is seen that as for 0.5 wt% Pd/UiO-66 (R = 15:1) operated in a trickle-bed reactor, the reaction temperature is the most significant one among the four factors. More experiment results in Table 3 indicate that the increase of reaction temperature from 80 °C up to 160 °C leads to an increase of the acetone conversion from 32.95% to 56.89%, accompanied by a volcanic-changed MIBK selectivity. The decline of MIBK selectivity at higher temperature is due to the formation of heavier condensation products (C_9^+) .³ In addition, it is noted that more MIBK is hydrogenated further to MIBC with the increase of temperature. Except the reaction temperature, the mole ratio of C₃H₆O/H₂ is the second significant factor on the catalyst performance. Considering the yield of MIBK + MIBC, the mole ratio of C_3H_6O/H_2 should be maintained at 1:2, consistent with the theoretic value. Therefore, the optimal reaction conditions of Pd/UiO-66 (R = 15:1) catalyst is reaction temperature of 120 °C, hydrogen pressure of 2 MPa, LHSV of 1.8 h^{-1} and mole ratio of C_3H_6O/H_2 of 1:2, which deliver the highest MIBK + MIBC yield of 51.95%.

The long-term stability of 0.5 wt% Pd/UiO-66 (R = 15:1) catalyst in the trickle-bed reactor under the optimal conditions was investigated (Fig. 1). After 1000 h run, the acetone conversion drops by 7.96% from 53.22% to 45.26%, and the MIBK + MIBC selectivity drops by 8.22% from 93.09% to 84.87%. Zhu⁴ reported that Pd/S (PS-GO) catalyst was tested for 1000 h and the acetone conversion dropped from 50.85% to 39.71% by 10.14% and the MIBK selectivity dropped from 77.86% to 73.47% by 4.39%. Duan³ showed that the TiO₂–SiO₂/SiO₂ (F)-1&Pd/Cor catalyst was active for 300 h with a MIBK selectivity of 83–93% at 26–36% acetone conversion. Comparatively, the catalyst in this paper displays the better stability than those reported in literature, presenting the prospect for industrial application.



Fig. 1 Stability of 0.5 wt% Pd/UiO-66 (R = 15 : 1) catalyst. Reaction conditions: volume of catalyst = 6 mL, T = 120 °C, P = 2.0 MPa, H₂ flow rate = 12 mL min⁻¹ and LHSV = 1.8 h⁻¹.

3.2 Characterizations of catalysts

The XRD patterns of UiO-66, UiO-66 (R = 15:1) and Pd/UiO-66 (R = 15:1) in Fig. 2 exhibit the characteristic peaks of the typical UiO-66 structure.¹⁶ No additional peaks can be observed, indicating that no impure crystalline phases are formed. These suggest that the variation of the Zr precursors and the Pd loading process have no influence on the bulk structure of UiO-66. In addition, it is noted that no characteristic peak of Pd at $2\theta = 40.1^{\circ}$ and 46.7° can be detected due to the quite low loading amount of Pd.

The SEM images of UiO-66 and UiO-66 (R = 15:1) are shown in Fig. 3. Obviously, two samples display different morphology. The UiO-66 granules display the cubic octahedron with the edge length size of around 150 nm (Fig. 3a and b). Comparatively, UiO-66 (R = 15:1) sample exhibits the amorphous morphology composed of flaky granules (Fig. 3c and d). It is noteworthy that although the macroscopic morphology of UiO-66 and UiO-66 (R = 15:1) seems quite different, both of them possess the same crystal structure revealed by XRD. It is known that the

Table 3	Catalytic performance of 0.5 wt% Pd/UiO-66 ($R = 15$: 1) at different temperature in the trickle-bed reactor ^a

Temperature (°C)	Acetone conversion (%)	Selectivity						
		MIBK	MIBC	МО	DAA	IPA	Others	MIBK + MIBC yield (%)
80	32.95	87.03	1.47	5.70	3.38	0.12	2.30	29.16
90	35.13	86.11	1.60	6.95	2.48	0.11	2.75	30.81
100	40.43	87.08	2.89	4.80	1.18	0.52	3.53	36.37
110	49.13	91.63	3.72	0.63	1.11	0.61	2.30	46.84
120	54.44	90.90	4.48	0.64	0.56	0.31	3.11	51.95
130	54.59	89.11	4.53	0.62	0.52	1.68	3.54	50.11
140	54.53	87.69	4.55	0.59	0.49	2.87	3.81	50.30
150	55.15	85.43	4.91	0.57	0.40	3.40	5.29	48.82
160	56.89	82.69	4.97	0.56	0.41	3.87	7.50	49.87

^{*a*} The catalyst volume is 6 mL, P = 2.0 MPa, H₂ flow rate = 12 mL min⁻¹, and time on stream = 2 h. IPA is the abbreviation of isopropanol. Others is mainly C₉⁺.



Fig. 2 XRD patterns of UiO-66, UiO-66 (R = 15: 1) and Pd/UiO-66 (R $= 15 \cdot 1$

crystalline process could be accelerated and small UiO-66 particles with several nanometers were formed when using ZrOCl₂·8H₂O as Zr-precursor.³⁷ Therefore, in this paper, the particle size of UiO-66 (R = 15:1) is much smaller than UiO-66. This is beneficial for the acetone hydrogenation reaction due to the readily accessibility of active sites on UiO-66 (R = 15:1).

The BET specific surface area and the porous structure of UiO-66 and UiO-66 (R = 15:1) are evaluated by N₂ adsorptiondesorption experiment (Fig. 4 and Table S4[†]). UiO-66 exhibits the type I isotherm in the region of low relative pressure, indicating the main existence of micropores. The BET specific surface area and the total pore volume are estimated to be 1677 m² g⁻¹ and 0.88 cm³ g⁻¹, respectively. The narrow distribution of the pore size justifies that the micropores come from the intrinsic structure of UiO-66. According to literature, the theoretical pore volume of UiO-66 is 0.77 $\text{cm}^3 \text{g}^{-1}$ and the surface area is 1160 m² g⁻¹.^{17,33,38} Comparatively, UiO-66 (R = 15:1) displays the type II isotherm with the hysteresis loop,



Fig. 4 N₂ adsorption-desorption isotherms and pore distribution curves of UiO-66 and UiO-66 (R = 15 : 1).

suggesting the main existence of mesopores, which probably are the secondary particle piled pores of formed by flaky granules. This was further confirmed by the pore size distribution calculated from the N₂ isotherms. Accordingly, the BET specific surface area of UiO-66 (R = 15:1) is 1068 m² g⁻¹, the total pore volume is 1.03 $\text{cm}^3 \text{g}^{-1}$ and the proportion of mesopores volume increases.

The thermal stability of UiO-66 and UiO-66 (R = 15:1)determined by TG measurement is shown in Fig. 5. The TG curve of UiO-66 reveal three broad areas representing weight loss. The first area is linked to the removal of water between 30 °C and 100 °C. And the second one is caused by the removal of the residual solvent N,N-dimethylformamide and/or of the unreacted terephthalic acid between 100 °C and 480 °C. The third one is assigned to the decomposition of the framework between 400 °C and 600 °C. The final solid product after thermal treatment in air is 6ZrO2.41 It is observed from Fig. 5 that UiO-66 and UiO-66 (R = 15:1) have almost the same TG profiles, suggesting their similar thermal stability.

The FT-IR spectra of UiO-66 and UiO-66 (R = 15 : 1) in Fig. 6 show that both samples have the characteristic peaks of UiO-66.



Fig. 3 SEM images of UiO-66 (a and b) and UiO-66 (R = 15: 1) (c and d).



Fig. 5 TG curves of UiO-66 and UiO-66 (R = 15: 1) in air.



Fig. 6 FT-IR spectra of UiO-66 and UiO-66 (R = 15 : 1). (a) Monodentate, (b) bidentate, (c) bridging.

The sharp peak at around 1400 cm⁻¹ represents the symmetric stretching of the O-C-O carboxylate group of BDC ligands, and the asymmetric stretching vibration is located at about 1587 cm⁻¹. The peak at 1670 cm⁻¹ is assigned to the stretching vibrations of C=O of the BDC ligands. The peak at 1503 cm⁻¹ represents the C=C vibrations of benzene rings.16 At the lower range of wavenumbers, the strong peaks at 848 cm⁻¹, 745 cm⁻¹, and 656 cm⁻¹ are attributed to the vibrations of O-H and C-H bonds in the BDC ligands. These peaks ranged in 848–656 cm⁻¹ overlap with the Zr-O modes. In addition, the gap between the two peaks of around 1400 cm⁻¹ and 1500–1700 cm⁻¹ can give us the information about the coordination modes of BDC and Zr atoms.^{39,40} When the gap is larger than 200 cm⁻¹, BDC acts as a monodentate ligand to Zr atoms. And the gap in the range of 130–200 cm⁻¹ indicates that BDC behaves as a bridging ligand. Alternatively, when the gap is in the range of $50-150 \text{ cm}^{-1}$, BDC acts as a bidentate ligand to Zr atoms.^{42,43} Fig. 6 shows that there is a strong splitting of 187 cm^{-1} with a weak one of 103 cm^{-1} for UiO-66 and UiO-66 (R = 15:1). This indicates that the coordination modes of BDC in UiO-66 and UiO-66 (R = 15:1) are mainly consisted of bridging ligands with a small amount of bidentate ligands.

The defects sites in the MOF materials could be identified by acid-base titration method.44,45 The acid-base titration curves of UiO-66 and UiO-66 (R = 15:1) are measured as shown in Fig. 7. To more accurately estimate the equivalence points in the titration curves, the first derivative curves are fit to Lorentziantype peaks. It is observed that both samples display the titration curves with three apparent inflection points, which correspond to three types of protons (µ₃-OH, Zr-OH₂, and Zr-OH).^{46,47} Among them, Zr-OH₂ and Zr-OH sites are generated as the defects sites by missing linker in the synthesis process. Therefore, the missing linker number can be further calculated from Fig. 7. The calculation process is shown in Tables S5 and S6.† And the missing linker numbers of UiO-66 and UiO-66 (R =15:1) are estimated to be 1.58 and 1.89, respectively. This suggests that more defect sites are introduced into UiO-66 (R =15:1) than UiO-66 via utilizing the precursor mixture of ZrCl₄



Fig. 7 Acid-base titration curve and first derivative curve for (a) UiO-66 and (b) UiO-66 (R = 15 : 1).

and $\text{ZrOCl}_2 \cdot 8\text{H}_2\text{O}$. In addition, according to the calculation results, the percentage of Zr–OH in defects increases from 65.9% to 83.4% in UiO-66 (R = 15:1) compared with UiO-66.

To correlate the nature of defect sites of UiO-66 and UiO-66 (R = 15:1) and the acid-base property required for acetone condensation, the NH₃-TPD and CO₂-TPD were carried out as shown in Fig. 8. It is well known that both Zr–OH and Zr–OH₂ sites belong to typical Brønsted acid.⁴⁸ The 6-fold-coordinated Zr sites on the Zr6 nodes of UiO-66 are Lewis acid. In Fig. 8a, the broad peak can be deconvoluted into three peaks. The peak centered at about 290 °C can be attributed to the Lewis acid of Zr sites, and the other two peaks can be assigned to the Brønsted acid sites of Zr–OH and Zr–OH₂. It is noted that the amount of Brønsted acid sites in UiO-66 (R = 15:1) is obviously increased, while the strength of Lewis acid sites is almost constant compared with UiO-66. It was previously reported that the change of specific coordination number of Zr has a minor effect on the Lewis acid strength of the active site.^{49–52}

As for base sites, the μ_3 -O on the Zr6 nodes was normally regarded as the Lewis base sites. When the missing linker number is higher, the unsaturated coordinative number of Zr atoms becomes higher and leads to the shortened length of Zr– O bond, which further results in the weakening of the Lewis



Fig. 8 NH₃-TPD (a) and CO₂-TPD (b) profiles of UiO-66 and UiO-66 (R = 15:1)

basicity. This can be reflected by the peak shift to lower temperature from 410 °C of UiO-66 to 355 °C of UiO-66 (R =15:1) in the CO₂-TPD profiles in Fig. 8b. In addition, the other broad peak shifting to lower temperature could be attributed to other kinds of basic sites related to oxygen in the framework.

In a word, compared with UiO-66, UiO-66 (R = 15:1)exhibits the obviously increased amount of Brønsted acid sites, mainly Zr-OH and Zr-OH₂ sites, caused by the increasing of missing linker number. For acetone condensation and dehydration reactions, except for base site, it is intensively reported that Brønsted acid is the typical catalytic site for the reaction, for instance sulfonate ion.⁵ In this paper, although the basicity of UiO-66 (R = 15:1) is weakened, the enhanced Brønsted acid amount leads to the enhancement of the cascade reaction.

The comprehensive contrast of the characterizations of UiO-66 and UiO-66 (R = 15:1) has been presented in the above text, which help us to understand the reason for the better performance of Pd/UiO-66 (R = 15:1). To further reveal the nature of the activity drop of Pd/UiO-66 (R = 15:1) after 1000 h run, the TG curves, FT-IR spectra, NH₃-TPD and CO₂-TPD profiles of the fresh and used catalysts are presented as follows. From the contrast of the TG curves of the fresh and used Pd/UiO-66 (R =15:1) in Fig. 9, it is seen that an increased weight loss 26% ranged in 100-500 °C is observed, suggesting the existence of the carbonaceous accumulation on the used catalyst. Except



Fig. 9 TG curves of Pd/UiO-66 (R = 15: 1) before and after 1000 h reaction in air

this, the crystal structure of UiO-66 (R = 15:1) has not changed even after 1000 h run, since the decomposition temperature of the framework at 500 °C is the same as the fresh one. In addition, the structure stability of UiO-66 (R = 15:1) can also been identified by the contrast of FT-IR spectra in Fig. S1.† The change of the acid-base property of fresh and used Pd/UiO-66 (R = 15:1) is revealed by NH₃-TPD and CO₂-TPD profiles in Fig. S2.† After 1000 h run, the Brønsted acid amount of the used Pd/UiO-66 (R = 15:1) decreases, and the acidity and basicity is strengthened as shown in Fig. S2b.[†] About the reason for the change of acidity and basicity, we assume that the long chain alcohol produced during the reaction may be co-adsorbed by the adjacent zirconium atoms, leading to the reduction of the acid sites. Then, the alcohol further deprotonate to form alkoxide, and the proton is transferred to the μ_3 oxygen to strengthen its basicity. Hence, compared with the fresh catalyst, the CO₂ desorption peak on the used Pd/UiO-66 (R = 15:1) shifts to a higher temperature.

In addition, the surface chemical state of Pd, Zr and O on Pd/ UiO-66 and Pd/UiO-66 (R = 15:1) was characterized by XPS as shown in Fig. S3.† The existence of C, O, Pd and Zr elements on Pd/UiO-66 and Pd/UiO-66 (R = 15:1) is identified in Fig. S3(a).† The Pd 3d and Zr 3p spectra are present in Fig. S3(b).[†] The two strong peaks in Fig. S3(b)^{\dagger} are assigned to Zr 3p_{3/2} and Zr 3p_{1/2}, respectively.⁵³ And the two weak peaks at 335.6 and 341.1 eV can be attributed to Pd 3d_{5/2} and Pd 3d_{3/2}, respectively, which are partially overlapped by the two peaks of Zr 3p. The similar XPS results were previously reported by Guan⁵³ and Yang.⁵⁴ It is observed in Fig. S3(b)[†] that the binding energy of Pd species on both samples is slightly higher than the standard value of 335.3 eV of Pd metal, probably attributed to the interaction between Pd and the UiO-66 framework.54 As for Zr 3d and O1s spectra in Fig. S3(c and d),† no difference is observed between Pd/UiO-66 and Pd/UiO-66 (R = 15:1).

Furthermore, the Pd mass percent and Pd dispersion of fresh Pd/UiO-66, fresh and used Pd/UiO-66 (R = 15:1) were characterized by ICP and CO chemisorption as shown in Table S7.†

Paper

The two samples present the close Pd mass percent. And the average particle size of Pd on Pd/UiO-66 is estimated to be 3.05 nm, slightly larger than that of 2.20 nm on Pd/UiO-66 (R = 15:1). Considering the contrast of the Pd average particle size and the surface chemical state between Pd/UiO-66 and Pd/UiO-66 (R = 15:1), we can regard that there is no obvious difference in view of the Pd particles loaded thereon. In addition, the condensation of acetone is normally regarded as the rate-determining step, instead of the hydrogenation step, for the overall cascade reactions. Therefore, the gap of catalytic performance between Pd/UiO-66 and Pd/UiO-66 (R = 15:1) can be mainly ascribed to the difference of acid and base sites.

As for the fresh Pd/UiO-66 (R = 15:1), the Pd mass percent is 0.486% with an average Pd particle size of 2.20 nm. Comparatively, as for the used Pd/UiO-66 (R = 15:1), the Pd mass percent drops to 0.442% (by 9.05%) accompanied by the slight aggregation of Pd particle to 2.73 nm. The change of the Pd mass and particle size is quite smaller than those reported in literature,^{3,4} This suggests that the phenomena of metal loss and aggregation of Pd particles can be efficiently inhibited by loading Pd in the small cavities of UiO-66 (R = 15:1), resulting in the excellent stability in long term run. Therefore, the reason for the deactivation of Pd/UiO-66 (R = 15:1) can be mainly attributed to the carbonaceous accumulation, instead of the Pd loss and aggregation. The used Pd/UiO-66 (R = 15:1) catalyst after 1000 h run was regenerated in our experiments. It was calcined in air flow with 10 mL min⁻¹ of at 150 °C for 1 h followed by reduced in H_2 flow with 10 mL min⁻¹ at 150 °C for 1 h. Then, the catalytic performance of the regenerated catalyst was tested again under the same reaction conditions. The regenerated catalyst exhibited an acetone conversion of 53.23% and a MIBK + MIBC selectivity of 94.19%, close to the initial performance. This indicates the deactivation of Pd/UiO-66 (R =15:1) is reversible and can be readily regenerated by calcination. This is valuable for the prolongation of the service life of Pd catalyst.

4. Conclusions

The cage structure, high thermal stability and the adjustability of acid-base property of UiO-66 allowed it to act as an excellent candidate to build multifunctional catalysts. In this paper, the defective UiO-66 was synthesized with the optimal ratio 15:1 of ZrOCl₂·8H₂O to ZrCl₄ as Zr-precursors firstly. Then, Pd was loaded thereon and tested in a typical cascade reaction, the onepot synthesis of methyl isobutyl ketone and methyl isobutyl methanol from acetone and hydrogen. The Pd/UiO-66 (R =15:1) catalyst exhibited a stable run for at least 1000 h with a MIBK + MIBC selectivity of 84.87-93.09% and acetone conversion of 45.26-53.22% in a continuous trickle-bed reactor. Compared with pure UiO-66, the improvement of the activity can be attributed to the increased Brønsted acid amount caused by the introduced defect sites. And the confinement of UiO-66 (R = 15:1) structure efficiently prevented the Pd loss and aggregation during the long term run. The contrast of the characterizations of the fresh and used Pd/UiO-66 (R = 15:1) indicated that the deactivation of the catalyst was attributed to

carbonaceous accumulation on the catalyst surface. This belongs to the reversible deactivation and the catalyst could be easily regenerated by calcination. This work supplied a new alternative for the design and utilization of industrial catalysts for MIBK and MIBC synthesis.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (Grant 21676303) and the Fundamental Research Funds for the Central Universities of Central South University (Grant 2020zzts415).

References

- 1 N. Takarroumt, M. Kacimi, F. O. Bozon-Verduraz and L. F. Liotta, J. Mol. Catal. A: Chem., 2013, 377, 42–50.
- 2 M. Mediavilla, L. Melo, Y. Díaz, A. Albornoz and A. Llanos, *Microporous Mesoporous Mater.*, 2008, **116**, 627–632.
- 3 D. Hui, W. Zhihui, C. Lifeng, L. Baining and Z. Yonghua, *Ind. Eng. Chem. Res.*, 2018, **57**, 12358–12366.
- 4 Y. Zhu, B. Lin, Y. Hu, Z. Cai and Z. Yonghua, *Mol. Catal.*, 2019, **478**, 110609.
- 5 W. Nicol and E. L. D. Toit, Chem. Eng. Process., 2004, 43, 1539–1545.
- 6 S. Talwalkar and S. Mahajani, *Appl. Catal. A.*, 2006, **302**, 140–148.
- 7 E. D. Toit, R. Schwarzer and W. Nicol, *Chem. Eng. Sci.*, 2004, **59**, 5545–5550.
- 8 L. M. Gandía, R. Malm, R. Marchand, R. Conanec, Y. Laurent and M. Montes, *Appl. Catal. A.*, 1994, **114**, L1–L7.
- 9 J. J. Gamman, S. D. Jackson and F. A. Wigzell, *Ind. Eng. Chem. Res.*, 2010, **49**, 8439–8443.
- 10 S. M. Yang and Y. M. Wu, Appl. Catal. A., 2000, 192, 211-220.
- 11 A. Chakrabarti and M. M. Sharma, *React. Polym.*, 1993, **20**, 1–45.
- 12 A. C. C. Rodrigues, J. L. F. Monteiro and C. A. Henriques, *C. R. Chim.*, 2009, **12**, 1296–1304.
- 13 R. Unnikrishnan and S. Narayanan, *Mol. Catal.*, 1999, 144, 173–179.
- 14 V. Chikán, á. Molnár and K. Balázsik, *J. Catal.*, 1999, **184**, 134–143.
- 15 G. Waters, O. Richter and B. Kraushaar-Czarnetzki, *Ind. Eng. Chem. Res.*, 2006, **45**, 6111–6117.
- 16 J. H. Cavka, S. Jakobsen, U. Olsbye, N. Guillou, C. Lamberti, S. Bordiga and K. P. Lillerud, *J. Am. Chem. Soc.*, 2008, **130**, 13850–13851.
- 17 H. Wu, Y. S. Chua, V. Krungleviciute, M. Tyagi, P. Chen and T. Yildirim, *J. Am. Chem. Soc.*, 2013, **135**, 10525–10532.
- 18 Y. Bai, Y. Dou, L. H. Xie, W. Rutledge, J. R. Li and H. Zhou, *Chem. Soc. Rev.*, 2016, **45**, 2327–2367.

- F. Vermoortele, M. Vandichel, B. V. d. Voorde, R. Ameloot, M. Waroquier, P. V. V. Speybroeck and P. D. Edition, *Angew. Chem. Int. Edit.*, 2012, **124**, 4887–4890.
- 20 F. G. Cirujano, A. Corma and F. X. L. Xamena, *Catal. Today*, 2014, 257, 213–220.
- 21 A. H. Valekar, K. H. Cho, S. K. Chitale, D. Y. Hong, G. Y. Cha, U. H. Lee, D. Hwang, C. Serre, J. S. Chang and Y. Hwang, *Green Chem.*, 2016, **18**, 4542–4552.
- 22 J. Hajek, M. Vandichel and B. V. Voorde, *J. Catal.*, 2015, **331**, 1–12.
- 23 H. Fei and S. Cohen, *Chem. Commun.*, 2014, **50**(37), 4810–4812.
- 24 L. Chen, R. Luque and Y. Li, *Chem. Soc. Rev.*, 2017, **46**, 4614–4620.
- 25 Q. Yang, Q. Xu and H. L. Jiang, Chem. Soc. Rev., 2017, 46, 4774-4808.
- 26 C. A. Trickett, K. J. Gagnon, S. Lee, F. Gándara and H. B. Bürgi, *Angew. Chem.*, 2015, **128**, 11314–11319.
- 27 O. V. Gutov, M. G. Hevia, E. C. Escudero-Adán and A. Shafir, *Inorg. Chem.*, 2015, **54**, 8396–8400.
- 28 Z. Hu, I. Castano, S. Wang, Y. Wang, Y. Peng, Y. Qian, C. Chi, X. Wang and D. Zhao, *Cryst. Growth Des.*, 2016, 16, 2295– 2301.
- 29 L. Yuan, M. Tian, J. Lan, X. Cao and X. Wang, *Chem. Commun.*, 2018, **54**, 370–373.
- 30 G. C. Shearer, S. Forselv, S. Chavan and S. Bordiga, *Top. Catal.*, 2013, **56**, 770–782.
- 31 Z. Niu, Q. Guan, Y. Shi, Y. Chen, Q. Chen, Z. Kong, P. Ning,
 S. Tian and R. Miao, *New J. Chem.*, 2018, 42, 19764–19770.
- 32 Y. Wu, Q. Li, D. J. Rutstrom, M. Zhuravleva and C. Melcher, *Phys. Status Solidi Rapid Res. Lett.*, 2017, **1700403**, 1–4.
- 33 D. Jiang, G. Fang, Y. Tong, X. Wu, Y. Wang, D. Hong, W. Leng, Z. Liang, P. Tu, L. Liu, K. Xu, J. Ni and X. Li, ACS Catal., 2018, 8, 11973–11978.
- 34 R. Ma, Y. Li, G. Wu, Y. He, J. Feng, Y. Zhao and D. Li, *Chinese. J. Catal.*, 2018, **39**, 1384–1394.
- 35 F. Al-Wadaani, E. F. Kozhevnikova and I. V. Kozhevnikov, *J. Catal.*, 2008, 257, 199–205.
- 36 D. Wang, Z. Wang, B. Zahng, H. Sun, P. Wu, Z. Gao and W. Yang, *Chem. React. Eng. Technol.*, 2017, 33(2), 89–297.

- 37 P. Wang, S. Bai, J. Zhao, P. Su, Q. Yang and C. Li, *ChemSusChem*, 2012 Dec, 5, 2390–2396.
- 38 L. Hao, X. Li, M. J. Hurlock, X. Tu and Q. Zhang, Chem. Commun., 2018, 54, 11817–11820.
- 39 L. Valenzano, B. Civalleri, S. Chavan, S. Bordiga, M. H. Nilsen, S. R. Jakobsen, K. P. Lillerud and C. Lamberti, *Chem. Mater.*, 2011, 23, 1700–1718.
- 40 P. Ghosh, Y. J. Colon and R. Q. Snurr, *Chem. Commun.*, 2014, 50, 11329–11331.
- 41 G. C. Shearer, S. Chavan, J. Ethiraj, J. G. Vitillo, S. Svelle, U. Olsbye, C. Lamberti, S. Bordiga and K. P. Lillerud, *Chem. Mater.*, 2014, 26, 4068–4071.
- 42 F. Verpoort, T. Haemers, P. Roose and J. P. Maes, *Appl. Spectrosc.*, 1999, **53**, 1528–1534.
- 43 K. B. Lausund and O. Nilsen, Nat. Commun., 2016, 7, 13578.
- 44 R. C. Klet, Y. Liu, T. C. Wang, J. T. Hupp and O. K. Farha, J. Mater. Chem. A., 2016, 4, 1479–1485.
- 45 T. J. Bandosz, M. Laskoski, J. Mahle, G. Mogilevsky and G. W. Wagner, *J. Phys. Chem. C*, 2012, **116**, 11606–11614.
- 46 T. G. Glover, G. W. Peterson, J. B. DeCoste and M. A. Browe, *Langmuir*, 2012, **28**, 10478–10487.
- 47 G. Ye, D. Zhang, X. Li, K. Leng, W. Zhang, J. Ma, Y. Sun, W. Xu and S. Ma, ACS. Appl. Mater. Inter., 2017, 9, 34937– 34943.
- 48 Y. Liu, R. C. Klet, J. T. Hupp and O. Farha, *Chem. Commun.*, 2016, **52**, 7806–7809.
- 49 L. Hao, X. Li, M. Hurlock and X. Tu, *Chem. Commun.*, 2018, 54, 11817–11820.
- 50 B. Bueken, N. V. Velthoven and T. Willhammar, *Chem. Sci.*, 2017, **8**, 3939–3948.
- 51 C. A. Trickett, K. J. Gagnon, S. Lee, F. Gándara, H.-B. Bürgi and O. M. Yaghi, *Angew. Chem.*, 2015, **127**, 11314–11319.
- 52 J. Hajek, B. Bueken and M. Waroquier, *ChemCatChem*, 2017, 9, 2203–2210.
- 53 Q. Guan, B. Wang, X. Chai, J. Liu, J. Gu and P. Ning, *Fuel*, 2017, **205**, 130–141.
- 54 Y. Yang, D. Deng, D. Sui, Y. Xie, D. Li and Y. Duan, *Nanomaterials*, 2019, **9**, 1698–1704.