# **Catalysis<br>Science & Technolog** Technology (SS)

## PAPER

Cite this: Catal. Sci. Technol., 2021, 11, 6257

Received 11th February 2021, Accepted 16th July 2021

DOI: 10.1039/d1cy00269d

rsc.li/catalysis

## Introduction

Olefin metathesis (OM) is a powerful and versatile method of organic chemistry. During OM, carbon atoms of two C=C double bonds are reorganized to new double-bond-containing molecules. These reactions are very selective and require mild reaction conditions. The atom economy of the synthetic procedures is often 100%, i.e., all the starting materials are incorporated into the products.

Nature is abundant in bio-based materials containing the olefin bond. Low-value olefins can also be found in vast amounts in petrochemical by-products. These chemicals often appear as an underutilized feedstock.

With the advent of alternative fuels and electric cars, the worldwide demand for mineral oil-based fossil fuels will

## Propylene synthesis via isomerization–metathesis of 1-hexene and FCC olefins†

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Conversion of 1-hexene or olefins obtained by fluid catalytic cracking (FCC) to propylene via isomerization–metathesis (ISOMET) was investigated using ethylene as a cross-coupling agent. Zeolite H-beta (HBEA) was applied as an isomerization catalyst. The olefin metathesis (OM) catalysts were about 12 wt% molybdena, supported on zeolite beta (MoO<sub>3</sub>/HBEA), and γ-alumina (MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>). HBEA-supported catalyst with a lower molybdena content (6 wt%) was also investigated. The catalysts were characterized by X-ray diffractometry (XRD), H<sub>2</sub>-temperature-programmed reduction (H<sub>2</sub>-TPR), and Fourier transform infrared (FT-IR), visible Raman, in situ ultraviolet-visible (UV/VIS) and XPS spectroscopy. It was shown that HBEA is a highly active and robust catalyst of double-bond isomerization. Applying a physical mixture of HBEA and  $12M$ o $O_3$ /Al<sub>2</sub>O<sub>3</sub> catalyst at 150 °C and 3 bar ethylene pressure, 60% conversion of 1-hexene to propylene was attained. Interestingly, quantitative conversion to propylene was achieved after reactivation of the deactivated catalyst in an argon atmosphere at 550 °C. It was found that the pre-treatment of the catalyst with olefins such as ethylene before inert gas activation resulted in significant catalyst activity improvement. This suggests that the adsorbed olefins may play a key role in the formation of active metal centers during the catalyst reactivation process. The catalyst mixture also had good performance in the conversion of FCC olefins to propylene. The MoO<sub>3</sub>/HBEA catalysts have rendered reasonable activity; however, the catalysts showed a significantly shorter lifetime than the alumina-containing catalyst mixture. **PAPER**<br> **CO** Check for undates<br> **CO** Check for undates<br>
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certainly decrease. However, the need for high-value innovative materials, including advanced polymers, steadily increases.<sup>1-3</sup> Unfortunately, at present, these demands cannot be met by using renewable feedstocks only. Therefore, there is still a constant need for the development of high atom-economic chemical procedures for the efficient and environmentally benign conversion of petrochemicals to high-value materials.

Propylene is an emerging bulk chemical, the key monomer of polypropylene plastic and other commodity chemicals. It is mainly obtained as a by-product of ethylene production in steam cracker units.4,5 Possibilities to control the ratio of ethylene to propylene in the cracked product mixture are limited. As a result, cracking units are currently producing excess ethylene, while the increasing demand for propylene remains uncovered.6–<sup>9</sup> Propylene can also be synthesized on a large scale by propane dehydrogenation as well as Fischer–Tropsch synthesis and methanol-to-olefin conversion.3,10–<sup>12</sup> Propylene can also be produced selectively via OM under moderate reaction conditions. The most general example is the ethenolysis of 2-butene giving two propylene molecules.<sup>1,4,5,7,9,11-14</sup>

There are several industrial examples for OM of olefinic hydrocarbon over heterogeneous catalysts such as the Shell

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<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: 10.1039/ d1cy00269d

Higher Olefin Process (SHOP) for detergent production,<sup>15</sup> the Philips Triolefin Process for synthesis of 2-butene<sup>16</sup> and the Olefin Conversion Technology used by ABB Lummus Global for propylene production.<sup>17</sup> The most widely used catalyst systems include Mo (ref. 11, 14 and 18–22) W (ref. 3, 4, 6, 8, 9, 12, 18 and 23–25) or Re (ref. 2, 26 and 27) oxides. Mo- and W-based catalysts are widely used in the petrochemical industry. Some new OM reactions, catalysed by metal oxide, have been reviewed recently.<sup>10,28</sup>

A special case of OM is isomerization–metathesis (ISOMET). During ISOMET, the olefinic bonds migrate along the hydrocarbon chain (isomerization), which is followed by a cross-metathesis reaction with another olefin, a crosscoupling agent. A special case of OM, when the crosscoupling agent is ethylene, is called ethenolysis.<sup>10,29</sup> In the case of linear mono-olefins, the complete ISOMET using ethylene as a cross-coupling agent theoretically ends up with propylene as the only end product (Scheme 1).

The ISOMET of C4 and C5 olefins producing propylene is one of the most widely investigated areas. Interestingly, industrially applied systems are mainly based on W (5–10% on silica) or Re  $(5-10\%$  on alumina) catalyst systems.<sup>4,11,30-32</sup> The ISOMET process of higher olefins (C4–C9) and the mixtures thereof has also been patended.<sup>33</sup> Re<sub>2</sub>O<sub>7</sub>, WO<sub>3</sub> and  $MoO<sub>3</sub>$  were used as OM catalysts, whereas  $RuO<sub>2</sub>$ , MgO and K2CO3 were applied as isomerization catalysts. Another patent application describes a process for converting an olefin feed containing butenes, diolefins and polyolefins to propylene with ethylene using molybdenum as a metathesis catalyst and MgO,  $K_2CO_3$ , and  $K_2O$  as isomerization catalysts.34 In both cases, the use of a fluid catalytic cracking



**Scheme 1** The conversion of  $C > 5$  olefins into propylene via ISOMET with ethylene.

(FCC) mixture is mentioned as a source of olefin feed, but only after appropriate refining.<sup>35,36</sup>

The FCC and FCC light fractions are abundant in  $C > 5$ olefin components. Furthermore, not only petrochemical streams but also materials derived from renewable feedstocks may contain  $C > 5$  olefins. For example, the ethenolysis of oleic acid produces 1-decene,<sup>29,37</sup> while the ethenolysis of linoleic acid gives 1-hexene. Both vegetable oils are highly abundant in nature.<sup>38</sup> Light olefins (propylene, butenes, butadiene) are also available from biomass-originated bioethanol<sup>5,39</sup> and biobutanol<sup>40</sup> and from biomass pyrolysis.<sup>6</sup> Both micro- and mesoporous silicates, aluminosilicates were shown to be effective as supports for molybdena in the OM reaction.20–<sup>22</sup>

In the present study, zeolite beta was chosen as the isomerization catalyst for the ISOMET of 1-hexene model compound and FCC fractions. Beta is a large-pore zeolite with a three-dimensional structure of 12-membered ring channels.41 Owing to its large pore size, strong acid sites and high chemical and thermal stability, it is used as a catalyst in the petrochemical industry and fine chemistry. It is also utilized as an adsorbent.<sup>42</sup> Zeolite beta has already been applied as a support itself<sup>43,44</sup> or mixed with  $Al_2O_3$  in the OM of 2-butene with ethylene to propylene.<sup>45</sup> However, it has not been used in the ISOMET of long-chain olefins yet, where the role of zeolite is the double-bond isomerization of the reactant and intermediate alpha-olefin products. The dispersed  $MoO<sub>x</sub>$ , the active component of the oxidesupported catalysts, was shown to be present as isolated or oligomeric surface species as well as crystalline particles on a high-surface-area oxide support.<sup>28</sup> **Paper** Catalysis Science & Technical Philip Formula Catalysis Article. Published on 2011. Downloaded on 2011. The published on 12-10/2021. Downloaded under the summary of the summary of the summary of the summary of the

Papers reporting isomerization metathesis using C5+ alkenes as raw materials for propylene synthesis are rare. $46,47$ In particular, there is no information in the literature about the synthesis of propylene from C5+ alkenes via ISOMET using HBEA as isomerization and  $MoO<sub>x</sub>/Al<sub>2</sub>O<sub>3</sub>$  as olefin metathesis catalysts. This paper describes a zeolite-supported molybdena (MoO3/HBEA) catalyst system having isomerization (HBEA) as well as metathesis  $(M_0O_3)$  activity. The mixed bed of alumina-supported molybdena and zeolite beta  $(MOO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub> + HBEA)$  ISOMET catalyst is also reported.  $MoO<sub>3</sub>$  loading of about 12 wt% was applied on both supports. Based on the work of Li and co-workers, for HBEA, a catalyst containing lower  $MoO<sub>3</sub>$  loading  $(6 \text{ wt\%})$  was also investigated.<sup>48</sup> The catalysts were characterized and investigated in the ISOMET of 1-hexene model compound with ethylene. Conversion of FCC having high olefin content to propylene was also studied. It has been found that the initial catalytic activity can be significantly improved by simple high-temperature heat treatment of the deactivated ISOMET catalyst in an inert gas.

### Results and discussion

The molybdena content of the catalysts, determined by ICP-OES, is shown in Table 1. The ISOMET activity of  $13\text{MoO}_3$ /

Table 1 Characterization of the supports and catalysts

Catalyst	$\text{MoO}_{3}^{a}$ (wt%)	$SSA^{b}$ (m <sup>2</sup> g <sup>-1</sup> )	$H_2$ uptake <sup>c</sup> (mmol $g^{-1}$ )	$H/Mo^c$
$Al_3O_3$	$\hspace{0.1mm}-\hspace{0.1mm}$	192		$\hspace{0.1mm}-\hspace{0.1mm}$
$12MO3/Al2O3$	11.7	184	2.38	5.9
HBEA $(Si/Al = 12)$		480		
13MOO <sub>3</sub> /HBEA	13.5	235	2.61	5.6
6MoO <sub>3</sub> /HBEA	6.1	476	1.17	5.6

 $a$  Determined by ICP-OES.  $b$  Specific surface area (SSA) determined by the Brunauer–Emmett–Teller method.  $c$  Calculated from the TPR curve measured up to 800 °C.

HBEA and the physical mixture of  $12\text{MoO}_3/\text{Al}_2\text{O}_3$  and HBEA was higher than that of  $6MoO<sub>3</sub>/HBEA$  (vide infra). The effect of the amount of  $MoO<sub>3</sub>$  loading on HBEA zeolite (70%)/  $Al_2O_3(30\%)$  has been investigated by Li *et al.* using the abovementioned solid support mixture; a 6–8% molybdena loading was found to be optimal.<sup>48</sup> The characterization of  $13\text{MoO}_3$ / HBEA and  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  is presented below, whereas the properties of  $6MoO<sub>3</sub>/HBEA$  are included in the ESI.<sup>†</sup>

#### Catalyst characterization

**Structure and texture.** The XRD patterns of  $Al_2O_3$  and HBEA supports and  $12\text{MoO}_3/\text{Al}_2\text{O}_3$  and  $13\text{MoO}_3/\text{HBEA}$ catalysts are shown in Fig. 1. Gamma-alumina is the only detectable phase of  $MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  containing about 12 wt%  $MoO<sub>3</sub>$ . This amount of molybdena is about two-thirds of the monolayer capacity.

The monolayer coverage of the used  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> having a specific surface area of 192 m<sup>2</sup> g<sup>-1</sup> corresponds to about 19.2 wt%  $MoO<sub>3</sub>$  content.<sup>49</sup> The absence of the  $MoO<sub>3</sub>$  XRD reflections suggests that  $MoO<sub>3</sub>$  crystallites are well dispersed on the alumina surface; thus they are X-ray amorphous species. The obtained results confirmed that aluminasupported  $MoO<sub>3</sub>$  catalysts of a high dispersion can be



Fig. 1 XRD patterns of supports and catalysts calcined at 500 °C.

obtained below monolayer coverage or even at  $MoO<sub>3</sub>$  contents higher than that corresponding to the monolayer coverage.<sup>50,51</sup> Upon Mo loading  $(13.5 \text{ wt\% MO}_3)$  the characteristic reflections of HBEA at  $2\theta$  = 7.8° and 22.6° significantly decreased, as was also shown by others. $43,52$ 

In addition, intense reflections typical of orthorhombic MoO3 (JCPDS 35-609) appeared in the XRD pattern of  $13\text{MoO}_3/\text{HBEA}$  at  $2\theta = 27.4^\circ$ ,  $23.4^\circ$  and  $25.7^\circ$ . The average MoO3 particle size, calculated from the Scherrer equation using the  $2\theta = 27.4^{\circ}$  reflection, was about 90 nm. At lower  $MO<sub>3</sub>$  loading (6MoO<sub>3</sub>/HBEA), no crystalline MoO<sub>3</sub> phase was detected by XRD and the diffraction lines of the zeolite support were more intense (Fig. S1†). No crystalline  $Al<sub>2</sub>(MoO<sub>4</sub>)<sub>3</sub>$  phase was detected in these samples. Formation of this phase was found in samples treated at high temperature (680 °C).<sup>43</sup> In our case, however, the MoO<sub>3</sub> content was not high and the decomposition temperature of the Mo precursor was lower, 500 °C, so the formation of the aluminium molybdate phase is unlikely in any of the supports. Catalysis Science & Technology<br>
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As expected, the specific surface area of the molybdenaloaded catalyst was smaller than those of the corresponding supports. The difference in the surface areas depends on the pore size of the support (Table 1). For  $\gamma$ -Al<sub>2</sub>O<sub>3</sub>, which is mesoporous, the specific surface area decreased only slightly (from 192 to 184 m<sup>2</sup>  $g^{-1}$ ). The specific surface area of the microporous zeolite catalyst with 12 wt%  $MoO<sub>3</sub>$  loading, however, was only half of that of the zeolite support (235 and 480 m<sup>2</sup> g<sup>−1</sup>). At 6 wt% MoO<sub>3</sub> content the SSA of HBEA did not decrease  $(476 \text{ m}^2 \text{ g}^{-1})$ .

#### Temperature-programmed reduction by hydrogen  $(H_2$ -TPR)

The reducibility of the molybdenum species in  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$ and  $13MoO<sub>3</sub>/HBEA$  was investigated by temperatureprogrammed  $H_2$  reduction (Fig. 2). Two reduction peaks were observed on the H<sub>2</sub>-TPR curve of the  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  catalyst. The low-temperature peak in the range of 300–600 °C with a maximum of 450 °C represents the reduction of multilayered and octahedral Mo(v<sub>I</sub>) to Mo(v<sub>I</sub>).<sup>50,53</sup> The tetrahedral Mo (v<sub>J</sub> species has stronger interaction with the  $Al_2O_3$  support, leading to a reduction temperature in the range of 600–800  $\rm{°C}$ . In this higher temperature region the reduction of Mo(IV) to Mo(0) takes place.



Fig. 2  $H_2$ -TPR curves of the  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and  $13MoO<sub>3</sub>/HBEA$ catalysts. The catalysts were treated in  $O<sub>2</sub>$  flowing at a rate of 30 ml min<sup>-1</sup> for 1 h at 500 °C, cooled to 25 °C and then heated at a rate of 10 °C min<sup>-1</sup> up to 800 °C for 1 h in a flow of a 9% H<sub>2</sub>/N<sub>2</sub> mixture.

For the  $13\text{MoO}_3/\text{HBEA}$  catalyst, two reduction peaks were observed in the temperature range of 300–650 °C and 650– 800 °C. The peak at lower temperature is assigned to reduction of Mo(v<sub>I</sub>) to Mo(v<sub>I</sub>) as in the case of the  $12\text{MoO}_3$ /  $Al_2O_3$  catalyst. The second peak represents the complete reduction of Mo( $iv$ ) to Mo( $0$ ).<sup>54</sup> A similar TPR curve was observed for  $6M_0O_3/HBEA$  (Fig. S2†). Table 2 shows that the hydrogen uptake, expressed in the H/Mo ratio, in the  $Al_2O_3$ supported sample was 5.9. This amount of hydrogen

Table 2 Propylene yield in ISOMET of 1-hexene (1-H) with ethylene over activated and reactivated 13MoO3/HBEA catalyst at different temperatures and 0.5 h and 3 h TOS, at 3  $g_{cat} g_{1-H}^{-1}$  h space time under  $\frac{g_{1-H} - g_{1}}{g_{1-H}}$  has a the under 3 bar ethylene pressure. The ethylene/1-hexene molar ratio was 10

	<b>TOS</b>	Propylene yield (%)		
Catalyst	(h)	Activated <sup><math>a</math></sup>	Reactivated <sup>b</sup>	
75	0.5	10	20	
	3	6	16	
100	0.5	15	59	
	3	12	19	
125	0.5	29	86	
	3	5	17	
150	0.5	19	82	
	3		10	

 $a$  Activation: the ex situ calcined catalyst was in situ pre-treated in a flow of Ar (50 ml min−<sup>1</sup> ) at 550 °C for 2 h. The activated catalyst was cooled to the target temperature in a flow of Ar and the ISOMET reaction was performed.  $\overline{b}$  Reactivation: after 3 h TOS, the reactant feed was stopped and the total pressure was reduced to atmospheric pressure. The catalyst was purged with a flow of Ar (50 ml min<sup>-1</sup>) for 30 min at the reaction temperature to remove olefins, then heated to 550 °C at a ramp rate of 10 °C min−<sup>1</sup> and maintained at this temperature for 2 h. The reactivated catalyst thus obtained was cooled to the target temperature in a flow of Ar and the ISOMET reaction was performed again.

consumption is near to the amount needed for the total reduction of  $MoO<sub>3</sub>$  to  $Mo(0)$  (6). Molybdena has been shown to react with the zeolitic protons during oxidative decomposition of the heptamolybdate precursor.<sup>55</sup> At 500  $^{\circ}$ C,  $MoO<sub>x</sub>$  oligomers migrate into the channels and react with the Brønsted acid sites of the zeolite to form ditetrahedral Mo species, *i.e.*  $(Mo<sub>2</sub>O<sub>5</sub>)<sup>2+</sup>$  cations. These cationic Mo species can only be reduced at temperatures above 800  $^{\circ}$ C.<sup>54,56</sup> Table 1 shows that the H/Mo ratio was 5.6 for both  $MO_3/HBEA$ catalysts, suggesting that some Mo may occupy cationic positions and cannot be reduced under our TPR conditions. In line with the above results, the loss of Brønsted acid sites was also observed for these samples (vide infra). Before the catalytic experiments, the calcined catalysts were activated in situ in a flow of Ar at 550 °C. Thermal autoreduction of  $Mo(vi)$  cannot be detected by  $H_2$ -TPR. **Catalysis Science Articles.**<br>
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#### FT-IR spectroscopy of adsorbed pyridine

For the ISOMET reaction, the catalyst must contain Brønsted acid sites that promote double-bond isomerization of terminal olefins. Pyridine adsorption, followed by FT-IR spectroscopy, was used to characterize the acidity of the supported molybdena catalyst (Fig. 3). Brønsted and Lewis acid sites of the catalysts can be distinguished by the characteristic 19b/8a ring vibrations of pyridinium ions and pyridine bound to Lewis acid sites.

These vibrations of the two species appear in different regions of the spectrum, *i.e.*, at 1545/1637 cm<sup>-1</sup> and around 1455/1620 cm−<sup>1</sup> , respectively. Fig. 3 shows that the HBEA support has both Brønsted and Lewis acid sites. In zeolites,



Fig. 3 FT-IR spectra of pyridine. The samples were activated in high vacuum at 550 °C for 1 h. Pre-treated samples were allowed to come in contact with Py vapour at 5 mbar pressure at 200 °C for 30 min, then evacuated for 30 min. After evacuation, the spectra were recorded at room temperature. Labels L and B indicate the characteristic bands for Py bonded to Lewis and Brønsted acid sites, respectively.

Brønsted acidity is due to protons compensating for the negative charge on the zeolite framework generated by tetrahedral framework aluminum. Extra-framework aluminum represents Lewis acid sites. A high number of defect sites caused by crystallographic faults is typical for the structure of zeolite beta.<sup>57</sup> Thus, trigonal framework aluminum near the silanol nests shows Lewis acidity. Upon Mo loading, the number of both Brønsted and Lewis acid sites decreased significantly. The FT-IR results suggest that molybdenum reacts with zeolitic protons and occupies cationic positions as ditetrahedral Mo ions  $(Mo_2O_5)^{2^+}$ , as proposed by Ding et  $al$ <sup>58</sup> The amount of Brønsted acid sites in the zeolite sample was 0.71 mmol  $g^{-1}$ , measured by the ammonium ion exchange capacity. Comparing the band intensities of the pyridinium ions around 1545  $cm^{-1}$ , the number of Brønsted acid sites decreased by 36%, indicating that 0.12 mmol  $g^{-1}$  Mo is in the cationic position, which is about 15% of the total Mo content. In addition, surface  $MoO<sub>x</sub>$ species can be bound in the zeolite silanol nests both in the micropores and on the outer surface of the zeolite crystallites. These species cause a significant decrease in the number of Lewis acid sites. Catalysis Science & Technology<br>
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In the spectrum of  $12\text{MoO}_3/\text{Al}_2\text{O}_3$  the intensities of the bands characteristic of Lewis acid sites slightly increase at 1453 and 1621 cm<sup>-1</sup> compared to the bare support (Fig. 3). Highly polymerized  $MoO<sub>x</sub>$  species were suggested to be responsible for the additional Lewis acid sites at this Mo content.<sup>59</sup> An approximately 10% decrease in Brønsted sites was observed on  $6M_0O_3/HBEA$  relative to the HBEA support (Fig. S3†).

#### Raman spectroscopy

The structure of supported molybdenum oxide was investigated by Raman spectroscopy under ambient conditions in the hydrated state. The Raman spectra of  $12M_0O_3/Al_2O_3$  and  $13\text{MoO}_3/\text{HBEA}$  are presented in Fig. 4. In the spectrum of  $12\text{MoO}_3/\text{Al}_2\text{O}_3$ , the bands observed at 953, 910 and 355 cm<sup>-1</sup> are assigned to the symmetric stretching, asymmetric stretching, and bending vibrations of the terminal Mo=O bond of octahedral MoO6 species in hydrated heptamolybdate, respectively.<sup>60</sup> In addition, the bands at 566 and 222 cm<sup>-1</sup> are due to the Mo–O–Mo symmetric stretching and deformation vibrations of these molybdena species, respectively. These results confirm that at 12 wt% molybdena loading octahedral species dominate on alumina. It should be mentioned that tetrahedral MoO<sub>4</sub> units are also present as the Raman band at 910 cm<sup>-1</sup> is typical of the symmetrical stretching vibrations of the terminal Mo=O in tetrahedral molybdena species.<sup>61</sup> This observation was also confirmed by UV-vis spectroscopy. The intense and narrow bands in the spectrum of 13MoO<sub>3</sub>/HBEA are due to crystalline  $MoO<sub>3</sub>,<sup>62</sup>$  indicating that this catalyst contains the MoO<sub>3</sub> phase, which is also supported by the XRD data (Fig. 1). At 6 wt% molybdena loading, the intensity of the 970 cm−<sup>1</sup> band increased compared to that of the zeolite support (Fig. S4†). This indicates that the catalyst contains tetrahedral  $MoO<sub>x</sub>$  species.



Fig. 4 Raman spectra of  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and  $13MoO<sub>3</sub>/HBEA$  under ambient conditions.

#### In situ UV-vis spectroscopy

In situ UV-vis experiments were performed to determine the type of  $MoO<sub>x</sub>$  species in the calcined  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and  $13MoO<sub>3</sub>/HBEA$  catalysts and to follow changes in the degree of polymerization of  $MoO<sub>x</sub>$  during in situ activation in an inert atmosphere (Ar) at temperatures increased up to 550 °C (Fig. 5). Both calcined catalysts show a strong absorption band below 330 nm. The bands at 250 and around 290 nm are due to the ligand–metal charge transfers (LMCTs) from  $oxygen to Mo(v)$  in highly dispersed and small linear-chained tetrahedral MoOx, respectively. LMCT in polymerized



Fig. 5 In situ UV-vis spectra of the  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and  $13MoO<sub>3</sub>/HBEA$ catalysts. The catalysts were heated at a rate of 10 °C min−<sup>1</sup> up to 550 °C for 2 h in an Ar flow.

octahedral  $MoO<sub>x</sub>$  gives rise to a band at around 330 nm.<sup>59,63–65</sup> For  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$ , the intensity of the absorption band at 250 nm becomes weaker above 200 °C, while the band at 330 nm gains strength. This result suggests that during thermal treatment in inert atmosphere tetrahedral  $MO<sub>x</sub>$  is transformed to polymerized octahedral  $MO<sub>x</sub>$  species over alumina-supported molybdena. Comparing the relative intensity of the bands below 300 nm and at 330 nm in the spectra of  $13\text{MoO}_3/\text{HBEA}$ , similar but smaller changes were observed, indicating that a smaller portion of tetrahedral  $MoO<sub>x</sub>$  species is transformed to octahedral  $MoO<sub>x</sub>$  species on the zeolite-supported molybdena.  $MoO<sub>3</sub>$  located in microporous channels is less capable of forming longer  $MoO<sub>x</sub>$  chains. They can be formed only on the outer surface of the zeolite crystals, but in small amounts only because the outer surface is only approx. 4–5% of the specific surface area of the zeolite. In the spectrum of the calcined  $13\text{MoO}_3/\text{HBEA}$ the presence of a weak absorption band at around 400 nm is due to the crystalline  $MoO<sub>3</sub>$  phase, which was also confirmed by XRD. Crystalline  $MoO<sub>3</sub>$  was shown to be catalytically inactive.<sup>66</sup> In the spectrum of  $6M_0O_3/HBEA$ , at higher temperatures, those tetrahedral  $MoO<sub>x</sub>$  species that are surrounded by other Mo atoms cannot be detected (Fig. S5†). This result suggests that during heat treatment in Ar,  $MoO<sub>x</sub>$ species migrate into the microporous channels and occupy cationic positions in the zeolite. **Paper**<br>
Catalysis Science & Technology<br>
nm:<sup>263</sup> - Epi 2046, 2047, this method of the silence of the

#### XPS measurements

Quantitative evaluation of the XPS data indicated that the Mo oxide content of the  $MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  catalyst was around 12 wt%, in good agreement with the nominal composition. Spectral features of Al (Al 2p binding energy of 74.7 eV, Al KLL kinetic energy of 1386.8 eV and the Auger parameter defined as their sum at 1461.5 eV) as well as the main O 1s peak around 531.4 eV all pointed to an oxidized/hydroxylated environment for the Al( $\text{III}$ ) cations,<sup>67,68</sup> in agreement with literature data on similar catalysts.<sup>69</sup> No significant changes in the chemical environment of Al were observed during the treatments. In addition, no sign of dissolution of Mo into the alumina was detected. The Mo  $3d_{5/2-3/2}$  spin orbit doublet of welldispersed  $MoO<sub>3</sub>$  on alumina has a rather broad line shape with a  $3d_{5/2}$  binding energy of around 233 eV,<sup>69-71</sup> which is somewhat higher than the value characteristic for bulk MoO<sub>3</sub>  $(232.5 \text{ eV})$ .<sup>72</sup> While the shift is attributed to the strong interaction of the oxidized Mo species with the alumina support, $70$  the broadening can be the result of a charging effect<sup>70</sup> but may also indicate the existence of a range of slightly different environments for the adsorbed  $Mo(v)$ ions.<sup>73</sup> As shown in Fig. 6a, these features are well reproduced in the Mo 3d spectrum of the calcined  $12\text{MoO}_3$ / Al2O3 sample exposed to air. The spectrum can be well modelled by a single peak pair with the  $3d_{5/2}$  component at 233.2 eV, indicating the exclusive presence of  $Mo(v)$  ions.

Treatment in Ar at 550 °C resulted in a marginal shift of the maximum of the spectrum towards lower binding



Fig. 6 Mo 3d spectra of the calcined  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  sample exposed to air (a), after activation in Ar at 550 °C for 1 h (b), subsequent  $C_2H_4$ exposure at 100 °C for 1 h (c) and after reactivation in Ar at 550 °C for 1 h (d).

energies and further apparent broadening (Fig. 6b). Spectral modelling revealed that these changes can be interpreted as the result of the appearance of a new Mo 3d doublet with its  $3d_{5/2}$  peak at around 231.7–231.9 eV.

According to its binding energy, this new component was attributed to Mo(v) species.<sup>72</sup> The abundance of the Mo(v) species clearly increased upon ethylene exposure (Fig. 6c) and some further increase was observed after the subsequent treatment in Ar at 550 °C (Fig. 6d). Although the combination of the  $Mo(v)$  and  $Mo(v)$  states adequately modeled the observed line shapes, the broad peaks can overlap the weak  $Mo(w)$  signal, so their presence, especially after the ethylene exposure and the subsequent re-activation, cannot be completely ruled out.

#### 1-Hexene isomerization over HBEA catalyst

Preliminary experiments were carried out to investigate the isomerization of 1-hexene model compound using the HBEA catalyst at 75 °C and atmospheric pressure in an inert Ar gas atmosphere. At nearly 90% 1-hexene conversion, the yields of cis- and trans-2-hexene and cis- and trans-3-hexene were 70% and 20%, respectively. (Fig. 7). This result indicates that double bond shift and cis–trans rearrangement are the main reactions in the conversion of 1-hexene over zeolite beta, consistent with earlier results on large- and medium-pore zeolites, i.e., H-Y and H-ZSM-5. Side reactions such as polymerization are negligible.74–<sup>76</sup> It was demonstrated that under these conditions, HBEA is highly robust. No catalyst deactivation was observed during a time on stream (TOS) of 10 h.

The ISOMET of 1-hexene with ethylene was performed at 3 bar total pressure of ethylene and at an ethylene/1-hexene



Fig. 7 Isomerization of 30% 1-hexene (1-H) solution using the HBEA catalyst at 75 °C and 3  $g_{cat} g_{1-H}^{-1}$  h space time under 3 bar ethylene pressure. The calcined catalyst was pretreated in situ in a flow of Ar (50 ml min−<sup>1</sup> ) at 450 °C for 2 h at atmospheric pressure.

molar ratio of 10. In order to make sure that the isomerization is not affected by ISOMET conditions the isomerization reaction was performed in the presence of 3 bar ethylene (optimal ethylene pressure for OM, Fig. S7†) as well. The results showed that neither the catalytic activity nor the product selectivity was affected. Similar conversion and reaction product distribution to that in the absence of ethylene were observed. The influence of reaction temperature on the isomerization conversion and product distribution has been investigated at 75 (optimal reaction temperature for OM), 100, 125 and 150 °C and 3  $g_{\text{cat}} \, g_{\text{1--H}}^{-1}$  h space time under 3 bar ethylene pressure. It could be concluded that even at 75 °C neither the conversion of 1-hexene nor the product distribution has been changed significantly.

By using the optimized OM (ethenolysis) condition, the catalyst activity did not decrease within ten hours of the reaction (Fig. 7). The yield of 2-hexenes achieved 66% (55% trans-2-hexene and 11% cis-2-hexene) after 1 hour TOS. HBEA is a highly robust isomerization catalyst showing high activity and stability under optimized metathesis reaction conditions.

The 1-hexene isomerization activity of the  $MoO<sub>3</sub>$ containing HBEA catalysts was also investigated. The catalysts were pretreated in an  $O_2$  flow at 550 °C for 2 h to keep Mo in the  $Mo(w)$  state. It was found that the isomerization activity of HBEA significantly dropped upon impregnating with  $MoO<sub>3</sub>$ . With increasing  $MoO<sub>3</sub>$  content, the number of Brønsted acid sites decreases (shown by FT-IR spectroscopy), and therefore the isomerization activity also decreases. Actually, at 6 wt% loading 80% 1-hexene isomerization was observed (versus 90% on neat HBEA), while at 12 wt% loading the isomerization activity was around 60% after one hour time on stream.

At 12 wt% molybdena content the isomerization activity was halved after 5 hours time on stream, whereas  $6MoO<sub>3</sub>/$ HBEA showed good stability in 1-hexene isomerization.

#### 1-Hexene ISOMET using MoO<sub>3</sub>/HBEA catalysts

Theoretically, (not only) terminal olefins such as 1-hexene can be completely converted to propylene by using ethylene in sequential isomerization and ethenolysis steps (ISOMET) (Scheme 2). Two types of sites are required, Brønsted sites for the C=C double-bond isomerization of 1-hexene and the intermediate alpha-olefins formed (1-pentene, 1-butene) and active  $MoO<sub>x</sub>$  sites for the conversion of isomerized olefins (2and 3- hexenes, 2-pentenes, 2-butenes) with ethylene to give propylene and lower alpha-olefins (1-pentene, 1-butene).

The pathway of the 3-hexene isomer to propylene is shorter, as its reaction with ethylene gives 1-butene, which is converted to propylene in the isomerization and ethenolysis steps (not shown in Scheme 2).

In the following series of experiments, HBEA was used not only as an isomerization catalyst but also as a support for  $MoO<sub>3</sub>$ . The reactions were carried out under 3 bar ethylene pressure and 3  $g_{cat} g_{1-H}^{-1}$  h space time in the temperature range of 75–150 °C. The catalyst was pre-treated in inert gas at 550 °C and the activated catalyst thus obtained was tested in the ISOMET reaction for 3 h time on stream (TOS). After activation, the catalyst was regenerated under the same conditions as the pre-treatment. The activity of the reactivated catalyst was also studied in the ISOMET reaction.

On the activated  $13\text{MoO}_3/\text{HBEA}$  catalyst, the propylene yield was low, ranging from 10% to 29% at 0.5 h TOS (Table 2). It was 10% at 75 °C; with increasing reaction temperature it reached its maximum (29%) at 125 °C and then at 150 °C decreased to 19%. Reactivation, i.e., heat treatment in an inert atmosphere at high temperature (550 °C) after the ISOMET reaction, however, resulted in a significant increase in the initial activity of the  $13MoO<sub>3</sub>/$ HBEA catalyst. At 75 °C, the propylene yield doubled, while at higher temperatures the amount of propylene was 3 and 4



Scheme 2 Subsequent isomerization and ethenolysis steps in ISOMET of 1-hexene with ethylene to propylene. (The pathway of the 3-hexene isomer is not shown here).

times as high on the reactivated catalyst as on the freshly activated catalyst. Thus, at 125 and 150 °C more than 80% propylene yield can be obtained over reactivated  $13\text{MoO}_3$ / HBEA at 0.5 h TOS. However, the catalyst is rapidly deactivated under all conditions. The rate of deactivation was higher at higher temperatures, and higher on the reactivated catalyst. During 3 h TOS, at 75 °C the propylene yield decreased from 10% to 6% and from 20% to 16% on the activated and reactivated catalyst, respectively. At 125 °C, the initial activity of the activated and reactivated catalyst decreased from 29% to 5% and from 86% to 17%, respectively, in three hours (Table 2). The  $13MoO<sub>3</sub>/HBEA$ catalyst, in which the active sites of both the isomerization and the OM reactions are located in a microporous aluminosilicate support, is rapidly deactivated due to the polymerization of olefins.<sup>77</sup> In particular, at 150 °C, after three hours TOS the propylene yield dropped significantly from 85% to 10%. **Paper** Catalysis Science & Packmost times as high on the results are on the freeholy . Complementary characterization techniques revealed and access are highly deprecised and the same in the same of the most simply depen

This catalyst contains 90 nm-size  $MoO<sub>3</sub>$  crystals, which block the micropores; therefore we have halved the  $MoO<sub>3</sub>$ content of the catalyst. No crystalline  $MoO<sub>3</sub>$  phase was detected by XRD in  $6MoO<sub>3</sub>/HBEA$  and only the low-intensity Raman bands indicate the presence of a small amount of  $MoO<sub>x</sub>$  species; however, the catalyst showed lower activity in 1-hexene ISOMET (Table S1†).

At 75 °C the activity of the two catalysts was about the same, but as the temperature increased, the propylene yield on  $6M_0O_3/HBEA$  hardly changed. The number of active sites is presumably decreasing during the reaction as carbonaceous deposits may gradually block the micropores; thus the reactant molecules cannot access the active sites located inside the pores. Not only coking but also the low oxidation state of Mo is a reason for the decreased activity (vide infra).

On MoO3/HBEA catalysts, the Brønsted acid sites of HBEA zeolite support protonate the 2-hexene to form the carbenium ion intermediate of double-bond isomerization.<sup>77-79</sup> Zhang et  $al^{11}$  reported that both the number of Brønsted acid sites and the high dispersion of  $MoO<sub>3</sub>$  species have an important role in determining the OM activity.

Brønsted acid sites not only catalyse double-bond isomerization of alpha-olefins but are also involved in the OM reactions, i.e., participate in the generation of active metal carbene species, which are the active site of the OM reactions. Previous work suggested that Brønsted acid OH groups interact with adjacent metal oxides, for example,  $MoO<sub>x</sub>$ ,  $WO<sub>x</sub>$ , or  $ReO<sub>x</sub>$ , for the generation of OM active carbene species. $80,81$  Recent results showed that using the  $MoO_x/SBA-15$  catalyst, surface Brønsted acidic OHs coordinated to Mo(vi) protonate propylene and isopropoxide species are formed upon propylene adsorption.82 Such species are further oxidized by the lattice oxygen of  $MoO<sub>x</sub>$  to gas-phase acetone, yielding reduced Mo( $iv$ ). This species reacts with gas-phase propylene to form the OM active Mo(vi) alkylidene species. When Brønsted acid sites and  $MoO<sub>x</sub>$  species were located on the microporous support, rapid deactivation occurred during the ISOMET reaction.

Complementary characterization techniques revealed that in the  $13MoO<sub>3</sub>/HBEA$  sample Mo exists in highly dispersed tetrahedral and polymerized octahedral  $MoO<sub>x</sub>$  species, which are active in the OM reaction. However, a fraction of Mo is present as catalytically inactive crystalline  $MoO<sub>3</sub>$  phase. Tentatively it is presumed that the lower activity of  $6M_0O_3$ / HBEA is most probably due to the formation of hardly accessible  $MoO<sub>x</sub>$  species located in the microporous channels.

In order to achieve a more active and stable catalyst for the ISOMET of 1-hexene with ethylene, molybdena was loaded onto a gamma-alumina support  $(12M_0O_3/Al_2O_3)$ because at 12 wt% loading only highly dispersed  $MoO<sub>x</sub>$  is formed. Further experiments were performed with a mixed bed of  $12\text{MoO}_3/\text{Al}_2\text{O}_3$  (50%) and zeolite HBEA (50%) catalysts.

#### 1-Hexene ISOMET using  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and HBEA catalyst mixture

The activity and stability of the physical mixture of  $12\text{MoO}_3/$  $Al_2O_3$  and HBEA was studied in the ISOMET reaction of 1-hexene at 75 °C, ethylene pressure of 3 bar and 6 and 12  $g_{cat}$   $g_{1-H}^{-1}$  h space time (Fig. 8A and A', B and B') and compared with  $MoO<sub>3</sub>/HBEA$ . In these experiments, we compared the properties of catalysts with the same molybdena content but different weights, i.e., a mixture of 1 g of  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub> + 1$  g of HBEA as well as 1 g of  $13MoO<sub>3</sub>/$ 



Fig. 8 Propylene yield in ISOMET of 1-hexene (1-H) with ethylene over activated and reactivated catalyst mixture of HBEA and  $MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  as a function of time on stream (TOS) at space time of 6 (A, A′) and 12 (B, B')  $g_{cat} g_{1-H}^{-1}$  h, at 75 °C, 3 bar ethylene pressure, and ethylene/1-<br>hovene ratio of 10, activation: the mixture of av situ calcined eatshet hexene ratio of 10. Activation: the mixture of ex situ calcined catalyst was in situ pre-treated in a flow of Ar (50 ml min<sup>-1</sup>) at 550 °C for 2 h.<br>Boastivation: after <sup>7</sup> h TOS in the ISOMET reastion, the satalyst was Reactivation: after 3 h TOS in the ISOMET reaction, the catalyst was purged with Ar flow (50 ml min<sup>-1</sup>) at 550 °C for 2 h (see Table 2 for detailed conditions). After activation and reactivation the catalyst was cooled to the reaction temperature in a flow of Ar (50 ml min<sup>-1</sup>) and the ISOMET reaction was performed again.

HBEA. Over the freshly activated catalyst, the propylene yield was 33% after 0.5 h TOS. However, after reactivation of the catalyst the propylene yield more than doubled and reached a level of nearly 80%.

Repeated reactivation gave similar high propylene yield (Fig. S9†). Not only the initial activity of the catalyst improved significantly but the lifetime of the catalyst also increased. (Fig. 8A and A′).

The propylene yield on the activated catalyst mixture decreased from 33% to 18% in 3 h TOS. Deactivation of the reactivated catalyst is slower; the propylene yield dropped from 79% to only 57% in 3 h. Significantly lower initial activity and faster deactivation were observed for  $13\text{MoO}_3$ / HBEA (Table 2). Over the activated and reactivated  $13\text{MoO}_3/$ HBEA catalyst in three hours TOS the propylene yield decreased from 10% to 6% and 20% to 16%, respectively. The higher activity of the  $12\text{MoO}_3/\text{Al}_2\text{O}_3$  catalyst compared to 13MoO3/HBEA is due to the higher concentration of active  $MoO<sub>x</sub>$  species on alumina than on zeolite (microporous aluminosilicate), as confirmed by XRD and Raman and UVvis spectroscopy. Higher space time resulted in a significantly higher propylene yield. The catalyst reactivation has shown significantly higher (>90%) propylene yield and longer catalyst lifetime (Fig. 8B and B′). Catalysis Science & Technology<br>
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Further experiments were carried out to investigate the influence of the reaction temperature on the overall propylene yield (Fig. 9). At 125 °C, the propylene yield was around 50%. The catalyst activity did not change as a function of TOS. After reactivation of the catalyst, a propylene yield of more than 90% can be obtained. Some catalyst deactivation was observed; however, after 3 h TOS still 80% propylene yield could be measured.

By increasing the reaction temperature up to 150  $\degree$ C, even higher yield and longer catalyst lifetime was observed. After the catalyst reactivation approximately 100% propylene yield was found, which was maintained up to two hours. From the third hour of the reaction the propylene yield slightly decreased; however, it was still high (>80%). As was observed after the first run and catalyst reactivation, the catalyst performance was always significantly higher.

#### Activation and reactivation of  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and HBEA catalyst mixture

Conventional pre-treatment of the heterogeneous OM catalysts includes high-temperature calcination and inert gas purging. It has also been shown that olefin pre-treatment at low or high temperatures can improve the initial activity. Amakawa et  $al$ <sup>82</sup> reported that heat treatment in inert gas at 550 °C for 2 hours after the room-temperature olefin adsorption doubled the catalytic activity of  $MoO<sub>3</sub>/SBA-15$ . Surface isopropoxide species were supposed to form and activate surface  $Mo(w)$  sites by reduction to  $Mo(w)$  and formation of C3 oxygenate (acetone).

This reduced Mo species react with propylene and give active  $Mo(w)$ -alkylidene species. Two orders of magnitude



Fig. 9 Propylene yield in ISOMET of 1-hexene (1-H) with ethylene over activated and reactivated catalyst mixture of HBEA and  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$ as a function of time on stream (TOS) at 125 °C and 150 °C, 6  $g_{\text{cat}}$  $g_{1-H}^{-1}$  h space time, 3 bar ethylene pressure, and ethylene/1-hexene<br>ratio of 10. Activation: the mixture of ex city calcined catalyst was in ratio of 10. Activation: the mixture of ex situ calcined catalyst was in situ pre-treated in a flow of Ar (50 ml min<sup>-1</sup>) at 550 °C for 2 h.<br>Peactivation: after 3 h TOS in the ISOMET reaction the catalyst was Reactivation: after 3 h TOS in the ISOMET reaction the catalyst was purged with an Ar flow (50 ml min<sup>-1</sup>) at 550 °C for 2 h (see Table 2 for detailed conditions). After activation and reactivation, the catalyst was cooled to the reaction temperature in a flow of Ar (50 ml min<sup>-1</sup>) and the ISOMET reaction was performed under the conditions described above.

increase in activity was observed over silica-supported  $MoO<sub>3</sub>$ and  $WO<sub>3</sub>$  when propylene adsorption was performed at high temperature, 550 °C and 700 °C, respectively.<sup>83</sup> The hightemperature activation was explained by a pseudo-Wittig mechanism.

Activation in  $H<sub>2</sub>$  or Ar: the *ex situ* calcined catalyst was in situ pre-treated in 5% H<sub>2</sub>/Ar or Ar flow (50 ml min<sup>-1</sup>) at 550 °C for 2 h. Reactivation in Ar: after 3 h TOS in 1-hexene ISOMET the catalyst was purged in a flow of Ar  $(50 \text{ ml min}^{-1})$ for 0.5 h at 75 °C to desorb olefins, then heated to 550 °C at a ramp rate of 10 °C min−<sup>1</sup> and maintained at this temperature for 2 h. Reactivation in  $O_2$ : after 3 h TOS in 1-hexene ISOMET the catalyst was purged in a flow of  $O<sub>2</sub>$  (50 ml min<sup>-1</sup>) for 0.5 h at 75 °C to desorb olefins, heated to 550 °C at a ramp rate of 10 °C min−<sup>1</sup> and maintained at this temperature for 2 h, and then purged with Ar  $(50 \text{ ml min}^{-1})$ for 0.5 h and cooled to the reaction temperature. Activation in ethylene: after Ar-activation ethylene was fed into the reactor for 1 h at 75 °C then changed to Ar flow for 0.5 h and heated to 550 °C at a ramp rate of 10 °C min−<sup>1</sup> and maintained at this temperature for 2 h.

Others state that the active species is  $Mo(v).^{84}$  Thus, there is still some debate in the scientific community as to which catalyst oxidation state results in high catalytic activity. The surface  $Mo(v)$  oxide species are known to be catalytically

inactive;  $Mo(v)$  and  $Mo(w)$  are shown to be active. However, lowering the oxidation state results in catalytically inactive species again; therefore the appropriate activation is crucial for the high catalyst activity.

Fig. 10 shows the effect of different activation and reactivation on propylene yield in 1-hexene ISOMET over the  $12\text{MoO}_3/\text{Al}_2\text{O}_3$ –HBEA catalyst mixture. No propylene was formed over the  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and HBEA mixture oxidized in situ at 550  $\degree$ C, confirming that surface Mo(v<sub>I</sub>) oxide species are not active in the OM reaction. The experiment was stopped after 0.5 h TOS without waiting for the catalyst to be activated in situ during the reaction, as proved by Amakawa et  $al$ .<sup>62</sup>

The catalyst pretreated in a 5%  $H_2/Ar$  mixture at 550 °C for 2 hours has the lowest activity (Fig. 10 and S8 $\dagger$ ). At 75 °C and 3 bar ethylene pressure the propylene yield was about 15%. Higher propylene yield (∼28%) was obtained over Aractivated catalyst (Fig. 8 and 10). However, after performing the ISOMET reaction for 3 h TOS and treatment again in inert gas flow at 550 °C for 2 h, improved activity was observed. The propylene yield increased to 80%. When the catalyst was reactivated in an  $O_2$  flow at 550 °C for 1 h followed by purging with an Ar flow for 2 h no catalyst activity improvement was observed. It remained the same as that of the Ar-activated catalyst. Fig. 10 shows that improved ISOMET activity was also observed when only the crosscoupling agent ethylene was fed into the reactor at 75 °C (actually, self-metathesis proceeds) and the catalyst was heattreated in Ar at 550 °C (ethylene-activated catalyst). These results suggest that after 1-hexene ISOMET or ethylene OM, during the Ar reactivation the remaining olefins may also participate in the generation of the catalytically active molybdenum species.<sup>83</sup> With calcination of the catalyst in  $O_2$ flow at 550 °C Mo–carbene species are decomposed and Mo is oxidized to  $Mo(w)$ . **Paper** Catalyis Science Catalyis Science Article. Published on 19 Access Articles Conserved United Un

The XPS results show that during inert gas treatment at 550 °C some of the Mo(v<sub>I</sub>) are reduced to Mo(v) species. The ethylene-activated catalyst contains more Mo(V) species. The high catalyst performance can also be explained by the partial reduction of Mo initiated by the hydrocarbon residues remaining on the catalyst surface.

#### Propylene synthesis via ISOMET of FCC fractions using  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  and HBEA catalyst mixture

Our further aim was to develop an OM-based chemical process for a mixture produced by FCC cracking (further named as crude FCC and FCC light). The analysis of FCC light and FCC fractions revealed that the olefin content is about 37 wt% FCC light and 22 wt% crude FCC and approximately 50% of the olefins are 2-olefins (Fig. S10†). The main olefin components of the FCC light fraction are C5–C7 olefins; however, some C4 and C8 components could also be detected. Experiments have been carried out to synthetize propylene from the crude FCC fraction. Theoretically, regardless of the nature of the linear mono olefins (either terminal or internal) all olefins can be converted to propylene via ISOMET using ethylene (Scheme 1). In terms of the weight of olefins, approximately 100 tons of C4–C8 olefin mixture can be converted to 200 tons of propylene by using 100 tons ethylene.

The ISOMET of FCC light and FCC fractions was carried out without any pretreatment at 75 °C and 3 bar ethylene pressure. The FCC light fraction contained mainly C5–C7 olefin components. Due to the relatively high olefin content



Fig. 10 Propylene yield over activated and reactivated  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$ HBEA catalyst mixture in ISOMET of 1-hexene at 75 °C for 1 h TOS, 6 g<sub>cat</sub> g<sub>1−H</sub><sup>-1</sup> h space time under 3 bar ethylene pressure and ethylene/1-<br>boxane ratio of 10 hexene ratio of 10.



Fig. 11 Propylene yield in ISOMET of FCC light and FCC over activated and reactivated HBEA and  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  catalyst mixture as a function of time on stream at 75 °C and 6  $g_{\text{cat}} g_{\text{olefin}}^{-1}$  h space time under 3 bar ethylene pressure. Activation: the mixture of ex situ calcined catalyst was *in situ* pre-treated in a flow of Ar (50 ml min<sup>-1</sup>) at  $550.8$  for 2 h, Boastivation: after 3 h, TOS in JSOMET reaction the 550 °C for 2 h. Reactivation: after 3 h TOS in ISOMET reaction the catalyst was purged with Ar flow (50 ml min<sup>-1</sup>) at 550 °C for 2 h (see Table 2 for detailed conditions). After activation and reactivation, the catalyst was cooled to the reaction temperature in a flow of Ar (50 ml min<sup>-1</sup>) and the ISOMET reaction was performed under the conditions described above.

(35–37%) and purity, high propylene yield (70%) and long catalyst lifetime were obtained (Fig. 11). Following the reactivation significantly higher (90%) propylene yield and longer catalyst lifetime were observed. After 8 hours TOS the catalyst remained still active and the propylene yield reached 50%. Compared to the  $12MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub> - HBEA$  catalyst mixture, significantly lower propylene yield could be obtained using the  $12MoO<sub>3</sub>/HBEA$  catalyst (Fig. S11†).

The ISOMET of FCC showed that after 0.5 h TOS slightly higher than 45% propylene yield was obtained, which declined steadily in the next two hours (Fig. 11). The catalyst reusability was also investigated. It was found that experiments carried out with reactivated catalyst resulted in significantly higher (75%) propylene yield and longer catalyst lifetime.

## **Experimental**

### Catalyst preparation

The  $MoO<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub>$  olefin MET catalyst was prepared by incipient wetness impregnation of the alumina support. Commercial  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> (Ketjen, CK 300) was first calcined at 500 °C for 8 h (at a heating rate of 10 °C min−<sup>1</sup> ) and then impregnated with ammonium heptamolybdate solution. The sample was air-dried at 120 °C for 6 h and then slowly heated (2 °C min<sup>-1</sup>) to 500 °C and kept at this temperature for 12 h. The  $MoO<sub>3</sub>$  content of the air-calcined catalyst was 11.7 wt%, so the catalyst is designated as  $12\text{MoO}_3/\text{Al}_2\text{O}_3$ .

MoO3/HBEA ISOMET catalysts were prepared by wet impregnation of H-Beta (HBEA; Si/Al = 12; SÜD-Chemie AG, München) with ammonium heptamolybdate solution. Before impregnation, zeolite HBEA was calcined at 500 °C for 8 h. After impregnation, the solvent was evaporated by keeping the preparation at 120  $\degree$ C for 6 h. The samples were then heated to 500 °C in air and kept at this temperature for 12 h. The air-calcined samples are designated as  $x\text{MoO}_3/\text{HBEA}$ . The  $MoO<sub>3</sub>$  contents of the catalysts were 13.5 and 6.1 wt%.

### Catalyst characterization

X-ray powder diffraction patterns of the catalysts were recorded by a Philips PW 1810/3710 diffractometer equipped with a graphite monochromator. CuK $\alpha$  radiation ( $\lambda = 0.1541$ nm) was used. The X-ray tube was set at 40 kV and 35 mA current. The scan step size was 0.02 degrees 2-theta, whereas the scan time was five seconds in each step.

The specific surface area (SSA) of the catalysts was calculated from the  $N_2$  adsorption isotherms using the Brunauer–Emmett–Teller (BET) method. The adsorption isotherms were measured at −196 °C using a Thermo Scientific Surfer automated gas sorption instrument. Before measurement, samples were evacuated at 250 °C for 2 h in high vacuum (∼10−<sup>6</sup> mbar).

The reducibility of the catalysts was studied using temperature-programmed  $H_2$ -reduction ( $H_2$ -TPR). About 100 mg of sample was treated in a 30 ml min<sup>-1</sup> O<sub>2</sub> flow at 500 °C for 1 h in a quartz reactor tube (6 mm ID). To obtain the TPR

The acidity of the catalysts was characterized using a Nicolet Impact Type 400 spectrometer applying the selfsupported wafer technique and using an *in situ* cell. The acid site concentrations were determined using transmission Fourier-transform infrared (FT-IR) spectra of the adsorbed pyridine (Py) on Brønsted- and/or Lewis acid sites of the catalysts. IR spectra of the sample were recorded at room temperature, averaging 32 scans at a resolution of 2  $cm^{-1}$ . Spectra were normalized to a wafer thickness of 5 mg  $cm^{-2}$ . Difference spectra of adsorbed Py were generated by subtracting the spectrum of the wafer before pyridine adsorption from the spectrum of the wafer loaded by pyridine. Catalysis Science & Technology<br>
Osea Technology (13-3-7%) and putity, high propylene yield (70%) and long<br>
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The in situ UV/vis spectra were collected using a Thermo Scientific Evolution 300 UV/vis spectrophotometer equipped with a Praying Mantis diffuse reflectance accessory and a high temperature and pressure reaction chamber. The catalysts were diluted 50–50 wt% with BaSO4.

Raman analyses were carried out at the Research and Industrial Relations Center of the Faculty of Science, Eötvös Loránd University of Budapest, using a HORIBA JobinYvon LabRAM HR 800 Raman microspectrometer. A frequency doubled Nd-YAG green laser with a 532 nm excitation wavelength was employed, displaying 10–20 mW on the sample surface. An OLYMPUS  $\times$  50 (N.A. = 0.6) objective was used to focus the laser. For Raman analyses, a 100 μm confocal hole, 600 grooves per mm optical grating, and 20– 120 s cumulated exposition time were used. The spectral resolution of measurements was  $3.0 \text{ cm}^{-1}$ .

The XPS measurements using self-supported wafers were performed by using an Omicron EA 125 electron spectrometer in the fixed analyser transmission mode; photoelectrons were excited by non-monochromatized MgKα (1253.6 eV) radiation. Detailed spectra were recorded with a pass energy of 30 eV, providing a resolution of around 1 eV. Spectra were collected in the as-received state of the catalysts, after 2 h annealing in 300 mbar Ar at 550 °C (simulated activation), after 3 h exposure to 100 mbar  $C_2H_4$  at around 100 °C (simulated reaction) and after repeated activation at 550 °C. The treatments were performed in a preparation chamber attached to the electron spectrometer so no exposure to the atmosphere was needed during sample transfer. Spectra were processed with the CasaXPS package [N. Fairley, "CasaXPS: Spectrum Processing Software for XPS, AES and SIMS," Cheshire, 2006, [http://www.casaxps.com\]](http://www.casaxps.com), while quantitative evaluation of the data was performed with the XPSMultiQuant package [M. Mohai, "XPS MultiQuant: Multimodel XPS Quantification Software," Surf. Interface Anal. 36 (2004) 828–832, M. Mohai, "XPS MultiQuant: Multimodel X-ray photoelectron spectroscopy quantification

program," 2011, [http://aki.ttk.hu/XMQpages/XMQhome.php\]](http://aki.ttk.hu/XMQpages/XMQhome.php). As the samples were insulating, charge compensation was performed by setting the binding energy of the main C 1s peak (arising from adventitious hydrocarbon) of the asreceived catalysts to 285.0 eV.

### Catalytic test

The catalytic experiments were carried out in a downstream, fixed-bed, stainless-steel tube reactor (12 mm ID). The reactor was loaded with 2 g of catalyst grains prepared by compression of the sample powder, crushing and sieving (0.315–0.63 mm) and the flow rate of the reactant liquid mixture  $(1,5-6 \text{ ml h}^{-1})$  was varied. The reactor temperature was controlled using an Omron E5CN controller. The liquid mixture (1-hexene in hexane solution (30%), FCC; FCC light) was fed into the reactor by a high-pressure micro pump (Teledyne Isco, Model 100DM). The gas was introduced by a mass flow controller (Aalborg GFC17). The total pressure and the reaction temperature were varied in the range of 1–50 bar and 25–150 °C, respectively. Using water cooling, the reactor effluent was separated to liquid and gas products. The liquid product mixtures were analysed by GC-MS (Shimadzu QP-2010) using a 60 m ZB-WAX PLUS capillary column. The gaseous reactor effluent was analysed for detection of propylene and light hydrocarbons using an on-line gas chromatograph (Varian 3300) with a flame ionization detector (FID) and applying a 30 m Supelco (alumina/chloride) capillary column. **Paper** Catalysis Science & we are included on 19 July 2021. The complete is determined on the theoretical on the specific terminal was the common about the specific is license to the common about the specific is licensed

The propylene yield as outcome parameter was determined; in some cases complete mass balance including the quantitative distribution of the intermediate species was calculated.

#### Catalyst activation/reactivation

Activation and reactivation of the catalysts were performed in an inert gas (Ar) flow at 550 °C for 2 h.

Activation: the ex situ calcined catalysts were in situ pretreated in a flow of Ar (50 ml min−<sup>1</sup> ) at atmospheric pressure and 550 °C for 2 h. The activated catalyst was cooled to the target temperature in a flow of Ar and the ISOMET reaction was performed.

Reactivation: after 3 h TOS, the reactant feed was stopped and the catalyst was purged with a flow of Ar  $(50 \text{ ml } min^{-1})$ for 30 min at reaction temperature and atmospheric pressure. Then it was heated to 550 °C at a ramp rate of 10 °C min−<sup>1</sup> and maintained at this temperature for 2 h. The reactivated catalyst thus obtained was cooled to the target temperature in a flow of Ar and the ISOMET reaction was performed again.

After the ISOMET reaction, the catalyst was purged with an Ar flow at 75 °C for 0.5 h and the effluent was analysed by GC-MS. It was found that the reactant and the intermediate olefins (ethylene, propylene, butenes, pentenes and hexenes) completely desorbed from the catalyst surface within 30 min. With increasing temperature longer olefins/paraffins  $(>C6)$  were detected by GC-MS. Above 300 °C there were no further organic species observed.

### General procedure for isomerization with HBEA

The isomerization catalyst (HBEA) was pre-treated in situ in Ar (50 ml min−<sup>1</sup> ) flow in the reactor at atmospheric pressure and 450 °C for 2 h. After catalyst pre-treatment the reactor was cooled to the target temperature under an Ar flow. The isomerization of 1-hexene was investigated at the temperature range of 75–150 °C, 6  $g_{cat} g_{1-hexene}^{-1}$  h space time in Ar flow at atmospheric pressure.

General procedure for ethenolysis. After catalyst pretreatment the reactor was cooled to the target temperature under Ar flow and then pressurized to the target pressure with ethylene. The ethenolysis of the reactants was investigated at different temperatures (25–150 °C), pressure (1–50 bar) and space time (6–12  $g_{cat} g_{reactant}^{-1}$  h). The ethylene/1-hexene molar ratio was 10.

## **Conclusions**

Heterogeneous molybdena catalysts were prepared using alumina and zeolite beta (HBEA) as support. The catalysts and their mixtures were characterized and studied in the ISOMET of 1-hexene and FCC olefins. It was shown that the Brønsted sites of HBEA catalyze the double-bond isomerization, while the  $MoO<sub>x</sub>$  species supported on alumina or HBEA are active in the ethenolysis of olefins to give propylene as the final product from both 1-hexene and FCC olefins. Over a mixture of  $12\text{MoO}_3/\text{Al}_2\text{O}_3$  and HBEA catalysts, quantitative conversion of 1-hexene to propylene could be achieved at 150 °C, 3 bar ethylene pressure and 6  $g<sub>cat</sub>$  $g_{1-\text{hexene}}^{-1}$  h space time. The catalyst mixture also had good performance in the ethenolysis of FCC olefins. Relative to the  $12\text{MoO}_3/\text{Al}_2\text{O}_3$ -HBEA system, the  $13\text{MoO}_3/\text{HB}$ EA catalyst rendered lower activity and significantly shorter catalyst lifetime presumably due to the lower amount of highly dispersed  $MoO<sub>x</sub>$  sites formed on the zeolite.

The deactivated ISOMET catalyst system could be reactivated by treatment in an argon flow at 550 °C for 2 hours. The reactivated catalysts showed higher initial activity and stability than the fresh catalysts pre-treated in the same way. It has been demonstrated that treatment of the catalyst with olefins such as ethylene before inert gas activation resulted in significant catalyst activity improvement. It indicates that the adsorbed olefins may play a key role in the formation of active metal centers during the catalyst reactivation process. Activating the catalyst either in a strongly reducing or in an oxidizing environment, however, led to the loss of activity.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors acknowledge the support of the project No. VEKOP-2.3.2-16-2017-00013 financed by the European Union and the State of Hungary, co-financed by the European Regional Development Fund. The financial support of the National Research, Development and Innovation (NRDI) Office (2019-2.1.13-TÉT\_IN-2020-00043) is also greatly acknowledged. Further thanks are due to COST Action CA17128 (LignoCOST) "Establishment of a Pan-European Network on the Sustainable Valorization of Lignin" for initiating this work. The authors also thank Dr. László E. Aradi for the Raman experiments and the anonymous reviewers for the fruitful suggestions. Catalysis Science & Technology<br>
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## References

- 1 R. Beucher, R. D. Andrei, C. Cammarano, A. Galarneau, F. Fajula and V. Hulea, ACS Catal., 2018, 8, 3636–3640.
- 2 S. Vorakitkanvasin, W. Phongsawat, K. Suriye, P. Praserthdam and J. Panpranot, RSC Adv., 2017, 7, 38659–38665.
- 3 G. Zuo, Y. Xu, J. Zheng, F. Jiang and X. Liu, RSC Adv., 2018, 8, 8372–8384.
- 4 G. Chen, M. Dong, J. Li, Z. Wu, G. Wang, Z. Qin, J. Wang and W. Fan, Catal. Sci. Technol., 2016, 6, 5515–5525.
- 5 R. D. Andrei, M. I. Popa, F. Fajula, C. Cammarano, A. Al Khudhair, K. Bouchmella, P. H. Mutin and V. Hulea, ACS Catal., 2015, 5, 2774–2777.
- 6 W. Jiang, X. Mo, S. Feng, F. Xu, G. Zhou, H. Zhou, C. Xu and B. Chen, Mol. Catal., 2017, 442, 49–56.
- 7 W. Jiang, R. Huang, P. Li, S. Feng, G. Zhou, C. Yu, H. Zhou, C. Xu and Q. Xu, Appl. Catal., A, 2016, 517, 227–235.
- 8 S. Lwin and I. E. Wachs, ACS Catal., 2017, 7, 573–580.
- 9 H. Liu, S. Huang, L. Zhang, S. Liu, W. Xin and L. Xu, Catal. Commun., 2009, 10, 544–548.
- 10 V. Hulea, Catal. Sci. Technol., 2019, 9, 4466–4477.
- 11 D. Zhang, X. Li, S. Liu, X. Zhu, F. Chen and L. Xu, Chem. Asian J., 2015, 10, 1647–1659.
- 12 K. Gayapan, S. Sripinun, J. Panpranot, P. Praserthdam and S. Assabumrungrat, RSC Adv., 2018, 8, 11693–11704.
- 13 M. Stoyanova, U. Bentrup, H. Atia, E. V. Kondratenko, D. Linke and U. Rodemerck, Catal. Sci. Technol., 2019, 9, 3137–3148.
- 14 X. Li, X. Zhu, D. Zhang, F. Chen, P. Zeng, S. Liu, S. Xie and L. Xu, J. Energy Chem., 2013, 22, 145–150.
- 15 J. C. Mol, J. Mol. Catal. A: Chem., 2004, 213, 39–45.
- 16 R. L. Banks, D. S. Banasiak, P. S. Hudson and J. K. Norell, Chem. Eng. Prog., 1979, 75, 73.
- 17 G. Parkinson, Chem. Eng., 2001, 108, 27–35.
- 18 T. Otroshchenko, O. Reinsdorf, D. Linke and E. V. Kondratenko, Catal. Sci. Technol., 2019, 9, 5660–5667.
- 19 O. V. Klimov, O. S. Alekseev and A. N. Startsev, React. Kinet. Catal. Lett., 1995, 56, 143–150.
- 20 M. A. Ibrahim, M. N. Akhtar, J. Čejka, E. Montanari, H. Balcar, M. Kubů and S. S. Al-Khattaf, J. Ind. Eng. Chem., 2017, 53, 119–126.
- 21 H. Balcar, D. Mishra, E. Marceau, X. Carrier, N. Žilková and Z. Bastl, Appl. Catal., A, 2009, 359, 129–135.
- 22 R. D. Andrei, M. I. Popa, C. Cammarano and V. Hulea, New J. Chem., 2016, 40, 4146–4152.
- 23 N. Poovarawan, K. Suriye, J. Panpranot, W. Limsangkass, A. Guntida, F. J. C. S. Aires and P. Praserthdam, Catal. Today, 2018, 309, 43–50.
- 24 S. Maksasithorn, P. Praserthdam, K. Suriye and D. P. Debecker, Microporous Mesoporous Mater., 2015, 213, 125–133.
- 25 D. P. Debecker, M. Stoyanova, U. Rodemerck, F. Colbeau-Justinc, C. Boissère, A. Chaumonnot, A. Bonduelle and C. Sanchez, Appl. Catal., A, 2014, 470, 458–466.
- 26 M. Selva, S. Guidi, A. Perosa, M. Signoretto, P. Licence and T. Maschmeyer, Green Chem., 2012, 14, 2727–2737.
- 27 L. Sang, S. L. Chen, G. Yuan, M. Zheng, J. You, A. Chen, R. Li and L. Chen, J. Nat. Gas Chem., 2012, 21, 105–108.
- 28 S. Lwin and I. E. Wachs, ACS Catal., 2014, 4, 2505–2520.
- 29 M. Sibeijn and J. C. Mol, J. Mol. Catal., 1992, 76, 345–358.
- 30 P. Schwab, R. Schulz, M. Schulz, B. Breitscheidel and G. Meyer, US6166279, 2000.
- 31 N. P. Christopher, E. Mazoyer, M. Taoufik, J. M. Basset, P. T. Barger and J. E. Rekoske, US8704029, 2014.
- 32 S. Choi and B. Ramachandran, US9611193, 2017.
- 33 M. Röper and J. Stephan, US2004/0242944A1, 2004.
- 34 V. A. Dang, L. Zhang, J. T. Ruszkay, E. Morales and S. T. Coleman, WO2017/079509Al, 2017.
- 35 C. Zhang, X. Liu, T. Liu, Z. Jiang and C. Li, Appl. Catal., A, 2019, 575, 187–197.
- 36 Y. Zhang, H. Wang, L. Zhao, J. Gao and C. Xu, Sep. Purif. Technol., 2019, 228, 115757.
- 37 C. P. Park, M. M. Van Wingerden, S. Y. Han, D. P. Kim and R. H. Grubbs, Org. Lett., 2011, 13, 2398–2401.
- 38 P. D. Nieres, J. Zelin, A. F. Trasarti and C. R. Apesteguía, Catal. Sci. Technol., 2016, 6, 6561–6568.
- 39 B. Szabó, G. Novodárszki, Z. May, J. Valyon, J. Hancsók and R. Barthos, Mol. Catal., 2020, 491, 110984.
- 40 S. Jeong, H. Kim, J. H. Bae, D. H. Kim, C. H. F. Peden, Y. K. Park and J. K. Jeon, Catal. Today, 2012, 185, 191–197.
- 41 J. B. Higgins, R. B. LaPierre, J. L. Schlenker, A. C. Rohrman, J. D. Wood, G. T. Kerr and W. J. Rohrbaugh, Zeolites, 1988, 8, 446–452.
- 42 B. Yilmaz and U. Müller, Top. Catal., 2009, 52, 888–895.
- 43 X. Li, W. Zhang, S. Liu, X. Han, L. Xu and X. Bao, J. Mol. Catal. A: Chem., 2006, 250, 94–99.
- 44 J. Guan, G. Yang, D. Zhou, W. Zhang, X. Liu, X. Han and X. Bao, Catal. Commun., 2008, 9, 2213–2216.
- 45 S. Liu, S. Huang, W. Xin, J. Bai, S. Xie and L. Xu, Catal. Today, 2004, 93–95, 471–476.
- 46 E. A. Buluchevskii, L. F. Saifullina, A. V. Lavrenov, T. R. Karpova and N. A. Glazov, Russ. J. Appl. Chem., 2017, 90, 1893–1899.
- 47 M. Röper, G. P. Schindler and J. Stephan, EP1379489A1, 2002.
- 48 X. Li, W. Zhang, S. Liu, L. Xu, X. Han and X. Bao, J. Phys. Chem. C, 2008, 112, 5955–5960.
- 49 E. Hillerová, H. Morishige, K. Inamura and M. Zdražil, Appl. Catal., A, 1997, 156, 1-17.
- 50 B. Wang, G. Ding, Y. Shang, J. Lv, H. Wang, E. Wang, Z. Li, X. Ma, S. Qin and Q. Sun, Appl. Catal., A, 2012, 431–432, 144–150.
- 51 I. E. Wachs, Catal. Today, 1996, 27, 437–455.
- 52 E. Mannei, F. Ayari, M. Mhamdi, M. Almohalla, A. Guerrero Ruiz, G. Delahay and A. Ghorbel, Microporous Mesoporous Mater., 2016, 219, 77–86.
- 53 S. Rajagopal, H. J. Marini, J. A. Marzari and R. Miranda, J. Catal., 1994, 147, 417–428.
- 54 Y. Song, C. Sun, W. Shen and L. Lin, Appl. Catal., A, 2007, 317, 266–274.
- 55 R. W. Borry, Y. H. Kim, A. Huffsmith, J. A. Reimer and E. Iglesia, J. Phys. Chem. B, 1999, 103, 5787–5796.
- 56 A. Martínez, E. Peris, M. Derewinski and A. Burkat-Dulak, Catal. Today, 2011, 169, 75–84.
- 57 P. A. Wright, W. Zhou, J. Pérez-Pariente and M. Arranz, J. Am. Chem. Soc., 2005, 127, 494–495.
- 58 W. Ding, G. D. Meitzner and E. Iglesia, J. Catal., 2002, 206, 14–22.
- 59 T. Hahn, U. Bentrup, M. Armbrüster, E. V. Kondratenko and D. Linke, ChemCatChem, 2014, 6, 1664–1672.
- 60 H. Knözinger and H. Jezlorowski, J. Phys. Chem., 1978, 82, 2002–2005.
- 61 H. Hu and I. E. Wachs, J. Phys. Chem., 1995, 99, 10897–10910.
- 62 G. Mestl and H. Knozinger, Langmuir, 1998, 7463, 3964–3966.
- 63 C. C. Williams, J. G. Ekerdt, J. M. Jehng, F. D. Hardcastle, A. M. Turek and I. E. Wachs, J. Phys. Chem., 1991, 95, 8781–8791.
- 64 C. C. Williams, J. G. Ekerdt, J. M. Jehng, F. D. Hardcastle and I. E. Wachs, J. Phys. Chem., 1991, 95, 8791–8797.
- 65 E. L. Lee and I. E. Wachs, J. Phys. Chem. C, 2007, 111, 14410–14425.
- 66 D. P. Debecker, B. Schimmoeller, M. Stoyanova, C. Poleunis, P. Bertrand, U. Rodemerck and E. M. Gaigneaux, *J. Catal.*, 2011, 277, 154–163.
- 67 J. T. Kloprogge, L. V. Duong, B. J. Wood and R. L. Frost, J. Colloid Interface Sci., 2006, 296, 572–576.
- 68 M. Harju, S. Areva, J. B. Rosenholm and T. Mäntylä, Appl. Surf. Sci., 2008, 254, 5981–5989.
- 69 B. M. Reddy, B. Chowdhury, E. P. Reddy and A. Fernández, Appl. Catal., A, 2001, 213, 279–288.
- 70 Y. V. Plyuto, I. V. Babich, I. V. Plyuto, A. D. Van Langeveld and J. A. Moulijn, Appl. Surf. Sci., 1997, 119, 11–18.
- 71 P. Novotný, S. Yusuf, F. Li and H. H. Lamb, J. Chem. Phys., 2020, 152, 044713.
- 72 J. Baltrusaitis, B. Mendoza-Sanchez, V. Fernandez, R. Veenstra, N. Dukstiene, A. Roberts and N. Fairley, Appl. Surf. Sci., 2015, 326, 151–161. **Paper Cathyis Science, A view Access Article on 2021.** This article is licensed on 2021. This article is licensed on 2021. This article is licensed under a method on 2021. Downloaded the simulation 2021. Downloaded the s
	- 73 W. Grünert, A. Y. Stakheev, W. Mörke, R. Feldhaus, K. Anders, E. S. Shpiro and K. M. Minachev, J. Catal., 1992, 135, 269–286.
	- 74 B. W. Wojciechowski and A.-N. Ko, Int. J. Chem. Kinet., 1983, 15, 1249–1274.
	- 75 J. Abbot, A. Corma and B. W. Wojciechowski, J. Catal., 1985, 92, 398–408.
	- 76 M. Pérez-Luna, A. Cosultchi, J. A. Toledo-Antonio and L. Díaz-Garcia, Catal. Lett., 2009, 128, 290–296.
	- 77 X. Li, W. Zhang, X. Li, S. Liu, H. Huang, X. Han, L. Xu and X. Bao, J. Phys. Chem. C, 2009, 113, 8228–8233.
	- 78 X. Li, A. Zheng, J. Guan, X. Han, W. Zhang and X. Bao, Catal. Lett., 2010, 138, 116–123.
	- 79 X. Li, J. Guan, D. Zhou, G. Li, X. Han, W. Zhang and X. Bao, J. Mol. Struct.: THEOCHEM, 2009, 913, 167–172.
	- 80 X. Xiaoding, C. Boelhouwer, D. Vonk, J. I. Benecke and J. C. Mol, J. Mol. Catal., 1986, 36, 47–66.
	- 81 A. Andreini, X. Xiaoding and J. C. Mol, Appl. Catal., 1986, 27,  $31 - 40.$
	- 82 K. Amakawa, S. Wrabetz, J. Kröhnert, G. Tzolova-Müller, R. Schlögl and A. Trunschke, J. Am. Chem. Soc., 2012, 134, 11462–11473.
	- 83 K. Ding, A. Gulec, A. M. Johnson, T. L. Drake, W. Wu, Y. Lin, E. Weitz, L. D. Marks and P. C. Stair, ACS Catal., 2016, 6, 5740–5746.
	- 84 B. Zhang, N. Liu, Q. Lin and D. Jin, J. Mol. Catal., 1991, 65, 15–28.