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Nickel-aluminosilicate catalysts for ethylene oligomerization: recent scientific progress

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Significant scientific effort has been made during the last decades to develop heterogeneous catalysts and processes for ethylene oligomerization. Among the reported catalysts, Ni-aluminosilicates are regarded as the most promising candidates. This paper reviews the recent advancements in ethylene conversion catalyzed by Ni-aluminosilicates. The main fundamental and practical aspects on this topic, including types of catalysts depending on the support, active nickel sites, oligomerization mechanism and kinetics and catalyst deactivation, are examined. The multi-reaction catalytic processes in which oligomerization is the key step have been also discussed.

1. Introduction

With an annual worldwide production of about 150 million tonnes, ethylene is a key molecule for producing other major platform molecules, end-use chemicals and polymers. Additionally, ethylene is one of the most promising bio-based chemicals, mainly because its high-volume production by bioethanol dehydration became economically feasible. The dimerization/oligomerization of ethylene is of considerable interest for the synthesis of butenes and higher olefins. This reaction can be efficiently catalysed by both homogeneous and heterogeneous catalysts containing metal species as active sites, as shown in recent reviews.^{1,2} Note that the first review article describes for the most part homogeneous oligomerization catalysts and only a little bit of heterogeneous catalysts,¹ while the second one focused on oligomerization on metal-containing zeolites.²

In line with sustainable chemistry principles, during the last few decades, research attention has been focused on the development of oligomerization processes based on heterogeneous catalysts. By far, Ni-based materials have been

developed and used as heterogeneous catalysts in these research studies.

Ten years ago, I co-authored an extended review, providing knowledge on this topic at the time.³ It has been shown that Ni-exchanged microporous and mesoporous aluminosilicates are very efficient catalysts for this important application. It was also specified that, despite the scientific progress achieved at that time, some key aspects, including the nature of the Ni species, the active sites involved in the catalytic act, the oligomerization mechanism, the effect of the catalyst texture or the effect of the reaction parameters on the oligomerization, were not yet clearly elucidated. To answer the questions that remained open, a notable research effort has been conducted during the last decade.

Based on the recent literature, I try here to identify the progress attained in this scientific area. The first section discusses the main families of Ni-aluminosilicate catalysts used in ethylene oligomerization, paying principal attention to their composition and texture. In the next two major sections, I examine the nature of the active sites and identify them, as well as the elementary reactions involved in the oligomerization mechanism. Then, the effect of the contaminants and the parameters on the oligomerization process are evaluated. Finally, the multi-reaction processes, such as fuel or propylene production, where ethylene

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Vasile Hulea has been full professor of heterogeneous catalysis at the National School of Chemistry of Montpellier (France) until 2023 and currently holds the position of Professor Emeritus at the “Charles Gerhardt” Institute of Montpellier. He has worked tirelessly in catalysis research for more than 40 years. His research activities focused on synthesis and characterization of microporous and mesoporous catalysts, their reactivity in acid–base and redox catalysis, and kinetic and mechanistic aspects. He developed original catalysts and friendly processes for valorisation of renewable/fossil sources and wastes (light olefin oligomerization & metathesis, methanol conversion to diesel and mild distillate, aromatics alkylation, mercaptan conversion, and mild oxidation of S-containing organic compounds). Prof. Hulea has (co)-authored more than 200 research articles, patents and book chapters in the fields of materials science and catalysis.



oligomerization is a crucial step, are discussed. The major contributions of my group in the field are underscored in this review.

2. Ni-aluminosilicate catalysts for ethylene oligomerization

Metal-containing microporous and mesoporous aluminosilicates are efficient catalysts for many industrial and academic chemical applications. Among them, Ni-containing aluminosilicates have proven their high catalytic potential in reactions such as selective hydrogenation and methane reforming or low olefin oligomerization. While the hydrogenation and reforming reactions require metallic nickel as catalytic centers,^{4,5} the oligomerization of olefins is activated by the isolated ionic Ni sites.^{1,3} Ni-aluminosilicates are considered the most promising heterogeneous catalysts for ethene oligomerization carried out at moderate temperatures, without any co-catalysts and activators. The catalysts are typically obtained by ion exchange or impregnation of different supports. Table 1 lists the main families of Ni-containing catalysts prepared, characterized and used during the last decade in ethylene oligomerization.

Ni-containing zeolites

Zeolites are ordered crystalline microporous aluminosilicates, which are widely used as catalysts for many industrial applications. Thanks to their properties, such as high thermal, mechanical and chemical stability, these materials were frequently used as carriers for preparing Ni-containing catalysts, which showed real abilities in ethylene oligomerization.

The major drawback exhibited by Ni-based zeolites was their low stability against deactivation.^{3,17} Indeed, they often suffered severe deactivation, mainly due to the blocking of micropores with heavy products. As shown in Table 1, Ni-Beta was the most used catalyst among the Ni-zeolites. Beta is a 3D zeolite, with pores of 12 MR (0.66 × 0.67; 0.56 × 0.56 nm). This topology is favorable for the transfer of bulky molecules, which are responsible for catalyst deactivation.

Usually, the nickel species were incorporated into the zeolite support using two post-synthesis procedures, namely ionic exchange^{9–11,13,15,16,23,29} and incipient wet impregnation.^{6,7,11,12,14,19,22,25,27} The general protocol applied for preparing Ni-zeolites by ionic exchange consists of three steps: (i) exchange of the as-synthesized Na-zeolite with aqueous NH₄NO₃ which results in NH₄-zeolite; (ii) exchange with aqueous Ni(NO₃)₂ and (iii) thermal treatment. Fig. 1 shows such a protocol, used by McCaig and Lamb¹⁴ for preparing the Ni-Beta catalyst. The authors found that [NiOH]⁺ and H⁺ were the primary charge-compensating cations in the uncalcined catalyst, as evidenced by TPR (Fig. 2). More generally, it is accepted that in the uncalcined Ni-exchanged catalyst the nickel ions are in a hydrated state, similar to isolated Ni hexaaqua ions.⁸³ To remove the water

ligands and thus release the Ni ions, heat treatment at temperatures above 500 °C is mandatory prior to catalytic application.⁸⁴

The impregnation approach was also used for incorporating the Ni species in zeolites. This is a simpler method, but it often produces a large amount of NiO, which is not active in the oligomerization of ethylene.³

Ni-containing mesoporous catalysts

Ni-mesoporous aluminosilicates, which possess large pore diameters, were also widely used in ethylene oligomerization. These catalysts have taken advantage of the increasing knowledge in materials science during the last decades. It allowed the design of tailored mesoporous materials, which are able to improve the catalytic activity and stability.

Catalysts with different topologies including Ni-ALMCM-41,^{12,17,48–54} Ni-ALSBA-15,^{36–42} and Ni-KIT-6 (ref. 43–47) revealed excellent behavior in terms of catalytic activity and deactivation stability. For example, productivities up to 175 g of oligomers per gram of catalyst per hour and high conversions during 80 h on stream were obtained over Ni-ALSBA-15, in both batch and flow modes by Andrei *et al.*⁴⁰ This result, obtained at 150 °C and 3.5 MPa, was superior to those exhibited by other Ni-based heterogeneous catalysts, without using alkylaluminum cocatalysts. The outstanding behavior exhibited by the Ni-mesoporous materials has been attributed to their pores, which are large enough to allow free diffusion of large molecules, resulting in a lower deactivation rate.¹⁷

The general protocol applied for preparing these catalysts starts either from Al-containing mesostructured silica (in the case of Ni-ALMCM-41) or from mesostructured silica (in the case of Ni-ALSBA-15 and Ni-AIKIT-6). Such a protocol is given in Scheme 1. The Al-containing sample was obtained from SBA-15 silica by grafting with sodium aluminate. The nickel ions were incorporated into aluminosilicates using an ionic exchange procedure, and after the thermal treatment at 550 °C, a bifunctional catalyst with both Ni and acid sites was obtained.

Although Ni-mesoporous aluminosilicates exhibit good catalytic activity, their preparation cost is high enough, which makes them difficult to apply on a commercial scale.

Ni-non-ordered aluminosilicates, also known as *Ni-amorphous aluminosilicates* (Ni-ASA), are also considered as promising catalysts for ethylene oligomerization, thanks to their merits of simple/easy synthesis and cheapness. Additionally, their catalytic properties, such as the moderate strength acid sites and mesoporous texture, are comparable to those exhibited by the Ni-based ordered mesoporous catalysts. Note that amorphous silica–alumina was the earlier support for nickel-based oligomerization catalysts.³

Inspecting the recent literature, we can find many sources of ASA supports used for the nickel species: commercial



Table 1 Microporous and mesoporous aluminosilicates-supported Ni catalysts for ethylene oligomerization

Catalyst	Support topology	Reaction conditions	Main aim of the study	Ref.
<i>Ni-microporous materials</i>				
Ni-Beta	Beta zeolite	180 °C, 0.1 MPa	Active sites, mechanism	6
Ni-Beta	Beta zeolite	120 °C, 0.1 MPa	Active sites, mechanism	7
Ni-Beta	Beta zeolite	180 °C, 0.1 MPa	Catalyst deactivation	8
Ni-Beta	Beta and ZSM-5	200 °C, 3.5 MPa	Beta vs. ZSM-5: crystal morphology effect	9
Ni-ZSM-5	zeolites			
Ni-Beta	Beta zeolite	30–300 °C, 0.1 MPa	Subcritical and supercritical conditions, coke formation	10
Ni-Beta	Beta zeolite	200 °C, 3.5 MPa	Active sites	11
Ni-Beta	Beta zeolite	120 °C, 3.5 MPa	Active sites, Beta vs. MCM-41 and SIRAL: effect of the support	12
Ni-Beta	Beta zeolite	120 °C, 3.5 MPa	Active sites, Beta vs. ASA and Al ₂ O ₃ effect of the support	13
Ni-Beta	Beta zeolite	225 °C, 1.1 MPa	Active sites	14
Ni-Beta	Beta zeolite	100 °C, 2.8 MPa	Active sites, mechanism	15
Ni-Beta	Beta and ZSM-5	300 °C, 0.1 MPa	Beta vs. ZSM-5: effect of pore topology, Ni sites, and acid sites	16
Ni-ZSM-5	zeolites			
Ni-Beta	Beta and FAU	–30, –15 °C,	Beta and FAU vs. MCM-41: Ni site deactivation, working	17
Ni-FAU	zeolites	0.1–2.4 MPa	at low <i>T</i>	
Ni-Beta	Beta zeolite	250 °C, 2.5–3.5 MPa	Ni–SiO ₂ –Al ₂ O ₃ vs. Ni-Beta, kinetic study	18
Ni-Beta	Beta zeolite	30–190 °C, 3.5–6.5 MPa	Fuel production	19
Ni-Beta	Beta zeolite	50–190 °C,	Effect of parameters	20
		0.85–2.56 MPa		
Ni-Beta	Beta zeolite	120, 250 °C,	Kinetic study	21
		3.0–3.5 MPa		
Ni-Beta	Beta zeolite	50–100 °C,	Kinetic study	22
		0.5–2.8 MPa		
Ni-Beta	Beta and ZSM-5	180, 200 °C, 0.25,	Ni-Beta vs. Ni-ZSM-5: active sites	23
Ni-ZSM-5	zeolites	3 MPa		
Ni-Beta	Beta zeolite	180 °C, 0.1 MPa	Beta heteroatom composition; X-Beta, X = Al, Ga, Fe	24
Ni-Beta	Beta zeolite	250 °C, 3 MPa	Beta heteroatom composition; X-Beta, X = Sn, Ge, Hf, Zr, Ti	25
Ni-Beta	Beta zeolite	30–120 °C,	Subcritical or supercritical ethylene	26
		1.1–5.3 MPa		
Ni-ZSM-5	ZSM-5 zeolite	300 °C, 3.5 MPa	Effect of the incorporation procedure on the nature of the active sites	27
Ni-ZSM-5	ZSM-5	250 °C, 2 MPa	Relationship between activity and acid site distribution	28
Ni-ZSM-5	ZSM-5	300–400 °C, 0.1 MPa	Deactivation mechanism	29
Ni-MgY	FAU zeolite	25 °C	1-Butene production, dimerization mechanism DFT calculation	30
Ni-FAU	FAU zeolite	300–350 °C, 3.5 MPa	Fuel production, cascade reactions	31
H-ZSM-5				
Ni-MCM-22	MWW zeolite	300 °C, 2 MPa	Synergetic effect of acid and nickel sites	32
		450 °C, 0.1 MPa		
Ni-ERB-1	MWW zeolite	300 °C, 2 MPa	Synergetic effect of acid and nickel sites	32
		450 °C, 0.1 MPa		
Ni-SSZ-24	SSZ-24 zeolite	No reaction	Mechanism, DFT	33
		150 °C, 3 MPa	Mechanism, kinetic, DFT-MD	34
Ni-ETS-10	ETS-10 zeolite	180 °C, 0.5 MPa	New catalysts	35
Ni-CIT-6	BEA zeolite			
<i>Ni-ordered mesoporous catalysts</i>				
Ni-ALSBA-15	ALSBA-15	80 °C, 3 MPa	ETP (ethylene to propylene), metathesis catalyst: MoO ₃ –SiO ₂ –Al ₂ O ₃	36
Ni-ALSBA-15	ALSBA-15	210 °C, 1 MPa	Fuel production, Ni-ALSBA-15 + Amberlyst-35	37
Ni-ALSBA-15	ALSBA-15	150 °C, 3.5 MPa		38
Ni-ALSBA-15	ALSBA-15	250 °C, 1.5 MPa	Ni-ALSBA-15 vs. Ni-ALSBA-15 diluted stream	39
Ni-ALSBA-15	ALSBA-15	150 °C, 3.5 MPa	Mechanism	40
Ni-ALSBA-15	ALSBA-15	150–350 °C, 0.1–2 MPa	Fuel production	41
Ni-ALSBA-15	ALSBA-15	30–120 °C,	Subcritical or supercritical ethylene	26
		1.1–5.3 MPa		
Ni-ALSBA-15	ALSBA-15	300 °C, 1.15 MPa	Effect of the catalyst morphology	42
Ni-ALKIT-6	ALKIT-6	60 °C, 3 MPa	ETP, metathesis catalyst: ReO _x /Al ₂ O ₃	43
Ni-ALKIT-6	ALKIT-6	60–120 °C, 3 MPa	ETP, metathesis catalyst: WO _x /KIT-6	44
Ni-ALKIT-6	ALKIT-6	60–120 °C, 0.1–3 MPa	ETP, metathesis catalyst: ReO _x /Al ₂ O ₃	45
Ni-ALKIT-6	ALKIT-6	40–120 °C, 0.1–2 MPa	Kinetic study	46
Ni-ALKIT-6	ALKIT-6	120 °C, 4 MPa	Effect of calcination temperature	47
Ni-ALMCM-41	ALMCM-41	250 °C, 2 MPa	Relationship between activity and acid site distribution	28
Ni-ALMCM-41	ALMCM-41	–30, –15 °C,	MCM-41 vs. Beta and FAU: Ni site deactivation, working	17
		0.1–2.4 MPa	at low <i>T</i>	
Ni-ALMCM-41	ALMCM-41	120 °C, 3.5 MPa	Beta vs. MCM-41 and SIRAL active sites, effect of the support	12



Table 1 (continued)

Catalyst	Support topology	Reaction conditions	Main aim of the study	Ref.
<i>Ni-microporous materials</i>				
Ni- AlMCM-41	AlMCM-41	120 °C, 0.1 MPa	Active sites	48
Ni- AlMCM-41	AlMCM-41	375 °C, 1 bar	ETP	49
Ni- AlMCM-41	AlMCM-41	75–475 °C, 0.1 MPa	ETP	50
Ni- AlMCM-41	AlMCM-41		Active sites, mechanism, DFT	51
Ni- AlMCM-41	AlMCM-41	–30 °C, 1.5 MPa	Active sites, low T , CO poison	52
Ni- AlMCM-41	AlMCM-41	–30, –20 °C, 2.6–3.5 MPa	Mechanism	53
Ni- AlMCM-41	AlMCM-41	180–300 °C, 1–4 MPa	Distribution of the active sites	54
Ni- AlMCM-41	AlMCM-41		DFT, ethylene adsorption on Ni sites	55
Ni- AlMCM-41	AlMCM-41		Theory, mechanism	56
<i>Ni-non-ordered mesoporous catalysts</i>				
Ni-ASA	ASA	150 °C, 3.5 MPa	ETP, metathesis catalyst: $\text{MoO}_x/(\text{Al})\text{SiO}_2$	57
Ni-ASA	ASA	170–230 °C, 1.5–3.5 MPa	Kinetic study	58
Ni-ASA	ASA	150–350 °C, 3 MPa	Nature of the Ni species	59
Ni-ASA	ASA	300 °C, 0.1 MPa	Nature of the Ni species, effect of the catalyst properties	60
Ni-ASA	ASA	300 °C, 3–4 MPa	Effect of the catalyst properties	61
Ni-ASA	ASA	50–350 °C, 0.35 MPa	Nature of the Ni species	62
Ni-ASA	ASA	375 °C, 0.1 MPa	ETP	49
Ni-ASA	ASA	80–360 °C, 3.5 MPa	Fuel production	63
Ni-ASA	ASA	200 °C, 1 MPa	Nature of the Ni species	64
Ni-ASA	ASA	300 °C, 0.1 MPa	Effect of the catalyst preparation	65
Ni-ASA	ASA	60 °C, 3 MPa	Nature of the Ni species	66
Ni-ASA	ASA	60 °C, 3 MPa	Structure of Ni active sites	67
Ni-ASA	ASA	275 °C, 4 MPa	Fuel production	68
Ni-ASA	ASA	350 °C, 0.1 MPa	ETP, mechanism	69
Ni- AlSiO_2	AlSiO_2	250 °C, 1.5 MPa	Ni- AlSiO_2 vs. Ni- AlSBA-15 diluted stream	39
Ni- SiAlO_x	SiAlO_x	375 °C, 0.1 MPa	ETP	70
Ni-ASA	SIRAL-30	200 °C, 1 MPa	Fuel production	71
Ni-ASA	SIRAL-30	50–200 °C, 4, 6.5 MPa	Subcritical and supercritical ethylene	72
Ni/Siralox-30	SIRALOX-30	120 °C, 3.5 MPa	Active sites, Beta vs. MCM-41 and SIRAL: effect of the support	12
Ni/Siralox-30	SIRALOX 40 HPV	120 °C, 5 MPa	Fuel production	73
Ni/Siralox-30	SIRALOX 40	120 °C, 4 MPa	Fuel production	74
Ni-ASA	SIRAL-30	200–350 °C, 1 MPa	Fuel production, cascade reactions	75
H-ZSM-5				
Ni-Siral-70	SIRAL-70	375 °C, 0.1 MPa	ETP	76
Ni-ASA composite	ZSM-5, MCM-41	250 °C, 2 MPa 450 °C, 0.15 MPa	Synergetic effect of Ni-ions and acid sites	77
<i>Other catalytic systems</i>				
Ni-clay	Montmorillonite	150–350 °C, 3 MPa	New catalysts	78
$\text{NiSO}_4/\text{Al}_2\text{O}_3$	Al_2O_3	70 °C, 3.5 MPa	Mechanism	79
$\text{NiSO}_4\text{-ReO}_x/\text{Al}_2\text{O}_3$	Al_2O_3	70 °C, 0.1 MPa	ETP	80
Ni-POM-WD ($\text{NiK}_{10}\text{P}_2\text{W}_{17}\text{O}_{61}$)	POM-WD	200 °C, 2 MPa	Ni^{2+} single sites	81
Ni-POM-WD ($\text{NiK}_{10}\text{P}_2\text{W}_{17}\text{O}_{61}$)	POM-WD	200 °C, 2 MPa	Kinetic study	82

products (e.g. SIRAL series),^{12,71–75} synthetic ASA and “natural” aluminosilicates (e.g. montmorillonite pillared clay).⁷⁷ The synthetic ASA were prepared by various methods: homogeneous co-precipitation,^{60,61,65–67} alumination of the non-order silica⁵⁶ and hydrolytic non-hydrolytic sol-gel synthesis.^{59,62}

The introduction of nickel into ASA supports was performed using either methods similar to those discussed above for zeolites and ordered mesoporous aluminosilicates (i.e. ion-exchange and impregnation), or by “one-pot” co-

precipitation, using tetraethyl orthosilicate, aluminium chloride/nitrate and nickel nitrate/bis(acetylacetonate) as sources of Si, Al and Ni.^{62,65}

3. Nickel state in oligomerization catalysts

Characterization techniques, including DRIFT, XPS, EPR, and H_2 -TPR, allowed the identification of the different Ni species contained in the oligomerization catalysts. Ionic



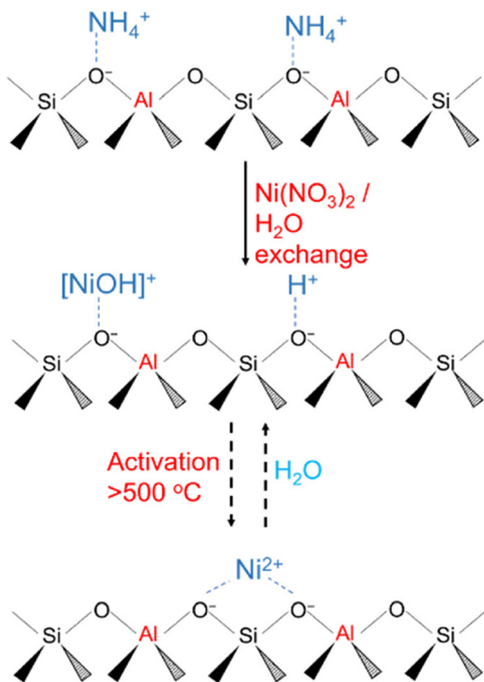


Fig. 1 General protocol for preparing the Ni-Beta catalyst. This figure has been adapted from ref. 14.

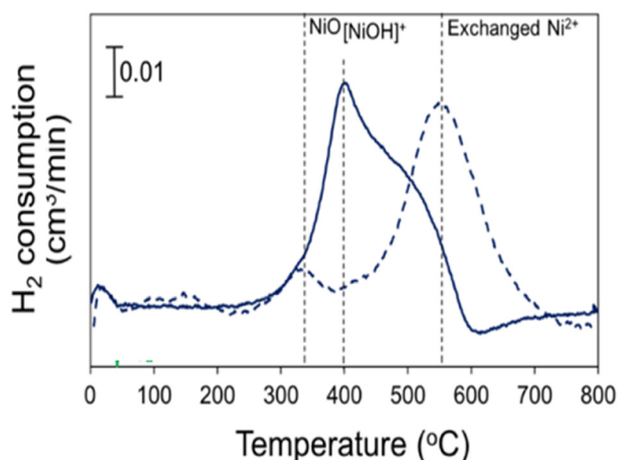


Fig. 2 H_2 TPR profiles of Ni-Beta catalysts after *in situ* pretreatment at $300\text{ }^\circ\text{C}$ (solid lines) and $500\text{ }^\circ\text{C}$ (dashed lines). This figure has been reproduced from ref. 14.

species, such as Ni^{2+} , Ni^+ , $[\text{Ni}-\text{O}-\text{Ni}]^{2+}$ and $[\text{Ni}-\text{O}-\text{H}]^+$, but also small NiO and metallic (Ni^0) particles were formed during the introduction of nickel into the carrier matrix and the heat treatment made before to the oligomerization process (Table 2). This diversity of sites is due to the fact that Ni can adopt a number of oxidation states and that it can bind to various atoms on the aluminosilicate support. The nature of the support, as well as the presence of residual Brønsted acid sites plays an important role in the formation of Ni species.

4. Nickel active catalytic centers: nature and oligomerization mechanism

The identification of the Ni-based active centers and of the elementary steps involved in the dimerization/oligomerization mechanism was the main challenge of the studies carried out in the last decade. But, as Table 3 shows, there is still controversy around the specific nickel species responsible for ethylene oligomerization and their contribution to the initiation of the reaction over nickel aluminosilicates.

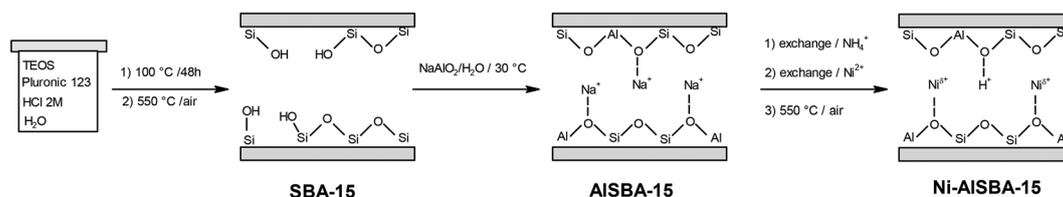
The nature of the nickel species mainly depends on the preparation method of the catalysts. Generally, nickel incorporation by an aqueous ion exchange or grafting method leads to higher contents of Ni^{2+} cations.^{13–15,27,46,76} On the other hand, solid-state ion exchange mainly seems to generate intrazeolitic mono(μ -oxo) dinickel ($[\text{Ni}-\text{O}-\text{Ni}]^{2+}$) species.¹¹ The post-synthesis impregnation leads to high contents of cationic Ni^{2+} in ion exchange positions, but some NiO clusters are frequently formed.^{23,27,76} The *one-pot* synthesis of Ni-aluminosilicates favours the formation of a large amount of NiO particles. Xu *et al.*⁶⁶ showed that in the case of Ni-ASA catalyst, the oxidation state of Ni can be regulated by varying the pretreatment atmosphere. Thus, the majority of the Ni species were found to exist in Ni^{2+} , Ni^+ , and Ni^0 states when pretreated in air, N_2 , and H_2 respectively.

As some research groups have shown, the carrier identity had an important influence on the nature of the nickel species. Moussa *et al.*¹² prepared bifunctional 5%Ni catalysts by impregnation of three acidic porous aluminosilicates: nanocrystalline Beta zeolite, mesoporous Al-MCM-41 and silica-doped alumina Siralox-30. According to characterization results, the authors distinguished significant differences in the nickel speciation depending on the support identity. Isolated Ni^{2+} cations in ion exchange positions were identified on Ni-Beta, while Ni^{2+} attached to weakly acidic silanol and aluminol functions and undercoordinated Ni^{2+} on the surface of small NiO nanoparticles prevailed on Ni-Al-MCM-41 and Ni-Siralox-30 catalysts. Comparable results were reported by Agirrezabal-Telleria and Iglesia⁵² for the Ni-MCM-41catalyst: isolated cations $[\text{Ni}-\text{OH}]^+$ were identified as active sites, besides inactive NiO clusters.

Unlike homogeneous catalysis, the ethylene oligomerization performed in the presence of Ni-aluminosilicate catalysts does not require the use of activators. It is also important to note that for the oligomerization catalyzed by Ni-based complexes, the activator is involved in the formation of the first Ni-C bond. However, the oligomerization mechanism proposals for the Ni-aluminosilicate catalysts are based on organometallic chemistry.

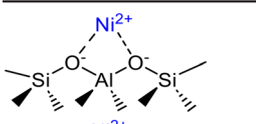
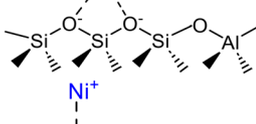
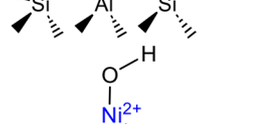
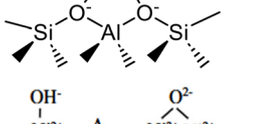
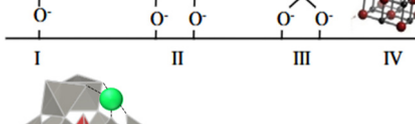
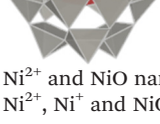
^{13}C NMR and FTIR spectroscopy undoubtedly proved that the conversion of ethylene over Ni-based sites starts





Scheme 1 General protocol for preparing the Ni-AISBA-15 catalyst. This scheme has been adapted from ref. 40.

Table 2 Proposed nickel species contained in Ni-aluminosilicate catalysts

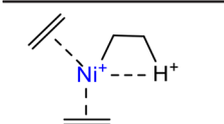
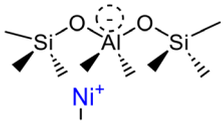
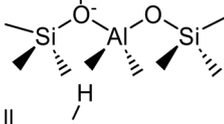
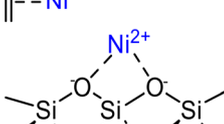
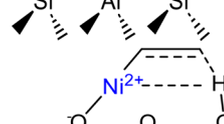
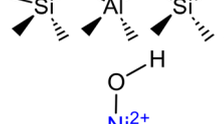
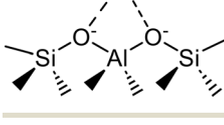
Nickel species	Catalysts	Refs.
	Ni-SSZ-24, Ni-Beta, Ni-ASA, Ni-MCM-41, Ni-ZSM-5	6, 13–15, 18, 23, 27, 34, 46, 48
	Ni-Beta, Ni-ASA, Ni-MCM-41	7, 48, 67
	Ni-SBA-15, Ni-Beta	40, 66, 67
	Ni-Beta	11
	Ni-MCM-41	51
	Ni-MCM-41	52
	Ni-POM	82
Ni ²⁺ and NiO nanoclusters cationic Ni ²⁺ , Ni ⁺ and NiO	Ni-Beta, Ni-ASA	23, 66
Ni ²⁺ and NiO	Ni-ZSM-5	27

with the formation of a π -complex.^{2,7} The subsequent modification of the hydrocarbon chain and its interaction with the metallic site can occur *via* diverse mechanisms, which can be classified according to the nature of the key intermediate species: alkyl-based, vinyl-based, allyl-based (referred to as coordination–insertion or Cossee–Arlman) and metallacycle (Scheme 2). For the coordination–insertion mechanism, largely accepted by

the research groups, the main unclear aspect is the formation of the primary nickel–carbon bond. On this point there are controversies and speculation. The metallacycle mechanisms have been suggested by analogy with the homogeneous catalysis. However, some theoretical research carried out on catalysts based on Ni-aluminosilicates has discouraged the realization of this mechanism.

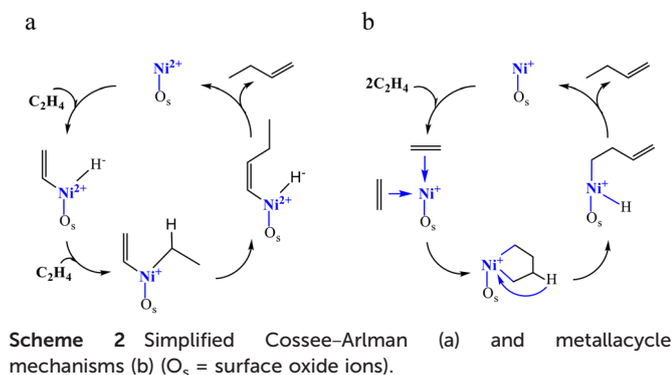


Table 3 Ni-active sites in Ni aluminosilicate catalysts

Active catalytic sites	Catalysts	Ref.	
	Mobile [(ethene) ₂ -Ni-alkyl] ⁺ species	Ni-SSZ-24	34
	Isolated Ni ⁺ cations	Ni-SBA-15 Ni-ASA	40, 66
	<i>In situ</i> ethene-assisted [Ni(II)-H] ⁺ species	Ni-Beta	6
	Isolated Ni ²⁺ cations grafted on acidic silanols	Ni-Beta Ni-MCM-41	7, 48
	Intrazeolitic mono(μ-oxo) dinickel species	Ni-Beta	11
	Isolated Ni ²⁺ cations in ion exchange state	Ni-Beta Ni-ZSM-5	13–15, 27
	Isolated nickel-hydride [Ni(II)-H] ⁺ centers	Ni-Beta	15
	Isolated hydroxylated Ni ²⁺	Ni-MCM-41	51, 52

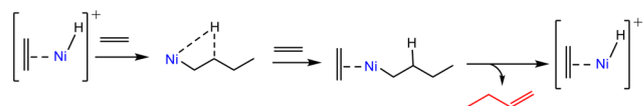
Cossee–Arlman mechanisms

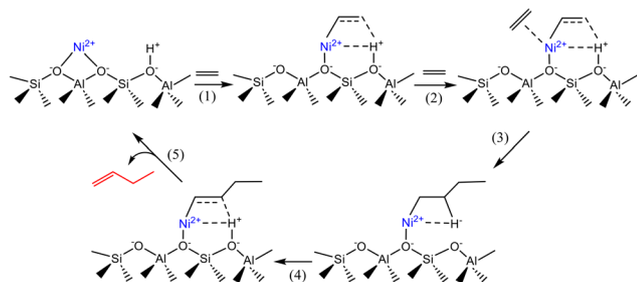
Using H/D isotope scrambling and H₂-D₂ isotope exchange experiments, Joshi *et al.*⁶ evidenced the H₂-assisted formation of [Ni(II)-H]⁺ species, which were the proposed active sites in the Cossee–Arlman mechanism (Scheme 3).



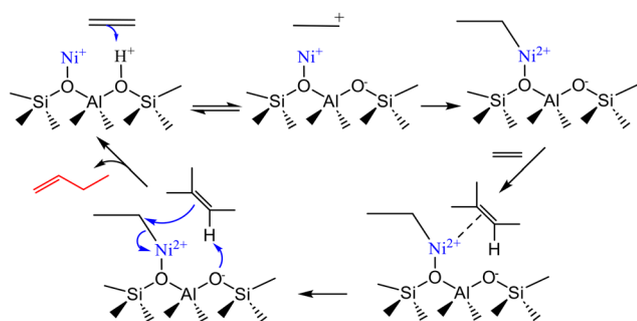
Seufitelli *et al.*¹⁵ suggested the [Ni(II)-H]⁺ active centers involved in the Cossee–Arlman mechanism were generated by ethylene adsorption on a Ni²⁺ site, assisted by an adjacent proton (Brønsted site) (Scheme 4).

The involvement of a neighboring Brønsted center in the Cossee–Arlman cycle was also proved by Rabeah *et al.*⁸⁵ They used *operando* electron paramagnetic resonance (EPR) and *in situ* X-ray absorption spectroscopy (XPS) for finding the active centers in Ni-ASA catalysts during butene oligomerization. Single Ni⁺/Ni²⁺ redox couples were identified as active sites and the reaction mechanism occurred as shown in Scheme 5.



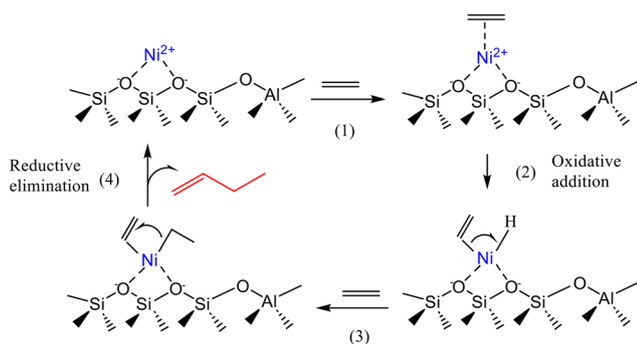


Scheme 4 Proposed mechanism for formation of a nickel-hydride site involving a Brønsted acid: (1) ethylene adsorption over a Ni^{2+} site and an adjacent Brønsted site, (2) formation of a $[\text{Ni}(\text{II})\text{-H}]^+$ center followed by coordination of another ethylene molecule, (3) insertion to form adsorbed butyl, (4) recovery of the Brønsted site, and (5) desorption to form a product followed by recovery of the Ni^{2+} site. This scheme has been adapted from ref. 15.

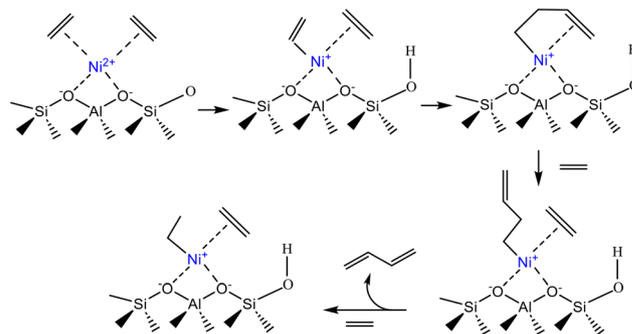


Scheme 5 Reaction mechanism proposed for olefin oligomerization over Ni-ASA catalysts. This scheme has been adapted from ref. 85 with permission from American Chemical Society, copyright 2025.

Based on the intermediates detected by FTIR-ethylene spectroscopy coupled to online MS analysis, Moussa *et al.*⁷ proposed the mechanism shown in Scheme 6. This mechanism starts with the oxidative activation of the C–H bond of ethylene at the Ni^{2+} site, leading to a nickel-ethenyl-hydride intermediate, which will then be involved in the coordination–insertion Cossee–Arlman cycle. According to



Scheme 6 Simplified catalytic cycle proposed for the activation and dimerization of ethylene on the active Ni^{2+} centers of Ni-Beta catalysts: (1) ethylene adsorption on Ni site; (2) oxidative addition; (3) ethylene insertion; (4) reductive elimination of butene. This scheme has been adapted from ref. 7 with permission from American Chemical Society, copyright 2025.



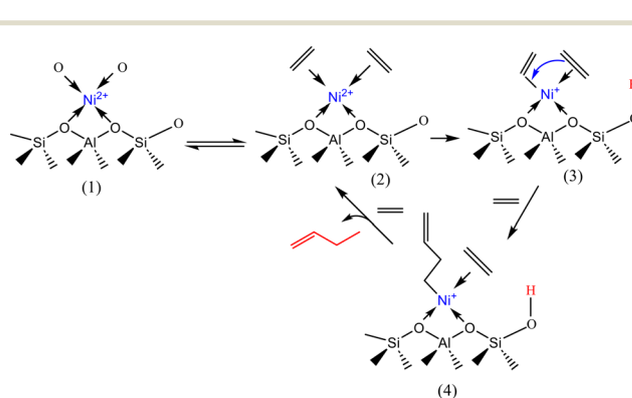
Scheme 7 Formation of the active center over the Ni-Beta catalyst. This scheme has been adapted from ref. 13 with permission from Elsevier, copyright 2025.

the experimental results, the Ni^{2+} ions grafted on acidic silanols of the Ni-Beta catalyst appeared to be the active species, rather than the more accepted ion-exchanged nickel ions.

Henry *et al.*¹³ also examined the nature of the Ni species in the Ni-Beta catalyst, using FTIR spectroscopy with CO as a probe molecule. They found that Ni^{2+} counterions, which were the predominant active sites, interacted with ethylene to form Cossee–Arlman oligomerization sites (Scheme 7). Moreover, the Ni ions grafted on silanol groups and NiO particles were considered as spectators in the oligomerization process.

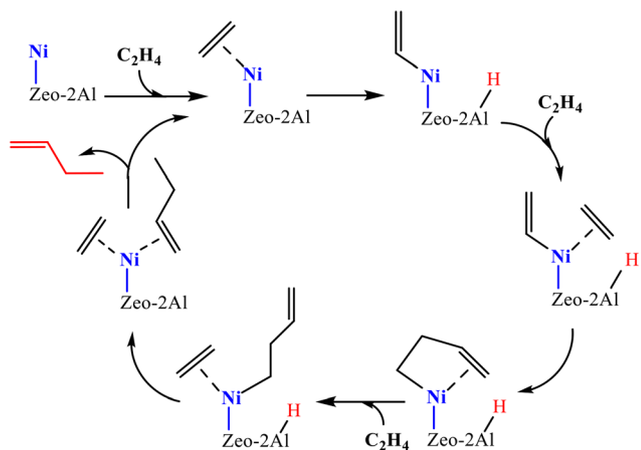
Beucher *et al.*⁴⁶ suggested that the coordination of the Ni ions on the KIT-6 surface is similar to that observed in the coordination chemistry. Thus, the dispersed nickel ions can reversibly bind ligands (L) such as C_2H_4 or surface oxide ions O^{2-} , leading to various $\text{Ni}(\text{L})_n^+$ species. Based on these results, the following mechanism for the formation of 1-butene is proposed (Scheme 8).

By combining advanced characterization techniques (*in situ* XAS, *in situ* FTIR, XPS, H_2 -TPR), DFT calculation and microkinetic simulations, Wang *et al.*³⁰ demonstrated that the *in-situ* generated Ni–vinyl motif is the intrinsic active site and ethylene dimerization proceeds *via* the Cossee–Arlman



Scheme 8 Proposed mechanism for the dimerization of ethylene by Ni-AIKIT-6. (1) Nickel complexation; (2) ethylene complexation; (3) metal–vinyl intermediate; (4) 1-butene formation. This scheme has been adapted from ref. 46.



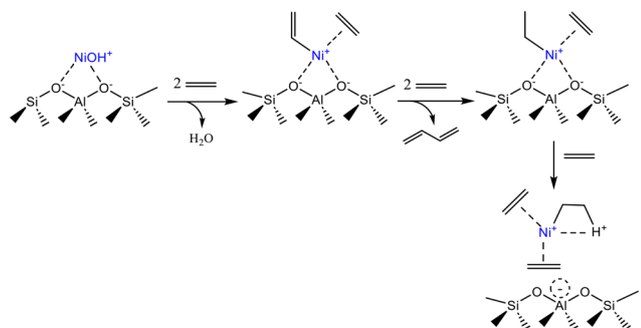


Scheme 9 Proposed Cossee-Arlman pathway for ethylene dimerization to 1-butene on isolated Ni^{2+} ; Zeo = Ni-FAU zeolite. This scheme has been adapted from ref. 30 with permission from Wiley-VCH GmbH, copyright 2025.

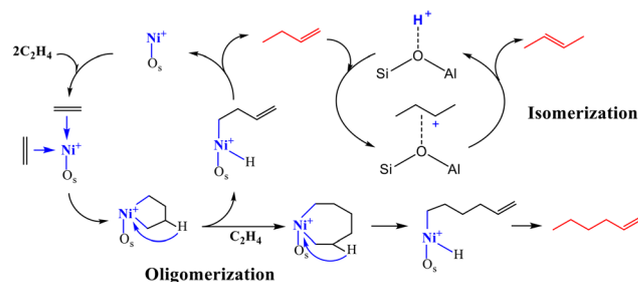
mechanism (Scheme 9). The dynamic hydrogen transfer between the ethylene/vinyl ligand and zeolite framework participated in the formation of the Ni active sites.

Brogaard and Olsbye³³ used DFT calculation for discriminating between Cossee-Arlman and metallacycle mechanisms involved in the ethene dimerization over Ni sites (Ni^{2+} and Ni^+) of SSZ-24 zeolite. They proved that the metallacycle mechanism was energetically unfavorable and Cossee-Arlman mechanisms prevailed for this catalyst. In order to identify the active site involved in the oligomerization mechanism, the same group³⁴ used both experimental and computational investigations. They suggested that the active species form as shown in Scheme 10.

The initial $[\text{NiOH}]^+$ ions served as a precursor for generating the Ni-alkyl species. This was followed by coordination of two ethylene molecules on Ni-alkyl species to form mobile $[(\text{ethene})_2\text{-Ni-alkyl}]^+$ active sites. The mobilization was reversible, as ethene dynamically exchanged with oxygens of the zeolite support as a ligand on Ni during the reaction.



Scheme 10 Formation of the active sites involved in the Cossee-Arlman mechanism. This scheme has been adapted from ref. 34 with permission from American Chemical Society, copyright 2025.



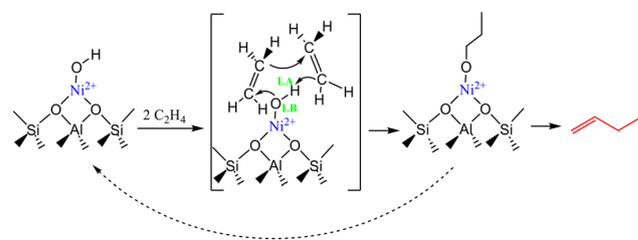
Scheme 11 Proposed catalytic cycles for linear ethylene oligomerization involving Ni^+ and H^+ as active sites on the Ni-AlSBA-15 catalyst (O_s^{2-} = surface oxide ions). This scheme has been adapted from ref. 40.

Metallacycle mechanisms

Andrei *et al.*⁴⁰ showed that on the Al-SBA-15 catalyst it is possible to produce Ni^+ ions and that the coordination of the Ni^+ surface species is similar to that observed in the coordination chemistry.¹ Accordingly, the dispersed nickel ions can reversibly bind ligands such as C_2H_4 or surface oxides. The growth oligomerization mechanism involves a metallacyclopentane intermediate resulting from a concerted coupling of two olefin molecules on each Ni^+ site (Scheme 11).

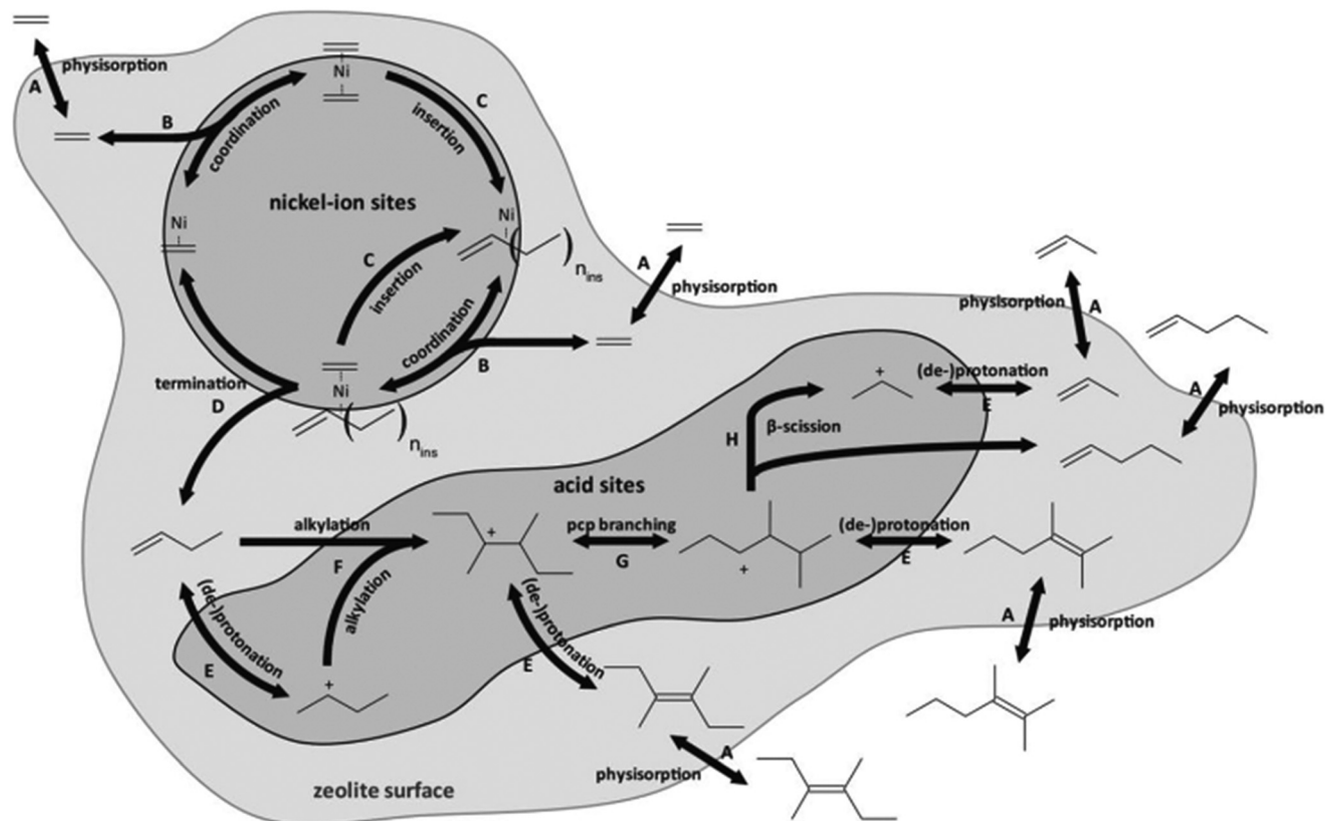
The cyclic intermediate then liberates 1-butene by β -hydride transfer. Insertion of a third molecule of ethylene leads to a metallacycloheptane species, which can release 1-hexene, regenerating the catalytic site. When the desorbed 1-butene and 1-hexene migrate on an Brønsted acid site, they are easily converted to internal double-bond olefins.

Among the oligomerization mechanisms proposed in recent years, that proposed by Jaegers and Iglesia⁵¹ clearly stands out. Their DFT study showed that alkene dimerization cannot occur on active sites consisting of Ni^+ (metallacycle mechanism) or $\text{Ni}^{2+}\text{-H}^-$ (Cossee-Arlman cycle), which would bind ethene very strongly, in contradiction with observed kinetic trends. Instead of classical Cossee-Arlman or metallacycle pathways, they proposed a concerted Lewis acid-base pathway on $(\text{NiOH})^+$ sites contained on Ni-Al-MCM-41 (Scheme 12).



Scheme 12 Probable ethene dimerization catalytic cycle on $(\text{Ni-OH})^+$ moieties acting as a Lewis acid-base pair proposed from a DFT analysis of ethene reactions on Ni-Al-MCM-41. This scheme has been adapted from ref. 51 with permission from American Chemical Society, copyright 2025.





Scheme 13 Schematic representation of the ethylene oligomerization network involving Ni-ion oligomerization and acid-catalyzed alkylation, isomerization and cracking: (A) ethylene physisorption; (B) ethylene coordination on Ni-ethylene species; (C) ethylene insertion between Ni and coordinated ethylene molecule; (D) release of butene and restoration of the active species; (E) butene protonation on an acid site; (F) butene alkylation; (G) carbenium ion isomerization; (H) carbenium ion cracking. This figure has been adapted from ref. 18 with permission from Elsevier, copyright 2025.

Ethylene oligomerization pathways involving both nickel and acid sites

As is known, Ni-aluminosilicates are bifunctional catalysts, containing both Ni and acid sites (see Scheme 1).³ The mechanisms examined in the previous section referred to the dimerization of ethylene to butenes on Ni centers, and show that the Ni sites are indispensable for ethylene oligomerization. Some research groups considered that the Brønsted acid sites also participate in the formation of the active species involved in oligomerization.^{15,30} Generally, the role of catalyst acidity in the process is more complex. They

catalyze a series of reactions that determine the final composition of the reaction mixture, which is usually quite complex. Typically, the products are C₄, C₆, C₈ and C₁₀ olefins, but under severe conditions (*i.e.* high temperature and high acidity of the catalyst), other hydrocarbons such as alkanes and odd-numbered alkenes are formed. To explain the formation of various molecules, several distinct reactions, involving both Ni and acid sites, have been considered (Scheme 13).^{3,6,18,40}

The reactions involving ethylene-ethylene and butene-ethylene couples to form linear olefins occur on nickel sites, *via* the Cossee-Arlman mechanism (called as *true*

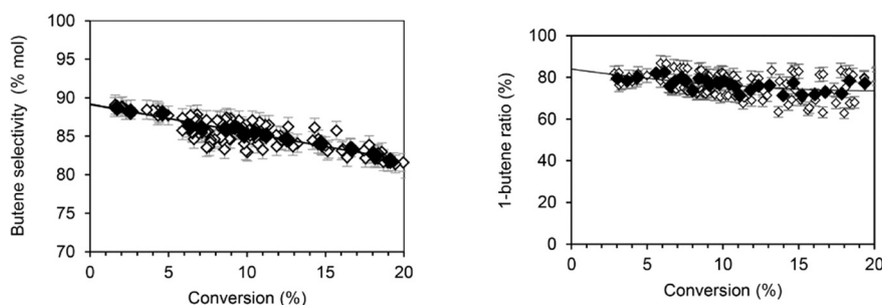


Fig. 3 Butene selectivity (left) and 1-C4 isomer ratio (right) vs. ethylene conversion. Catalyst: 0.1 g of Ni-AIKIT-6, $T = 40\text{--}80\text{ }^{\circ}\text{C}$, $P_{\text{ethylene}} = 0.4\text{--}2\text{ MPa}$, $P_{\text{total}} = 3.0\text{ MPa}$, contact time = 15–98 s. This scheme has been reproduced from ref. 46.



oligomerization). Once formed, these olefins are involved in further acid-catalyzed oligomerization, referred as the *hetero-oligomerization* pathway, including isomerization, alkylation, cracking reactions and H transfer reactions. Following an ionic mechanism, these reactions mainly lead to heavy branched olefins, aromatics and alkanes. The acid site involvement increases at high conversion, temperature, acid concentration and strength.^{3,12,18}

5. Kinetics on Ni-aluminosilicates

Despite the fact that a large number of studies have been devoted during the last decade to the oligomerization of ethylene over Ni-aluminosilicate catalysts, only a few of them have focused on determining the kinetic models. Among the extensive kinetic studies there is that performed by Seufitelli *et al.*²² with the Ni-Beta catalyst. Their micro-kinetic model obeying an Eley–Rideal-type mechanism allowed the rate expressions for the consumption of ethylene and production of butene, hexene and octene to be obtained accurately. Working at various partial pressures of ethylene, the authors found that the reaction order for butene and hexene formation was consistent with a coordination–insertion Cossee–Arlman mechanism, while the reaction order for octene formation points to a cascade co-oligomerization reaction of butene and hexene. The reaction constants for the formation of butene, hexene and octene were 1.8, 0.2 and 2.4 g g_{cat}⁻¹ h⁻¹, respectively. In this model, the Ni²⁺ ions were the precursors for the formation of the [Ni–H]⁺ active sites, while the involvement of the acid sites in the process was not considered. The experimental activation energies for the formation of butene, hexene and octene were 45, 79, and 60 kJ mol⁻¹, respectively.

Combining experimental observation and single-event microkinetic modeling, Toch *et al.*^{18,58} have evaluated the kinetic parameters for ethylene dimerization on Ni-ASA and Ni-Beta catalysts. The reaction rate increased linearly with ethylene pressure (0.15 to 0.35 MPa) and the reaction was found to be first order on ethylene converted. The activation energy for the insertion and termination step was 76 and 74 kJ mol⁻¹, respectively. The kinetic model obtained at low ethylene conversion (*i.e.* high selectivity to butenes) has been extrapolated to different parameters for predicting the effect of conversion, temperature, acid concentration and strength.

Agirrezabal-Telleria and Iglesia⁵² examined the kinetics of ethylene dimerization over the Ni-MCM-41 catalyst, at sub-ambient temperatures (–30–15 °C) and 1.5 MPa. Under these conditions, ethylene condensed within ordered mesopores of the catalyst. A second-order dimerization rate with respect to ethylene pressure was reported. The same reaction order was observed for ethylene dimerization on Ni-SSZ-24 zeolite.³⁴

More recently, Beucher *et al.*⁴⁶ performed a kinetic study of ethylene oligomerization on Ni-ALKIT-6. As shown in Fig. 3, and as reported in many studies,^{6,10,22,25,40} 1-butene is the primary ethylene dimer formed in the processes catalyzed

by the Ni-aluminosilicates. The selective formation of butenes (90%) and hexenes (9%) as primary products has been explained considering a Cossee–Arlman mechanism.

The experimental data obtained in ethylene oligomerization fitted with a first order kinetics, while the activation energy was of 15.2 kJ mol⁻¹. The authors suggested that the kinetic-limiting step is the insertion of ethylene into the Ni–alkyl bond.

6. Effect of the reaction parameters

Effect of the temperature

Oligomerization studies of ethylene in the presence of Ni-aluminosilicates were performed in various temperature ranges, placed between –30 °C and 350 °C. Each working temperature has advantages and disadvantages, depending on the purpose of the study. Fig. 4 shows the representative behavior of a Ni-based catalyst like Ni-SBA-15, obtained at various temperatures.⁴⁰ The ethylene conversion strongly increased, from 14 to 90% in the temperature range of 50 to 150 °C. After that, the ethylene conversion increases slowly up to 300 °C.

As shown in Fig. 4, there is a change in product distribution as a function of ethylene conversion.

Typically, at low conversion, C₄ is the major oligomerization product, while at higher conversion, the oligomerization was directed toward the formation of C₆ and C₈ olefins.^{19,41,51} Among the butenes, with increasing temperature and conversion, the proportion between 1-C₄ and 2-C₄ decreases.^{40,73}

Even if most recent studies have been carried out at moderate temperatures, *i.e.* 50–120 °C, some of them were conducted at high or very low temperatures.^{19,20,40,43} Jan and Resende¹⁹ worked at high temperatures, with the aim of converting ethylene into jet fuel range hydrocarbons. They found that the maximum liquid yield over Ni-Beta can

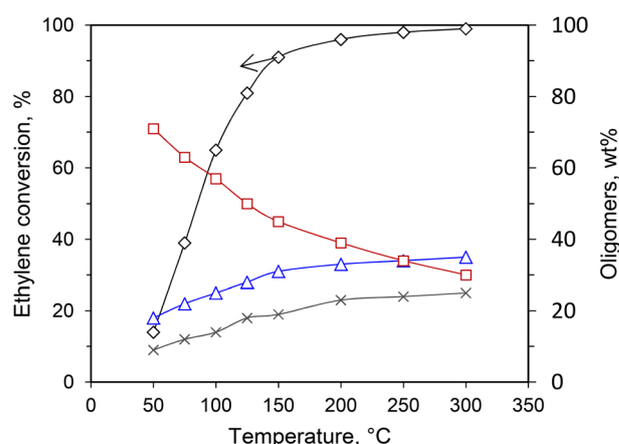


Fig. 4 Ethylene conversion and oligomer distribution at various temperatures; Ni-SBA-15 catalyst; (◇) % ethylene conversion, (□) % C₄, (Δ) % C₆, (x) C₈; conditions: 3.0 MPa, WHSV = 10 h⁻¹. This scheme has been reproduced from ref. 40.



be obtained at 190 °C, 52 bar, and WHSV of 2.0 h⁻¹. Attanatho *et al.*⁴¹ showed that the optimum temperature for the formation of C₈₊ hydrocarbons was in the range of 275–300 °C.

Agirrezabal-Telleria and Iglesia^{51,52} studied the dimerization of ethylene over Ni-Al-MCM-41 at an ethene pressure of 1.5 MPa and sub-ambient temperatures (between -30 °C and -15 °C). The unique performances in terms of reactivity, selectivity and stability obtained under these conditions were attributed to the ethene liquid phase formed in the pores of the catalyst. The liquid phase solvates the reaction transition states, promoting the desorption of butenes and thus preventing their isomerization and their growth towards bulky oligomers which can block the Ni sites and the pores of the catalyst. A similar behavior was described by Jan *et al.*²⁶ for the ethylene oligomerization carried out over Ni-Beta and Ni-SBA-15 under supercritical conditions. The increase in ethylene conversion was attributed to the high solubility of C₄₊ products in supercritical ethylene and the easy desorption of large molecules from the active catalytic centers. Keeping the supercritical state, the catalytic activity increased when the temperature increased from 30 to 120 °C.

Effect of the pressure

As expected, for a reaction involving a gaseous reactant, *i.e.* ethylene, the pressure played a major role in the process. Because the experiments carried out at low pressure led to low ethylene conversions, in most studies the reaction pressure was set between 1 and 4 MPa. The experimental data showed that the effect of the ethylene pressure in the reactor is analogous to that observed for the temperature: the ethylene conversion and the amount of higher olefins (C₆₊) increased with increasing ethylene pressure from 1 to 4 MPa.^{20,40,41,46} For example, over the Ni-SBA-15 catalyst, the specific activity increased, from 214 to 336 mmol g_{cat}⁻¹ h⁻¹ in this range of pressure.⁴⁰ Over Ni-ASA, the ethylene conversion was 10% at 1.0 MPa, but it increased up to 99% when the pressure increased at 3.5 MPa.⁷³ In the presence of Ni-Beta, the steady-state ethylene conversion increased from 38 to 57% as the pressure increased from 0.85 to 2.6 MPa.²⁰ Seufitelli and Gustafson showed that over Ni-ASA the production of liquid products reached a maximum under supercritical conditions, *i.e.* 6.5 MPa of ethylene.⁷²

In addition to these expected results, the recent studies revealed new aspects linked to the effect of pressure in the oligomerization reaction catalyzed by the Ni-aluminosilicates. First, it was shown that the deactivation of the catalytic centers of Ni-based mesoporous aluminosilicates during the oligomerization process can be suppressed by operating at high ethylene pressure.^{17,52,53} The high pressure led to capillary condensation of liquid-like ethylene within mesoporous voids, which solvates and facilitates the transport to external fluid phases of the bulky hydrocarbons that cause catalyst deactivation.

On the other hand, Rabeah *et al.*⁸⁵ showed that the nature and the stability of the Ni active centers in the dimerization of the lower olefins depended on the ethylene pressure. Using *operando* EPR and *in situ* XAS techniques, they identified the Ni^I/Ni^{II} couples as active sites, which were stable only at pressure higher than 1.2 MPa. In contrast, at low pressure (<0.2 MPa), Ni^I sites form inactive Ni⁰ aggregates.

Effect of the space velocity (WHSV)

Typically, over both microporous and mesoporous Ni-aluminosilicates, the ethylene conversion and the amount of the higher hydrocarbons (C₆₊) decreased, while the C₄ fraction significantly increased with increasing WHSV (*i.e.* with decreasing contact time).^{12,20,21,40,41} Fig. 5 shows such a dependence. These results suggest that C₄-hydrocarbons are the primary molecules formed in the ethylene process, while the higher olefins, *i.e.* C₆₊, are produced in a second step, by oligomerization and co-oligomerization, involving the ethylene and butenes.

Jan and Resende¹⁹ evaluated the effect of WHSV on the formation of liquid fraction and coke in the presence of the HNi-Beta catalyst. At WHSV of 0.5 h⁻¹, the yield of coke was nearly 52 wt%, along with a 3.7 wt% yield of liquid hydrocarbons. At 2.0 h⁻¹, there was a noticeable shift in the product distribution, with the yield of the liquid product increasing three times to 10.6 wt% and a minimum coke yield of 6.2 wt%.

7. Effect of the contaminants in ethylene oligomerization

As shown above, ethylene oligomerization over Ni-containing heterogeneous catalysts follows either a Cossee–Arlman or metallacyclopentane mechanism. Whatever the mechanism, ethylene first coordinates *via* π-bonding to the nickel ion site.

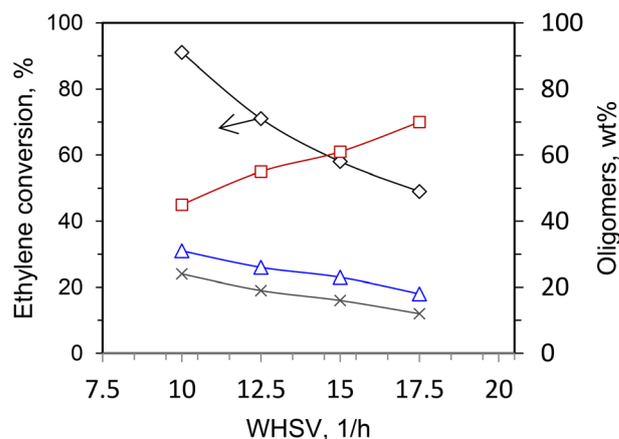


Fig. 5 Ethylene conversion and oligomer distribution at various WHSV; Ni-SBA-15 catalyst; (○) % ethylene conversion, (□) % C₄, (Δ) % C₆, (x) C₈₊; conditions: 3.0 MPa of ethylene, T = 150 °C, TOS = 1 h. This scheme has been reproduced from ref. 40.



But, the nickel ions can also form complexes with other unsaturated organic compounds or with O- and S-containing compounds. During the oligomerization process, these molecules become contaminants for the catalyst, as ligands competing with ethylene.

It is known that the Ni^{2+} cations can be easily hydrated by strong electrostatic interactions with n H_2O molecules as ligands ($n \leq 6$).⁸⁶ Some studies showed that the Ni-aluminosilicate catalysts are extremely air- and moisture-sensitive.^{39,85} For example, when the Ni-ASA catalyst was prepared under aerobic conditions, no catalytic activity in the oligomerization reaction was observed, because of the blockage of active Ni sites by H_2O and/or the formation of stable Ni^{II} oxide species, in which the Ni^{II} coordination sphere is saturated by O ligands and, therefore, is not prone to interact with olefin molecules.⁸⁵

On the other hand, with CO, nickel cations easily form mono- and polycarbonyl complexes. For this reason, CO has been usually used in FTIR spectroscopic studies in order to characterize the Ni species.⁸⁷ CO interacts with Ni^+ and Ni^{2+} sites of low coordination.¹² Exposing Ni-MCM-41 samples to 1 kPa CO at 263 K led to an infrared band at 2200 cm^{-1} , corresponding to $\text{Ni}^{2+}(\text{CO})$.⁸⁸ Fig. 6 shows dimerization rates (1.5 MPa, $-30\text{ }^\circ\text{C}$) as a ratio to that before contact with CO pulses as a function of the cumulative CO uptakes.

The rate decrease as a function of cumulative CO per total Ni is similar for Ni-MCM-41 samples, leading to CO uptakes (per Ni atom) that are similar at these two $\text{Ni}^{2+}/\text{H}^+$ ratios.

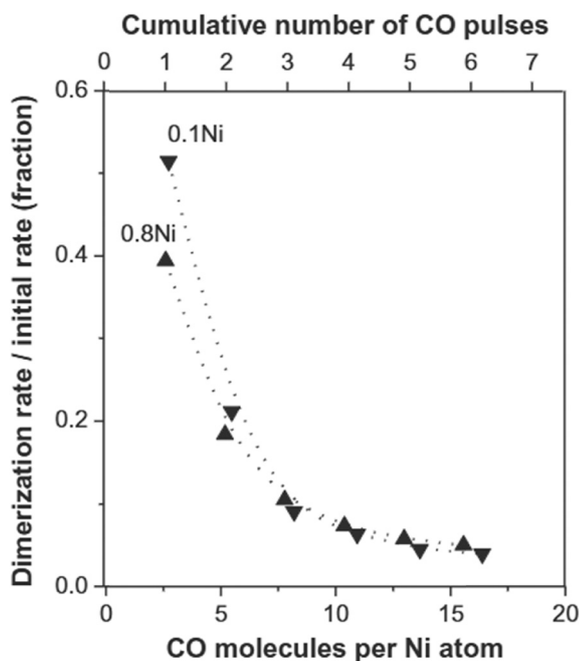


Fig. 6 Ratio of rates after each CO pulse introduction (per Ni) at $-30\text{ }^\circ\text{C}$, 1.5 MPa ethene. This figure has been reproduced from ref. 52 with permission from Elsevier, copyright 2025.

To remove possible contaminants coordinated on Ni sites, thermal activation of the Ni-aluminosilicate catalysts at high temperature ($>300\text{ }^\circ\text{C}$) before the reaction is a crucial step.⁸⁴

Another way to keep the catalytic activity, even in the presence of contaminants, is to work at high temperatures. In one of the earlier studies, Kimura *et al.*⁸⁹ reported that the dimerization of the ethylene was completely prevented when CO or H_2O was pre-adsorbed over the NiO-SiO_2 catalyst. The catalyst was activated by evacuation at temperatures higher than $200\text{ }^\circ\text{C}$.

In another study, Zhang *et al.*⁶⁸ showed that at $280\text{ }^\circ\text{C}$ and 4.0 MPa the oligomerization behavior of the Ni-ASA catalyst was enough high, despite the presence of other molecules, including C_3H_6 , C_4H_8 , CH_4 , CO, CO_2 , H_2 and N_2 together with the ethylene in the feed.

More recently, Andrei *et al.*³⁹ evaluated the effect/role on the Ni-SBA-15 and Ni-ASA of some potential contaminants, such as H_2O , CO, and H_2 , during ethylene oligomerization. The catalytic tests were carried out at $250\text{ }^\circ\text{C}$ and 15 bar, while the concentrations of CO, H_2O and H_2 in the reactor feed were 90, 300 and 1000 ppmv, respectively. Fig. 7 compares the ethylene conversions obtained in the absence and in the presence of contaminants. While carbon monoxide has not affected the catalyst activity, water has only a minor negative effect. In contrast, in the presence of hydrogen, the ethylene conversion remained at higher values all throughout the catalytic test. Most likely, the presence of hydrogen limits the formation of large unsaturated species, which are responsible for catalyst deactivation.

The oligomer distribution also depended on contaminants, in particular CO and H_2O . In both cases, the amount of C_4 was higher compared to that produced in other tests (Fig. 8).

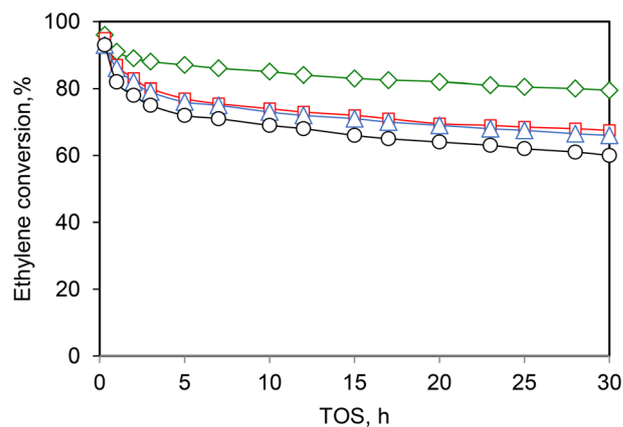


Fig. 7 Ethylene conversion vs. time on stream for various feed compositions (\square) $\text{C}_2\text{H}_4 + \text{N}_2$, (\diamond) $\text{C}_2\text{H}_4 + \text{N}_2 + \text{H}_2$ and (Δ) $\text{C}_2\text{H}_4 + \text{N}_2 + \text{CO}$ and (\circ) $\text{C}_2\text{H}_4 + \text{N}_2 + \text{H}_2\text{O}$, over Ni-ASA-15 catalyst. Reaction conditions: $T = 250\text{ }^\circ\text{C}$, $P = 15\text{ bar}$, $\text{WHSV} = 1.1\text{ h}^{-1}$. This scheme has been reproduced from ref. 39.



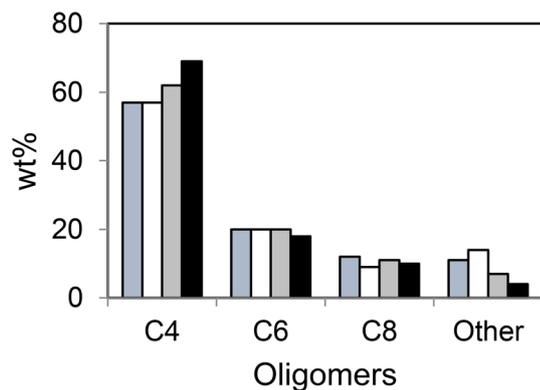


Fig. 8 Effect of the contaminants on the product distribution; blue = C₂H₄ + N₂, white = C₂H₄ + N₂ + H₂, grey = C₂H₂ + N₂ + CO, black = C₂H₂ + N₂ + H₂O; TOS = 30 h, T = 250 °C, catalyst: Ni-ALSBA-15. This scheme has been reproduced from ref. 39.

8. Catalyst deactivation in ethylene oligomerization

Deactivation is one of the main drawbacks of porous heterogeneous catalysts. In the case of the Ni-aluminosilicate catalysts used in the oligomerization reaction, the deactivation is mainly due to heavy hydrocarbons that can bind strongly to Ni sites and/or obstruct the porous voids. The experimental data showed that the amount of hydrocarbons trapped in the catalyst pores and the deactivation rate of the Ni centers mainly depended on the catalyst porosity, but also on its surface acidity and the reaction conditions.

Generally, the Ni-based microporous zeolites deactivate more rapidly than mesoporous materials.^{12,13,23,29} Moussa *et al.*¹² examined the influence of the support identity in the oligomerization of ethylene on three acidic Ni-aluminosilicates: Ni-Beta zeolite, mesostructured Ni-ALMCM-4 and Ni-Siralox-30. As shown in Fig. 9, both mesoporous catalysts were significantly more active than zeolite Ni-Beta.

The authors considered that the heavy oligomers formed during the reaction decreased the catalytic activity by hindering the access of the reactant to the internal Ni sites in Ni-Beta.

Similar differences in terms of activity and deactivation rate between the mesoporous Ni-ASA and microporous Ni-Beta catalysts were identified by Henry *et al.*¹³ Additionally, the authors showed that the crystal size of the Ni-Beta catalysts also had an impact on deactivation. The larger micro-crystallites (which present a longer diffusion path) exhibited a lower activity and higher deactivation rate than the nano-crystallites because of diffusion limitations inside the pores of the long chain products.

Martínez Gómez-Aldaraví *et al.*²³ compared the behavior of Ni-ZSM-5 and Ni-Beta, two zeolite type catalysts with medium and large pores, respectively. The higher deactivation rate of Ni-ZSM-5 has been attributed to rapid blocking by the heavy oligomers of the Ni active sites placed in the small pores.

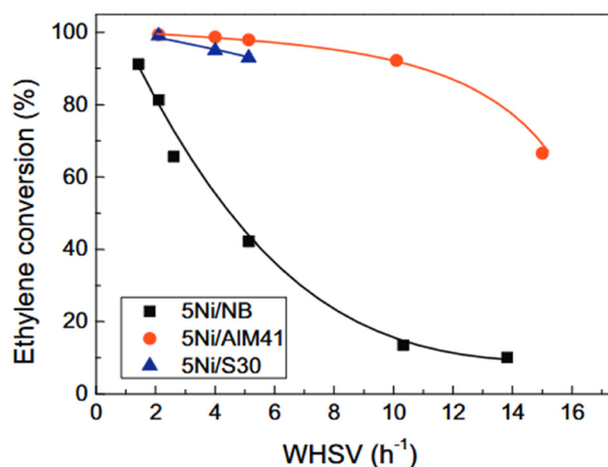


Fig. 9 Ethylene conversion over Ni-Beta (5Ni/NB), Ni-MCM-41 (5Ni/AlM41) and Ni-ASA (5Ni/S30) catalysts. Reaction conditions: T = 120 °C, P_{tot} = 3.5 MPa, P_{C₂} = 2.6 MPa. This figure has been reproduced from ref. 12 with permission from Elsevier, copyright 2025.

In contrast to Ni-zeolites, Ni-mesoporous catalysts such as Ni-ASBA-15,^{40,41} Ni-ALKIT-6 (ref. 42) and Ni-ALMCM-41,^{17,52} are very stable catalysts against deactivation in oligomerization processes. For example, in an experiment performed during 80 h on stream, Andrei *et al.*⁴⁰ showed that the catalytic activity of Ni-ALSBA-15 (pore size of 7.9 nm) declined only smoothly (deactivation rate of $1.6 \times 10^{-3} \text{ h}^{-1}$). This behavior has been related to the SBA-15 topology, with large interconnected mesopores, facilitating the diffusion of the large molecules. For a Ni-KIT-6 catalyst (pore size of 5.4 nm), Hwang *et al.*⁴⁷ found an apparent 2nd order deactivation rate constant of 2 h^{-1} .

On the other hand, the mesoporous supports have another advantage over the microporous ones. At high pressure and low temperature, they allow the formation of a liquid phase (consisting in ethylene and oligomers) within mesoporous voids, which solvates, desorbs and facilitates the transport to external fluid phases of the bulky intermediates that are

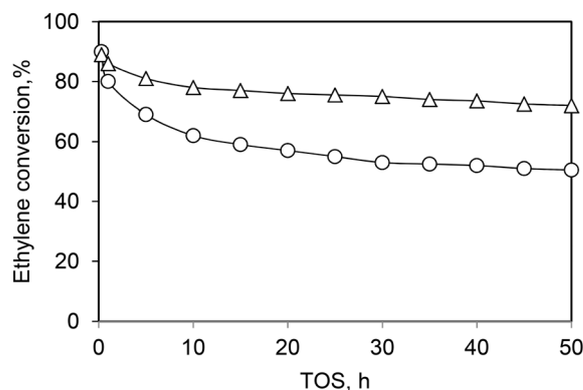


Fig. 10 Ethylene conversion vs. time on stream in the presence of (O) C₂H₄ + N₂ and (Δ) C₂H₄ + N₂ + H₂ over the Ni-ASA catalyst. Reaction conditions: T = 250 °C, P = 1.5 MPa, WHSV = 1.1 h⁻¹. This scheme has been reproduced from ref. 39.



precursors to deactivation.^{51–53} By contrast, the capillary condensation of ethylene within the spatial constraints of microporous voids of Ni-zeolites is not possible.¹⁷

Andrei *et al.*³⁹ proved that the initial deactivation rate and the lifetime of the Ni-ALSBA-15 and Ni-ASA catalysts can be improved by adding hydrogen in the reaction feed (Fig. 7 and 10). The presence of hydrogen limits the formation of large unsaturated species, which are responsible for catalyst deactivation.

Some recent studies have focused on the kinetics of catalyst deactivation. They showed that the kinetic parameters depend on the reaction parameters or the density of the Ni sites in the catalyst. Agirrezabal-Telleria and Iglesia⁵² modeled the deactivation of the Ni-ALMCM-41 catalyst using a first-order deactivation constants (k_d), defined as follows:

$$r_t/r_0 = \exp[-k_d(t - t_0)]$$

where r_t and r_0 represent ethene dimerization rates at each time t and at the initial time t_0 .

Fig. 11 shows the dependence between this constant and the ethene pressure, at $-30\text{ }^\circ\text{C}$ and $-20\text{ }^\circ\text{C}$. An abrupt increase of k_d as the ethene pressure decreases below 1.2 MPa at $-30\text{ }^\circ\text{C}$ and below 1.8 MPa at 253 K was observed.

Caulkins *et al.*¹⁷ studied the effect of ethylene pressure on the deactivation of Ni-FAU, Ni-Beta and Ni-ALMCM-41 catalysts at sub-ambient temperature ($-15\text{ }^\circ\text{C}$). The apparent deactivation constants have been calculated using the following model:

$$r = \frac{r_0}{(1 + (n-1)k_d r_0^{n-1} t)^{\frac{1}{n-1}}}$$

where k_d is the apparent deactivation constant, r is the ethene oligomerization rate at a given time t , r_0 is the initial

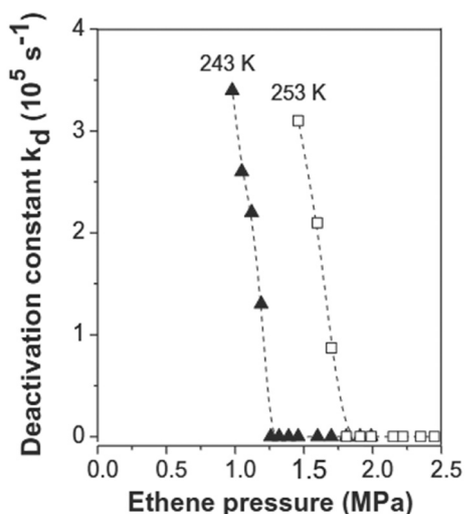


Fig. 11 First-order ethylene dimerization deactivation constant (k_d) as a function of ethylene pressure. This figure has been reproduced from ref. 52 with permission from Elsevier, copyright 2025.

rate at time 0, and n is the deactivation order. The experimental results are summarized in Table 4. Both k_d and n parameters strongly depend on the catalyst type and reaction pressure. The first-order deactivation kinetics suggests a deactivation by the formation of heavier oligomers adsorbed at Ni sites according to a single-site mechanism. The second-order deactivation suggests a dual-site a deactivation mechanism that involves two Ni sites.

Based on the experimental results and DFR calculations, Saxena *et al.*⁸ showed that the deactivation mechanisms and kinetics depended on the Ni site density in the Ni-Beta catalyst. An exponential deactivation (*i.e.* single-site and deactivation rates of first-order in Ni) was observed for a low-site density sample (Ni/Al = 0.06), while a hyperbolic deactivation (*i.e.* dual-site and deactivation rates of second-order in Ni) was found for the high-site density catalyst (Ni/Al = 0.25).

In the first case, the deactivation was due to inhibition and poisoning by strongly bound alkyl groups formed from heavy oligomers, while in the second case the deactivation is due to the formation of bridging alkyl groups between two Ni species to form unreactive centers.

Catalyst regeneration

It is well known that the regeneration ability of solid catalysts is of vital importance for commercialization. Typically, regeneration involves thermal treatment to remove surface coatings and/or adsorbed species. In most cases, the spent catalyst is treated at $500\text{--}550\text{ }^\circ\text{C}$ under air, then treated under an inert gas. Using this regeneration way, Andrei *et al.*³⁹ examined the behavior of the Ni-ALSiO₂ catalyst in two reaction cycles, at $250\text{ }^\circ\text{C}$. The ethylene conversion *versus* time on stream of 75 h is presented in Fig. 12. The conversion profiles of the fresh and regenerated samples were almost the same.

The efficiency of the regeneration process was also evaluated for the used Ni-ALSBA-15 (ref. 40) and Ni-SIRAL-30.⁷¹ The regenerated catalysts exhibited catalytic properties similar to those of the original catalysts. Another regeneration method is to treat the used catalyst in an inert gas. Thus, Agirrezabal-Telleria and Iglesia⁵² demonstrated

Table 4 Deactivation constants (k_d) and orders (n) measured at different ethylene pressures (P/P_0)

Catalyst	P/P_0	k_d^a	n
Ni-MCM-41	0.36	3.15	1.9
	0.71	0	0
	0.89	0	0
Ni-Beta	0.04	6.07×10^{-4}	2
	0.24	8.49×10^{-3}	1.8
	0.84	3.90×10^{-2}	2.2
Ni-FAU	0.04	3.04×10^2	2
	0.36	2.13×10^3	2.2
	0.84	1.52×10^{-2}	1.2

^a Units for k_d : $\text{mol Ni}^{n-1} (\text{mol ethene})^{1-n} \text{s}^{-n-2}$.



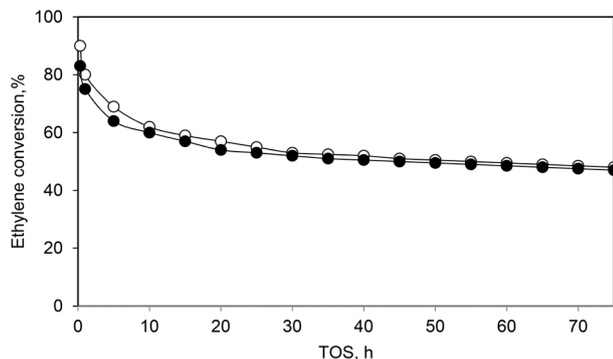


Fig. 12 Ethylene conversion vs. time on stream, over (○) fresh and (●) regenerated Ni-ALSiO₂ catalyst; reaction conditions: C₂H₄/N₂ = 1/5 (vol), *T* = 250 °C, *P* = 15 bar, WHSV = 1.1 h⁻¹. This scheme has been reproduced from ref. 39.

that the activity of the Ni-Al-MCM-41 catalyst can be fully restored by treating sample in He at 550 °C.

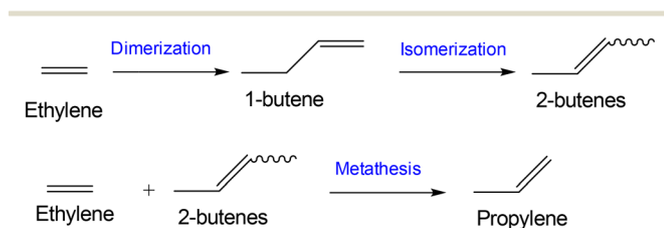
These experimental results show that nickel aluminosilicates used in ethylene oligomerization can be efficiently regenerated using recognized methods.

9. Potential applications based on Ni-aluminosilicate catalysed oligomerization

The previous sections have dealt with aspects closely related to the oligomerization reaction catalyzed by nickel aluminosilicates. However, in order to obtain products of high commercial interest, some research groups have developed in recent years' applications that involve ethylene oligomerization. Two examples, concerning the production of propylene and fuels, will be analyzed below. In both cases, multi-reaction catalytic processes, in which oligomerization is the key step, have been used.

Ethylene to propylene process

Starting from the selective dimerization of ethylene over Ni-aluminosilicates, a number of recent studies have focused on the direct conversion of ethylene to propylene (ETP). This process has been carried out using only ethylene as the reagent, over heterogeneous catalysts which were able to work without activators or co-catalysts. The ETP process,



Scheme 14 Ethylene to propylene in a catalytic cascade process (2-butenes mean *cis*- and *trans*-2-butenes).

carried out in flow mode, consisted in 3 cascade reactions, *i.e.* dimerization–isomerization–metathesis (Scheme 14).

Each step requires a specific catalytic site. Dimerization and isomerization are catalysed by nickel and acid (H⁺) sites, respectively. Both types of catalytic sites are provided by the Ni-aluminosilicates. The metathesis reaction is catalysed by Mo, W or Re oxides. Experimentally, to produce propylene from ethylene according to Scheme 13, two catalyst beds consisting of Ni- and W/Re-based materials were placed in one^{36,45,57,80} or two consecutive reactors^{43,44} (Fig. 13). Table 5 summarizes data from relevant recent studies.

Andrei *et al.*³⁶ explored for the first time the direct conversion of ethylene into propylene using two catalysts, in a single flow reactor. Under identical conditions, ethylene was first selectively dimerized/isomerized over the Ni-ALSBA-15 catalyst to form 2-butene, which reacted then with the excess of ethylene over MoO₃-SiO₂-Al₂O₃ to produce propylene. At 80 °C and 3 MPa, specific activities up to 48 mmol of propylene per gram of catalyst per hour were obtained. The authors extended the research on other catalyst couples: Ni-ALSBA-15/MoO_x-SBA-15 and Ni-ASA/MoO_x-ASA.⁵⁷

In a similar experimental mode, but working at a lower temperature (60 °C) and using Ni-ALKIT-6 and ReO_x/Al₂O₃ as catalysts, Beucher *et al.*⁴³ showed that ethylene can be simultaneously and selectively converted into propylene and 1-butene (Fig. 14).

The global selectivity to C₃ and 1-C₄ olefins was about 86%, while the yield of C₃ and 1-C₄ was 26 and 13 mmol g_{catal}⁻¹ h⁻¹, respectively. Despite their notable initial activity, the supported MoO_x and ReO_x metathesis catalysts used in these studies suffered significant deactivation in the process. In order to circumvent this inconvenience, Beucher *et al.*⁴⁴ combined the dimerization/isomerization catalyst (Ni-ALKIT-6) with a highly efficient metathesis catalyst, *i.e.*, WO_x, placed in two reactors (Fig. 13). In the first one, at 60 °C and 3 MPa, ethylene has been partially converted into 2-butene, which reacted with the unconverted ethylene in the second reactor, operated at 450 °C and 0.1 MPa. Under these conditions, an ethylene conversion of 85% and a selectivity to

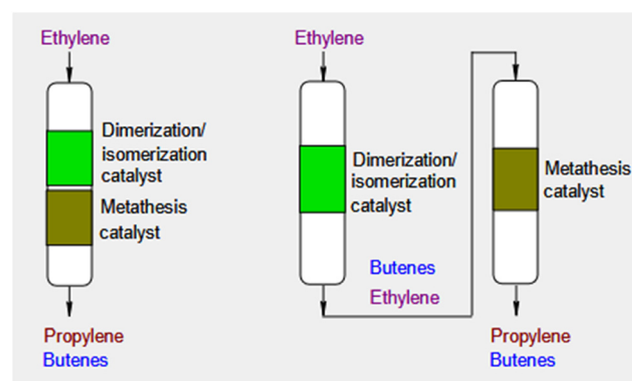


Fig. 13 ETP processes with one or two reactors.



Table 5 Direct conversion of ethylene into propylene over multifunctional catalysts

Dimer–isomer catalyst ^a	Metathesis catalyst	Reactor (s)	Conditions	C ₃ ^b	C ₂ conv. ^c	C ₃ in prod. ^d	Ref
Ni-ALSBA-15	MoO ₃ –SiO ₂ –Al ₂ O ₃	One	80 °C, 3 MPa	48	40	73	36
Ni-ALSBA-15	MoO _x /SBA-15	One	80 °C, 3 MPa	54	43	71	57
Ni-ASA	MoO _x /ASA	One	80 °C, 3 MPa	28	41	38	57
Ni-ALKIT-6	ReO _x /Al ₂ O ₃	One	60–120 °C 0.1–3 MPa	29	52	50	45
Ni-ALKIT-6	ReO _x /Al ₂ O ₃	One	60 °C, 3 MPa	26	73.5	61	43
Ni-ALKIT-6	WO _x /KIT-6	Two	1. 60–120 °C 2. 450 °C	39	48.8	59.4	44
NiSO ₄ –ReO _x /γ–Al ₂ O ₃		One	50 °C, 0.1 MPa	13	60		80

^a Dimerization–isomerization. ^b C₃ productivity (mmol g_{catal}^{−1} h^{−1}). ^c Ethylene conversion (%). ^d Propylene concentration in products.

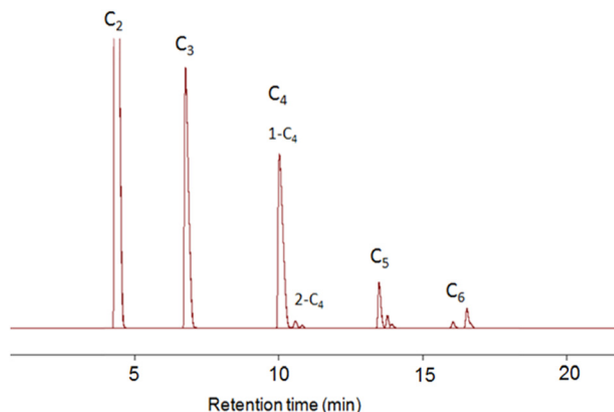


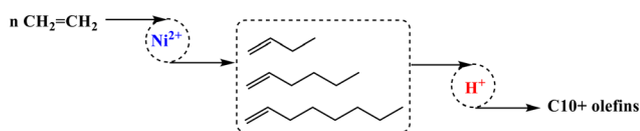
Fig. 14 Selective production of propylene and 1-butene from ethylene by catalytic cascade reactions. This scheme has been reproduced from ref. 43.

propylene of 55% were maintained during 24 on reaction stream.

In a recent study, Li *et al.*⁸⁰ performed the ETP process at 50 °C and 0.1 MPa, over a single multi-site catalyst, *i.e.* NiSO₄/ReO_x/γ–Al₂O₃. The two catalyst functions worked independently for dimerization/isomerization (NiSO₄) and metathesis (ReO_x).

Ethylene to fuels

Most experimental studies showed that, under optimized reaction conditions, ethylene oligomerization is very selective to linear C₄ and C₆ olefins. To produce hydrocarbons in the distillate range (>C₁₀ carbon atoms), the Ni-catalysed ethylene oligomerization must be assisted by an acid catalysis.⁹⁰ In other words, to produce jet fuels from ethylene, a multi-stage reaction is required. Thus, the dimerization/trimerization step is followed by the co-



Scheme 15 Main reaction pathways of ethylene oligomerization into C₁₀₊ olefins by catalytic cascade reactions.

oligomerization of the C₄–C₆ primary molecules (Scheme 15).

Practically, the process can be performed using either a dual-bed catalyst (in a continuous fixed-bed reactor),^{31,75} two catalysts in batch mode (autoclave),³¹ or by combining a fixed bed reaction system with an autoclave reactor.³⁷ Babu *et al.*³⁷ proposed an integrated process for the production of jet-fuel range olefins using Ni-ALSBA-15 and Amberlyst-35 catalysts. In a fixed bed reaction system, the Ni-based catalyst led to C₄–C₈ olefins, at 99% ethylene conversion, during 60 h on stream, at 200 °C and 1 MPa. Co-oligomerization reactions of the as-synthesized oligomers over the Amberlyst-35 catalyst (at 100 °C, 3 MPa N₂, 24 h reaction time, batch mode) produced 98% liquid hydrocarbons, with about 42% C₁₀₊.

Kwon *et al.*⁷⁵ demonstrated that the Ni–Siral-30 + H-ZSM-5 one-pot cascade catalysis in a continuous flow fixed-bed reactor can be an efficient method for producing jet-fuel range hydrocarbons from ethylene. Compared to the results obtained over a single catalyst (Ni–Siral-30), the cascade process with two catalysts exhibited higher ethylene

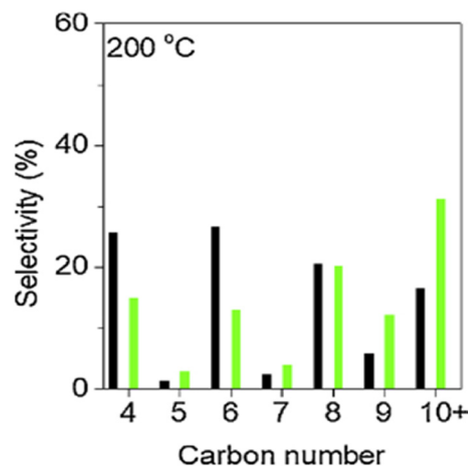


Fig. 15 Carbon distribution of products obtained after ethylene reaction for 16 h on stream over Ni/Siral-30 (black) and Ni/Siral-30 + H-ZSM-5 (green) in the continuous fixed-bed reactor. This figure has been reproduced from ref. 75 with permission from Elsevier, copyright 2025.



conversion (close to 100%) and a completely reversed Schulz-Flory type product distribution $C_{10} > C_8 > C_6 > C_4$, Fig. 15).

In a more recent study, Mohamed *et al.*³¹ used Ni-Y and H-ZSM-5 as catalysts for oligomerization and co-oligomerization, respectively. Under optimized dual-bed conditions (300 °C, 3.5 MPa, 0.68 g_{cat} h g_{C₂}⁻¹), the process produced 64 wt% of jet-fuel at the beginning of the reaction and maintained a 50 wt% selectivity to this fraction for 20 h on stream.

Conclusion and future outlook

On the basis of an important quantity of research results, this contribution clearly shows that ethylene dimerization/oligomerization catalyzed by Ni-aluminosilicates is a dynamic topic, of great fundamental and practical interest. The scientific progress achieved in the last decade can be summarized as follows:

1. Ni-containing zeolites, due to their highly ordered crystalline structure, have served as excellent model supports for fundamental studies in terms of mechanism, active sites and DFT simulation. On the other hand, the well-tailored Ni-mesoporous aluminosilicates, due to their high catalytic stability, were widely used for developing oligomerization processes, under optimized reaction conditions.

2. Concerning the dimerization/oligomerization mechanism, the experimental and theoretical arguments clearly favor the Cossee-Arlman mechanism over the metallacyclic pathway.

3. Unfortunately, there is still controversy surrounding the nature of the nickel active site, how it forms, its location on the aluminosilicate matrix and its contribution to the initiation of the Cossee-Arlman mechanism.

4. Studies specifically dedicated to the effects of the contaminants in ethylene oligomerization have been conducted for the first time during the last decade. It has been shown that the adsorption competition on the Ni centers between ethylene and contaminants, *i.e.* H₂O, CO, olefins, can be circumvented by working at temperatures higher than 200 °C.

5. The conversion of ethylene in the liquid phase, *i.e.* at sub-ambient temperature or under supercritical conditions, highlighted the favorable effect of intrapore solvation on the catalyst activity and stability, as well as the selectivity to 1-butene.

6. Based on the remarkable results obtained in the oligomerization of ethylene, some research groups enriched the potential of the Ni-mesoporous aluminosilicates by developing multi-reaction/catalyst processes in which oligomerization is the key step. The direct ethylene-to-propene conversion and the ethylene conversion in jet fuels are the most representative applications investigated.

7. Thanks to their robust structures, Ni-aluminosilicates allow their complete regeneration by thermal treatments in an inert or oxidizing environment when deactivation occurs.

In my opinion, the main challenges regarding this topic are the following:

1. Unlike the nickel-based organometallic catalysts, Ni-aluminosilicates do not require activators or cocatalysts. Further research is needed to understand this behavior.

2. The interaction between Ni and matrix atoms and the role of the support in the formation of active species require more investigation.

3. The selective catalytic conversion of ethylene into higher hydrocarbons could be incorporated as a stage into an integrated system that includes bioethanol production from biomass and bioethanol dehydration to form ethylene (biomass → bioethanol → ethylene → higher hydrocarbons). Although promising progress has been obtained for each way, further research is still needed in order to produce the knowledge necessary to design the “ideal” catalysts and large-scale processes.

4. Developing large-scale processes capable of producing propylene and fuels from ethylene remains a challenge. Research is also needed to increase the selectivity to 1-C₄ or long-chain linear hydrocarbons.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

Conflicts of interest

There are no conflicts to declare.

References

- 1 H. Olivier-Bourbigou, P. A. R. Breuil, L. Magna, T. Michel, M. F. Espada Pastor and D. Delcroix, *Chem. Rev.*, 2020, **120**, 7919–7983.
- 2 Z. N. Lashchinskaya, A. A. Gabrienko and A. G. Stepanov, *ACS Catal.*, 2024, **14**, 4984–4998.
- 3 A. Finiels, F. Fajula and V. Hulea, *Catal. Sci. Technol.*, 2014, **4**, 2412–2426.
- 4 S. Nishimura, *Handbook of heterogeneous catalytic hydrogenation for organic synthesis*, John Wiley & Sons, Inc., New York, 2001.
- 5 *Hydrogen production by steam reforming of natural gas and other nonrenewable feedstocks*, in Compendium of hydrogen energy, Hydrogen production and purification, ed. L. Garcia, Woodhead Publishing, Kidlington, 2015, vol. 1, pp. 83–107.
- 6 R. Joshi, G. Zhang, J. T. Miller and R. Gounder, *ACS Catal.*, 2018, **8**, 11407–11422.
- 7 S. Moussa, P. Concepción, M. A. Arribas and A. Martínez, *ACS Catal.*, 2018, **8**, 3903–3912.
- 8 A. Saxena, R. Joshi, R. R. Seemakurthi, E. Koninckx, L. J. Broadbelt, J. Greeley and R. Gounder, *ACS Eng. Au*, 2022, **2**, 12–16.
- 9 S. Moon, H. J. Chae and M. B. Park, *Appl. Catal., A*, 2018, **553**, 15–23.



- 10 G. V. S. Seufitelli and F. L. P. Resende, *Appl. Catal., A*, 2019, **576**, 96–107.
- 11 K. Lee and S. B. Hong, *Appl. Catal., A*, 2021, **615**, 118059.
- 12 S. Moussa, M. A. Arribas, P. Concepción and A. Martínez, *Catal. Today*, 2016, **277**, 78–88.
- 13 R. Henry, M. Komurcu, Y. Ganjkhanlou, R. Y. Brogaard, L. Lu, K.-J. Jens, G. Berlier and U. Olsbye, *Catal. Today*, 2018, **299**, 154–163.
- 14 J. McCaig and H. H. Lamb, *Catalysts*, 2022, **12**, 824.
- 15 G. V. S. Seufitelli, J. J. W. Park, P. N. Tran, A. Dichiarà, F. L. P. Resende and R. Gustafson, *Catalysts*, 2022, **12**, 565.
- 16 O. Abed, H. O. Mohamed, I. Hita, V. Velisoju, N. Morlanés, O. El Tall and P. Castaño, *ChemCatChem*, 2024, **16**, e202301220.
- 17 R. Caulkins, R. Joshi, R. Gounder and F. H. Ribeiro, *ChemCatChem*, 2022, **14**, e202101478.
- 18 K. Toch, J. W. Thybaut, M. A. Arribas, A. Martínez and G. B. Marin, *Chem. Eng. Sci.*, 2017, **173**, 49–59.
- 19 O. Jan and F. L. P. Resende, *Fuel Process. Technol.*, 2018, **179**, 269–276.
- 20 O. Jan, K. Song, A. Dichiarà and F. Resende, *Ind. Eng. Chem. Res.*, 2018, **57**, 10241–1025021.
- 21 E. Koninckx, R. Gounder, J. W. Thybaut and L. J. Broadbelt, *Ind. Eng. Chem. Res.*, 2022, **61**, 3860–3876.
- 22 G. V. S. Seufitelli, J. J. W. Park, P. N. Tran, A. Dichiarà, F. L. P. Resende and R. Gustafson, *J. Catal.*, 2021, **401**, 40–53.
- 23 A. Martínez Gómez-Aldaraví, C. Paris, M. Moliner and C. Martínez, *J. Catal.*, 2023, **426**, 140–152.
- 24 M. Meloni and R. C. Runnebaum, *Catal. Sci. Technol.*, 2021, **11**, 3393–3401.
- 25 Y. Bai, T. Cordero-Lanzac, A. Nova, U. Olsbye, E. Taarning and J. S. Martínez-Espin, *Catal. Sci. Technol.*, 2024, **14**, 1991–2002.
- 26 O. Jan, K. Song, A. Dichiarà and F. L. P. Resende, *Chem. Eng. Sci.*, 2019, **197**, 212–222.
- 27 H. O. Mohamed, V. K. Velisoju, I. Hita, O. Abed, R. K. Parsapur, N. Zambrano, M. Ben Hassine, N. Morlanes, A. H. Emwas, K. W. Huang and P. Castano, *Chem. Eng. J.*, 2023, **475**, 146077.
- 28 C. Wang, L. Wang, G. Wu, F. Jin, X. Zhan and Y. Ding, *Catal. Lett.*, 2019, **150**, 429–437.
- 29 Y. Ganjkhanlou, G. Berlier, E. Groppo, E. Borfecchia and S. Bordiga, *Top. Catal.*, 2017, **60**, 1664–1672.
- 30 L. Wang, J. Ke, Y. Chai, G. Wu, C. Wang and L. Li, *Angew. Chem., Int. Ed.*, 2025, e202502563.
- 31 H. O. Mohamed, O. Abed, N. Zambrano, P. Castaño and I. Hita, *Ind. Eng. Chem. Res.*, 2022, **61**, 15880–15892.
- 32 K. Lu, F. Jin, G. Wu and Y. Ding, *Sustain. Energy Fuels*, 2019, **3**, 3569–3581.
- 33 R. Y. Brogaard and U. Olsbye, *ACS Catal.*, 2016, **6**, 1205–1214.
- 34 R. Y. Brogaard, M. Komurcu, M. M. Dyballa, A. Botan, V. Van Speybroeck, U. Olsbye and K. De Wispelaere, *ACS Catal.*, 2019, **9**, 5645–5650.
- 35 J. Thakkar, X. Yin and X. Zhang, *ChemCatChem*, 2018, **10**, 4234–4237.
- 36 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.
- 37 B. H. Babu, M. Lee, D. W. Hwang, Y. Kim and H. Chae, *Appl. Catal., A*, 2017, **530**, 48–55.
- 38 R. D. Andrei, M. I. Popa, F. Fajula and V. Hulea, *J. Catal.*, 2015, **323**, 76–84.
- 39 R. D. Andrei, E. Borodina, D. Minoux, N. Nesterenko, J. P. Dath, C. Cammarano and V. Hulea, *Ind. Eng. Chem. Res.*, 2020, **59**, 1746–1752.
- 40 R. D. Andrei, M. I. Popa, F. Fajula and V. Hulea, *J. Catal.*, 2015, **323**, 76–84.
- 41 L. Attanatho, S. Lao-ubol, A. Suemanotham, N. Prasongthum, P. Khowattana, T. Laosombut, N. Duangwongsa, S. Larpkiattaworn and Y. Thanmongkhon, *SN Appl. Sci.*, 2020, **2**, 971.
- 42 T. W. Kim, J. W. Jun, S. I. Hong and C. U. Kim, *J. Nanosci. Nanotechnol.*, 2018, **18**, 2026–2031.
- 43 R. Beucher, R. D. Andrei, C. Cammarano, A. Galarneau, F. Fajula and V. Hulea, *ACS Catal.*, 2018, **8**, 3636–3640.
- 44 R. Beucher, C. Cammarano, E. Rodríguez-Castelló and V. Hulea, *Catal. Commun.*, 2020, **144**, 10609.
- 45 R. Beucher, C. Cammarano, E. Rodríguez-Castellón and V. Hulea, *Ind. Eng. Chem. Res.*, 2020, **59**, 7438–7446.
- 46 R. Beucher, C. Cammarano and V. Hulea, *React. Chem. Eng.*, 2022, **7**, 133–141.
- 47 A. Hwang, S. Kim, G. Kwak, S. K. Kim, H. Park, S. C. Kang, K. Jun and Y. T. Kim, *Catal. Lett.*, 2017, **147**, 1303–1314.
- 48 S. Moussa, P. Concepción, M. A. Arribas and A. Martínez, *Appl. Catal., A*, 2020, **608**, 117831.
- 49 M. Stoyanova, M. Schneider, M. M. Pohl and U. Rodemerck, *Catal. Commun.*, 2017, **92**, 65–69.
- 50 L. A. Perea, M. Felischak, T. Wolff, C. Hamel and A. Seidel-Morgenstern, *Chem. Ing. Tech.*, 2017, **89**, 903–914.
- 51 N. R. Jaegers and E. Iglesia, *J. Am. Chem. Soc.*, 2023, **145**, 6349–6361.
- 52 I. Agirrezabal-Telleria and E. Iglesia, *J. Catal.*, 2017, **352**, 505–514.
- 53 I. Agirrezabal-Telleria and E. Iglesia, *J. Catal.*, 2020, **389**, 690–705.
- 54 W. Li, C. Zhou, W. Li, L. Ge, G. Yu, M. Qiu and X. Chen, *New J. Chem.*, 2022, **46**, 9461–9469.
- 55 M. Ghambarian, M. Ghashghaee, Z. Azizi and M. Balar, *Phys. Chem. Res.*, 2019, **7**, 235–243.
- 56 M. Ghambarian, M. Ghashghaee, Z. Azizi and M. Balar, *Struct. Chem.*, 2019, **30**, 137–150.
- 57 R. D. Andrei, M. I. Popa, C. Cammarano and V. Hulea, *New J. Chem.*, 2016, **40**, 4146–4152.
- 58 K. Toch, J. W. Thybaut and G. B. Marin, *Appl. Catal., A*, 2015, **489**, 292–304.
- 59 A. A. Khudhair, K. Bouchmella, R. D. Andrei, A. Mehdi, P. H. Mutin and V. Hulea, *Microporous Mesoporous Mater.*, 2021, **322**, 111165.



- 60 K. Shimura, S. Yoshida, H. Oikawa and T. Fujitani, *Microporous Mesoporous Mater.*, 2022, **338**, 111955.
- 61 K. Shimura, S. Yoshida, H. Oikawa and T. Fujitani, *Mol. Catal.*, 2022, **528**, 112478.
- 62 A. Al Khudhair, K. Bouchmella, R. D. Andrei, V. Hulea and A. Mehdi, *Molecules*, 2024, **29**, 4172.
- 63 L. Chen, G. Li, Z. Wang, S. Li, M. Zhang and X. Li, *Catalysts*, 2020, **10**, 180.
- 64 J. S. Yoon, M. B. Park, Y. Kim, D. W. Hwang and H. Chae, *Catalysts*, 2019, **9**, 933.
- 65 K. Shimura, S. Yoshida, H. Oikawa and T. Fujitani, *Catalysts*, 2023, **13**, 1303.
- 66 J. Xu, R. Wang, L. Zheng, J. Ma, W. Yan, X. Yang, J. Wang, X. Su and Y. Huang, *Catal. Sci. Technol.*, 2021, **11**, 1510–1518.
- 67 J. Xu, R. Wang, Y. Zhang, L. Li, W. Yan, J. Wang, G. Liu, X. Su, Y. Huang and T. Zhang, *Chin. J. Catal.*, 2021, **42**, 2181–2188.
- 68 Q. Zhang, T. Wanga, Y. Lia, R. Xiaoe, T. Vitidsant, P. Reubroycharoen, C. Wang, Q. Zhang and L. Ma, *Fuel Process. Technol.*, 2017, **167**, 702–710.
- 69 Z. Chen, S. R. Docherty, P. Florian, A. Kierzkowska, I. B. Moroz, P. M. Abdala, C. Copéret, C. R. Müller and A. Fedorov, *Catal. Sci. Technol.*, 2022, **12**, 5861–5868.
- 70 M. Stoyanova, M. Schneider, M. M. Pohl and U. Rodemerck, *Catal. Commun.*, 2017, **92**, 65–69.
- 71 M. Lee, J. Yoon, Y. Kim, J. Yoon, H. Chae, Y. Han and D. Hwang, *Appl. Catal., A*, 2018, **562**, 87–93.
- 72 G. V. S. Seufitelli and R. Gustafson, *Ind. Eng. Chem. Res.*, 2022, **61**, 4286–4299.
- 73 M. Betz, C. Fuchs, T. A. Zevaco, U. Arnold and J. Sauer, *Biomass Bioenergy*, 2022, **166**, 106595.
- 74 C. Fuchs, U. Arnold and J. Sauer, *Chem. Ing. Tech.*, 2023, **95**, 651–657.
- 75 M. H. Kwon, J. S. Yoon, M. Lee, D. W. Hwang, Y. Kim, M. B. Park and H. J. Chae, *Appl. Catal., A*, 2019, **572**, 226–231.
- 76 M. Stoyanova, U. Bentrup, H. Atia, E. V. Kondratenko, D. Linke and U. Rodemerck, *Catal. Sci. Technol.*, 2019, **9**, 3137–3148.
- 77 F. Jin, Y. Yan and G. Wu, *Catal. Today*, 2020, **355**, 48–161.
- 78 A. Aid, R. D. Andrei, S. Amokrane, C. Cammarano, D. Nibou and V. Hulea, *Appl. Clay Sci.*, 2017, **146**, 432–438.
- 79 S. Forget, H. Olivier-Bourbigou and D. Delcroix, *ChemCatChem*, 2017, **9**, 2408–2417.
- 80 L. Li, S. Chavan, Y. Ganjkhanelou, E. Groppo, E. Sagstuen, S. Bordiga, U. Olsbye and K.-J. Jens, *Appl. Catal., A*, 2022, **637**, 118598.
- 81 Y. Cho, J. A. Muhlenkamp, A. G. Oliver and J. C. Hicks, *Chem. Commun.*, 2021, **57**, 13772–13775.
- 82 Y. Cho, A. G. Oliver and J. C. Hicks, *Appl. Catal., A*, 2023, **666**, 119391.
- 83 F. P. Rotzinger, *J. Am. Chem. Soc.*, 1996, **118**, 6760–6766.
- 84 M. Lallemand, A. Finiels, F. Fajula and V. Hulea, *J. Phys. Chem. C*, 2009, **113**, 20360–20364.
- 85 J. Rabeah, J. Radnik, V. Briois, D. Maschmeyer, G. Stochniol, S. Peitz, H. Reeker, C. La Fontaine and A. Brückner, *ACS Catal.*, 2016, **6**, 8224–8228.
- 86 E. Dooryhee, C. R. A. Catlow, J. W. Couves, P. J. Maddox, J. M. Thomas, G. N. Greaves, A. T. Steel and R. P. Townsend, *J. Phys. Chem.*, 1991, **95**, 4514–4521.
- 87 H. A. Aleksandrov, V. R. Zdravkova, M. Y. Mihaylov, P. S. Petkov, G. N. Vayssilov and K. I. Hadjiivanov, *J. Phys. Chem. C*, 2012, **116**, 22823–22831.
- 88 K. Góra-Marek, A. Glanowska and J. Datka, *Microporous Mesoporous Mater.*, 2012, **158**, 162–169.
- 89 K. Kimura, A.-I. Hideo and A. Ozaki, *J. Catal.*, 1970, **18**, 271–280.
- 90 A. Lacarriere, J. Robin, D. Swierczynski, A. Finiels, F. Fajula, F. Luck and V. Hulea, *ChemSusChem*, 2012, **5**, 1787–1792.

