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A review of methanol-to-olefins conversion over SAPO-34: catalyst design, mechanisms, and kinetics

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The methanol-to-olefins (MTO) process has gained significant attention as a promising alternative for light olefin production, particularly because traditional methods are energy-intensive and rely on a limited number of feedstocks. Methanol, on the other hand, is both abundant and can be sustainably derived, supporting the concept of a closed carbon cycle. This literature review combines three decades of key developments in the MTO process, with a focus on the role of SAPO-34, the catalyst which plays a critical part in achieving high selectivity for ethylene and propylene. The review begins by comparing the MTO process with existing olefin production technologies, followed by an overview of how the reaction mechanisms have developed over time, from early-stage concepts to the widely accepted hydrocarbon pool model. Kinetic models developed to describe the complex reactions involved in the MTO process are also examined. A significant part of this review is placed on catalyst design, including strategies for enhancing SAPO-34 performance through structural modifications, metal incorporation, and shaping techniques aimed at extending the catalyst's lifespan. This comprehensive review provides insights intended to inform future advancements in the efficiency and sustainability of MTO-based industrial applications.

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1 Introduction

Light olefins (ethylene and propylene) are often considered one of the key feedstocks for the petrochemical industry. These components take on the role as building blocks and are normally polymerized to form polyethylene and polypropylene, respectively.¹ The significant value of light olefins in today's society is due to the wide range of their derivatives' applications, including packaging, construction, synthetic fibers and coatings.² In terms of market trends, the demand for light olefins has escalated significantly in recent years (see Fig. 1(A)). Fortifying this surge in the light olefins market, future projections seem to portray an even further intensified expansion, which further highlights the need to generate these valuable petrochemical products. Studies estimate that this market will achieve a compound annual growth rate of 5.58% by 2030, capable of generating a revenue of roughly 475.8 million US dollars in this timeframe.³

Light olefin production can occur through a wide array of different processes and technologies, however, traditional methods such as thermal (steam) cracking (SC) and fluid

catalytic cracking (FCC) of hydrocarbons still remain the pillars of today's light olefin production industry (as seen in Fig. 1(B), in the case of propylene production).⁸ Steam cracking, the dominant technology in light olefin production (contributing to 30–50% of total global demand)^{6,7,9–11} can have various feedstocks that are diluted with steam and undergo thermal decomposition/pyrolysis at ~850 °C (at normal pressure of ~1.65 bar) in a tubular furnace and in the absence of oxygen.^{12,13} The products subsequently formed *via* steam cracking are largely dependent on the choice of feedstock (as seen in Fig. 2(A)) as well as reaction conditions. Typically, the conversion lies between 70 and ~95% for the different available feedstocks,¹⁴ where light compounds, like ethane, are more difficult to be cracked and possess a lower conversion. Pyrolysis gas (which contains primarily H₂, CH₄, CO and CO₂) contributes to a roughly 10–25 wt% yield amongst products;¹⁵ hence, steam cracking units remains a heavy threat to the environment, as they are responsible for the emission of more than 300 million tons of CO₂ per annum.¹⁶

Fluid catalytic cracking (FCC), on the other hand, requires operating conditions of lesser severity (a lower reaction temperature of roughly 500 to 650 °C) due to the use of catalytic materials.¹⁹ The process can be manipulated to maximize gasoline, middle distillate, liquified petroleum gas (LPG), or lighter compounds by alteration of the feedstock used, type of catalyst, and operating conditions.²⁰ Nevertheless, it should be

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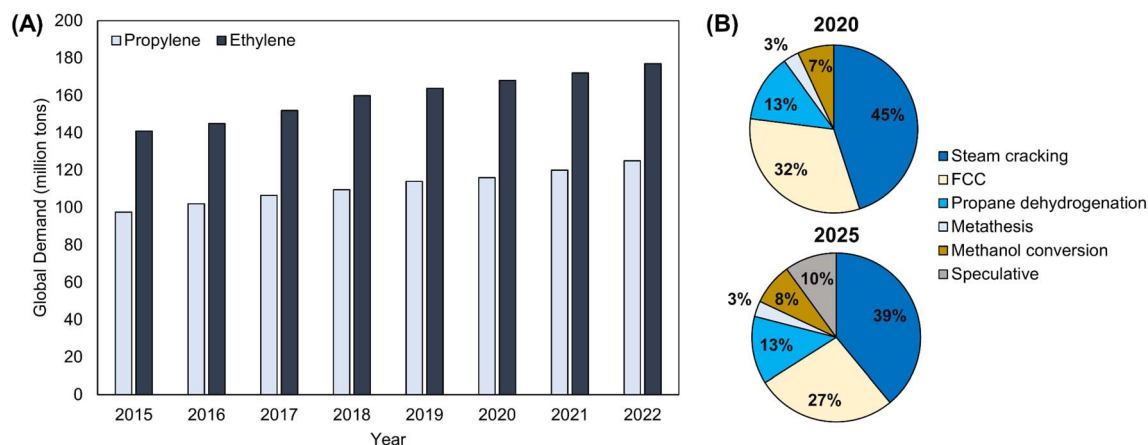


Fig. 1 (A) Global ethylene and propylene demand (consumption) from 2015 to 2022. Adapted from ref. 4 and 5 with permission from Statista, Copyright 2025. (B) Contribution of different processes towards global propylene production in early 2020 and end of 2025 (projected).^{6,7}

noted that, even if the process was adjusted to maximize the production of light compounds (*e.g.*, by using technology licensed by UOP termed PetroFCC), the light olefin yield would still be in the range of 20 to 30 wt% (ref. 21) (see Fig. 2(B)). Besides the low yield of light olefins, another disadvantage associated with FCC could be substantial coke formation (4 to 6 wt%) that instigates rapid catalyst deactivation.²² Evidently, coke removal in the regenerator to reactivate the catalyst coincides with substantial CO₂ emissions.

Another process is dehydrogenation of ethane and propane, which requires the use of a hybrid catalyst – one that possesses both Brønsted and Lewis acidic sites (*e.g.* promoted platinum-based zeolite catalysts and promoted chromium-oxide).^{23,24} These processes typically take place between 535 and 650 °C, under pressures of 0.3 to 6 bar.²⁵ The maximum conversion of propane and ethane in industry are typically around 65% and 20%, respectively, with a selectivity of roughly 90% for propylene and ethylene (notable side products are methane, CO, and CO₂).^{26,27} Overall, the constraints of these processes include the use of costly and environmentally harmful catalysts, high energy consumption, thermodynamic limitations leading

to low single-pass conversion, and restricted availability of feedstocks.²⁸

In general, other than the disadvantages enumerated, since the available resources for these traditional cracking and light alkane dehydrogenation processes are limited and non-renewable (relying on either crude oil fractions or gas condensate (C₂⁺)), these production routes would eventually have to be substituted. As a result, researchers are continuing to look for alternative routes to synthesize light olefins, particularly those that possess a greater abundance of resources and even renewable ones. Methanol to light olefins (MTO) process has started to become more commercially adopted worldwide in recent years, as the feed, methanol, can be generated from a wide range of different carbonaceous resources.^{29,30} Natural gas reserves (154.7 billion metric tons available as of 2017,³¹ which contain ~85 to 97% methane) and gas hydrate reservoirs (3000 billion metric tons, with an assumed 17 to 20% recovery rate, containing roughly 95 to 99% methane^{32–34}), can be converted to syngas, and then ultimately to methanol for use in the MTO process. Reflecting on only methane as a feedstock (derived from either hydrate or natural gas), there are 10 times

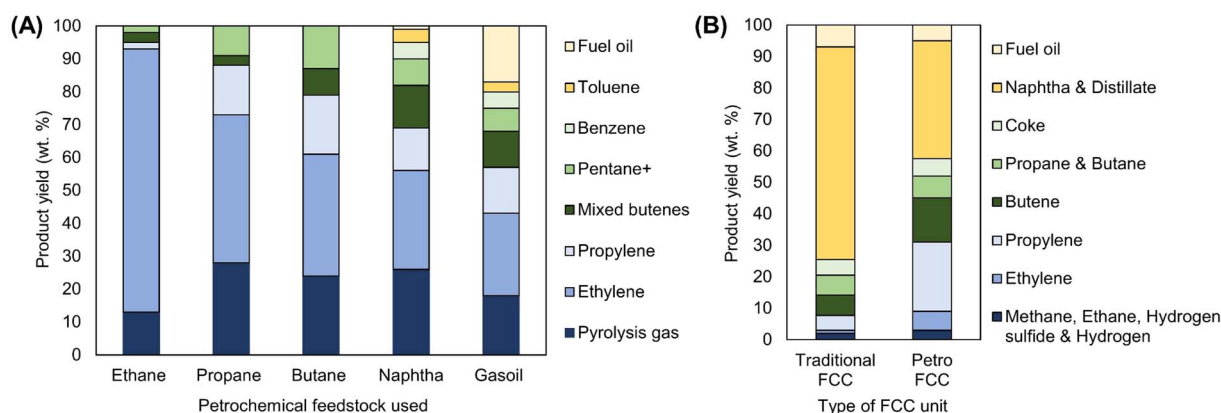


Fig. 2 (A) Steam cracking product distribution (yield) for different petrochemical feedstock. Adapted from ref. 17 with permission from Springer Nature, Copyright 2015. (B) Traditional FCC and PetroFCC product yields using VGO feedstock.¹⁸



greater available resources for MTO compared to the ones for cracking and dehydrogenation processes.^{35,36} Recently, much research has also been allotted towards investigating sustainable methanol production through both the direct and indirect conversion of carbon dioxide.^{37–40} The indirect conversion of CO₂ to methanol has even been commercialized, with the first plant (George Olah) being introduced in Iceland in 2011.^{41,42}

Other than CO₂-derived methanol production, the development of green technologies revolving around a closed-loop carbon cycle has also added to the appeal of the MTO process. Carbon recycling in this context pertains to the indirect conversion of plastic wastes to methanol. Initially, plastic wastes are converted to synthetic gas *via* steam-reforming or partial oxidation, followed by the subsequent conversion of syngas to methanol, which ultimately can be utilized in the MTO process.⁴³ Furthermore, the promotion of carbon recycling can in turn help address the global waste management dilemma (especially with ever rising landfill sites and the negative environmental effects associated with incineration). It should be noted that a multitude of start-up firms have begun to take on the initiative of converting municipal solid waste (and residual biomass) into biofuels, methanol, and other value-added renewable chemicals.⁴⁴ With the aid of these feasible pathways for converting abundant resources to methanol, this value-added chemical has experienced an upsurge in production worldwide (Fig. 3(A)).⁴⁵ Subsequently, to further convert methanol to chemicals in even higher demand, Fig. 3(B) demonstrates that the MTO process ranked as the highest consumption pathway of produced methanol globally (in 2021). This has led to the establishment of the MTO process in the petrochemical industry within the last two decades, contributing ~7% of global light olefin generation (see Fig. 1(B)).

The commercial introduction of the MTO process was first achieved by Union Carbide in 1981 (ref. 47) by alteration of the catalyst of the methanol to gasoline process (MTG) (introducing exceptional shape-selective SAPO-34 as opposed to ZSM-5). The technology adopted by this research group, which is now patented by the Honeywell UOP/Norsk Hydro joint venture, remains the current most prevalent industrially-implemented

MTO technology.⁴⁷ As of yet, SAPO-34 remains the most frequently used catalyst for the process, since it seems to drive peak catalytic performance relative to all other examined silicoaluminophosphate catalysts (as observed in Fig. 4).⁴⁸ Under optimal MTO reaction conditions, SAPO-34's relatively small pores (3.7 Å),⁴⁹ medium acidity (~1 mmol g⁻¹)^{50–52} and high thermal stability (up to 1000 °C)^{53,54} have been recognized as the key reasons for high initial feed conversion (~100%) and high selectivity towards light olefins (~80% total of C₂= and C₃=).^{55–57} The rather small pore size of SAPO-34 inhibits the diffusion of large hydrocarbon species out of the framework, with only smaller sized hydrocarbons (mainly light olefins) being capable of diffusing out of SAPO-34's cages, evidently explaining the selectivity towards these species.

In terms of drawbacks, one major issue that still exists is the rather short catalyst lifetime that SAPO-34 encounters. Over time on stream, it has been experimentally validated that large multi-ring aromatic species (termed coke) build up inside and on the outer surfaces of the framework. These species block catalyst pores and in turn inhibit the diffusion of molecules, resulting in catalyst deactivation.^{60,61} However, a variety of strategies at the lab scale (such as those discussed in Section 4) have been investigated to try and resolve this problem of pore blockage.

SAPO-34 is generally described as having “good hydrothermally stability”, however studies show that prolonged exposure to steam at elevated temperatures gradually weaken its framework through hydrolysis of Si–O–Al and P–O–Al bonds.⁶² Initially, the hydrolysis begins reversibly even at ambient temperature and extends into the MTO range (300 to 500 °C).⁶³ Long-term exposure to moisture eventually causes irreversible hydrolysis, leading to dealumination or desilication, loss of acidity, the formation of silanol defects, and in severe cases partial collapse or amorphization of the structure. Interestingly, coke deposits that form during the MTO cycle can temporarily protect the framework by shielding Brønsted acid sites from water, although they also block the pores and cause deactivation.^{62,64}

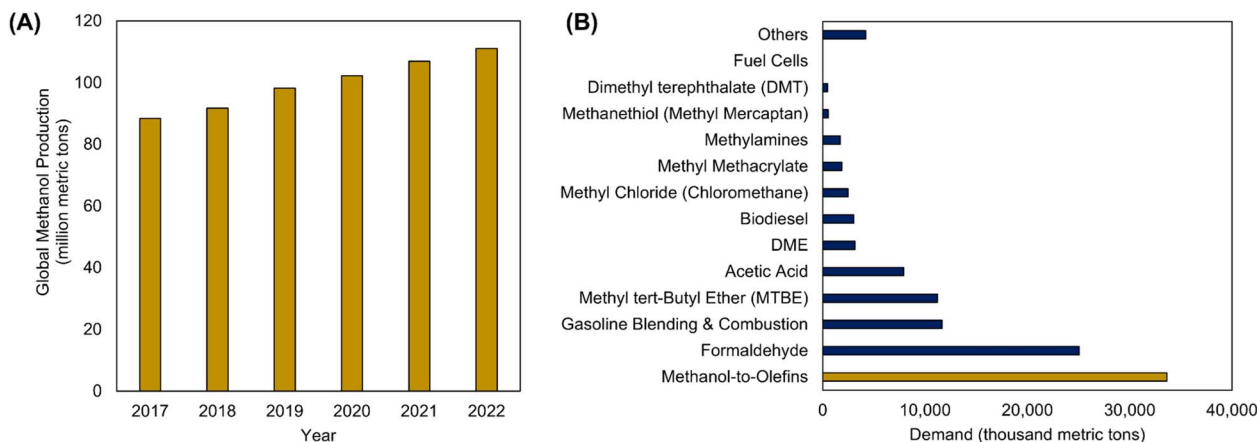


Fig. 3 (A) Global methanol production between 2017 and 2022. (B) Global methanol demand by application in 2021. Adapted from ref. 45 and 46 with permission from Statista, Copyright 2025.



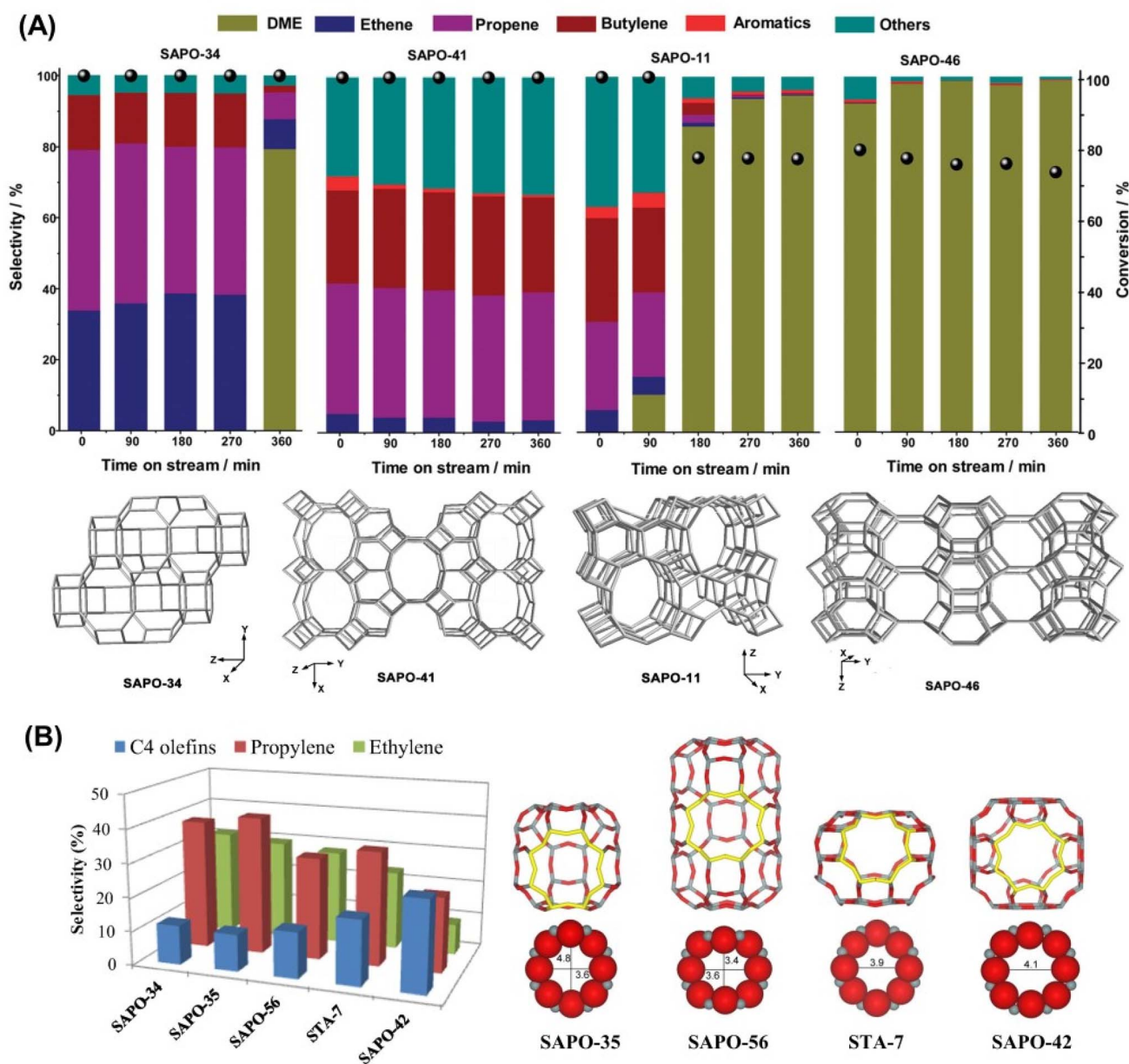


Fig. 4 Product distribution obtained by application of different silicoaluminophosphate catalysts (alongside their topology structure) in the MTO reaction. (A) Parent SAPO-34 compared to SAPO-41, SAPO-11, and SAPO-46. Reaction conditions: $T = 450\text{ }^{\circ}\text{C}$, $p = 1\text{ atm}$ and Weight Hourly Space Velocity (WHSV) = $1\text{ g}_{\text{MeOH}}\text{ g}_{\text{Cat}}^{-1}\text{ h}^{-1}$. Reproduced from ref. 58 with permission from American Chemical Society, Copyright 2015. (B) Parent SAPO-34 compared to SAPO-35, SAPO-56, STA-7 and SAPO-42. Reaction conditions: $T = 400\text{ }^{\circ}\text{C}$, $p = 1\text{ atm}$, WHSV = $0.35\text{ g}_{\text{MeOH}}\text{ g}_{\text{Cat}}^{-1}\text{ h}^{-1}$. Reproduced from ref. 59 with permission from Elsevier, Copyright 2017.

Multiple studies identify critical temperatures where SAPO-34 stability rapidly decline. At around $400\text{ }^{\circ}\text{C}$, which is already within the MTO operating window, reversible hydrolysis sets in, and with time the framework starts to degrade. At higher temperatures, however, controlled steaming can have a stabilizing effect. Treatments at or above about $550\text{ }^{\circ}\text{C}$ cause silicon to migrate within the lattice, which preserves microporosity and strengthens resistance to hydrolysis.⁶⁵ In contrast, steaming at lower temperatures such as at $400\text{ }^{\circ}\text{C}$ can severely damage nanocrystalline SAPO-34. Beyond $700\text{--}800\text{ }^{\circ}\text{C}$, dealumination becomes extensive and the microporous structure collapses,

while above $1000\text{ }^{\circ}\text{C}$ the framework completely amorphizes into a glass-like state.^{66,67}

These structural changes directly affect catalytic performance. Hydrolysis and dealumination reduce the number of active Brønsted sites, lowering conversion activity, while pore collapse hinders diffusion and accelerates coke-induced deactivation.⁶² To address these issues, several strategies have been developed. Controlled high temperature steaming at $550\text{--}600\text{ }^{\circ}\text{C}$ can improve hydrothermal stability by restructuring the silicon environment. Acid-site passivation techniques such as silylation or temporary ammonia treatment reduce the



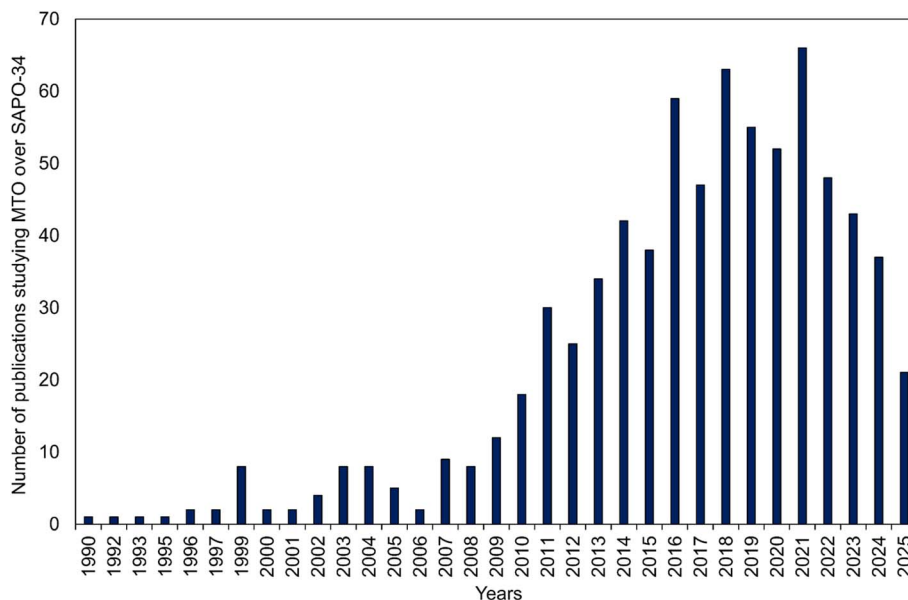


Fig. 5 Number of publications related to methanol conversion to light olefins over SAPO-34 published per year from 1990 to 2025. Data collected from the Web of Science Core Collection.⁷¹

vulnerability of framework bonds to hydrolysis.^{68,69} Incorporating metal oxides like indium oxide, ceria, or magnesium into SAPO-34 has proven effective in redistributing acidity, creating mesoporosity, suppressing coke, and thereby extending catalyst lifetime and olefin selectivity.⁷⁰ Morphological engineering, such as synthesizing hierarchical or nanosized SAPO-34 (discussed in Section 4.4) can alleviate diffusion limitations and slow coking, though this often comes at the expense of intrinsic

framework stability. On the operational side, strategies like mild water co-feeding and carefully controlled regeneration are used to moderate coke deposition and limit steam-induced degradation.⁶²

To date, research is continuously being allocated towards the redesign and modification of the catalyst in the hope of mitigating its short lifetime. Fig. 5 demonstrates the surge in research that has been allocated towards MTO process over the

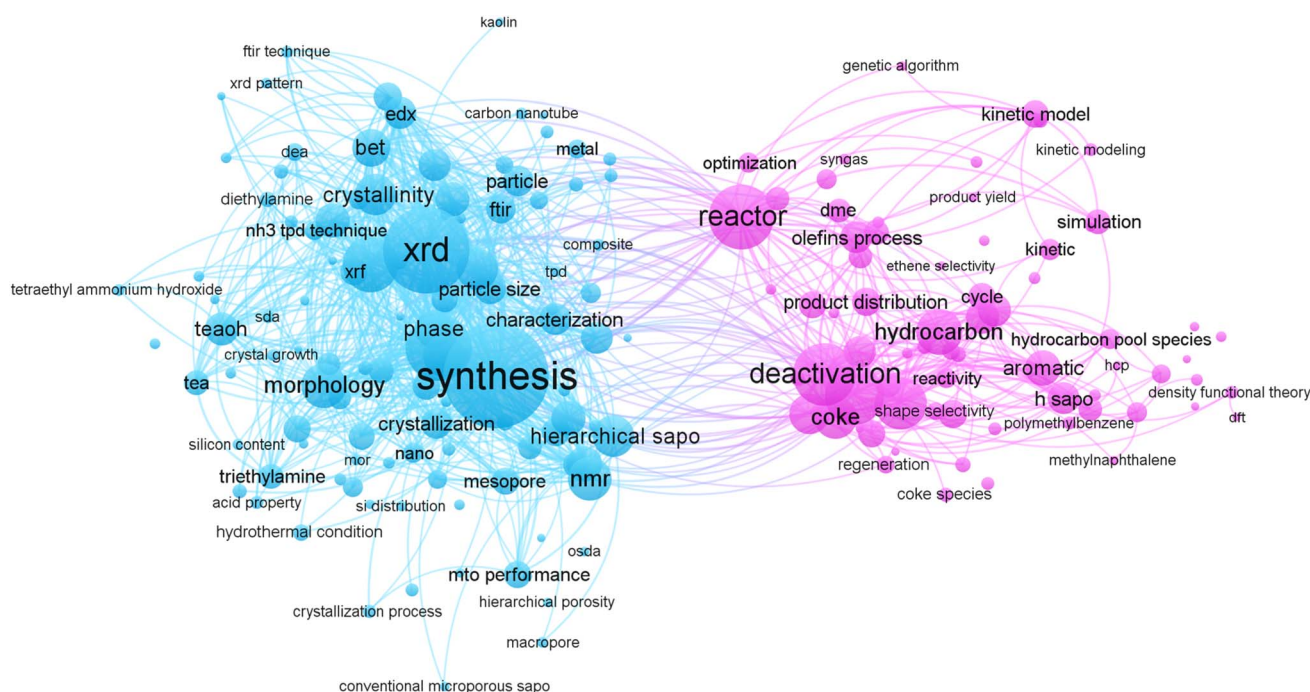


Fig. 6 Bibliometric map generated by VOSviewer,⁷² based on 669 articles (keywords "MTO" and "SAPO-34") from the Web of Science Core Collection.⁷¹



last 30 years, which can project both the importance and emergence of this process as a means for sustainable light olefins production.

A bibliometric mapping image (Fig. 6) has been formulated to decipher the most prominent terms and themes present in MTO-related articles between the years of 1999 and mid-2025. The size of the circles in the schematic correlates with the number of occurrences of a term. It seems there are two distinct clusters that correspond to (i) SAPO-34 synthesis and characterization (blue) and (ii) reaction and mechanistic studies (purple). The “reactor” seems to act as the interlinking concept between these cluster topics. The rationale lies in the fact that no matter whether a group is developing the catalyst's synthesis or studying the reaction mechanism and kinetics, conducting the activity test using a reactor is a vital step.

Considering all the justifications mentioned in the introduction regarding the importance of this process, alongside the extensive attention that has been brought to this topic from researchers worldwide in the last three decades (Fig. 5), we were encouraged to conduct a comprehensive literature review on all the important aspects of this process. In the following Sections, the literature review will be dedicated specifically to methanol conversion to light olefins over SAPO-34 catalyst. In detail, we will commence with a description of the process and the operating conditions that influence it, followed by an analysis of the reaction mechanism and deactivation pathways, reaction kinetics, and catalyst design and modification.

2 MTO process

2.1 Process description & process flow diagram

The simplified process flow diagram of the methanol conversion to light olefins process, based on the technology developed

by the Honeywell UOP/Norsk Hydro joint venture, has been presented in Fig. 7. Methanol diluted with 20 wt% water is being warmed and vaporized by recovering heat from reactor effluent in E-101 heat exchanger. The methanol feed then passes through the feed distributor and fluidizes the fresh catalyst coming back from the regenerator. The regenerated catalyst (having a temperature of about 600 °C) heats up the feed to the reaction temperature (around 410 °C). The MTO reaction takes place in the reactor riser, and gaseous products along with the fluidized catalyst particles enter the disengagement section of the reactor. The higher cross-sectional area of this section causes an instant drop in the gas velocity, which in turn triggers the fall of most of the catalyst into the dense phase catalyst section known as the catalyst inventory. A small portion of the catalyst, that is entrained in the gas phase, will be separated using multiple successive cyclones. Simultaneously, a portion of the catalyst is being transported to the generator to burn the partial coke formed *via* air. The reactor effluent is quenched (to remove the water and unreacted methanol) and then compressed to be directed to products separation and purification units. After caustic wash (to remove carbon dioxide) and drying (to remove the moisture), the gas passes through the de-ethanizer where ethylene, ethane, and lighter compounds are being separated from heavier ones. After the acetylene converter, the fuel gas (methane and hydrogen) will be removed, followed by ethylene purification. In the heavy compounds section, high-purity propylene is being produced, where C_4^+ species are being sent to the olefin cracker to enhance the overall light olefin yield of the process.⁷³ The basic MTO process can yield 75–80% (carbon-based) ethylene plus propylene, 2–5% coke, 2–5% alkanes, and 0.5–1.0% Dimethyl Ether (DME), where the olefin cracker can improve the light olefin yield up to an extra 10%.

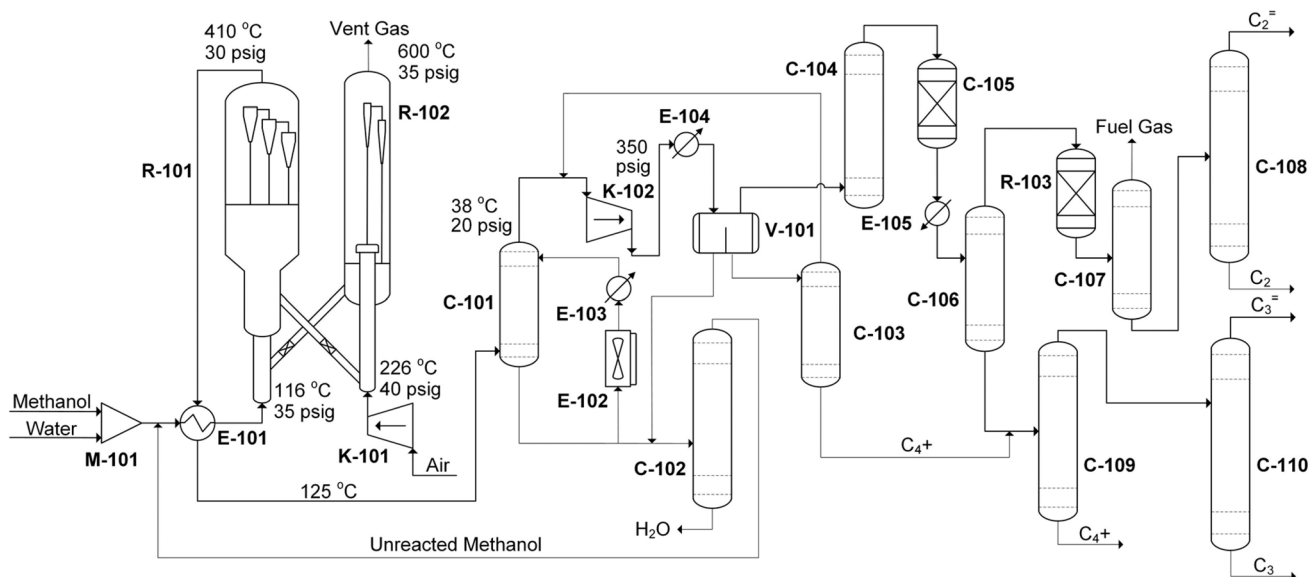


Fig. 7 Process flow diagram of MTO process adapted from UOP Co.⁷⁴ Unit identification: M-101 mixer; E-101 heat exchanger; R-101 fluidized reactor; R-102 regenerator; K-101 air compressor; C-101 quench tower; E-102 air-cooled exchanger; C-102 water stripper; K-102 product compressor; V-101 phase separator; C-103 condensate stripper; C-104 caustic scrubber; C-105 gas dryers; C-106 De-ethanizer; R-103 acetylene converter; C-107 De-methanizer; C-108 C_2 splitter; C-109 De-propanizer; C-110 C_3 splitter.



2.2 MTO products and coke evolution

In an industrial MTO unit, by employing a fluidized bed reactor with continuous regeneration of the catalyst, the coke content of the overall catalyst inventory within the reactor is maintained constant, resulting in a stable and unchanged product distribution. In contrast, in a fixed bed reactor, due to gradual coke buildup, the products evolve over time on stream, shedding light on important phenomena. As illustrated in Fig. 8(A and B), gradual coke formation narrows the pores of the catalyst, which in turn makes the production of smaller molecules more favorable.^{75,76} As a result, selectivity towards methane, ethylene, and ethane is increasing, while the formation of heavier compounds such as propylene, butene, and C₅⁺ olefins is being hindered. This continues until the formed coke completely blocks the pores of the catalyst, causing catalyst deactivation, where dimethyl ether (an intermediate) and unreacted methanol emerge. Interesting findings were revealed by the analysis conducted *via* Ghavipour *et al.*,⁷⁶ who transformed a thermogravimetric analyzer into a TG reactor in the pursuit of performing live/*in situ* measurements of coke generation during the MTO reaction over SAPO-34, in turn relating this to the

products' evolution. As seen in Fig. 8(C and D), after weight reduction due to moisture desorption, the MTO reaction commenced by feeding 7 mol% methanol in argon. Opposed to the speculation that the coke formation rate might be higher in the early stage of the reaction due to the higher number of active sites available, in reality, the coke formation gradually increases until it reaches the complete deactivation stage. According to Fig. 8(C), the group recorded a minor weight gain in the early stages of the reaction due to formation of entrained species such as hydrocarbon pool (HCP) autocatalytic intermediates. This was followed by moderate coke formation rate and subsequently enhanced coke formation towards the advanced times on stream (close to deactivation) due to the formation of poly-cyclic aromatic hydrocarbons. The group was able to approve this phenomenon using ¹³C NMR over partially and fully coked catalyst, explaining this through the schematic shown in Fig. 8(E and F). In the beginning of the reaction, as methanol molecules penetrate the catalyst pores, all the products, regardless of their sizes, can diffuse out. Conversely, as partial coke restricts the pores of the catalyst in advanced times on stream, methanol molecules still enter and light products

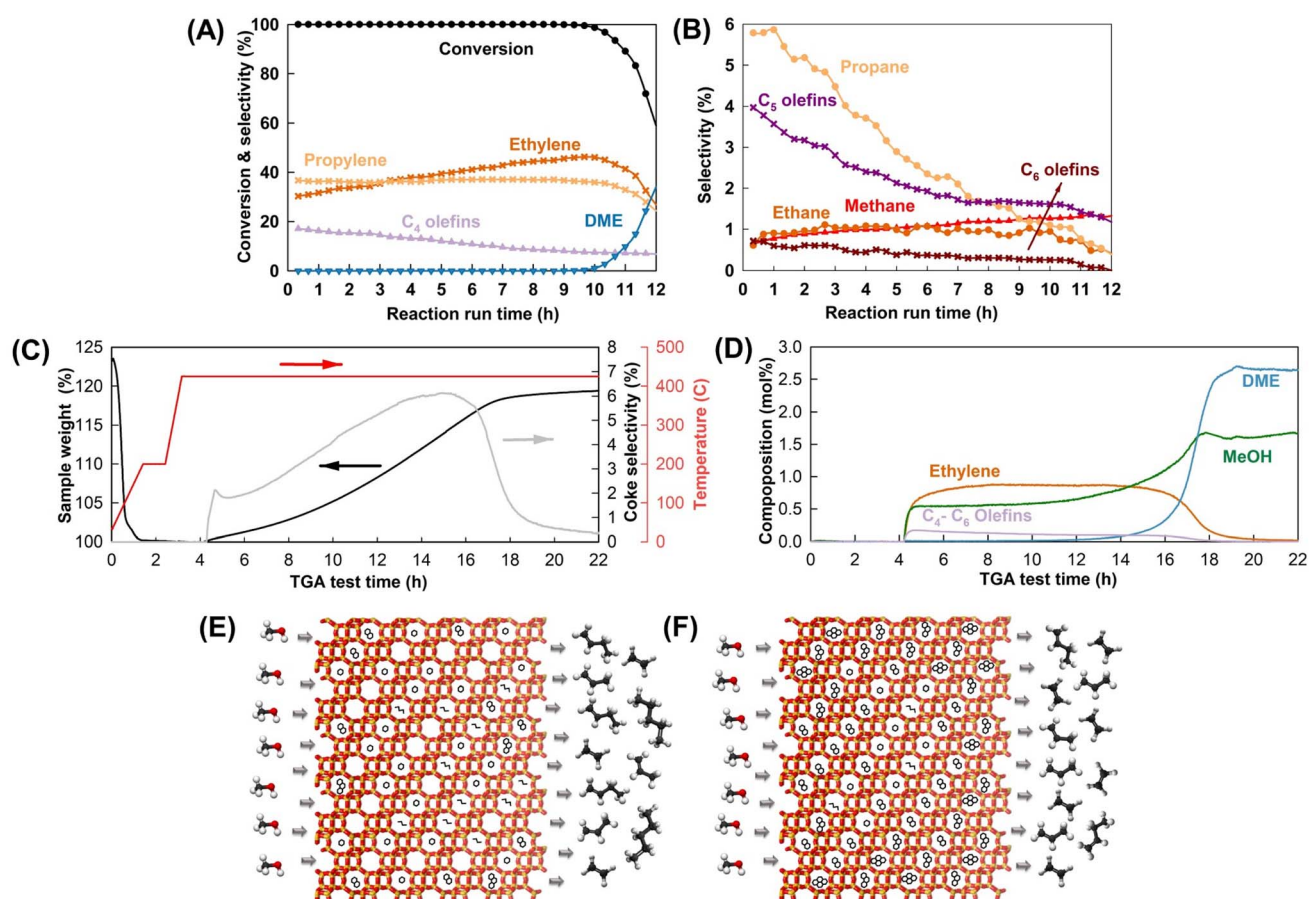


Fig. 8 (A and B) Products distribution *versus* time on stream over SAPO-34 catalyst in the MTO reaction. Reaction conditions: temperature of 400 °C, atmospheric pressure, WHSV of 1 g_{MeOH} g_{Cat}⁻¹ h⁻¹, and methanol diluted with 20 wt% water. (C and D) TGA-MS analysis: coke formation measurement using TGA and products distribution acquisition by MS. Reaction conditions: temperature of 400 °C, atmospheric pressure, WHSV of 1 g_{MeOH} g_{Cat}⁻¹ h⁻¹, and methanol diluted with 93 mol% argon. (E and F) Coke evolution in the pores of the SAPO-34 catalyst in (E) the beginning of the activity run, and (F) in enhanced times on stream close to catalyst deactivation. Reproduced from ref. 76 with permission from American Chemical Society, Copyright 2023.



exit, while heavier products become entrapped in the cavities and further grow. This causes faster coke formation in the later stages of the MTO reaction.

2.3 Reactor operating conditions

The influence of MTO reaction conditions on SAPO-34 catalytic performance is an area of research that has been well investigated. One important factor that researchers have considered is the operating temperature, which plays a large role in establishing product distribution. It has been found that the optimal temperature exists in the range of 400 to 460 °C (by weighing the combined values of both light olefins selectivity and lifetime, as can be seen in Fig. 9(A)).^{77–80} Reactions taking place at a temperature higher than the optimum range sustain an acceptable total average light olefins selectivity but suffer from lower catalyst active lifetime (due to enhanced rate of coke generation) and heightened light alkane (primarily methane) formation.^{78,81,82} At lower temperatures (smaller than 400 °C), on the other hand, the heat required to overcome the activation energy of the reaction seems to be too minimal, resulting in incomplete conversion of methanol and overall lower active lifetime.^{78,82} In general, higher temperature results in greater light compounds production as $C_2=C_3$ increases from 1.2 to 2.7 within temperature alteration of 370 °C to 490 °C.⁷⁸ Researchers have related the dependency of products distribution on reaction temperature to the type of activating and deactivating species present at specific reaction temperature ranges.⁸²

Other than operating temperature, one should consider the role that weight hourly space velocity (WHSV) may impose on catalytic activity. It has been found that the MTO reaction performs optimally under WHSVs of approximately 1 to 2 $g_{MeOH} g_{Cat}^{-1} h^{-1}$ (as seen in Fig. 9(B)).^{80,85} A WHSV below 1 $g_{MeOH} g_{Cat}^{-1} h^{-1}$ has also been proven to reduce the total average selectivity of $C_2=C_4$ species. This has been attributed to the longer residence time of species in the reactor, which can permit secondary reactions to take place and ultimately alter the product distribution.⁸⁰ The use of WHSVs larger than the aforementioned optimal range results in shortened catalyst active lifetime and, in even higher values, causes incomplete methanol conversion.⁸³

Assuming this stipulated ideal WHSV has been employed, groups have also investigated the effect that reactor pressure may impose on MTO product distribution. Generally, it has been found that the optimal pressure, taking into account the combination of both LOS and feed conversion, lies between 1 and 2 bar.^{86,87} A reactor pressure greater than this has been shown to negatively impact the average selectivity towards light olefins (see Fig. 9(C)) and encourage greater deposition of coke instead.^{87,88} This has been justified by the consequential increased residence time of reactants in the reactor, which encourages a greater likelihood of light olefins undergoing successive reactions to form byproducts. Furthermore, raising the pressure would favor the backward reaction in the conversion of, for instance, methanol to ethylene (from 2 to 3 moles of gas) or methanol to propylene (from 3 to 4 moles of gas),

according to Le Chatelier's principle. It should be noted that, on the other hand, a pressure too low (keeping other factors, such as space velocity, constant) would result in reduced residence time of reactants, thus resulting in incomplete conversion of methanol over the catalyst active lifetime.⁸⁷

Another important consideration is the impact feed composition may have on product distribution and catalyst lifetime. Studies have demonstrated that the use of diluents in the feed, such as water, decreases the partial pressure of methanol, which has correspondingly boosted selectivity towards light olefins and delayed catalyst deactivation.^{89,90} Although nitrogen has been utilized as an alternative diluent, water is readily available, possesses a relatively higher heat capacity as a vapor (to carry away the heat of reaction), and has been shown to more effectively reduce the rate of coke formation and further improve catalyst utilization.^{91,92} Since water will tend to reversibly adsorb on catalyst acidic sites (primarily strong ones) due to its basic properties, it has been deemed to reduce the residence time of coke precursors adsorbed on such active sites as well as inhibit the coke-generating steps involving alkene monomers undergoing oligomerization and cyclization.^{91,93,94} Furthermore, problems involving product separation may arise when diluents other than water are employed.⁹² Employing 70–80 mol% of water in the feed has proven to process roughly eight times more methanol (within the catalyst active period) than if the feed had consisted of only pure methanol.⁹² Exceeding this range (90 mol% water) has not been shown to significantly improve performance.⁹⁵ It is important to note that addition of substantial amounts of water will result in occupation of the reactor volume, which consequently lowers the reactor's nominal processing ability. Besides, heating up the water to the reaction temperature and condensing the outlet water, despite the application of heat recovery methods, can result in heat loss and decreased energy efficiency. Hence, in the industrial application, feed water content is usually kept around 30 mol% (20 wt% water in methanol).⁷⁴

Other than the conventional use of water/nitrogen diluent in the feed, groups have also investigated the potential effects that other co-fed gases like hydrogen may impose on catalytic activity.^{84,96} According to Fig. 9(D), Zhao *et al.*⁸⁴ reported that the SAPO-34 fed with pure methanol lasted for less than an hour; however, following hydrogen addition in the $H_2/MeOH$ ratios of 8.3, the lifetime was substantially enhanced to around 80 hours. However, the light olefins were inevitably hydrogenated to alkanes, leading to a considerable drop in light olefins selectivity, from around 70% using pure methanol, to around 50% in the presence of hydrogen. Another group⁹⁶ claimed milder reduction in light olefins selectivity in response to hydrogen addition (LOS of 76% for $0H_2/MeOH$ molar ratio compared to LOS of 70.5% for $231H_2/MeOH$ molar ratio) and over 70 times longer lifetime. Nonetheless, the reaction conditions that they employed (WHSV of 40 $g_{MeOH} g_{Cat}^{-1} h^{-1}$ over only 4 mg of catalyst) seem to be far from the optimal range. Interestingly, the simultaneous presence of water and hydrogen in the feed seems to trigger a synergistic effect, leading to long lifetime without compromising LOS (see Fig. 9(D)). Preceding studies have claimed that the fed H_2 may play a role in hydrogen



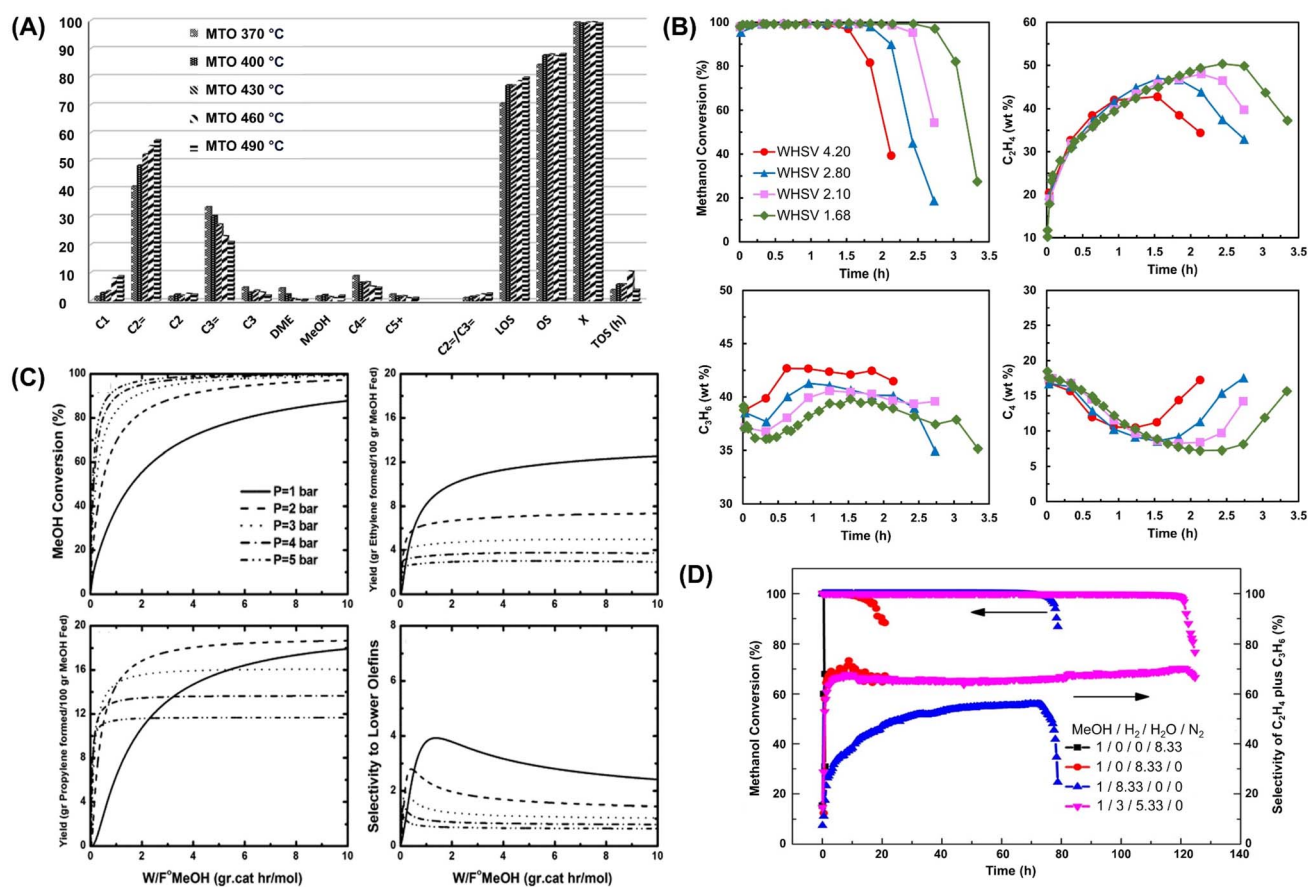


Fig. 9 (A) Product distribution of MTO reaction over SAPO-34 catalyst at different T . Notation: LOS = light olefin selectivity, OS = olefin selectivity, X = conversion, TOS = time on stream. Reaction conditions: 0.1 barg pressure, WHSV of $1 \text{ g}_{\text{MeOH}} \text{ g}_{\text{Cat}}^{-1} \text{ h}^{-1}$, and methanol diluted with 20 wt% water. Reproduced from ref. 78 with permission from Royal Society of Chemistry, Copyright 2016. (B) Effect of WHSV (in $\text{g}_{\text{MeOH}} \text{ g}_{\text{Cat}}^{-1} \text{ h}^{-1}$) on conversion of methanol, selectivity to ethylene, propylene, and butylene over time on stream reproduced from ref. 83 with permission from John Wiley and Sons, Copyright 2018. Reaction conditions: T of 450 °C, atm pressure, and methanol diluted with 20 wt% water. (C) The effects of pressure alteration on methanol conversion, ethylene yield, propylene yield and selectivity towards light olefins. Reproduced from ref. 79 with permission from Elsevier, Copyright 2011. Reaction conditions: T of 400 °C, WHSV of $12 \text{ g}_{\text{MeOH}} \text{ g}_{\text{Cat}}^{-1} \text{ h}^{-1}$, and methanol diluted with 69 wt% water. (D) Synergistic effect between H₂O and H₂ on the catalytic performance of SAPO-34 in the MTO process. Reaction conditions: T of 450 °C, 40 bar, and WHSV of $4.0 \text{ g}_{\text{MeOH}} \text{ g}_{\text{Cat}}^{-1} \text{ h}^{-1}$. Reproduced from ref. 84 with permission from American Chemical Society, Copyright 2019.

transfer reactions and seemingly intercept the pathways that lead to the generation of deactivating multi-ring aromatic species. It was proclaimed that, under combined feeding of H₂ and H₂O,⁸⁴ heavy aromatic deposits that formed were likely to undergo hydrogenation catalyzed by SAPO-34 Brønsted acidic sites,⁹⁷ followed by possible cracking (depending on the temperature employed). These species were declared to have likely reverted back to active intermediates and in turn decelerate the overall evolution of coke species.⁸⁴ Periodic density functional theory (DFT) modelling that ensued focused on investigating these hydrogenation-based mechanistic claims that revolved around hydrogen co-feeding.⁹⁸ It was uncovered that potential deactivation-inducing precursors such as formaldehyde and dienes, which have been claimed to drive the link between olefinic and aromatic dual cycles (see Section 4.3.1 on polyalkylaromatics formation), are selectively hydrogenated (*i.e.*, at a higher rate) relative to desired light alkene species. Hence, the elimination of such potential aromatic precursors by

hydrogenation, *via* high-pressure H₂ co-feeding, may be the plausible cause for the extended lifetime observed in the studies mentioned previously. Recently, the potential of undertaking MTO catalysis in the presence of syngas reagents (cofeeding a mix of H₂ and CO under high pressure) has been examined.⁹⁹ However, under elevated methanol partial pressure, which begins to more closely resemble the MTO process, co-feeding CO (in both presence and absence of H₂) resulted in no clear improvement in any catalytic performance metric.

3 MTO reaction and deactivation pathways

The uncovering of the MTO mechanism has been and continues to be a challenge for researchers. Over the last 60 years, a vast number of mechanistic proposals have been put forward. The pathways that have gained increasing popularity over the years are depicted as a simplified schematic in Fig. 10. In general, the



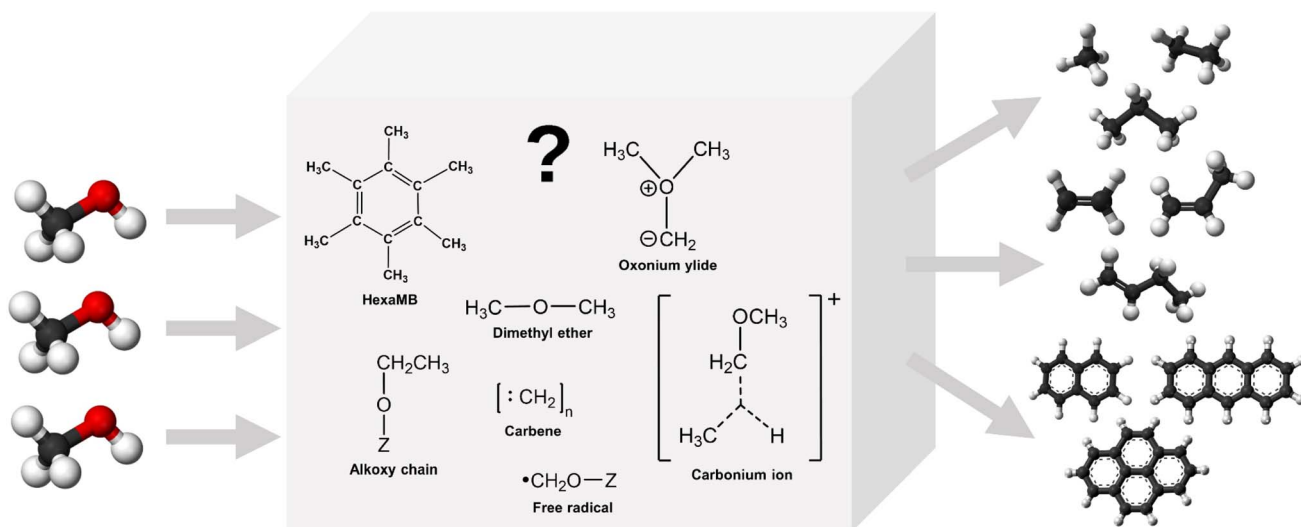


Fig. 10 Simplified schematic of the MTO overall reaction.

elementary steps of the first C–C coupling (in the formation of light olefins) are still under debate. However, it has been widely accepted that early stage constructed light olefins are likely to undergo further reactions. These include hydrogen transfer, alkylation, cyclization, and aromatization to form the following species: paraffins, larger olefins, naphthenes, and aromatics. The generation of large multi-ring aromatic species can, in turn, lead to the “coking” and hence the deactivation of the zeolite catalyst.

That being said, there still exists extensive discussion with regards to the reaction intermediates or active species leading to the previously mentioned hydrocarbon products. Some experimental evidence has ratified the hypothesis of dimethyl ether (DME) being an intermediate of the MTO reaction.^{100–102} For instance, DME is the sole product observed following catalyst deactivation, which means it is produced over weaker acidic sites and may be formed and consumed over the active catalyst. In addition, feeding the reactor with DME instead of methanol results in almost the same product distribution, which raises the idea that DME can act as an active intermediate.¹⁰³ However, there are also mechanistic proposals that introduce other species as active intermediates, such as the free radical mechanism, the alkoxy chain growth route, and the hydrocarbon pool concept (which suggests polyalkylaromatics as active components). In the following sections, the most widely utilized mechanisms of the MTO reaction will be explored.

3.1 Early studies: initial stages of light olefin production

The most probable path behind the first C–C bond formation has been the main topic of discussion for earlier studies conducted on the MTO process, lying primarily between the years of 1970 and 1990. A large number of mechanistic proposals have been put forward; however, the most widely discussed and relevant mechanisms have been relayed as follows.

3.1.1 Oxonium ylide mechanism. During early mechanistic research, the oxonium ylide mechanism (as seen in Fig. 11) has

received recognition as one of the viable paths for the formation of the initial C–C bond.^{104–106} It has been suggested that, in the presence of Brønsted acidic sites (H^+) of zeolite catalysts, the initial step (A) consists of the dehydration of two methanol molecules to a singular dimethyl ether (DME). This DME intermediate may interact with a Brønsted acidic site of the zeolite catalyst, to in turn form a dimethyl oxonium ion. These two preceding species can react further to produce a trimethyl oxonium ion (*i.e.*, step B). This is then followed by a deprotonation step (C) of one of the methyl groups by a basic site, to in turn formulate a surface-associated dimethyl oxonium methyl ylide species. The subsequent step comprises of two options, either a Stevens rearrangement (step D) or an intermolecular methylation (step D'). *Via* the Stevens rearrangement, the ylide species is converted to a methylethyl ether (as well as a singular methanol molecule), which is then ultimately transformed to ethylene *via* a β -elimination step (E). On the other hand, with the intermolecular methylation (the second option), an ethyldimethyl oxonium ion forms. This is once again followed by a β -elimination step (E') to produce the ethylene species, while regenerating a DME molecule in this case. An isotope labelling study¹⁰⁷ reached a conclusion that the intermolecular methylation (step D') is the preferred pathway in comparison to the potential ‘Stevens rearrangement’ (step D).

3.1.2 Carbene mechanism. Another proposed route that may be responsible for early-stage light olefin production is commonly referred to as the ‘carbene mechanism’ (Fig. 12).^{108,109} This mechanism commences *via* an initial α -elimination step (A), in which water and a carbene species form from the methanol reactant. This can then be followed by either polymerization of the carbene monomers to olefins (steps B and C), or *via* sp^3 insertion of the carbene species in dimethyl ether or methanol molecules (step D). The newly generated intermediates can then undergo cleavage to form olefins, as well as regenerate water or alcohol species (step E).



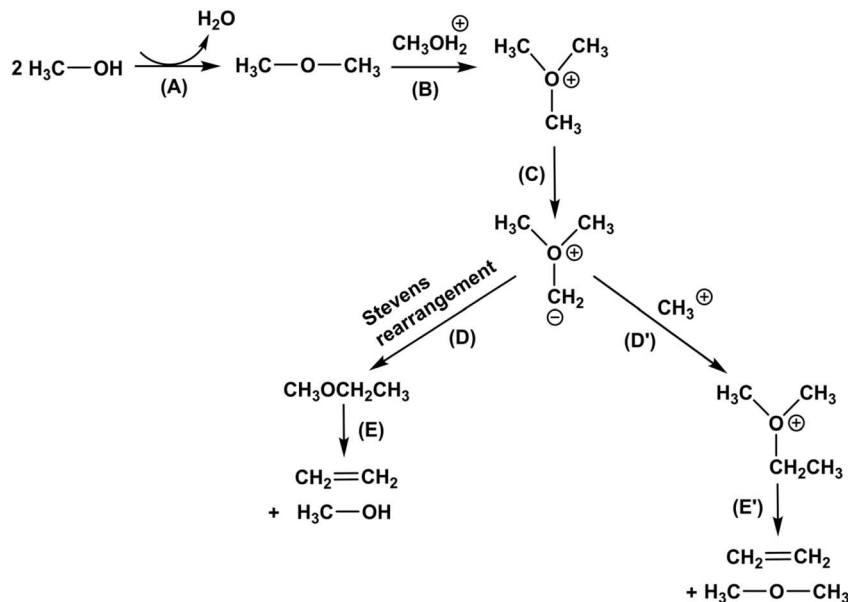


Fig. 11 Oxonium ylide mechanism. Adapted from ref. 104 with permission from the Royal Society of Chemistry, Copyright 2023.

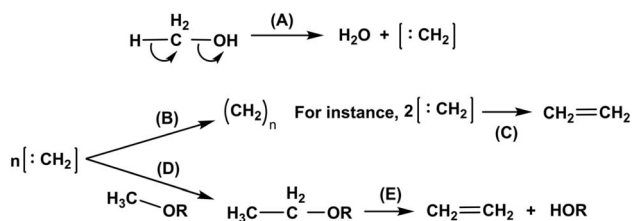


Fig. 12 Carbene mechanism. Adapted from ref. 108 with permission from Elsevier, Copyright 1981.

The search for supporting evidence behind carbene intermediacy was undertaken by isotope-labeling experiments involving ^{13}C methanol, in the presence of propane, over HZSM-5 zeolites.¹¹⁰ Here, the resulting ratio of iso-over normal butane produced was substantially low (*i.e.*, ~ 1 compared to 3.8 for the control run, which comprised a feed of only methanol and helium instead). This can provide evidence, as suggested by the authors, that the nature of the C_1 attacking species must be carbene-like. This is because insertion into a sp^3 C-H bond (a mode of attack by carbene species) statistically leads to a low *i/n* butane ratio. This low ratio can also act to disprove the existence of both the 'oxonium ylide' and the to-be-introduced 'carbocationic' mechanisms (as the suggested paths for initial C-C bond formation), since such routes have been characteristically shown to display higher *i/n* butane ratios instead. It was also revealed that the produced butanes were singly labeled, which validated the direct methylation of propane *via* methanol.

3.1.3 Carbocationic mechanism. Some research groups have looked to the carbocationic mechanism as a possible path for initial C-C bond formation and early-stage light olefin production.¹¹¹⁻¹¹³ Fig. 13 portrays the elementary steps that comprise this mechanism. The pathway commences *via* the formation of dimethyl ether (from methanol) through

a reversible dehydration step (A). *Via* protonation, dimethyl ether (*i.e.*, a surface methoxyl) may be the provider of free methyl cations (step B), which could in turn react with another dimethyl ether molecule by addition to its C-H bond (step C). In turn, a carbonium transition state can form; its subsequent deprotonation (by the anionic zeolite catalyst) can lead to the production of methylethyl ether (methoxyethane) while regenerating the catalyst Brønsted acidic site (step D). The methylethyl ether can then ultimately transform to ethylene *via* a β -elimination step (E), like that seen in the final step for the 'oxonium ylide mechanism'. However, the more likely path instead involves the methoxyethane species further reacting with free methyl cations (from dimethyl ether in its protonated form) to in turn generate a pentavalent carbonium transition state (step F). This can then follow *via* deprotonation to produce methyl isopropyl ether (step G), which could finally eliminate methanol to produce propylene (step H). It could, though unlikely, repeat the cycle of methyl carbonium ion addition to form even larger olefins.

It could be claimed that the C-H bond of DME (or methanol) may not be of sufficient nucleophilicity to undergo an attack by the electrophilic methyl cations (derived from dimethyl ether in its protonated form), as seen in step (C). Smith and Futrell¹¹⁴ embarked on an investigation that assisted in providing an answer to this claim. The group undertook experiments in which they reacted methylcarbonium ion (CH_3^+) with methanol. It was concluded that the reaction took place majorly (*i.e.*, 80%) through intermediates structured as dimethyl-oxonium ion or protonated DME. However, according to the tandem ion cyclotron resonance mass spectrometer, C_2 species (such as ethylene) were not detected as part of the product distribution; rather, C_1 species were present. Hence, these observations can question the overall validity of the carbocationic mechanism in its capability for ethylene production.



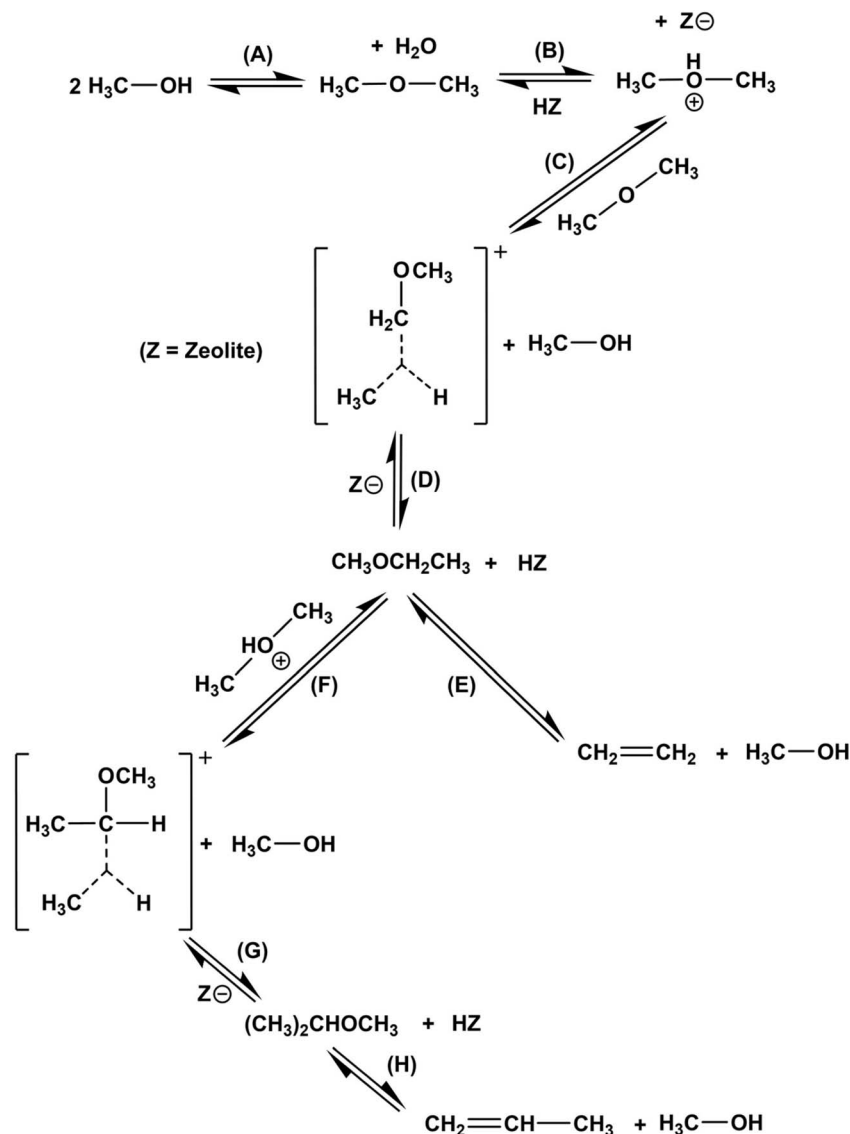


Fig. 13 Carbocationic mechanism. Adapted from ref. 112 with permission from Elsevier, Copyright 1981.

3.1.4 Free radical mechanism. A small portion of articles (during early mechanistic research) have proposed the participation of free radicals in the initial-stage production of light olefins,¹¹⁵⁻¹¹⁷ which have been depicted in Fig. 14. Initially, a methanol molecule approaches a Brønsted acidic site of the zeolite catalyst, resulting in a dehydration reaction in which water and a surface methoxy group are formed (step A). Simultaneously, it has been declared that the interaction of methanol molecules on radicogenic sites of the catalyst lead to the generation of free methyl radicals.¹¹⁷ Consecutive reactions lead to the carbon chain buildup of these radical species. Subsequently, the methoxy group on the zeolite surface interacts with these radical carbonaceous species (such as an alkyl, alkoxy, alkoxyalkyl) to in turn form a radical intermediate (step B). Further interaction between this intermediate and the zeolite surface ultimately leads to the production of surface-bound carbene species *i.e.*, methylene (step C). This can then

be followed by polymerization of the carbene monomers to olefins (step D). A final step (F) involves the regeneration of the zeolite acidic site (ZOH).

Experimental evidence in support of the existence of free radical intermediates was presented by Clarke *et al.*¹¹⁶ The group conducted experiments in which they reacted DME over

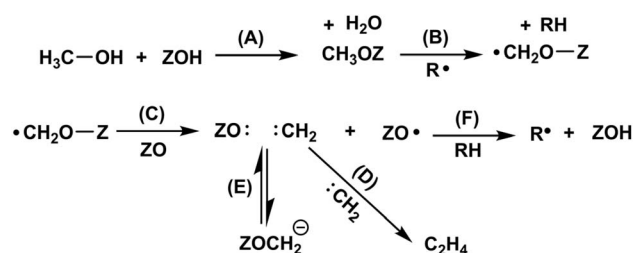


Fig. 14 Free radical mechanism. Adapted from ref. 115 with permission from Elsevier, Copyright 1989.



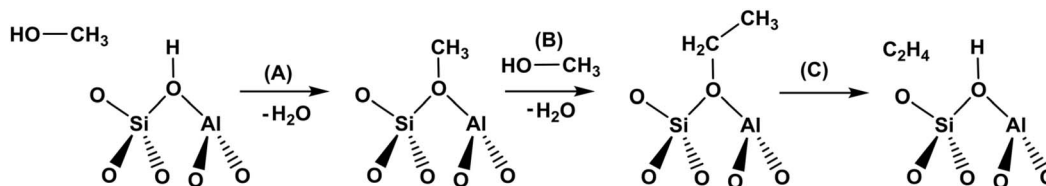


Fig. 15 Alkoxy chain growth route. Adapted from ref. 119 with permission from American Chemical Society, Copyright 2003.

H-ZSM-5 zeolite; using a spin trapping reagent with electron spin resonance (e.s.r.) analysis, free radicals were said to be detected during the reaction. Clarke *et al.* were also able to demonstrate that the source of these free radicals stemmed from small quantities of solid-state paramagnetic defects present in the zeolite, which were able to activate C–H bonds in DME (on ZSM-5 catalyst at ≥ 200 °C). Other tests were also performed using methanol fed over a catalyst comprising of 10 wt% MgO with 1 wt% Li₂O supported on a silica-alumina (13% wt Al₂O₃). This was done to observe how methanol would interact with a catalyst that contained an abundance of both paramagnetic centers (Li⁺O⁻) and active sites (Brønsted acidic sites). The results demonstrated that the catalyst was able to readily convert methanol feed into both ethene and aromatics, with regards to tests in the range of 300 to 500 °C. According to the authors, these findings can provide evidence to support the notion that the ‘free radical mechanism’ and ‘carbocationic mechanism’ may be working in tandem in the generation of the initial C–C bond.

3.1.5 Alkoxy chain growth route. Considering all previously discussed widespread mechanistic proposals presented during early research, the alkoxy chain growth route still remains the most simplified approach for the formation of the initial C–C bond.^{118,119} According to Fig. 15, the mechanism commences with a methanol molecule approaching an acidic site (*i.e.*, the active site) of a zeolite catalyst. This results in a dehydration reaction in which a surface-associated methoxy group is formed in place of the active site, with water being eliminated (step A). A subsequent methanol molecule approaches the methoxy group, resulting in methylation of this species to generate a surface-associated ethoxy group instead (step B). Detachment of this ethyl group would ultimately form ethylene and regenerate the acid site in the process (step C). It should be noted that, despite the apparent simplicity associated with this approach, both research groups^{118,119} declared that the growth of the alkoxy chain had not been previously observed or proven experimentally, which generates doubt as to its existence, relative to the previously discussed early stage proposals (Sections 3.1.1 to 3.1.4). To the best of our knowledge, there exists only limited research regarding experimental works (in the form of spectroscopic evidence) in support of the formation of surface alkoxide groups.¹²⁰

3.2 Current studies: the ‘hydrocarbon pool’ concept

As mechanistic research in the methanol conversion to olefins process developed, much research during the 1990s was allocated towards the understanding of a more novel parallel-type reaction concept *i.e.*, the notion of the hydrocarbon pool (HCP). The initial formal suggestion of this concept was

pronounced by Dahl and Kolboe.^{121,122} Through the use of isotope labelling experiments, the authors were able to provide experimental evidence to back up the carbon pool idea they proposed. The experiments involved cofeeding ¹³C-methanol and ¹²C ethanol and propanol, simulating ethylene and propylene adsorbed on the surface of the SAPO-34 catalyst. In the case of the methanol and ethanol co-fed reaction, in view of the product isotopic distribution, propene and butene species were generated primarily from ¹³C, coming from methanol, rather than a combination of ¹³C and ¹²C. This can demonstrate that the produced hydrocarbons were not generated *via* successive methylations of ethylene, but instead *via* a sort of ambiguous ‘carbon pool’ mechanism. As research to uncover the details expanded into the later 2000s, the concept began to receive much recognition as the major contributor for both olefin formation as well as coke production for advanced times on stream.^{30,123–129} With analysis of the species confined in catalyst pores during the reaction, substantial amounts of polyalkylaromatics were detected. This finding spawned the idea that a pool of intermediate carbonaceous species exists, in which polyalkylaromatics (generally polymethylbenzenes) are the active centers. In this pool, a variety of elementary reaction steps, including chain extension, cyclization, and aromatization take place on linear chain aliphatic hydrocarbons (detected during early stages of MTO),¹³⁰ in turn leading to the generation of aromatic species. As these aromatic species begin to alter and evolve, ethylene and propylene have been suggested to ‘split off’ as products. This in turn offers a path for light olefin generation, as opposed to the previously discussed ordinary formulation of these species (see Section 3.1). The HCP concept possesses two underlying mechanisms which are said to be generally responsible for light olefin formation. These are termed the exocyclic methylation route (*i.e.*, the side chain mechanism) and the pairing mechanism (also commonly referred to as the expansion–contraction mechanism),^{124–126} which will be discussed, along with other noteworthy HCP means of light olefin formation, in the following section.

3.2.1 The aromatic cycle: the formation of polyalkylaromatics. As previously discussed, polyalkylaromatics are generally referred to as the core/active intermediates of the MTO hydrocarbon pool. Studies^{127,131} have sought to experimentally validate the existence of these species over time on stream by investigating the organic deposits entrapped inside the pores and cages of SAPO-34 during both the reaction and deactivation periods. Specifically, the authors utilized a gas chromatograph equipped with a mass spectrometer detector (*i.e.*, GC-MS analysis) and *in situ* UV-visible spectroscopy, as well as ¹H magic-angle-spinning (MAS) and ¹³C cross-polarization MAS Nuclear



Magnetic Resonance (NMR) spectroscopy. Analysis of the results proved that polymethylbenzenes (polyMBs) with 4–6 methyl groups, alongside polymethylnaphthalenes (polyMNs) with 2–4 methyl groups, are the key active components of the reaction period over SAPO-34 (as illustrated in Fig. 16(A)). Although such species are typically related to their capabilities in coke generation, groups^{132,133} have used DFT modelling calculations to portray the potential of these species in assisting in the formation of light olefins within zeolite cages. In fact, one of the groups¹³² provided a strategy to convert coke species to aforementioned naphthalenic species through steam cracking (*i.e.*, passing hydrogen and steam over SAPO-34 catalyst) and uncovered a subsequent restoration of catalytic activity through boosted total light olefins selectivity.

In order to investigate the role that polyalkylbenzene (the first suggested active component) may play towards light olefin formation, it is crucial to first uncover how it could be initially generated. Wang *et al.*¹³³ took on this task by using density functional theory (DFT) calculations to propose an energetically feasible reaction pathway from early-stage-constructed propylene into hexamethylbenzene (see Fig. 16(B)–(D)). The group suggested that a singular propene molecule may undergo protonation (P1(a)) using an acid site of the zeolite catalyst in close proximity, in turn generating 1-propyl carbenium ion.

This intermediate can subsequently oligomerize (O1(a)) by reacting with another propene molecule, to form 2-hexyl carbenium ion. Deprotonation (D1(a)) of this ionic species can result in the production of 1-hexene. This may then be followed by a hydride transfer step (HT1(a)) between a zeolite surface-associated alkoxy (*i.e.*, a methoxy) as the hydride acceptor, and 1-hexene as the hydride donor. This leads to the generation of a 1-hexene carbenium ion and an alkane (*i.e.*, methane in this case). Subsequent cyclization (C1(a)) and deprotonation of the carbenium ion results in the formation of cyclohexene. A repetition of deprotonation and hydride transfer reactions (HT2(a) and HT3(a)) results in the production of benzene. Recurring methylation steps (M1–6(a)) ultimately leads to the formulation of hexamethylbenzene, which falls under the criteria as the active component of the HCP concept. The task of uncovering the route for polyalkylbenzene formation was also undertaken by Dai *et al.*,¹²⁷ with the employment of hexene as the starting species for the mechanistic route (stating that their generation came about *via* the methylation of early-stage light olefins). As part of the proposed reaction pathway, initially, hexene species are protonated to form C_6^+ ions alongside C_6^- alkoxy. The skeletal isomerization of these carbenium ions could also be undertaken *via* alkylcyclopropane intermediates. However, the observed alkylcyclopropanes were found to be

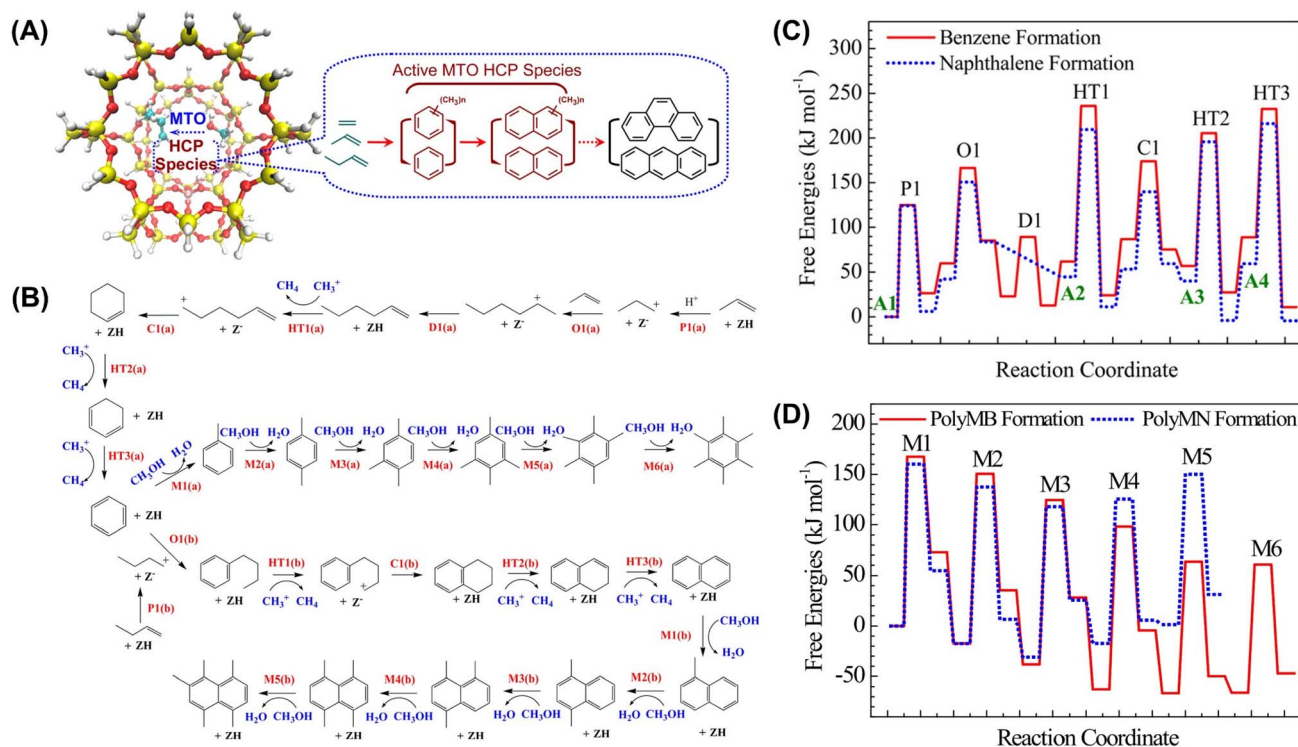


Fig. 16 (A) The development of species present in the cages of the zeolite framework during the MTO reaction (atom coloring: yellow (Si), red (O), white (H), and pink (Al)). (B) Proposed mechanistic path for the formation of polymethylbenzene (polyMB) and polymethylnaphthalene (polyMN) during the MTO reaction over zeolite catalyst (ZH). (C) Free energy profiles for the generation of benzene and naphthalene (at 400 °C) during MTO over zeolite catalyst. In addition to the zeolite framework, adsorbed propene as well as benzene and butene (A1) are taken as the reference states for benzene and naphthalene production, respectively. States A2, A3, and A4 represent the methoxyl formation at the active site prior to hydride transfer reactions. (D) Free energy profiles for the generation of polyMB and polyMN (at 400 °C) during MTO over zeolite catalyst. In addition to the zeolite framework, adsorbed benzene and naphthalene are taken as the reference states for polyMB and polyMN, respectively. Reproduced from ref. 133 with permission from American Chemical Society, Copyright 2016.



slightly less stable than hexenes, which could discourage the occurrence of such a route. The reaction pathway continues *via* a hydride transfer step, in which propene was selected to be the hydride-acceptor, leading to the formation of alkadienes from 3-methyl-1-pentene. The chain is then further propagated *via* subsequent methylation of the generated alkadienes. A repetition involving deprotonation and hydride transfer steps ultimately leads to the generation of polymethylbenzenes.

Referring to Fig. 16(B)–(D), the formation of polymethylnaphthalenes (*i.e.*, the other suggested active component of the hydrocarbon pool) has also been well documented and proven to be energetically feasible by the employment of DFT modelling. Taking benzene as the commencement point, this species has been suggested to initially react (step O1(b)) with a 1-butyl carbenium ion (which would have been previously generated *via* the protonation (P1(b)) of early-stage generated butene species). This would result in the formation of butylbenzene. Subsequently, a hydride transfer step (HT1(b)) may occur between a surface-associated methoxy group and butylbenzene, resulting in the production of a butylbenzene carbenium ion and a methane molecule. A 1,6-cyclization and deprotonation step would lead to the generation of phenylcyclohexane (step C1(b)). A repetition of the deprotonation and hydride transfer reactions (HT2(b) and HT3(b)), followed by methylation steps (M1-5(b)), would ultimately result in the production of polymethylnaphthalene.

3.2.2 The aromatic cycle: mechanisms for light olefin formation. The ‘exocyclic methylation route’, also commonly referred to as the ‘side chain mechanism’, is one out of the 2

central mechanisms under the HCP concept. To the best of our knowledge, this pathway was first proposed by Mole *et al.*¹³⁴ during the early 1980s and has been subsequently modified.¹¹⁹ Over recent years, the side chain mechanism has received recognition as a means of explaining how light olefins may form within the MTO reaction.^{124–126,135,136} Fig. 17(A) provides a close examination of this mechanistic route over zeolite catalyst utilizing either polymethylbenzene (top row) or polymethylnaphthalene (bottom row), *i.e.*, the previously mentioned active components of the hydrocarbon pool concept.¹³³ Nonetheless, the mechanism proceeds in the same manner for both scenarios. Studying the case of the PMB cycle (top row), commencing with hexamethylbenzene, this species first undergoes gem methylation with methanol (M1) to form 1,1,2,3,4,5,6-heptamethylbenzenium cation (heptaMB⁺). Its subsequent deprotonation (D1) generates hexamethylmethylene-cyclohexadiene (HMMC). The exocyclic double bond of HMMC reacts with an electrophilic methenium species (derived from a methanol molecule) to form an ethyl group on the benzenic ring (*i.e.*, an extension of the benzenic species side chain) (M2). As suggested by a different group,¹²⁴ dealkylation of this ethyl group may take place, resulting in the formation of ethylene. However, there is the alternative possibility that deprotonation occurs instead (D2), which would regenerate the exocyclic double bond, forming hexamethyl-ethylene-cyclohexadiene. Further reaction with a methenium species (M3) leads to the formation of an isopropyl group attached to the benzenium cation. Dealkylation of this propyl group takes place *via* an internal hydrogen shift (E1), resulting in the

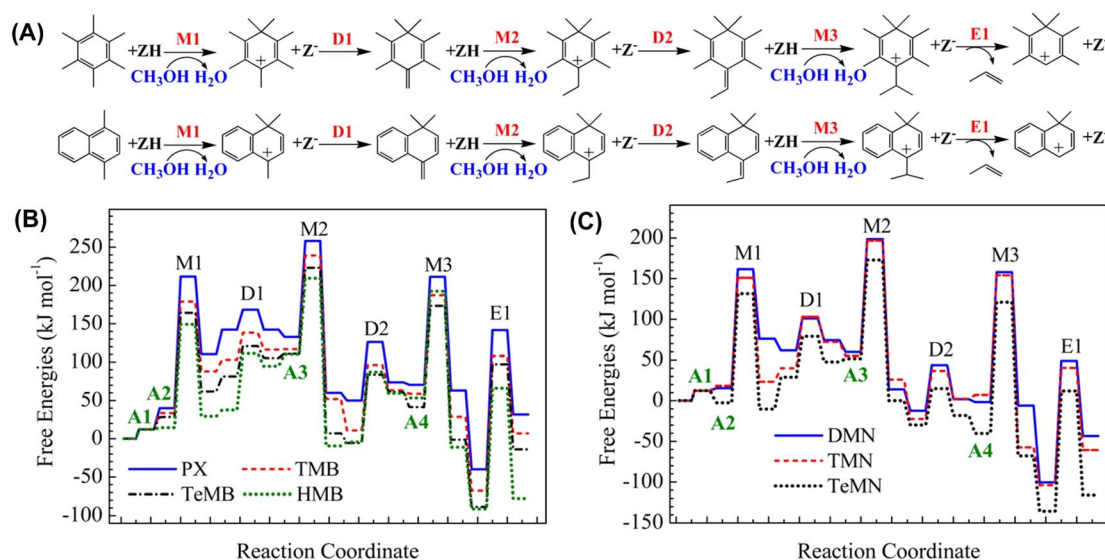


Fig. 17 (A) Side chain mechanism *via* the top: polyMB cycle (commencing with hexamethylbenzene) and bottom: polyMN cycle (commencing with dimethylnaphthalene) for the MTO reaction over zeolite catalyst. (B) Free energy profiles of the polyMB based side chain mechanism (at 400 °C) for different commencing polyMBs in the MTO reaction over zeolite catalyst. The commencing polyMBs studied are *p*-Xylene (PX), 1,2,4-trimethylbenzene (TMB), 1,2,4,6-tetramethylbenzene (TeMB), and hexamethylbenzene (HMB). The zeolite framework, methanol and different polyMBs in gaseous phase are taken as the reference state. A1, A3, and A4 are the adsorption of methanol, while A2 is the additional adsorption of polyMB. (C) Free energy profiles of the polyMN based side chain mechanism (at 400 °C) for different commencing polyMNs in the MTO reaction over zeolite catalyst. The commencing polyMNs studied are 1,4-dimethylnaphthalene (DMN), 1,4,5-trimethylnaphthalene (TMN) and 1,4,5,8-tetramethylnaphthalene (TeMN). The zeolite framework, methanol and different polyMNs in gaseous phase are taken as the reference state. A1, A3, and A4 are the adsorption of methanol, while A2 is the additional adsorption of polyMN. Reproduced from ref. 133 with permission from American Chemical Society, Copyright 2016.



formation of propylene. It should be noted that butene formation is highly unlikely to occur with respect to this mechanism due to steric hindrance.¹²⁴ Overall, groups^{126,133} have backed up this mechanistic proposal as a plausible route for light olefin formation *via* the use of theoretical modelling DFT calculations (as seen in Fig. 17(B) and (C)).

The second major mechanistic route that falls under the HCP concept is commonly termed the expansion–contraction mechanism, or the paring mechanism. The introduction of this pathway came about in 1961 when Sullivan *et al.*¹³⁷ sought to explain the formation of light olefins from polyalkyl benzenium ions. Since then, research fixating on this mechanistic proposal took flight, with many research groups agreeing on the viability of this path.^{124–126,136,138} The generalized paring route can be observed in Fig. 18(A). The mechanism suggests alkyl side chain growth by ring contraction and expansion (as opposed to the alkylation steps involving methenium species of the side chain mechanism). Specifically, the six-membered ring (heptamethyl benzenium ion) is contracted to a five-membered ring (step A), followed by subsequent dealkylation of the isopropyl and isobutyl groups (which leads to the formation of propylene and isobutene, respectively). This can be seen in steps B', B, and C. The five-membered ring undergoes deprotonation and carbon atom interchange to in turn expand to a six-membered ring (steps D and F). Methylation involving methanol leads to the ultimate formation of hexamethylbenzene (hexaMB) (steps E and G). This process is then repeated as hexaMB is converted back to the starting species (heptamethyl benzenium ion) *via* gem methylation with methanol (step H). Isotope labelling experiments conducted by Bjørgen *et al.*¹³⁹ promoted the idea that the paring mechanism may be the dominant route in olefin formation over zeolite catalysts. Precisely, the group reacted ¹³C-methanol and ¹²C-benzene over H-beta zeolite catalyst. In turn, methylbenzene species (comprised of a ¹²C-benzene ring) with ¹³C methyl groups were generated. The gas-phase product isotopic distribution provided evidence in support of the existence of the paring mechanism. A great portion of the propene product molecules

contained carbon atoms (¹²C) from the benzene ring, which is in line with the notion of the ring contraction–expansion of the paring mechanism. Furthermore, as seen in Fig. 18(B), one study¹³⁶ utilized DFT-based modelling to analyze the most crucial reaction steps of the paring mechanism (*i.e.*, up to ethylene formation (step B')). It was uncovered that the paring mechanism may be an energetically feasible pathway for certain zeolite catalysts (such as H-ZSM-5), but it was argued that in the case of H-SAPO-34, the side chain mechanism seems to be more favorable in comparison (due to both the less favorable kinetics and thermodynamics of the B' step).

Hence, overall, in terms of the two key mechanisms ('side chain' and 'expansion–contraction'), the articles studied have suggested that both are likely to take place, but that it is largely dependent on the zeolite's acid site strength, leading to the favoring of one over the other.^{124–126} From a theoretical perspective, as dictated by Olsbye *et al.*,¹²⁴ the side chain mechanism could be favored over the expansion–contraction mechanism by utilizing a zeolite catalyst with active sites of relatively low acidic strength.

3.2.3 Evolution of mechanistic understanding & developments. Overall, with consideration of Table 1, the following conclusions can be made as to the preferred mechanistic pathway adopted by SAPO-34 in the MTO conversion process. To commence, it is important to reiterate that the MTO reaction is a complex interplay of multiple mechanisms rather than a single, dominant pathway (see Fig. 19). These mechanisms may operate concurrently or sequentially, and their relative dominance can depend on several factors such as reaction conditions, time on stream (*i.e.*, induction period, steady-state operation, or deactivation phase), and catalyst structure or modification. During the induction period, the formation of light olefins primarily proceeds through a combination of five proposed direct mechanisms (as outlined in Table 1). Although the exact contribution of each remains debated, all five direct mechanisms are generally accepted to be responsible for initiating olefin production in the early stages of the reaction. As the

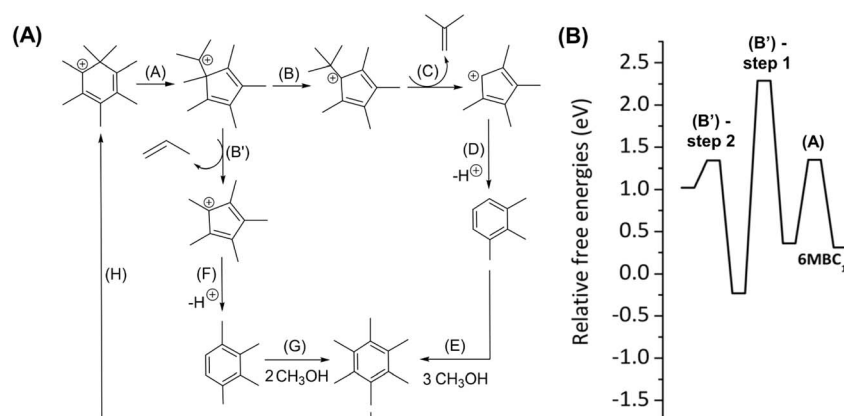


Fig. 18 (A) Paring mechanism *via* the polyMB cycle for the MTO reaction over zeolite catalyst. Adapted from ref. 124 and 125 with permission from Elsevier, Copyright 2005 and John Wiley and Sons, Copyright 2009 respectively. (B) Free energy profile of the most crucial steps of the polyMB based paring mechanism (at 452 °C), initiated from the most favorable gem-methylated carbenium intermediate (6MBC₁⁺) over H-SAPO-34. The energy reference is the free energy of the adsorbed gem-methylated carbenium intermediate (ZO⁻-6MBC₁⁺) alongside three gas-phase methanol molecules. Adapted from ref. 136 with permission from American Chemical Society, Copyright 2021.



reaction progresses into the steady-state phase (bulk reaction period), olefins formed initially begin to undergo secondary reactions-growth, cyclization, and aromatization-leading to the formation of key intermediates, including polyalkylaromatics and larger olefinic species. This marks the onset of the hydrocarbon pool (HCP) mechanism, which operates *via* a dual-cycle model: one cycle based on aromatic intermediates and another on olefinic species. The dominance of each cycle is strongly influenced by operating parameters such as temperature, pressure, and feed composition, and will be discussed in more detail. This, in turn, leads to the production of later-stage light olefins and exists as the main mechanistic concept encompassing the MTO process through to deactivation. Ultimately, the process concludes with catalyst deactivation, primarily due to the transformation of reactive intermediates-especially methylated aromatic species (*e.g.*, polymethylbenzenes)- into larger polycyclic aromatic hydrocarbons. These species accumulate and block the SAPO-34 micropores, hindering reactant diffusion and access to active sites, thereby terminating catalytic activity.

In recent years, researchers have focused their efforts on understanding the factors that influence the mechanistic pathways involved in methanol conversion over zeolite catalysts, particularly with regard to schemes that fall under the HCP concept. The dominant route, in terms of the likelihood of formation of specific HCP intermediates (*e.g.*, polymethylbenzenes (PMBs) *vs.* polymethylnaphthalenes (PMNs)), and subsequent production of light olefins utilizing these intermediates, seems to depend highly on the zeolite catalyst acidity, topology, and reaction conditions in place.

Commencing with acidity, researchers¹⁵¹ have placed great emphasis on the tuning of hydrocarbon pool intermediates by alteration of this parameter. Using GC-MS and NMR methods, it

has been uncovered that SAPO-34 with lower Brønsted acid site density (*i.e.*, by utilizing a SiO₂/Al₂O₃ molar ratio of 0.2 in the starting gel) were found to favor the promotion of PMBs as active hydrocarbon pool species as opposed to PMNs. This was justified by the suppression of intermolecular hydrogen transfer reactions, which are needed for the growth of PMBs into polycyclic aromatics, such as PMNs and beyond. It should be noted that researchers¹⁵¹ have discerned that alterations in SAPO particle size (*i.e.*, morphological changes), unlike acidity changes, has a much smaller impact on the product distribution. Hence, this indicates that particle size as a parameter holds little to no impact on the favoring of a PMB or PMN-based HCP route for olefin formation.

Researchers¹⁵² have also investigated the effects of zeolite topology on the mechanistic route adopted for olefin formation from methanol. DFT and *ab initio* calculations conducted on the paring mechanism (a key mechanism of the aromatic cycle of the HCP concept) for different zeolite topologies, such as MFI (H-ZSM-5), AFI (H-SSZ-24) and CHA (H-SAPO-34), demonstrated a clear favoring of this mechanism over the CHA (H-SAPO-34) zeolite. Differences in pore size amongst the different topologies were ascribed to play a major part in determining the likelihood of the paring mechanism to take place. The researchers justified that zeolites with larger pore sizes (such as ZSM-5) were able to better accommodate the mobility of heptaMB+ (a key intermediate of the paring mechanism), encouraging the adsorbate to rearrange and achieve more favorable transition state structures, which possessed generally higher free energy of adsorbed states. Hence, higher overall energy barriers for the paring mechanism were observed over larger pore zeotypes like ZSM-5, making it less favourable to occur relative to small-pore ones like SAPO-34.

Lastly, researchers have also demonstrated the effect that chosen reaction conditions can impose on the mechanistic route adopted for methanol conversion over SAPO-34 catalyst. As discussed in Chapter 2 (the MTO Process), it is well established, through experimental validation, that alterations in process operating parameters (such as temperature) play a large role in determining the product distribution.¹⁵³⁻¹⁵⁵ However, some researchers have taken the next step of investigating how this effect occurs. This has been achieved by both analyzing the species developed during the reaction period (upon changes in reaction conditions) *via in situ* and/or *operando* experimental techniques, as well as by applying modelling/computational-based methods. Commencing with one of the most important operating parameters, researchers have¹⁵⁶ demonstrated (*via* the use of *operando* UV-vis spectroscopy and methanol pulse experiments) that alterations in the process temperature can highly impact the nature of active and deactivating species formed during MTO over SAPO-34, thereby shifting the mechanistic pathway adopted to form olefins. Researchers¹⁵⁶ have observed that for the lower MTO process temperature range of 300 °C and 325 °C, the active HCP species are primarily highly methylated benzene carbocations (and low amounts of methylated naphthalene carbocations species), which have been found to favour the selectivity towards propylene formation. At this low temperature range, these methylated naphthalene

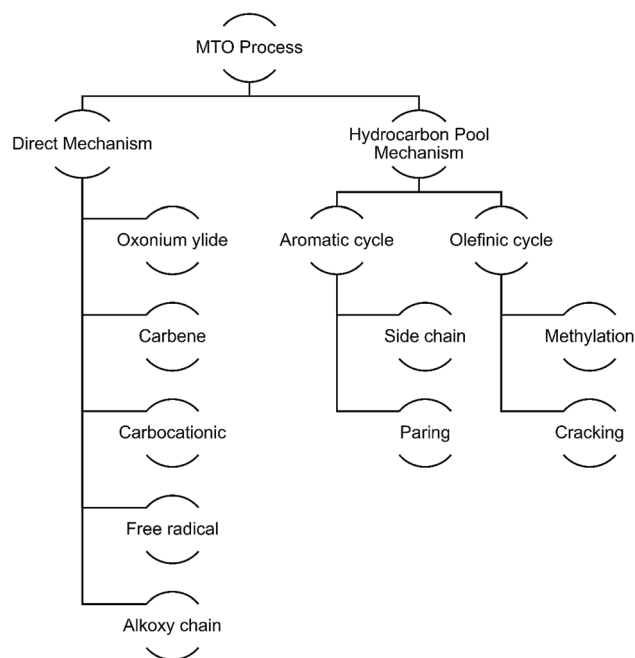


Fig. 19 MTO process classification chart of mechanistic concepts.



Table 1 Supporting and opposing evidence for major proposed MTO mechanistic pathways

Mechanistic pathway	Supporting evidence	Opposing evidence
Direct mechanism 1: oxonium ylide mechanism	<p>In the study conducted by Sayed,¹⁴⁰ IR spectroscopy (1000–4000 cm⁻¹) of CH₃OH over HZSM-5 was used to support an oxonium-based mechanism. This involved higher ethers (methylethyl, methylisopropyl, methyl-<i>t</i>-butyl) that decompose to ethene, propene, and isobutene. Two observations were observed by the author that provide the key evidence: (1) after DME consumption, methanol bands reappear (O–H stretch at 3675 cm⁻¹, C–O stretch at 1033 cm⁻¹) despite a DME feed, suggesting reversible oxonium intermediates; and (2) a loss of DME bands (1102 cm⁻¹ C–O stretch, 1178 cm⁻¹ CH₃ rock), which coincides with the growth of a C=C band at 1650 cm⁻¹, consistent with DME-to-alkene conversion</p> <p>In the study presented by Olah <i>et al.</i>¹⁴¹, GC and GC-MS analysis were carried out to investigate the mechanism of the conversion of methanol/dimethyl ether, with particular emphasis on how the initial C–C bond forms. Labeling experiments strongly indicated the formation of initial trimethyloxonium ions, followed by the production of oxonium ylides (<i>i.e.</i>, methylenedimethyloxonium ylide) <i>via</i> proton abstraction. The authors claimed that, ultimately, the cleavage of oxonium ion intermediates results in the formation of ethylene and dimethyl ether</p>	<p>The study by Sayed¹⁴⁰ was conducted for MTG/HZSM-5 conversion. Considering the key differences between ZSM-5 and SAPO-34 (ZSM-5 being more acidic in nature, holding a larger average pore size, and possessing MFI topology relative to CHA topology), it could be argued that the oxonium ylide mechanism, while applicable for ZSM-5, may not be as dominant in MTO over SAPO</p> <p>The focus of the study by Olah <i>et al.</i>¹⁴¹ is on bifunctional acid-base catalyst systems, which possess no shape selectivity of zeolites. Hence, the mechanistic details presented by this study on the oxonium ylide may also not be fully applicable to MTO over SAPO, due to this clear discrepancy in catalyst system utilized</p>
Direct mechanism 2: carbene mechanism	<p>Chang <i>et al.</i>¹⁴² provided evidence obtained from ¹³C isotope-labeling experiments showing the existence of carbenoid C₁ species during methanol reactions over HZSM-5 zeolites. It was ultimately concluded that these species are partly involved in the initial C–C bond formation process in the methanol conversion process. The study presented by Minova <i>et al.</i>¹²⁰ proposes a carbene coupling reaction pathway for olefin formation from surface methoxy groups. The group claimed that insertion of surface carbene-like species into adjacent surface methoxy groups is a key step in the induction period of MTO over SAPO-34. This was supported by spectroscopic evidence utilizing synchrotron FTIR microspectroscopy coupled with mass spectrometric analysis</p>	<p>In the study by Chang <i>et al.</i>¹⁴², it was claimed that the reactions of olefins over zeolites can be explained partly by classical carbenium ion mechanisms. The group claimed this led to alkanes and aromatics and in turn, catalyst deactivation. However, the precise mechanism of initial C–C bond formation from methanol, involving carbene species, was not presented, which makes the claimed involvement of these species still subject to speculation. Another point of consideration is the fact that this paper was conducted over a more acidic, larger pore size zeolite (ZSM-5) as opposed to SAPO-34</p>
Direct mechanism 3: carbocationic mechanism	<p>In the study presented by Nagy <i>et al.</i>,¹⁴³ ¹³C-NMR and gas chromatography analysis presented data suggesting that a carbenium ion mechanism was highly probable in methanol conversion reactions. The group claimed that the involvement of carbenium ions led to the presence of alkanes and ethylene under the reaction conditions reported. The report conducted by Minova <i>et al.</i>¹²⁰ presents experimental evidence in the form of spectroscopic studies, stating that methoxy cations also play an important role (alongside previously discussed carbene-like species) as intermediates during the MTO process over SAPO-34. A computational study by Lesthaeghe <i>et al.</i>¹⁴⁴ also claims the involvement of carbenium ions in methanol to olefins conversion. The group reports that the zeolite framework is able to assist in stabilizing carbenium ion species, thereby reducing the energy barriers associated with mechanistic steps involving their formation and presence. Furthermore, the group</p>	<p>The study by Nagy <i>et al.</i>¹⁴³ focuses on the conversion of methanol on H-ZSM-5 in the presence of carbon monoxide. Hence, it could be argued that the discrepancy in the feed (<i>i.e.</i>, not pure methanol fed) and the choice of catalyst system (ZSM-5 as opposed to SAPO-34) makes the evidence less reliable when applied to the traditionally conducted MTO process over SAPO-34. Although the study by Lesthaeghe <i>et al.</i>¹⁴⁴ provided valuable insight into the role of carbenium ions in methanol to olefins conversion, the group does not provide experimental findings to back up the computationally-directed claims presented in the paper</p>



Table 1 (Contd.)

Mechanistic pathway	Supporting evidence	Opposing evidence
Direct mechanism 4: free radical mechanism	<p>reported that the CHA topology of SAPO-34 and SSZ-13 was best able to stabilize cationic species as opposed to other studied zeolite topologies (<i>i.e.</i>, MFI)</p> <p>Clarke <i>et al.</i>¹⁴⁵ studied the mechanistic pathway of dimethyl ether over H-ZSM-5 zeolite, and in turn detected the presence of free radicals by using a spin trapping reagent with e.s.r. analysis. The group thereby provided evidence that free radicals are involved in the initial C–C bond forming step and exist as important intermediates for the ultimate production of both alkenes and aromatics</p>	<p>The study by Clarke <i>et al.</i>¹⁴⁵ remains valuable for its mechanistic insights into the involvement of free radical species in hydrocarbon conversion. However, the focus of the study is on the reaction of dimethyl ether over H-ZSM-5 zeolite, with the goal to produce alkene and aromatic hydrocarbons. Hence, as with other previously mentioned studies, the applicability of these mechanistic details to MTO over SAPO-34 could be questioned</p> <p>To the best of our knowledge, there exists limited research papers¹²⁰ containing experimental evidence in support of this direct route. The majority of MTO mechanistic studies mention the possible existence of these species as intermediates during the induction period, but relative to the HCP concept, the alkoxy chain growth route is of much lower significance</p>
Direct mechanism 5: alkoxy chain growth route	<p>Researchers¹²⁰ have validated the presence of surface alkoxy species during the methanol conversion process, through the use of spectroscopic evidence. The group claims that such alkoxy species (primarily methoxy species) are the core intermediates during the induction period of the MTO process, and exist amongst the pool of other pre-mentioned hydrocarbon species (<i>e.g.</i>, carbocationic species). Computational research by Ke <i>et al.</i>¹⁴⁶ has also provided partial evidence for the likely existence of alkoxy species during the MTO process. The group utilized microkinetic simulations in a three-site model to claim the existence of these species as intermediates in the reaction</p>	<p>In the study by Mole <i>et al.</i>,¹⁴⁷ the authors state that the experimental evidence collected implies alternative or additional pathways within the aromatic cycle (such as ring expansion/contractions in addition to the side chain mechanism). Specifically, it was stated that the group's interpretations should be regarded as a working hypothesis rather than a rigorously established mechanism. Hence, the complexity of the HCP mechanistic scheme can make it difficult to establish an exact and dominant path</p>
HCP mechanism 1: aromatic cycle	<p>A study by Mole <i>et al.</i>¹⁴⁷ provided evidence (in the form of ¹³C and ²H isotope labelling experiments) for the existence of the aromatic cycle of the hydrocarbon pool concept. The group targeted their efforts on validating the side chain mechanism of the aromatic cycle for light olefin formation. It was ultimately revealed <i>via</i> these experiments that the side chain mechanistic scheme is viable. The mechanism implies that the carbon of the methyl group of a methyl-aromatic compound should be eventually incorporated into ethylene. This was validated by the labelling experiments where conversion of methanol, in the presence of toluene (with ¹³C situated in the toluene methyl group), led to the capture of ¹³C labelled ethylene</p> <p>A study by Lesthaeghe <i>et al.</i>¹⁴⁴ also provided valuable insight on the aromatic cycle of the HCP concept, focusing on methanol to olefins conversion over ZSM-5 zeolite. The group's theoretical computations verified that the side chain mechanism on methylbenzenes over zeolite catalysts is a plausible reaction scheme in the formation of ethene from methanol. On the other hand, the paring mechanism of the aromatic cycle (focusing on ring expansion/contraction) has been shown to favour the production of propene or isobutene. Overall, the group claims that the aromatic cycle is a dominant mechanistic concept of the MTO process</p> <p>A study by Li <i>et al.</i>,¹⁴⁸ involving ¹³C-labeling pulse experiments, indicated that the aromatic based hydrocarbon pool mechanism was the most dominant reaction route of methanol conversion</p>	<p>In the study by Lesthaeghe <i>et al.</i>¹⁴⁴ it is important to note that the group faced some bottlenecks in uncovering the most energetically favourable route for ethene formation <i>via</i> the side chain mechanism. It was found that the main issue (energetically speaking) existed in the elimination steps of ethene. Hence, determining the exact route with lowest energy barriers for olefin formation is not always a straightforward process, especially given the pool of simultaneous reactions taking place in the aromatic cycle. Moreover, it should be noted that this study was conducted over H-ZSM-5 and not SAPO-34; hence, as with previously mentioned studies, the applicability of certain mechanistic details may be questioned given the discrepancies in acidity and pore size between the two zeotypes</p> <p>The study by Li <i>et al.</i>¹⁴⁸ demonstrates that zeolite type plays a large role in determining the route adopted within the hydrocarbon pool mechanism. Due to the space confinement discrepancies that</p>



Table 1 (Contd.)

Mechanistic pathway	Supporting evidence	Opposing evidence
HCP mechanism 2: alkene cycle	<p>over SAPO-34. The group studied a variety of zeolites, including SAPO-34, H-ZSM-5 and H-ZSM-22, concluding that SAPO-34 in particular favours the aromatic cycle, with the zeolite holding enough spaces in its framework to accommodate large aromatic intermediates</p> <p>A study by Gong <i>et al.</i>¹⁴⁹ presents the existence of the alkene (olefinic) cycle of the hydrocarbon pool concept. Spectroscopic evidence conducted by the group elucidates the fact that both the aromatic and olefinic cycle exist in the hydrocarbon pool concept as a dual cycle mechanism that simultaneously occurs. It was concluded that the extent to which each of the two cycles dominate depends on the treatment of the zeolite. A study by Ke <i>et al.</i>¹⁴⁶ performed simulations in a three-site model to study the MTO conversion in industrially relevant H-SAPO-34 zeolite under a wide range of operating conditions. <i>Via</i> a combination of DFT calculations and microkinetic simulations, the group confirmed the presence of a dual cycle concept (involving both the olefinic cycle and aromatic cycle) within the global hydrocarbon pool framework in the MTO process. The group concluded that the reaction conditions implemented affected the extent to which each cycle dominates. Specifically, it was uncovered that the aromatic-based cycle dominated in MTO conversion as operating temperature increased, but as pressure and water content (in the feed) decreased. An article by Yu <i>et al.</i>¹⁵⁰ provided experimental validation of the existence of the dual cycle concept over SAPO-34 in the MTO process. <i>Via</i> the employment of a ¹²C/¹³C-methanol isotope switching experiment, the group were able to acknowledge the existence of both an aromatic-based cycle and alkene-based cycle taking place during the bulk of the MTO reaction process. The detailed reaction route was highly dependent on types of organic species most dominant in the catalyst framework at different times on stream. It was uncovered by the group that during the efficient reaction period, the aromatics-based cycle existed as the dominant of the two. However, during both the induction and deactivation periods, the olefinic-based cycle contribution grew larger</p>	<p>exist between zeolites of different structural topologies, this alters the type of hydrocarbon pool species that are most dominant. Overall, this once again highlights the complexity involved in determining the exact aromatic-based path adopted for the MTO process</p> <p>The study conducted by Gong <i>et al.</i>¹⁴⁹ focuses on the effect of steamed ZSM-5 zeolite on the reaction mechanism for the methanol-to-hydrocarbons process. Hence, this discrepancy in catalyst system and treatment approach implies that the mechanistic details may not be directly applicable. The study by Ke <i>et al.</i>¹⁴⁶ provides valuable insight into the mechanistic details of the hydrocarbon pool framework- specifically capturing the dual cycle concept and the relative dominance of each cycle under different operating conditions. However, it should be noted that this study is still highly theoretical/computational. Hence, there still exists a greater need for more complementary studies bringing experimental validation of the recently proposed dual cycle notion</p>

species were found to act as deactivating species, since catalyst deactivation was caused by the gradual filling and eventual blocking of micropores by these species. At a greater temperature condition of 350 °C, the active species consist of a mixture of both highly methylated benzene carbocations and methylated naphthalene carbocations. A higher MTO process temperature range of 400 °C and 500 °C has been found to preferentially promote methylated naphthalene carbocations as the primary active HCP species (and low amounts of highly methylated benzene carbocations), which has been found to favour the formation of ethylene. At this high temperature

range, the methylated naphthalene carbocation species only act as active species. However, these species do eventually give rise to the production of phenanthrene/anthracene (polycyclic) aromatic deposits (both neutral and carbocationic species), which are then mainly responsible for the deactivation by pore blockage at the higher temperature range.

Besides temperature, researchers¹⁵⁷ have also investigated how other operating conditions shape the mechanistic pathways of the MTO process over SAPO-34. Microkinetic simulations, involving more than 300 elementary steps, have revealed that pressure (*p*) and water-to-methanol feed ratio (*x*) also play



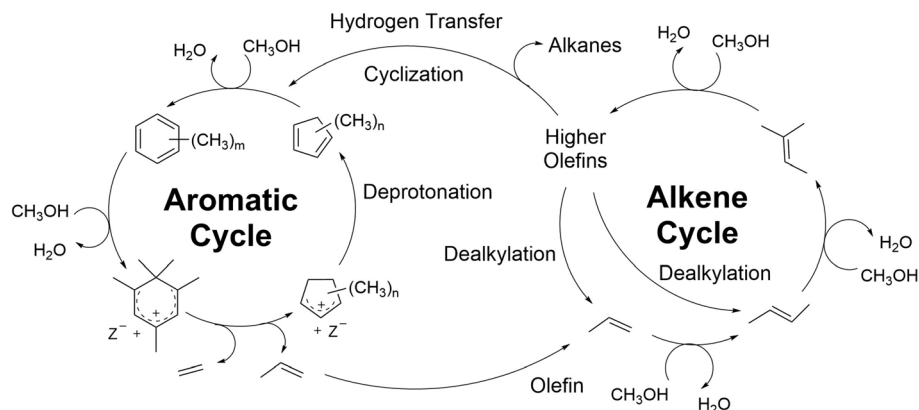


Fig. 20 Simplified mechanistic scheme of the dual cycle concept of the hydrocarbon pool mechanism. Reproduced from ref. 158 from Académie des Sciences, Copyright 2018.

significant roles in determining the route taken by the hydrocarbon pool (HCP). Before addressing these effects, it is important to note that recent studies on the HCP concept have begun to support the existence of a dual-cycle mechanism taking place (see Fig. 20). This involves the well-documented aromatic-based cycle (driven by polymethylbenzenes (PMBs) and polymethylnaphthalenes (PMNs) as key active intermediates, producing light olefins *via* side-chain and paring mechanisms) as well as the existence of an olefinic-based cycle (in which heavier olefins act as reactive intermediates and form lighter olefins through successive methylation and cracking steps). This represents a rather recent development to the original claims of a stand-alone, single aromatic-based cycle within the HCP concept. Hence, considering this, researchers are now beginning to determine how operating conditions affect both (i) the extent to which these two major cycles of the HCP occur, as well as (ii) the specific trends observed within each cycle. Commencing with the former, it can be seen that the overall rate of methanol conversion is dominated by the aromatic-based cycle at high temperatures, low pressures, and low water-to-methanol feed ratios. In contrast, the inverse trends cause the olefinic-based cycle to become dominant, often proceeding 1–5 orders of magnitude faster than the aromatic-based cycle. Regarding the latter, it can be seen that distinct selectivity trends emerge within each cycle.¹⁵⁷ For the aromatic-based cycle, the paring mechanism is favoured at high temperatures, low pressures, and low water content, leading to higher propylene selectivity relative to ethylene, whereas the side-chain mechanism is promoted under the opposite conditions, increasing ethylene formation. On the other hand, in the olefinic-based cycle, low temperatures and high pressures favor propylene formation (*i.e.*, the opposite trend to the aromatic-based paring mechanism). However, high water content favors ethylene formation, mirroring the aromatic side-chain mechanism. Overall, considering that the reaction conditions in place both affect the relative dominance of each cycle and the cycle-specific selectivity trends, the following two conclusions can be made. Firstly, propylene formation is more consistently associated with the dominant cycle under its optimal

conditions. Under higher T , lower p , and lower x , the dominant aromatic-based cycle favors propylene formation *via* the paring mechanism. Inverse reaction conditions dominates the olefinic cycle, which then tends to also favor propylene formation under those conditions. Secondly, with regards to ethylene formation, it seems that although ethylene can be favored within either cycle under certain conditions, those conditions in place do not usually coincide with those in which the ethylene-producing cycle dominates overall conversion. This demonstrates the overall dominance of propylene formation, relative to ethylene formation, under almost all operating conditions set.

Although it has been appropriate to uncover how the MTO mechanism (and henceforth, product distribution) alters as a function of the set operating conditions, it is of even greater value to understand how the mechanism takes place under specific, industrially relevant conditions. As outlined in Chapter 2 (MTO Process), the commercial fluidized-bed reactors typically operate at ~ 400 °C, near-atmospheric pressure (~ 1 bar), and with a water-to-methanol feed ratio of about 1 : 1 (molar). Under these conditions, researchers¹⁵⁷ have shown that the olefinic-based cycle of the hydrocarbon pool (HCP) strongly dominates, with the simulated methanol conversion rate exceeding that of the aromatic-based cycle by more than forty-fold, closely reproducing experimental conversion values. With regards to the cycle-specific selectivity, under these conditions, microkinetic simulations demonstrate that the dominant olefinic cycle favours propylene formation relative to ethylene, with a roughly one order of magnitude higher formation rate. Altogether, these findings highlight the critical role of the olefinic-based HCP cycle in governing MTO conversion over H-SAPO-34 under industrially relevant conditions.

3.3 Deactivation pathways

The consensus amongst articles regarding coke formation is that the mechanism commences with polyalkylnaphthalene. This group of molecules are said to undergo molecular rearrangement, ring expansion and aromatization, which leads to the formation of multi-ring aromatics, as seen in Fig. 22. Ultimately, these heavy aromatics (so-called coke) block zeolite



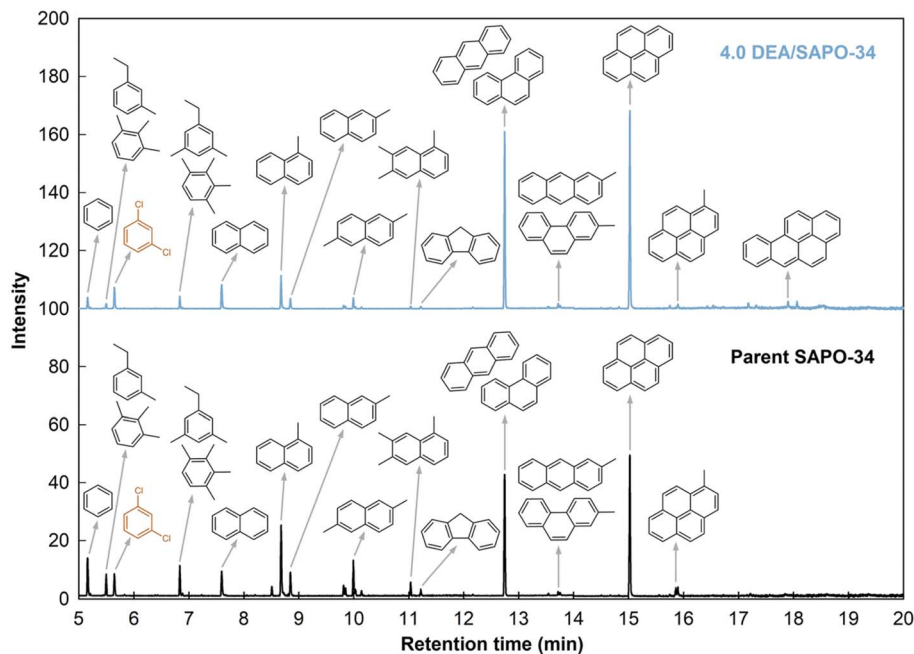


Fig. 21 GC-MS analysis of coke species formed over deactivated SAPO-34 catalyst (directed with tetraethylammonium hydroxide (TEAOH)) and deactivated SAPO-34 catalyst post-treated with 4.0 molar diethylamine (DEA) aqueous solution. Reproduced from ref. 160 with permission from Royal Society of Chemistry, Copyright 2023.

catalyst pores, preventing the diffusion of reactant molecules (methanol) towards active sites and resulting in catalyst deactivation.^{133,159} The nature of such coke species can be visually observed in Fig. 21, which demonstrates the results of a gas chromatography-mass spectrometry (GC-MS) analysis conducted over deactivated SAPO-34 catalyst, with the species beyond 11 minutes retention time indicative of the type of compounds responsible for deactivation.

Interestingly, as multi-ring aromatics begin to form, the necessary hydride transfer reactions between intermediate species and surface-associated methoxy groups leads to the formation of methane molecules (see Fig. 22). Thus, this can provide a justification as to why methane and coke production seem to rise concurrently during the MTO reaction, since the hydride transfer step (which generates methane) is necessary for the building up of aromatic rings to form coke species. Fig. 22 also illustrates the free energy profile associated with the mechanistic scheme developed for the formation of anthracene and phenanthrene (*i.e.*, types of multi-ring aromatics that constitute coke).¹³³ The conducted DFT modeling indicated that anthracene and phenanthrene production in zeolite cages is kinetically more difficult relative to the generation of benzene and naphthalene. Moreover, the free energy barriers of methylation and olefin elimination steps in the previously discussed polymethylnaphthalene (poly(MN)) based side chain mechanism are much lower in comparison to the hydride transfer reactions for anthracene and phenanthrene formation. Hence, it can be concluded that (poly)MB and (poly)MN are more likely to assist in the formation of olefins, the more favourable pathway, rather than simply build up to form polycyclic aromatics. In other words, these species take on more of a role as the active

components of the hydrocarbon pool mechanism, rather than being the precursors for inactive species.

3.4 Limitations of mechanistic studies

Although much progress has been made towards uncovering the mechanisms of the methanol conversion to olefins reaction, the individual steps amongst a majority of these mechanisms are somewhat vague, and do not normally possess the detail needed to be termed as elementary steps. Furthermore, a majority of the mechanisms presented have been generalized for different types of zeolites (such as mordenite, H-ZSM-5, and SAPO-34), which may not be accurate considering differences in acidity, topology, and pore sizes. In reality, these factors that distinguish one zeolite catalyst from another could largely affect the activation energies associated with elementary steps, and in turn influence the mechanistic route adopted. Research to explain how some zeolites may favor one path over another is quite limited and has only recently emerged.¹³⁶ With DFT calculations of van der Waals dispersive corrections, one group¹³⁶ concluded that the 'side chain mechanism' acts as the major reaction pathway over H-SAPO-34, whereas the H-ZSM-5 catalyst was shown to favor the paring route instead. Furthermore, according to the generalized HCP mechanism, poly-alkylaromatics, which are deemed the active intermediates for light olefin production, are themselves being produced from light olefins, which seems counterintuitive. To the best of our knowledge, there still exists a lack of comprehensive studies that compare the elementary steps and energy activation barriers of initial light olefin production and desorption *i.e.*, referring to early time on stream, with that of further olefin



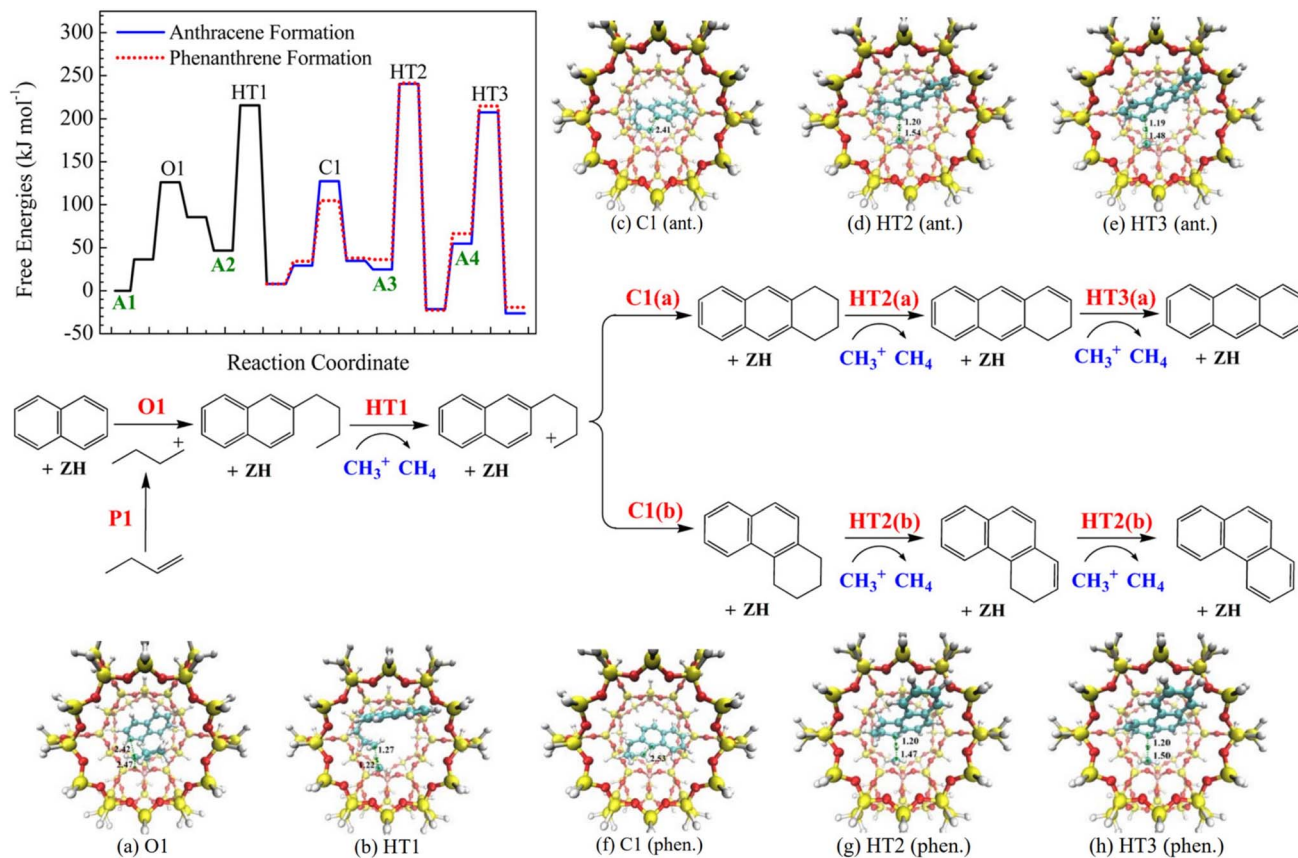


Fig. 22 Proposed mechanistic path for the formation of polycyclic aromatic species (anthracene and phenanthrene) during the MTO reaction over zeolite catalyst. The top left plot presents the free energy profile for both anthracene (ant.) and phenanthrene (phen.) production (at 400 °C) in the MTO reaction over zeolite catalyst. In addition to the zeolite framework, naphthalene and adsorbed butene (A1) are taken as the reference state for the formation of anthracene and phenanthrene. (A2, A3, and A4) represent methoxyl generation at the active site prior to the hydride transfer reactions. For each of the elementary steps, the optimized transition states have been presented schematically (from (a)–(h)), which correspond with both the free energy profile and the proposed mechanistic path. Notation: O1 refers to oligomerization, HT1–HT3 refer to hydride transfer reactions, and C1 is for the 1,6-cyclization. Atom colorings: yellow (Si), red (O), white (H), pink (Al) and blue (C). Bonding distances are presented in Å. Reproduced from ref. 133 with permission from American Chemical Society, Copyright 2016.

reconstruction to create polyalkylaromatics and late-stage light olefins. Furthermore, as has been touched upon in this section, it is clear that research groups are in agreement that aromatic species such as benzene and naphthalene play a dual role in both olefin and coke formation. However, up to the present day, mechanistic studies struggle to uncover the ways to fine-tune the mechanism in order to prevent or largely limit the pathway of coke production, while instead promoting olefin formation from these active intermediates. This remains at the forefront of future work in the field of MTO and its reaction mechanism. Lastly, research groups have begun to reveal superior ways to modify the catalyst synthesis procedure in order to achieve superior catalytic performance in the MTO reaction, involving the pre-treatment, post-treatment, and mixed templating of the catalyst (which will be discussed further in Section 4). As of now, mechanistic studies have largely focused on detailing the reaction route associated with the traditionally prepared SAPO-34 catalyst; hence, investigations into the potential mechanistic alterations associated with catalyst modification are yet to be fully revealed.

3.5 MTO reaction and deactivation kinetics

As seen by the highly varied and complex mechanistic scheme that constitutes the MTO reaction (discussed in Section 3), no kinetic model from research until present day has been able to fully encapsulate MTO's extensive reaction network. Research groups that have delved into the field of MTO kinetics (with SAPO-34 as the catalyst) have largely presented simplified lumped kinetic models that are derived from reaction schemes comprising a small number of steps. Iterative approaches were tried and tested until model predictions of MTO performance were comparable to that of the experimental data. Different approaches at developing these models have been utilized, ranging from the use of simple power laws to the application of the Langmuir–Hinshelwood (LH) approach. For a detailed insight into the specific reaction and deactivation kinetic models utilized, kinetic parameters, and rate expressions used by different research groups, consult the SI. One example can be taken from Fatourehchi *et al.*¹⁶¹ who utilized both LH and power law kinetic models to derive their rate equations. The developed simple mechanism consisted of five reaction steps (which can



Table 2 Reaction mechanism and final reaction rate expressions used to describe the MTO process over SAPO-34 catalyst, based on both Langmuir–Hinshelwood and power law kinetic models¹⁶¹

Reaction pathway ^a	Rate expressions ^b
$\text{MeOH} + Z \xrightleftharpoons[k'_m]{k_m} \text{MeOH} \cdot Z$	—
$2\text{MeOH} \cdot Z \xrightleftharpoons[k'_d]{k_d} \text{DME} \cdot Z + \text{H}_2\text{O} + Z$	—
$\text{DME} \cdot Z \xrightarrow{k_r} \text{DME} + Z$	$r_r = \frac{k_r K_d K_m^2 p_{\text{MeOH}}^2}{p_w (1 + K_m p_{\text{MeOH}} + (K_d K_m^2 p_{\text{MeOH}}^2 / p_w))}$
$\text{DME} \cdot Z \xrightarrow{k_c} \text{C}_2\text{H}_4 + \text{H}_2\text{O} + Z$	$r_c = \frac{k_c K_m^2 p_{\text{MeOH}}^2}{P_w (1 + K_m p_{\text{MeOH}} + (K_d K_m^2 p_{\text{MeOH}}^2 / p_w))}$
$\text{DME} \cdot Z + \text{MeOH} \cdot Z \xrightarrow{k_p} \text{C}_3\text{H}_6 + 2\text{H}_2\text{O} + 2Z$	$r_p = \frac{k_p K_d K_m^3 p_{\text{MeOH}}^3}{p_w (1 + K_m p_{\text{MeOH}} + (K_d K_m^2 p_{\text{MeOH}}^2 / p_w))^2}$
—	$r_{\text{Paraffin}} = k_{\text{Paraffin}} p_{\text{Paraffin}}^n$

^a Reaction rate constants (k_j) are in units of $\text{mol g}^{-1} \text{h}^{-1}$, except for k_{Paraffin} , which is in units of $\text{mol g}^{-1} \text{h}^{-1} \text{bar}^{-n}$ (where $n = 0.31$). Z in the reaction expressions refer to the zeolite active site. ^b r_i refers to the rate of species i production ($\text{mol g}^{-1} \text{h}^{-1}$). K_j refers to the equilibrium constant for a reaction j . p_w and p_{MeOH} refer, respectively, to the partial pressure (bar) of water and methanol in the feed.

be seen in Table 2), involving the adsorption of methanol on the surface of the zeolite (in place of a Brønsted acidic site), followed by the formation of a surface-associated dimethyl ether, which could subsequently desorb or instead convert to light olefins. With use of the LH formulation, it was assumed that the reactant methanol molecules were able to adsorb on the surface of the catalyst, diffuse in close proximity to the surface, and in turn react with surface associated molecules. Evidently, the generated products on the surface are assumed to desorb into the fluid phase without constraint. Other than the LH mechanism, all other possible paraffinic products were lumped together as a singular species, and the corresponding reaction rate expression was formulated as a simple power law. Reaction steps 3 to 5 were presumed to be rate-determining. The group established this *via* an iterative approach of altering the assumed rate-determining steps, and in turn comparing the predictions from a variety of suggested rate models with experimental data.

With regards to the detailed studies seen in the SI (*e.g.*, Table S5), some groups considered coke formation as part of their reaction rate expressions, as it influences the products distribution substantially.^{100,162–164} Accordingly, one group¹⁶² formulated a kinetic model and included the coke-dependent parameter by using a thermogravimetric reactor and measuring the coke content over time on stream. Both rate expressions, involving feed consumption (methanol and dimethyl ether, lumped as MDOH) and rate of production of different species i , contains the parameters a' and α_i , which, respectively, are indicative of the catalyst activity and the dependency of each species production on the coke content (see Table 3). The group tested a range of catalyst activity functions (a') and found that the exponential function provided the best fit for the experimental data, justified by its higher correlation coefficient (0.9965). This catalyst activity expression is a function of only the coke content (C), determined by the TGA. The coke content C in turn is a function of the temperature T , time t ,

weight hourly space velocity WHSV ($\text{g}_{\text{MeOH}} \text{g}_{\text{Cat}}^{-1} \text{h}^{-1}$), and methanol partial pressure p .

Overall, considering the dependence of product distribution on catalyst coke content, feed dilution, and temperature (as explained in Section 2), a universal and acceptable kinetic model should consider all the aforementioned influential parameters. In addition, the activity tests used to estimate the kinetic rate constants ought to be conducted under conditions that closely resemble industrial or realistic settings. Regrettably, some of the kinetic studies overlooked the influence of catalyst coke content on products' distribution,^{56,161,165} while others conducted the experimental tests under unrealistic conditions such as with a heavily diluted feed with water⁵⁶ (as opposed to the industrial case of 20 wt% dilution with water⁷³). A few points need to be highlighted here concerning the derivation of a kinetic model. Firstly, to simulate a differential reactor with limited conversion, a very high WHSV must be employed, but one should be aware that SAPO-34 catalyst tends to deactivate quite rapidly upon exposure to these conditions. Hence, utmost attention should be given to not collect data over partially deactivated catalyst. Intermittent (pulse) flow of feed could be a possible solution. Secondly, since the methanol conversion to light olefins reaction is not thermodynamically limited and backward reactions are not probable, the dependence of the methanol conversion rate on its concentration can be mirrored by different levels of dilution using an inert gas instead of products. However, the diluent should not block the active sites or alter the reaction mechanism; thus, water is not a suitable fit, as one group¹⁶⁵ varied the water content in the feed from 0 to 75 wt% and observed an influence of water dilution on the rate of each specific species' production. Besides this, to accurately mimic the actual reaction conditions, the diluent should impose similar molecular diffusion resistance as the actual products of the reaction (*i.e.*, with regards to methanol penetrating into the pores). For instance, methanol diffusion in the presence of dominant MTO products such as ethylene, propylene, and butylene is not similar to methanol in argon.



Table 3 MTO reaction pathways and rate expressions, with inclusion of a coke dependency term¹⁶²

Reaction pathway ^a	Rate expressions ^b	
	Rate of consumption of species A (MDOH)	Rate of production of species i ^c
MDOH $\xrightarrow{k_1}$ CH ₄	$r_A = a' \cdot r_{A0} = a' \cdot k_A^0 e^{-\left(\frac{E_{dA}}{RT}\right)} p_A$	$r_i = a' \cdot r_{A0} \cdot \alpha_i = a' \cdot k_A^0 e^{-\left(\frac{E_{dA}}{RT}\right)} p_A \cdot \alpha_i$
MDOH $\xrightarrow{k_2}$ C ₂ H ₄	$r_A = a' \cdot k_A^0 e^{-\left(\frac{E_{dA}}{RT}\right)} p_0 y_A$	$r_i = a' \cdot k_A^0 e^{-\left(\frac{E_{dA}}{RT}\right)} p_0 y_A \cdot \alpha_i$
MDOH $\xrightarrow{k_3}$ C ₃ H ₆	$r_A = a' \cdot k_A^0 e^{-\left(\frac{E_{dA}}{RT}\right)} p_0 \cdot \frac{1 - X}{1 + X \cdot \sum_i \alpha_i + \frac{N_{N_2}}{F_A \cdot t_p}}$	$r_i = a' \cdot k_A^0 e^{-\left(\frac{E_{dA}}{RT}\right)} p_0 \cdot \frac{1 - X}{1 + X \cdot \sum_i \alpha_i + \frac{N_{N_2}}{F_A \cdot t_p}} \cdot \alpha_i$
MDOH $\xrightarrow{k_4}$ C ₄ H ₈	where, catalyst activity function : $a' = e^{-\beta C}$	
MDOH $\xrightarrow{k_5}$ C _n H _m ^c	$C = C_0 \left(1 - e^{\left\{ \left[a+b \left(\frac{T}{1000} \right) + c \left(\frac{T}{1000} \right)^2 + d \left(\frac{T}{1000} \right)^3 \right] + f \cdot p \cdot \text{WHSV} \right\} \cdot t} \right)$	
	where, stoichiometric coefficient function : $\alpha_i = x_i (e^{u_i \cdot C} + z_i) \cdot e^{v_i/T}$	

^a Reaction pathway presented as simplified unbalanced reactions (side products omitted). MDOH refers to the lumped species of MeOH and DME.
^b N_{N_2} refers to the molar flow rate of N₂ (mol h⁻¹). F_A is the molar flow rate of methanol feed (mol min⁻¹). t_p is the pulse duration (min). p_A and p_0 respectively refer to the partial pressure of compound A (MDOH) and the initial total system pressure (both in MPa). R is the gas constant (8.314 J mol⁻¹ K⁻¹). C is the weight percentage of coke on the catalyst in g_{coke}/100 g_{cat}. C_0 is the final coke content on the catalyst in g_{coke}/100 g_{cat}. ^c C_nH_m refers to rest of hydrocarbons (*i.e.*, C₅⁻ and paraffins).

Table 4 summarizes the parameters and specifications of proposed kinetic models so once can assess the alignment of these models with these considerations.

It should be noted that some research groups have also begun to fixate on the kinetics surrounding the steam/air regeneration process of SAPO-34 in MTO,¹⁶⁹⁻¹⁷¹ as a means of de-coking the spent catalyst. Although this lies beyond the scope of the present review, it remains an interesting topic of discussion from an industrial perspective.

3.5.1 Limitations of the current kinetic approaches.

Considering the diverse variety of kinetic models discussed (see Table 4), the next step is to understand why such discrepancies exist and determine which models best capture the MTO process under realistic operating conditions. Commencing with the former, discrepancies among the models stem largely from one of the main areas: the considered assumptions, the mathematical simplifications made, and the experimental setup utilized. The first major discrepancy exists in the extent of lumping of the feed (*i.e.*, assumptions regarding the feed composition). Some models treat methanol and DME as separate reactants,^{172,173} while others have taken the approach of lumping the two species.^{174,175} The discrepancy lies in the fact that some articles choose to pursue a mathematically simpler approach with the consideration that DME exists in quasi-equilibrium with methanol. However, the realistic and more accurate approach, yet more complex one, is to still model the two species independently, since the operating conditions in place may shift the equilibrium, resulting in the lumping method being an oversimplification. Overall, this discrepancy exists depending on how researchers choose to weigh accuracy

relative to simplicity of the model. The second major discrepancy exists in the reaction mechanistic scheme utilized, in the development of the reaction rate expressions, and henceforth, the kinetic model. Some groups have used a simple power law approach for their model formulation,^{172,174-179} while others follow a more complex Langmuir-Hinshelwood (L-H) model.^{180,181} Some groups have also adopted a mixture of the two approaches.¹⁸² The reason behind differences in kinetic model type is primarily due to alterations in objectives regarding mathematical simplicity and data availability. For instance, adopting the power law approach allows for simple empirical fitting, with the goal of achieving fast, reactor-scale applicability. However, adopting approaches such as L-H or microkinetic-based models is more computationally expensive and complex, but allows for greater fundamental accuracy and mechanistic insight. The third major discrepancy lies in the level of product species lumping that takes place in formulation of the kinetic model. Research groups either take the route of lumping product species into groups such as light olefins or paraffins,^{174,175,177,178,182} whereas others choose to model products completely unlumped and individual from one another.^{172,180,181} As with the previously discussed discrepancy, these lumping differences arise from differences in research group objectives regarding simplicity and mechanistic reliability. The lumping approach allows for easier modelling, mathematically speaking, but simultaneously sacrifices on detailing fundamental mechanistic insights. The fourth discrepancy between different kinetic models is whether researchers have addressed the issue of catalyst deactivation *via* coke formation. Some researchers include deactivation



Table 4 Evaluation of the proposed kinetic models for the MTO reaction over SAPO-34^a

T (°C)	WHSV ($\text{g}_{\text{MeOH}} \text{g}_{\text{Cat}}^{-1} \text{h}^{-1}$)	Dilution (wt% of diluent in the feed)	Rate equation type	Produced and lumped species considered	Coke formation or deactivation consideration	Ref.
400–450	10.7–37.2	69 wt% water	Power law	DME, CH ₄ , C ₂ H ₄ , C ₃ H ₆ , C ₄ H ₈ , C ₅ H ₁₀ , CO, CO ₂ , C ₂ H ₆	The effect of coke was not considered	56
400–550	57–2558	Helium as diluent, N/A wt%	Power law	C ₂ H ₄ , C ₃ H ₆ , C ₄ H ₈ , C ₅ H ₁₀ , C ₆ H ₁₂ Lumped species: oxygenates (MeOH + DME), paraffins	Coke was included as a parameter C , referring to the wt% of coke on the catalyst	100
375–425	3.2–12.8	36 wt% water	Mainly Langmuir–Hinshelwood approach, minority being power law	MeOH·Z, DME·Z, C ₂ H ₄ , C ₃ H ₆ Lumped species: paraffins	The effect of coke was not considered	161
375–475	7.08–35.91	Nitrogen as diluent, N/A wt%	Power law	CH ₄ , C ₂ H ₄ , C ₃ H ₆ , C ₄ H ₈ Lumped species: MDOH (MeOH + DME), and rest of hydrocarbons (paraffins, except CH ₄ , and C ₅ ⁺)	Coke was included as a function of the final coke content, temperature, time, WHSV, and methanol partial pressure	162
350–475	2.27–100	0–75 wt% water	Power law	C ₂ H ₄ , C ₃ H ₆ , C ₄ H ₈ Lumped species: oxygenates (DME + MeOH) and rest of hydrocarbons (paraffins and C ₅ H ₁₀)	The effect of coke was not considered	165
450	1–500	~0.02 wt% helium	Power law	CH ₄ , C ₂ H ₄ , C ₃ H ₆ , C ₃ H ₈ , coke Lumped species: C ₄ (C ₄ H ₈ + C ₄ H ₁₀), C ₅ (C ₅ H ₁₀ , C ₅ H ₁₂ , C ₆ H ₁₂ and paraffins)	Coke was included as a function of time	163
400–470	31.7–339.4	~69 wt% water	Power law	CH ₄ , C ₂ H ₄ , C ₃ H ₆ , C ₃ H ₈ , C ₄ H ₈ Lumped species: MDOH (MeOH + DME), C ₄ (C ₄ H ₁₀ + 1,3-butadiene), and C ₅ ⁺ (includes C ₂ H ₆)	A deactivation rate was introduced (in terms of grams of catalyst that deactivates per unit time)	166
450	1.68–8.40	20–60 wt% water	Power law	C ₂ H ₄ , C ₃ H ₆ Lumped species: C ₄	Coke was included as a function of time, the maximum coke content, methanol feed concentration, WHSV, and the density of the catalyst bed	164
400–490	1.5–3	40–70 wt% water	Langmuir–Hinshelwood–Hougen–Watson approach	DME, CH ₄ , C ₂ H ₄ , C ₂ H ₆ , C ₃ H ₆ , C ₃ H ₈ , C ₄ H ₈ , C ₄ H ₁₀ , C ₅ H ₁₀ , H ₂ , CO, CO ₂	The effect of coke was not considered	167
400–500	3–6	N/A	Langmuir–Hinshelwood–Hougen–Watson approach	DME, CH ₄ , C ₂ H ₄ , C ₂ H ₆ , C ₃ H ₆ , C ₃ H ₈ , C ₄ H ₈ , H ₂ , CO, CO ₂	A catalyst activity function was introduced as a function of temperature, time and WHSV	168

^a Note on commercial conditions: WHSV of 2–3 $\text{g}_{\text{MeOH}} \text{g}_{\text{Cat}}^{-1} \text{h}^{-1}$; T of 400–450 °C; methanol dilution with 20 wt% water.⁷³

dynamics in the form of coke deposition in the model,^{175–178} while others choose to neglect it.^{172,174,181,182} The reasons for use *vs.* lack of integration of deactivation lies primarily in the researcher's objectives. One such objective exists in the complexity *vs.* simplicity trade-off with regards to inclusion *vs.* exclusion, respectively, of coke-based parameters. Another objective lies in the focus of the study, if the goal is to have fundamental kinetic insight (focusing on early stages of the process before the onset of deactivation) or if the goal is to have greater practical relevance (in which coke formation becomes inevitable). The fifth and last discrepancy of kinetic models lies in the chosen operating range in which these models are applicable to. The choice of operating condition depends on the researcher's purpose once again for formulating the model, *i.e.*,

trying to create models suitable for fixed-bed, lab-scale conditions, or to instead derive the model applicable for industrially based conditions (fluidized bed with a wider pressure and temperature range). This explains the wide range of operating conditions employed by different research groups.^{174,177}

Now having considered the discrepancies that exist between model types and the reasons therefore, it is hereby appropriate to critically assess which type of models are most transferable and applicable under realistic MTO conditions. From our understanding, it is clear that highly mechanistic microkinetic-based models (*i.e.*, founded on microkinetic schemes that detail hundreds of steps) are able to provide thorough fundamental insight. However, such models are highly complex, which limits their industrial relevance and makes them impractical for large-



scale reactor simulations. On the other hand, the choice of a simple power law approach lacks the capabilities to accurately predict and replicate experimental results, especially under wide or varying conditions. Hence, it seems that the most transferable approach is that of a Langmuir–Hinshelwood-type model, which possesses some detail of mechanistic insight but isn't as complex and computationally expensive as microkinetic models. The most transferable LH kinetic approach would incorporate deactivation kinetics within their model in the form of coking parameters, to reflect the reality of commercial MTO operation. The model should also be applicable to operating conditions typical of realistic MTO conditions, which means accounting for water co-feeding, and under operating ranges close to 400 °C and 1 bar. In terms of the degree of lumping of the feed and products, a semi-lumped LH based model is recommended. However, it is essential to note that a great degree of lumping can lead to a model that is able to describe overall conversion trends reasonably well, but is unlikely to be fully reliable for predicting detailed product evolution. This is because coke formation in SAPO-34 does not affect all reaction pathways for the different products equally. Thus, simple, heavily lumped models (which often treat deactivation as a single decay factor) are only able to capture the decline in activity, but cannot accurately predict the product distribution over time on stream (especially given the true complexity of the hydrocarbon pool mechanism for MTO over SAPO-34). Hence, for lumped models to be more transferable, they need to include a variety of deactivation effects explicitly stated, instead of a single empirical decay multiplier. This should be done in order to capture and validate experimental data over long time periods on stream, the route-specific nature of coking (*e.g.*, the olefin-based *vs.* aromatic-based cycle of the HCP concept), and any alterations in product distribution due to coke buildup in micropores. Thus, lumped models that include activity factors tied to coke mass and are route-specific (*e.g.*, separate activities for olefin-based *vs.* aromatic-based hydrocarbon-pool pathways) would be most transferable to commercial MTO application, but validity must still be confirmed by comparing model predictions to long-time experimental industrial datasets across a wide range of operating conditions.

Reflecting the need for catalyst deactivation dynamics in achieving a transferable kinetic model, one should consider the difficulties of including coking parameters and in turn seek solutions to mitigate such issues. Hence, to begin, the main challenges most models face in integrating deactivation dynamics in kinetic models will be discussed as follows. The first challenge in modeling these reactions is due to the complexity of the hydrocarbon pool concept, *i.e.*, referring specifically to the dual-cycle mechanism in which the MTO process takes place. This mechanism consists of two parallel, autocatalytic cycles: an olefin-based cycle and an aromatic-based cycle, which operate simultaneously, and in turn complicate the modeling of catalyst deactivation. This is because the aromatic cycle exhibits both activating and deactivating roles. While certain aromatic species, *i.e.*, polymethylbenzenes, are active intermediates, others, such as polymethylnaphthalenes, while also active for light olefin

formation, can grow into larger, polyaromatic species that cause deactivation. This dual role of some key intermediates makes it difficult to model deactivation dynamics accurately.¹⁸³ Overall, this complexity often necessitates oversimplifying the process by reducing the number of HCP-based elementary steps. A second major challenge in modeling deactivation dynamics is the nature of coke formation and deposition. Coke is composed of multi-ring aromatics of varying sizes, from simple two-ring structures to extensive multi-ring species. Hence, this diversity in size leads to differences in the sites of coke deposition. Smaller coking molecules may develop within the internal cages of the catalyst, whereas larger coke molecules may accumulate more on the external surface. This variability in deposition location, which is directly affected by the size of coke, can result in a spectrum of diffusional effects that are both location-dependant and product-dependant, presenting one of the primary challenges in accurately modeling this coke forming process. It is clear from research¹⁸³ that the accumulation of coke on the catalyst, and the type of coke species that forms, plays a crucial role in both methanol conversion and product selectivity. However, existing kinetic models still struggle to represent mathematically the effects of coke deposition, location and coke type associated with the MTO process over SAPO-34. A third major challenge for modelling deactivation dynamics is the rapidity and transient nature in which SAPO-34 catalyst deactivates. Specifically, deactivation of SAPO-34 in MTO is a highly transient process, which exhibits a strong dependence on time rather than progressing at a steady state. Unlike reactions that can be adequately described with time-independent rate constants under steady-state assumptions, in SAPO-34, product distribution evolves continuously as a function of both coke composition and distribution within the zeolite. Since it is difficult to track in real time, this rapid and fluctuating deactivation nature complicates its accurate integration into a kinetic model.¹⁸³ Hence, taking the prementioned into account, most kinetic models struggle to properly include all the pertinent factors in a way that accurately describes deactivation tendencies, resorting to lumped or simple models.

Addressing the problems of accuracy and oversimplification in MTO kinetic modelling, modern studies have begun to incorporate Machine Learning (ML) techniques to achieve a more reliable approach. One way researchers¹⁸⁴ have done so is by design of an Artificial Neural Network (ANN) to model catalyst deactivation. After validation of the fidelity of the hybrid model, by comparing simulated results with experimental kinetic data, it was observed that ANNs enhance the simulation of deactivation dynamics as they are able to more accurately capture the nonlinear relationships between activity and altering operating conditions. Rather than relying on direct coke deposition data over time on stream, the ANN implicitly accounts for coke formation and pore blockage by learning from experimental trends of activity decay. This way, the ANN works as a surrogate model for catalyst activity decline, replacing detailed mechanistic coke models. Its output triumphs that generated by simple lumped kinetic models, in turn yielding a hybrid model that is mechanistically interpretable and flexible enough to capture real deactivation behaviour. Another



promising ML approach is the use of Relevance Vector Machine (RVM) modelling, which widely uses industrial data sets for optimal operation of the MTO process at a commercial level.¹⁸⁵ RVM assists in predicting the yield distribution of main products given operating conditions, providing both mean and uncertainty estimates. The connection between kernel parameters and the predicted mean and variance is analyzed to make the model interpretable. While research acknowledges that coke deposition and catalyst deactivation affect yields, RVM is used primarily as a probabilistic yield predictor for process optimization, not as a direct mechanistic model of SAPO-34 deactivation.

Overall, although research on enhancing our knowledge of the kinetics of the MTO process has been promising, especially with the recent emergence of machine-learning-assisted approaches,^{184,185} there still exist some limitations to provide a clearer fundamental understanding. One such factor is the consideration of how diffusion limitations, transport restrictions, and mass transfer phenomena take place inside the small CHA cages of SAPO-34 during the MTO process, which in turn would affect the apparent kinetics, deviating from the intrinsic kinetics of elementary steps. The growth of the hydrocarbon pool and aromatic species within these small cages can introduce diffusion limitations, leading to slower-than-expected reaction rates. Such transport restrictions can lead to coke precursors building up near pore openings and promoting secondary reactions, which is usually not accounted for by intrinsic kinetic models. Hence, kinetic models that ignore such diffusion limitations, due to coke buildup in pores, tend to underestimate deactivation rates. Some studies have begun to delve into the effects of diffusion explicitly within their kinetic models by coupling intracrystalline mass transport with reaction networks, thereby forming a simulated dataset better in agreement with experimental data, especially during later stages of time on stream (*i.e.*, the deactivation period). Researchers have employed reaction-diffusion models using Maxwell–Stefan transport (intraparticle) theory, for MTO on SAPO-34, using the dual-cycle mechanism, with the effect of coke formation on diffusion and adsorption accounted for.¹⁸⁶ Generally, studies^{186–188} have concluded that both reaction site loss and internal diffusion limitations (which increase with coke accumulation in pores over time on stream) slow down apparent kinetics, reducing rates observed experimentally relative to computed intrinsic elementary-step rates (where diffusion effects are neglected). Specifically, the reduced effective diffusivities due to coke growth create a recurring loop that accelerates deactivation, in turn further slowing down the observed kinetics of the process. As deactivating species build up in pores, this slows down diffusion and increases local residence of such species, which leads to the buildup of even more coke and even slower diffusion as a result (*i.e.*, a positive feedback loop). Studies have shown that effective diffusion coefficients reduce by up to four orders of magnitude during deactivation as pores are blocked—especially for larger hydrocarbons.¹⁸⁶ These larger intermediates get trapped within the CHA cages, causing intraparticle resistances and in turn shifting product selectivity primarily to smaller compounds such as ethylene, which can diffuse out of the crystal more smoothly.¹⁸⁸ These pre-mentioned

diffusion-controlled, apparent product selectivity shifts cannot be easily captured by lumped kinetic models, which generally neglect such effects. Overall, despite the progress within this research direction of the kinetics of the MTO process, it seems that the development of multiscale models, comprising of both intrinsic reaction rates and transport phenomena inside SAPO-34's cages, still remains open for development in order to obtain a highly accurate model to describe the MTO process under industrially relevant conditions.

4 Catalyst design and modification

4.1 Introduction to SAPO-34

SAPO-34 is a microporous silicoaluminophosphate that possesses a topology analogous to that of the naturally occurring silico-aluminate chabazite (CHA) zeolite. The structure consists of a three-dimensional pore system of alternating D6R (double-six rings) cages and large CHA cavities (~9.4 Å in diameter) that are accompanied by smaller eight-ring (8r) windows/pore openings (3.8 Å) (Fig. 23). The 3D structure of SAPO-34 has been depicted in a variety of manners. Its molecular sieving capabilities makes SAPO-34 suitable for applications in the MTO process, allowing for the diffusion of the primarily desired C₂ and C₃ olefinic products out of its framework. In comparison to the neutral aluminophosphate (AlPO₄-34) structure, which predominantly consists of alternating Al–O–P bonds, SAPO-34 possesses a similar framework, one in which primarily phosphorous atoms (of PO₄ tetrahedra) are substituted by silicon.^{189–192} This is considered the first (and well-acknowledged) substitution mechanism (SM1), which forms Si (4Al) (isolated) structures; however, the second potential mechanism (SM2) involves the double substitution of neighboring Al and P by Si atoms, generating island-based entities *i.e.*, Si (*n*Al) (*n* ranging from 3 to 0).^{190–192} Overall, the incorporation of Si⁴⁺ for P⁵⁺ (SM1 route) yields a negatively charged framework; hence, protons (as additional cations) with Brønsted acid properties are introduced into the structure as part of bridging hydroxyl groups (*i.e.*, Si–OH–Al) for total charge compensation.^{190–192} These ultimately play a major role as the active sites for the MTO reaction.

SAPO-34 molecular sieves are primarily synthesized using a hydrothermal method due to its effectiveness and simplicity. However, alteration of the types of precursors utilized in synthesis (as well as the overall procedure used) has shown to have varied effects on particle shaping, capable of generating particles that are sheet-like^{196,197} and micro-spherical.^{196,198}

Overall, since SAPO-34 suffers from relatively rapid deactivation as a result of pore blockage by organic species, there exists the need to uncover methods to prolong catalyst lifetime.^{60,61} Hence, Table 5 summarizes the key scientific and technological challenges in SAPO-34-based MTO catalysis, with mitigation strategies and future directions for such work. Furthermore, as will be seen in the following sections, research groups have begun to improve catalyst performance by employing techniques during synthesis as a means of reducing particle size (to shorten the molecular diffusion pathway) as well as introducing secondary meso- and macro-pores, resulting



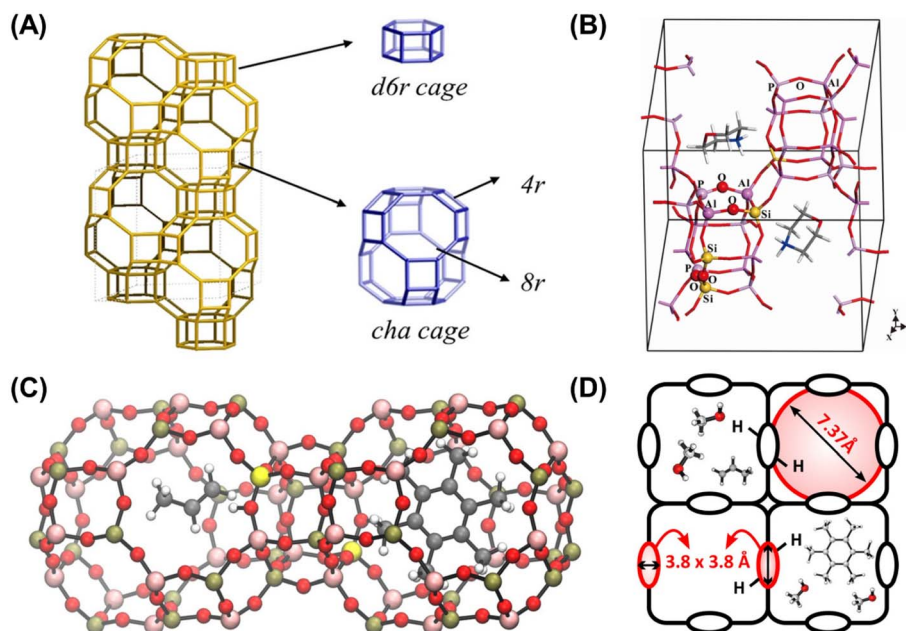


Fig. 23 (A) SAPO-34 (CHA type framework) topology structure alongside the composite building units. Reproduced from ref. 193 with permission from John Wiley and Sons, Copyright 2018. (B) Geometry-optimized SAPO-34 structure, with the four O connectivities indicated (Si–O–Si, Si–O–Al, Si–O–P, and Al–O–P). Reproduced from ref. 194 with permission from American Chemical Society, Copyright 2018. Here, the Si–O–Si species exists due to the presence of an Si island configuration. The illustrated Si–O–P linkage, which may be generated via the SM I mechanism, has not been experimentally observed. (C) H-SAPO-34 cell with two adjacent cages containing hydrocarbon pool species adapted from. Color code: Al (pink), P (green), Si (yellow), O (red), H (white), C (gray). (D) H-SAPO-34 pore system demonstrating the large cages connected via 8-ring windows. Reproduced from ref. 195 with permission from American Chemical Society, Copyright 2025.

in a hierarchical SAPO structure (to facilitate intra-particle diffusion). Furthermore, procedures involving template/OSDA (organic structure directing agent) alteration alongside modifications in silicon content (which directly impacts active site content) have also been utilized to improve catalytic activity. Fig. 24 summarizes the activity performance of optimum catalysts developed by over 35 research groups *via* utilization of the aforementioned techniques. Light olefins' selectivity and catalytic lifetime, the two most important metrics of acceptable catalytic performance, have been depicted. Since the methanol weight hourly space velocity can largely affect the catalyst's lifetime and varies largely from one article to another, to draw a fair comparison amongst the articles, a universal term 'methanol processing ability' was defined. This term describes the amount of methanol (in grams) that can be processed over 1 gram of catalyst over its lifetime (until conversion drops to 90%). Overall, the highest average light olefin selectivity that can be achieved generally falls in the range of 80–85 mol%, while the greatest methanol processing ability was recorded to be close to 30 g_{MeOH} g_{Cat}⁻¹. Although the datapoints related to each technique are scattered throughout the graph, in summary, the creation of a hierarchical structure and sub-micron synthesis seem to be the most efficient routes for SAPO-34 activity improvement.

4.2 Comparison of SAPO-34 vs. other zeolites

Given that the MTO process can employ a wide variety of microporous catalysts, each material's framework and

composition can strongly influence its performance. Silicoaluminophosphate molecular sieves like SAPO-34 have been identified as benchmark MTO catalysts due to their exceptional light olefin selectivity.²⁰⁵ By contrast, traditional aluminosilicate zeolites such as ZSM-5 exhibit different product distributions and stabilities. To contextualise SAPO-34's performance against the other readily available options, Table 6 summarises key characteristics against some representative MTO catalysts.

As depicted in the table above, SAPO-34 stands out for its high selectivity for light olefins (primarily propylene and ethylene) while suppressing the formation of heavier hydrocarbons.^{206–208} This arises from its small chabazite (CHA) cages, which favour C₂–C₄ olefin products and exclude larger molecules; consequently, SAPO-34 produces minimal aromatics or C₅⁺ by-products under MTO conditions.²⁰⁷ In addition to its structure, the milder acidity associated with SAPO-34's silicoaluminophosphate composition also provides mild acid sites that limit secondary reactions like oligomerisation or hydrogen transfer (that could form paraffins or aromatics).²¹⁰

On the other hand, the ZSM-5 zeolite is a medium-pore catalyst with larger channels (10-membered ring).²⁰⁷ Although it can achieve notable methanol conversion with a fair amount of C₂–C₃ olefins, its selectivity to light olefins is lower than SAPO-34's as a direct consequence to its more dispersed MFI channels, serving as a gateway to the formation of bulkier hydrocarbons.²⁰⁷ As such, ZSM-5 permits the generation of aromatics (such as benzene, toluene, and xylenes)²¹¹ that cannot



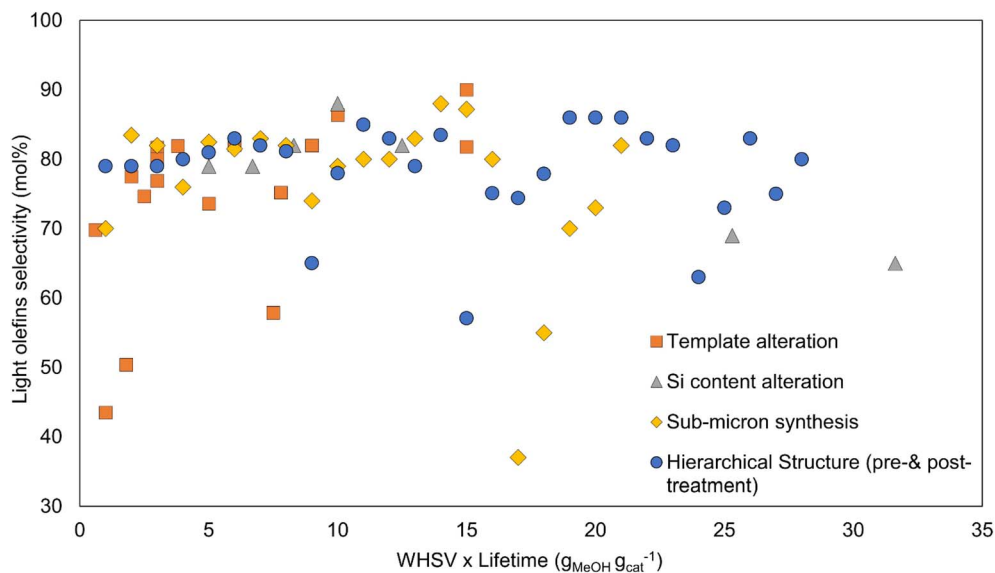


Fig. 24 Catalytic performance of SAPO-34 catalysts synthesized by employing different techniques for the MTO reaction. The data points refer to all references within Sections 4.4 to 4.5.

form in SAPO-34's constrained cages. Consequently, a significant fraction of the carbon in the MTO reaction is converted into aromatic and gasoline-range byproducts.^{207,211} Therefore, although ZSM-5 can be utilized for methanol-to-olefins, it compromises olefin yield for the generation of heavier products due to selectivity constraints.

One crucial consideration is that of catalyst stability (reported as the time-on-stream before deactivation) which markedly varies between these materials. For instance, as reported previously, the opportunity cost of employing SAPO-34 for its high olefin selectivity is its limited catalyst lifetime.²¹² SAPO-34's CHA cages tend to trap hydrocarbon intermediates as core reactions are being undergone, and the buildup of polymethylbenzene species (otherwise known as the "hydrocarbon pool") eventually leads to rapid coke deposition inside the cages, which then blocks available active sites for subsequent cycles.²¹² Consequently, SAPO-34 tends to deactivate relatively quickly in fixed-bed applications (without regeneration). As discussed in Section 2.1, this is alleviated in industry by using SAPO-34 in fluidized-bed reactors that have continuous regeneration. Nevertheless, at the particle level, SAPO-34's active lifetime per cycle is relatively short. On the other hand, ZSM-5 generally offers a longer operational time under MTO operating conditions.²¹³ This is because its larger MFI channels allow for more polyaromatic coke precursors to migrate out of its micropores; as a result, coke then forms on the external surface of the ZSM-5 crystals rather than blocking off its internal pore network (as is the case with SAPO-34).²¹¹ This prolongs lifetime and delays catalyst deactivation when compared to SAPO-34, even if the coke deposits on ZSM-5's outer surface eventually degrade the catalyst.²⁰⁷

Researchers have also opted to explore other small-pore SAPO catalysts in the pursuit of improved MTO performance, such as SAPO-18.²⁰⁹ With structural similarities to SAPO-34, the

AEI framework is reminiscent of the CHA structure due to its large cages that can be accessed through 8-ring windows, which similarly yield high selectivity towards light olefins. However, the orientation of the double six-ring unit in SAPO-18 is different, and it often features slightly lower acidity (depending on silicon content), which can influence diffusion and coking behaviour. In a study by Huang *et al.*²⁰⁹ SAPO-18 demonstrated excellent MTO performance with high olefin yield and stability (maintained over 6000 minutes on stream with co-fed syngas before deactivation). While this test involved a bi-functional setup (and the presence of hydrogen in syngas), it suggests that inducing modifications in SAPO-18 can prolong catalyst life.²⁰⁹ On the other hand, SAPO-35 (LEV framework) is another 8-ring small-pore silicoaluminophosphate that is a direct example of topology affecting stability. Similar to SAPO-18, SAPO-35 has cages accessible through 8-MR windows of similar size. However, unlike SAPO-18 (which has a three-dimensional structure), SAPO-18's LEV structure forms a 2D channel system with more restricted connectivity. This poorer diffusivity in SAPO-35 leads to faster coke saturation, which was observed in MTO testing (contrasted with SAPO-18).²⁰⁹ These comparisons are critical to elucidate how not all small-pore frameworks are made equal, characteristics like cage connectivity and surface openness can impact the delicate balance between olefin selectivity and catalyst longevity.

The case of ZSM-22 is one that depicts how necessary an appropriate pore structure is to sustain MTO catalysis. ZSM-22 is a 10 MR zeolite just like ZSM-5, though it has a one-dimensional channel system with no inherent structural inter-sections.²⁰⁷ Its shape-selective environment is so hindered that the classic hydrocarbon pool mechanism, which is a necessary intermediary step to arrive at the MTO reaction, cannot easily be sustained.²⁰⁷ In practice, only dimethyl ether (DME) is obtained as the primary product, where any light olefins obtained from



Table 5 Summary of core scientific and technological challenges in SAPO-34-based MTO catalysis, with mitigation strategies and future directions

Category	Challenges	Description & impact	Current mitigation strategies	Potential research direction
Scientific (catalyst-level)	Rapid deactivation <i>via</i> coke formation	Large multi-ring aromatics (coke) accumulate in pores and on surfaces, blocking active sites and reducing lifetime to hours in lab tests. This lowers conversion/selectivity over time and correlates with unwanted methane production <i>via</i> hydride transfer	Hierarchical structures <i>via</i> templating or desilication to improve coke tolerance (2–6× lifetime extension); metal promotion (<i>e.g.</i> , Zn) to alter acidity and suppress coke precursors (see Sections 4.4–4.5)	Explore CO ₂ co-feeding to inhibit coke deposition (<i>e.g.</i> , <i>via</i> ZnCeZrO _x /SAPO-34 dual catalysts); integrate AI-driven modeling for predictive coke management ¹⁹⁹
	Diffusion limitations in small pores	SAPO-34's 3.7 Å CHA cages enable high light olefin selectivity (~80%) but restrict larger hydrocarbon diffusion, leading to uneven coke buildup (faster at pore mouths) and mass transfer issues	Sub-micron/nanosheet synthesis to shorten diffusion paths; hierarchical meso/macropore introduction (see Sections 4.3.4–4.4)	Particle size control <i>via</i> advanced synthesis (<i>e.g.</i> , reducing to <100 nm for better acid site utilization); combine with <i>in situ</i> spectroscopy for real-time diffusion monitoring ²⁰⁰
	Acidity imbalance and silicon distribution	Medium acidity (~1 mmol g ⁻¹) is optimal, but high Si content creates islands promoting side reactions/aromatization; low Si reduces active sites, affecting initial conversion (~100%)	Tune Si/Al ratio (0.2–0.3) <i>via</i> templates (<i>e.g.</i> , TEAOH); metal doping (<i>e.g.</i> , Zn ²⁺) for Brønsted/Lewis acid synergy (see Sections 4.3.2–4.3.3, 4.5)	Theoretical studies on Zn-modified SAPO-34 for diene formation; hybrid catalysts with oxides like In ₂ O ₃ –ZrO ₂ for acidity optimization ^{201,202}
Technological (process- & industry-level)	Short lifetime requiring frequent regeneration	Industrial fluidized beds need continuous coke burn-off, but this is energy-intensive, emits CO ₂ , and risks framework damage (hydrothermal instability)	Shaping into microspheres/extrudates for better fluidization; mild steam regeneration (400–500 °C) to gasify coke (see Sections 2.1–4.6)	H ₂ co-feeding or alternative regenerants for lower-energy cycles; scale up hydrothermal-stable variants ²⁰³
	Scalability of catalyst synthesis/modifications	Lab-scale hierarchical/metal-modified SAPO-34 improves performance but faces reproducibility issues, high template costs, and inconsistent Si distribution at scale	Solvent-free/microwave-assisted synthesis; binder optimization in shaping (see Sections 4.3–4.4)	Commercial-scale templating (<i>e.g.</i> , soft/hard) for cost reduction; integrate with upstream methanol from CO ₂ /plastics for closed-loop economics ²⁰⁴
	Energy intensity and economic viability	High temperatures (400–460 °C) promote coking/alkanes; integration with feed production (<i>e.g.</i> , from CO ₂) adds separation costs, limiting <i>vs.</i> steam cracking	Diluents like water to reduce partial pressure; process optimization (WHSV 1–5) (see Section 2.3)	Direct syngas-to-olefins to bypass methanol step; techno-economic analyses for CO ₂ -derived feeds ¹⁹⁹

ZSM-22 originated from either impurity phases or reactions on external acid sites, rather than through a sustained, stable catalytic cycle in the micropores of the zeolite.²⁰⁷ Furthermore, ZSM-22's products are skewed more towards non-aromatic C₆⁺ olefins when activity is present, and virtually no aromatics are produced due to its severe spatial constraints.²⁰⁷ Thus, this inherent inability of ZSM-22 to sustain any MTO conversion illustrates the importance of having at least a 2D or 3D channel system to form methanol. In addition, the structure must not only be balanced between small pore openings in a way to allow for olefin production but also host a sufficiently connected network to alleviate coke buildup.

The alternative catalyst choices presented above exhibit some advantages over SAPO-34 such as catalyst longevity and stability, though this is at the expense of producing heavier hydrocarbons and unwanted by-products. Thus, with process-driven intentions, it is evident that SAPO-34 is the most pertinent for the MTO application.

4.3 Characterization of SAPO-34

X-ray diffraction (XRD) is typically the first characterization tool employed for crystalline-structured catalysts to verify that the desired phase has been formed *via* peak matching of the sample with that of its reference pattern. Depending on the choice of



Table 6 Comparison of selected MTO catalysts (structure, acidity, selectivity, products, and stability)^a

Catalyst & topology	Acid strength	Light olefin selectivity	Main byproducts	Lifetime categorization	Ref.
SAPO-34 (CHA, 3D small-pore, 8 MR)	Mild (silicoaluminophosphate)	High (primarily C ₂ -C ₃ olefins)	Minimal heavies (no aromatics or C ₃ ⁺ , only minor C ₄ hydrocarbons)	Short (rapid coke build-up in cages; deactivates in minutes-hours)	206–208
SAPO-18 (AEI, 3D small-pore, 8-MR)	Mild, weaker than SAPO-34 (silicoaluminophosphate)	High (comparable C ₂ -C ₃ olefins to SAPO-34)	Minimal heavies (no aromatics, similar to SAPO-34)	Moderate (improved diffusion and slightly weaker acidity extend lifetime vs. SAPO-34)	209
SAPO-35 (LEV, 2D small-pore, 8-MR)	Moderate, stronger than SAPO-18	High (comparable C ₂ -C ₃ olefins to SAPO-34)	Mainly light olefins initially; rapid formation of heavy coke/aromatics later	Short (deactivates quickly; conversion drops sharply within ~100 min)	209
ZSM-5 (MFI, 3D medium-pore, 10-MR)	Moderate (aluminosilicate)	Moderate (significant C ₂ -C ₃ olefins, but lower than SAPO-34)	Substantial aromatics and C ₅ ⁺ olefins (gasoline-range byproducts from larger 10 MR channels)	Long (hours of operation; larger pores streamline coke exit, so deposits are mainly on external surface)	207
ZSM-22 (TON, 1D medium-pore, 10-MR)	Strong (aluminosilicate)	Low (mostly heavier C ₃ ⁺ olefins)	Predominantly C ₆ ⁺ olefins; negligible aromatics (DME formed instead of olefins after prolonged operation)	Negligible (cannot sustain MTO activity; 1D pores prevent hydrocarbon pool; olefin production only observable in brief initial period)	207

^a MR = membered ring (aperture size of micropore); C₂-C₃ = ethylene/propylene range. AEI, LEV, TON is a framework type code assigned by International Zeolite Association.

synthesis procedure and template/structure directing agent utilized, competing phases (SAPO-5, -11, and -18) may present in combination with the main SAPO-34 structure.^{214–216}

Another common characterization technique is physisorption analysis, widely used tools to measure surface area and pore volume, as well as the relevant contribution of macro, meso and micropores. SAPO-34 normally exhibits an adsorption isotherm akin to that of the type I curvature.²¹⁷ Recent attempts to enhance mass transfer, by introducing meso- and macropores into the SAPO-34 structure (subsequently termed hierarchical), has led to isotherm patterns that are a combination of both type I and IV.^{218,219}

Scanning electron microscopy (SEM), a technique used to observe morphology, has shown that SAPO-34 generally possesses a cube-like particle morphology,^{220,221} with particle sizes ranging from the hundreds of nanometers^{222,223} to tens of microns.^{224,225} However, research groups have demonstrated that by alteration of synthesis precursors as well as the procedure used, different types of SAPO-34 particle morphologies are possible (such as spherical and sheet-like particle).^{196–198} SEM images may be utilized in tandem with an image-analyzing software to visually determine and quantify the particle size distribution.^{226,227} The morphological characterization of multi-phase materials is generally conducted *via* the analysis of the back scattered electrons (BSE) emitted by the sample in a scanning electron microscope.²²⁸ Heavier elements (*i.e.*, elements possessing a greater number of protons/Z-number) are able to generate more BSEs, and in turn appear brighter, relative to lighter elements.²²⁹ Hence, this presents as a useful tool when performing the morphological analysis of SAPO-34 catalyst modified with secondary metal species to assess the distribution of the promoted metal in the SAPO-34 structure.

In order to investigate the bulk elemental composition of a produced catalyst, the most commonly employed methods are inductively coupled plasma (ICP) spectroscopy and X-ray fluorescence (XRF) spectroscopy. The ICP-based technique can operate in a variety of manners depending on the spectroscopy method utilized (*e.g.*, mass spectrometry (MS) and atomic emission spectroscopy (AES)). Conducting an analysis of the bulk elemental composition can be highly valuable, as the acidity of the catalyst is a function of the number of silicon atoms present within the sample.²³⁰ Moreover, with respect to *in situ* metal incorporation as well as promotion *via* wet impregnation (which involves washing steps as opposed to dry impregnation), it is important to uncover the quantity of secondary metal species that were able to retain in the final solid catalyst.²³¹ When choosing the type of bulk elemental analysis tool to employ on SAPO-34, it is important to consider the difficulties involved in sample preparation. In terms of XRF, although there are a few sample preparation methods available such as fusion and pelletizing (to assure the homogeneity of the sample), it is often seen that direct XRF analysis of SAPO-34 powder, without any preparation, is sufficiently accurate. ICP-based methods, on the other hand, are destructive and do in fact require quite harsh sample preparation in the case of SAPO-34 catalyst. Since SAPO-34 is a silica-containing material, digestion involving hydrofluoric acid (HF), a highly toxic chemical, as well as high temperatures, are simultaneously required to destroy the framework structure.^{232–234} However, a recently published study⁷⁶ revealed that high concentration sulphuric and phosphoric acid solutions, at room temperature, are also able to digest SAPO-34, which makes ICP analysis more appealing.



X-ray photoelectron spectroscopy (XPS) and scanning electron microscopy with energy dispersive X-ray spectroscopy (SEM-EDXS) are the techniques typically utilized in the revelation of a catalyst's surface elemental composition. With the aid of these techniques, it is possible to locally specify the composition of visually-distinguished phases of the sample^{235,236} while bulk composition analysis gives an average composition of all phases presented in the specimen. In the case of metal-modified SAPO-34, depending on the procedure of secondary metal promotion (*i.e.*, impregnation based *vs.* *in situ* metal incorporation), the distribution of these participating metals may be different on the outer surface as opposed to the bulk, and could play a different role with regards to catalytic capabilities.^{237,238}

Fourier transform infrared spectroscopy (FTIR) is often used to further assist in deciphering and verifying the elemental environments present in a sample. This technique can be used to identify the chemical bonds present in the SAPO structure; however, it may fail to reveal the entire environment narrative. For instance, in the case of comparing secondary metal promoted SAPO-34 to its parent H-SAPO-34, no substantial differences are visible between the samples' spectra which could set them apart.²³⁹ The same peaks are observable for these calcined samples, present at roughly 499 cm⁻¹, 637 cm⁻¹, 725 cm⁻¹ and 1098 cm⁻¹ in the fingerprint region, as well as peaks at 1638 cm⁻¹ and 3400 cm⁻¹. These are respectively assigned to T-O bending of Si tetrahedra, T-O bending in 6-membered rings, T-O-T symmetric stretching, asymmetric stretching of TO₄ tetrahedra, -OH vibration (water absorbed on SAPO samples), and bridging hydroxyl (Si-OH-Al) groups (*i.e.*, the active sites).²³⁹⁻²⁴² Minor peaks at 3675 cm⁻¹ and 3743 cm⁻¹ have also been observed, which, respectively, can correspond to P-OH and Si-OH weak hydroxyl groups.^{240,242,243}

Nuclear magnetic resonance (NMR) spectroscopy is also another commonly employed tool in the field of catalysis, where it can determine the types of environments prevalent in a catalyst structure. With respect to ²⁹Si-NMR, five different peaks close to -90.5, -94.4, -99.6, -104.3 and -109.8 ppm are often observable, respectively corresponding to environments of Si(4Al) (*i.e.*, isolated Si), and silica islands such as Si(3Al,Si), Si(2Al,2Si), Si(Al,3Si) and Si(4Si).²⁴⁴⁻²⁴⁶ Generally, it has been observed that the greater the amount of silicon incorporated into the SAPO framework, the more likely the formation of Si islands in comparison to isolated Si environments.²⁴⁷ A general consensus amongst research groups exists in that the formation of primarily isolated silicon environments (as opposed to Si islands) is more desirable from a catalyst activity standpoint, which is often justified by the lower acidity strength associated with isolated Si sites.^{248,249} ²⁷Al-NMR spectroscopy is also another technique generally employed in studies, where two peaks are visible near roughly 35 to 39 ppm and -7 to -21 ppm, which respectively correspond to tetrahedral framework aluminum (main source of Brønsted acid sites) and octahedral aluminum species (main source of Lewis acid sites).^{51,250-253} Some research groups have also reported a weak signal close to 15 ppm due to penta-coordinated Al atoms,^{253,254} which these groups have also attributed to being Lewis acid sites.²⁵⁵ With

respect to ³¹P-NMR, a singular peak is usually present at roughly -28 ppm, which has been suggested to correspond to tetrahedral phosphorus in the SAPO framework.^{256,257}

As mentioned previously, the primary drawback associated with the MTO reaction is catalyst deactivation due to coke formation, which occurs inevitably over time on stream. The development of coke species leads to blockage of catalyst pores, resulting in rapid decline in methanol feed conversion and reduced light olefins selectivity.^{60,61,258-260} Ample research has focused on deciphering the mechanistic pathways responsible for coke formation,^{133,261} as well as uncovering ways to limit its generation rate.^{262,263} Therefore, tools such as thermogravimetric analysis-temperature programmed oxidation (TGA-TPO) have been utilized by researchers as a means to quantify the coke content over deactivated SAPO catalysts.^{222,264} Generally, the weight reduction observed in the temperature range of 50 to 200 °C corresponds to the desorption of water from the sample, whereas a second weight depletion is observed gradually between 200 and 600 °C and can be attributed to the combustion of coke species from the catalyst sample.^{222,233,265}

Other than identifying the coke generation rate, deciphering the types of species that constitute coke can provide a broader perspective on coke analysis. For instance, the identification of coke species that are distinctly responsible for ultimate pore blockage may provide further insight into the deactivation mechanistic sequence. One method to accomplish this involves analysis of coke species in deactivated catalyst *via* ¹³C or ¹H nuclear magnetic resonance (NMR) spectroscopy.^{54,266,267} Although this technique does not necessarily analyze the detailed structure of the species contained in coke, it can still provide an indication of the type of functional groups present. Groups who have utilized this method have indicated that coke species are largely aromatic-based as opposed to linear chain aliphatic compounds.^{266,267}

For a more in-depth analysis of the structure of coke species, a technique involving gas chromatography-mass spectrometry (GC-MS) may be utilized instead, which depicts coke species as ranges of multi-ring aromatics.^{84,268,269} Here, the deactivated catalyst may be digested in hydrofluoric acid, with the hydrocarbon phase extracted using dichloromethane (CH₂Cl₂).⁸⁴ The organic compounds can be subsequently injected into the GC as a means of separating the different coke species by their retention time, which can then be analyzed *via* MS successively. Although this technique may seem promising with regards to structural identification, there lies some inconsistencies between the types of coke species entrained in deactivated SAPO-34 reported by research groups. One group was able to analyze and identify insoluble hydrocarbons (up to 55 C atoms in size) within coke species,²⁷⁰ while another suggested that pyrene (16 C atoms) *i.e.*, a soluble hydrocarbon, was the largest identifiable coke species.⁵⁰

Since the active site for the MTO reaction is well known to be the Brønsted acid site (*i.e.*, the bridging hydroxyl groups, Si-OH-Al) over the catalyst, acidity measurements play an important role in catalyst characterisation. The easiest and most commonly used tool in the determination of acidic site quantity and strength is NH₃ temperature programmed desorption



(TPD).^{252,271,272} This tool generally provides two peaks that correspond to weak sites (combination of Lewis and Brønsted sites) and strong sites (primarily Brønsted) in ranges of 150 to 200 °C and 350 to 450 °C, respectively.^{273–275} FTIR spectroscopy goes beyond by distinguishing between the two types of acidic sites (Lewis and Brønsted) over the catalyst using a probe molecule. Choosing the probe molecule regarding SAPO-34 is quite challenging. Pyridine, having a molecular kinetic diameter of 5.33 Å, is not able to penetrate the small pores of SAPO-34 (pore size of 3.7 Å) and cannot be used as the probe molecule.²⁷⁶ NH₃-FTIR is not viable either, as the region where the vibrational modes of ammonia adsorbed on Lewis centers and Brønsted ones (LAS at 1625 cm⁻¹, and BAS related peaks at 1457 and 1405 cm⁻¹)²⁷⁷ interferes with the characteristic peaks related to SAPO-34 framework in the range of 500–1700 cm⁻¹. Therefore, deuterated acetonitrile (CD₃CN) was employed by one research group.⁷⁶ The C≡N stretching vibration of CD₃CN, interacting with BAS and LAS, appears within the 2100–2500 cm⁻¹ where no interference exists. The CD₃CN-FTIR test provides a qualitative analysis. In order to study the acidic sites quantitatively, CD₃CN-TPD was also conducted and related to the CD₃CN-FTIR test.⁷⁶ According to Fig. 25, since the LAS-related peak almost disappeared at around 100 °C, the low temperature peak that has appeared in the CD₃CN-TPD test can be attributed to both LAS and BAS while the other two peaks at higher temperatures are almost solely related to Brønsted acidic sites. Overall, it can be said that the majority of acidic sites presented over SAPO-34 are BAS rather than LAS.

4.4 Synthesis of SAPO-34

4.4.1 Standard synthesis approach. The synthesis of SAPO-34 zeolites typically involves mixing the precursors of the main constituent elements, specifically silicon, aluminum, and phosphorus. In addition to these primary components, other chemicals such as organic structure-directing agents (OSDAs), commonly referred to as templates, alkaline metals used as

mineralizing agents, and water as solvent play a crucial role in the synthesis process. The selection, type, and concentration of these chemicals are critical factors that significantly influence the physicochemical properties of the resulting SAPO-34, including its crystallinity, particle size, morphology, and catalytic performance. These variables directly affect the catalytic efficiency and stability of SAPO-34 in the MTO reaction, highlighting the importance of optimizing the synthesis parameters. The following sections will discuss and summarize the different parameters influencing the synthesis of SAPO-34, while in this section we focus on the basic steps involved in the synthesis of SAPO-34.

The synthesis of SAPO-34 was first reported by the Union Carbide Corporation (UCC) in the early 1980s as part of a broader study on SAPO (silicoaluminophosphate) molecular sieves.²⁷⁸ UCC researchers successfully developed a series of SAPO materials, including SAPO-34, by incorporating silicon into aluminophosphate frameworks, which led to the creation of novel molecular sieves with unique properties. This breakthrough was significant as it expanded the range of available microporous materials, combining the benefits of both zeolites and aluminophosphates. The original synthesis method involved the hydrothermal crystallization of a gel containing sources of silicon, aluminum, and phosphorus, along with organic structure-directing agents (OSDAs), which played a crucial role in defining the framework and pore structure of SAPO-34. After this innovative research, many different versions of the synthesis protocol have been developed. These variations have allowed for further adjustment of SAPO-34 properties, such as pore size, acidity, and thermal stability. As a result, SAPO-34 has become a versatile catalyst in different applications, including the MTO process.

To this day, the hydrothermal synthesis approach exists as the most commonly used procedure for the generation of SAPO-34 molecular sieves.^{189,279} The synthesis commences by gel formation *via* the mixing of different precursors (*i.e.*, sources of

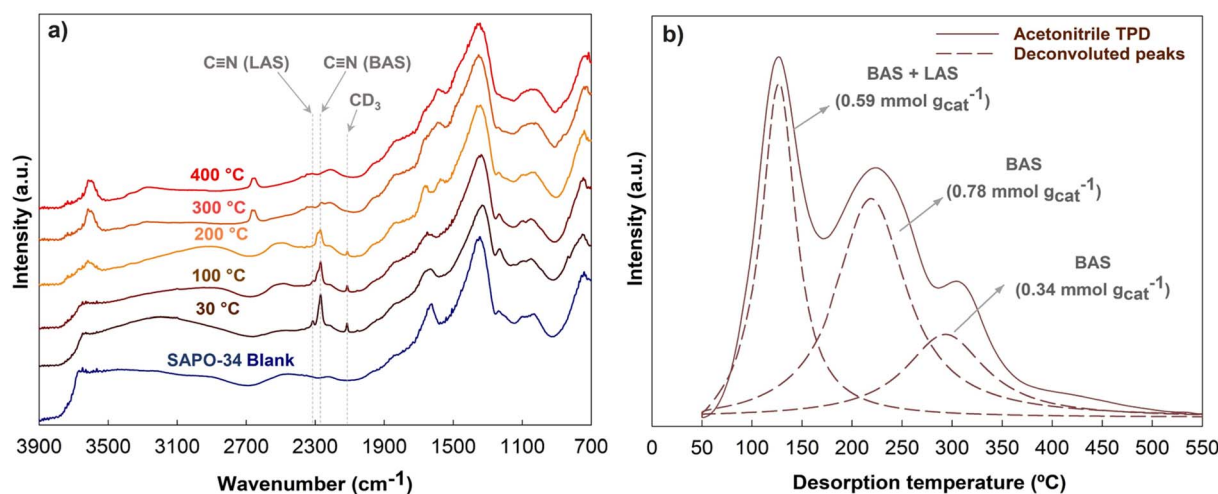


Fig. 25 (a) Temperature-dependent DRIFT spectra of SAPO-34 catalyst saturated with CD₃CN (blank sample is without CD₃CN). (b) Deuterated acetonitrile temperature programmed desorption (CD₃CN-TPD) over SAPO-34 and deconvolution of the peaks. Reproduced from ref. 76 with permission from American Chemical Society, Copyright 2023.



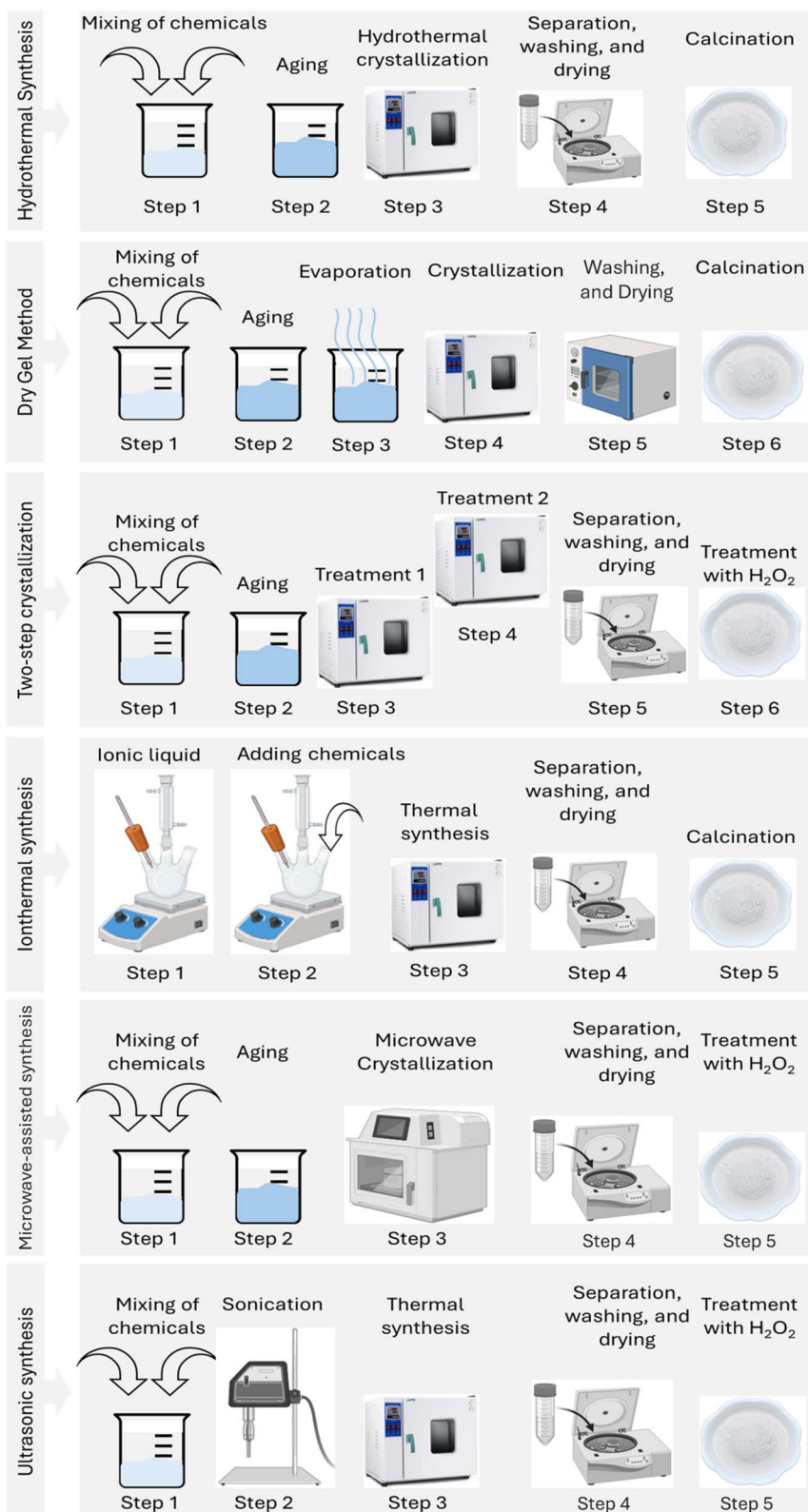


Fig. 26 Illustration of the summary of the main steps involved in the synthesis of SAPO-34 using different synthesis protocols.

Al, P and Si) and the template (commonly tetraethylammonium hydroxide (TEAOH)). This is followed by the crystallization step, typically conducted in a Teflon-lined stainless-steel autoclave at

175–215 °C, with the total period of crystallization typically lasting ~24 hours.^{189,280} The subsequent steps utilized in the final formulation of SAPO-34 are the separation of the solid



Table 7 Summary of the common synthesis protocol with the physical properties of the produced SAPO-34

Synthesis protocol	Gel composition	Crystallization temperature (°C)	Crystallization time (h)	Particle size (μm)	Morphology	Ref.
Hydrothermal	0.5(TEA) ₂ O : 0.6SiO ₂ : Al ₂ O ₃ : P ₂ O ₅ : 52H ₂ O	200	168	—	—	278
Dry gel	1.0Al ₂ O ₃ : 1.0P ₂ O ₅ : 0.6SiO ₂ : 1.8TEAOH : 77H ₂ O	180	3–24	0.045–0.075	Cube-like shape	286
Two-step crystallization	Al ₂ O ₃ : 0.6SiO ₂ : P ₂ O ₅ : 0.5TEAOH : 65H ₂ O	Step 1: 130–150 Step 2: 180	Step 1: 4 Step 2: 10	0.55–0.32	Cube-like shape	282
Ionothermal synthesis	—	130–200	3–48	1–10	Spherical and cubic ^a	283
Microwave synthesis	1.0Al ₂ O ₃ : 1.0P ₂ O ₅ : 0.6SiO ₂ : 6TEAOH : 110H ₂ O	150	7–20	0.01–0.1	Plate-like shape	285
Ultrasonic synthesis	1.0Al ₂ O ₃ : 1.0P ₂ O ₅ : 0.6SiO ₂ : 2.0TEAOH : 70H ₂ O	200	1.5 (15) ^b	0.05	Sphere-like shape	284

^a The morphology varies with the type of ionic liquid. ^b Sonication time in minutes.

from the liquid phase (*via* filtering or centrifugation), followed by drying and calcination (550–600 °C for 4–5 h to remove volatile organic compounds, *i.e.*, the template).^{189,280} Typically, the resulting catalyst particles exhibit a cube-like morphology^{220,221} (as shown in the SEM images in Fig. 28), with sizes ranging from hundreds of nanometers^{222,223} to tens of microns.^{224,225}

Other synthesis approaches have been explored for the preparation of SAPO-34, each offering unique advantages and challenges. Among these, the dry-gel conversion method involves using a minimal amount of solvent, making it an eco-friendly approach that reduces waste and allows for the precise control of crystal size and morphology.²⁸¹ The two-step crystallization method involves an initial low-temperature aging step, followed by crystallization at higher temperatures. This staged process helps in controlling the nucleation and growth phases separately, leading to better control over the final crystal size and shape.²⁸² Ionothermal synthesis, which utilizes ionic liquids as both solvents and structure-directing agents, offers a unique pathway to synthesizing SAPO-34 without the need for traditional organic templates.²⁸³ Ultrasonic synthesis uses ultrasonic waves to induce cavitation in the precursor solution, promoting rapid nucleation and crystallization. This technique not only shortens synthesis time but also enhances the

dispersion of reactants, leading to more uniform crystal growth.²⁸⁴ Microwave synthesis uses microwave radiation to directly heat the precursor mixture, significantly speeding up the crystallization process compared to conventional hydrothermal methods. This approach allows for rapid heating and uniform temperature distribution, leading to faster reaction kinetics and the formation of well-defined crystalline structures. The ability to fine-tune the microwave power and reaction time provides additional control over the final properties of SAPO-34.²⁸⁵

Fig. 26 summarizes the main steps involved in these different synthesis protocols, highlighting the unique characteristics of each method. Subsequently, Table 7 provides a detailed comparison of the gel compositions and other important details pertaining to each synthesis procedure. This summary overview serves as a valuable resource for researchers, guiding them in selecting the most suitable synthesis route based on the desired properties of SAPO-34. By understanding these synthesis methods, researchers can replicate the procedures with greater precision, reducing the time and effort required while ensuring consistent and high-quality SAPO-34 synthesis, thus allowing them to focus on advancing their core research objectives.

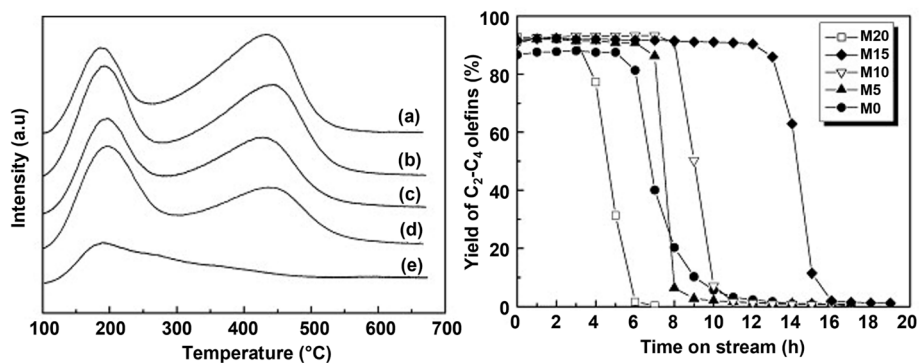


Fig. 27 (Left) NH₃-TPD (a) M20, (b) M15, (c) M10, (d) M5, and (e) M0, and (right) yield of C₂–C₄ olefins with time on stream over SAPO catalyst prepared at different compositions of MOR and TEAOH templates. “M” stands for mixed template, and the number following refers to the degree of mixing, where M0 and M20 mean only TEAOH and MOR were used, respectively. Reprinted from ref. 290 with permission from Elsevier, Copyright 2007.



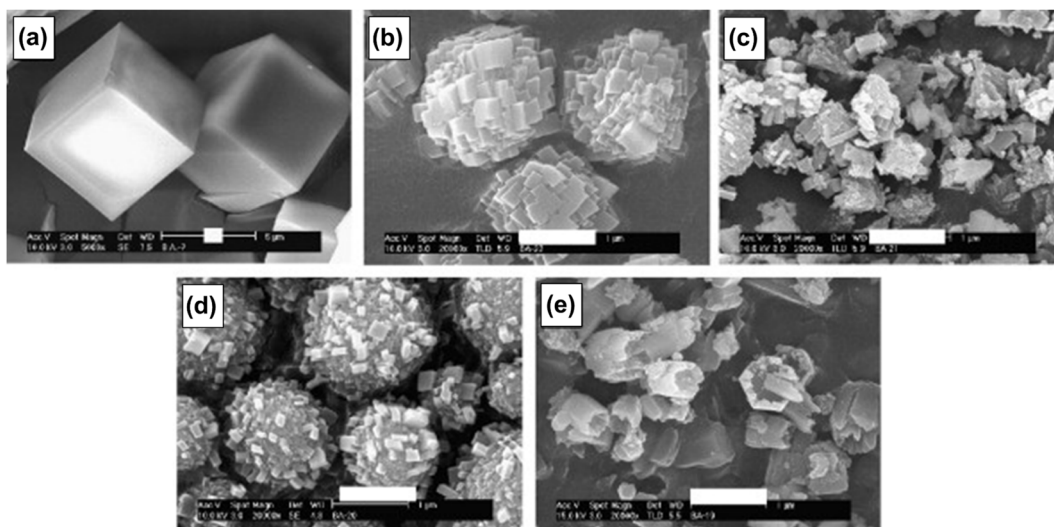


Fig. 28 SEM images elaborating the influence of the combined templates on the morphology and SAPO-34 (a) M20, (b) M15, (c) M10, (d) M5, and (e) M0. Bars correspond to 1 μm . Reprinted from ref. 290 with permission from Elsevier, Copyright 2007.

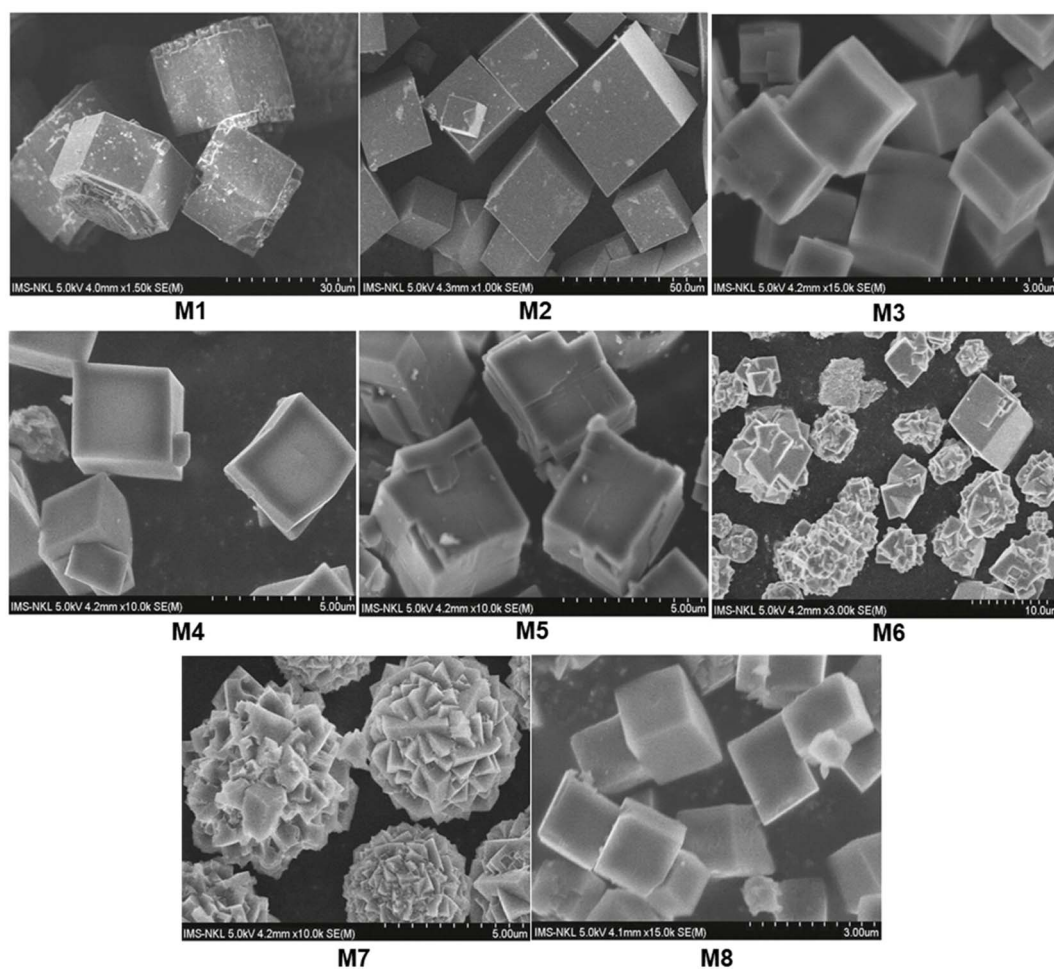


Fig. 29 Influence of different types of templates and their combination on morphology and purity of SAPO-34: (M1) TEA, (M2) MOR, (M3) TEAOH, (M4) 3MOR:1TEAOH, (M5) 3TEA:1TEAOH, (M6) 3MOR:1TEA, (M7) 1MOR:1TEA, (M8) 3MOR:3TEA:1TEAOH. Reproduced from ref. 288 with permission from John Wiley and Sons, Copyright 2019.



Table 8 Summary of the effect of different templates on phase purity, morphology, and catalytic activity

Sample	Templates	Phase	Operating conditions			Morphology		Activity		
			T (°C)	p (atm)	WHSV ($\text{g}_{\text{MeOH}} \text{g}_{\text{Cat}}^{-1} \text{h}^{-1}$)	Shape	Particle size (μm)	Stability (h)	Olefins selectivity ($\text{C}_2\text{-C}_4$)	Ref.
S-1	TEA	CHA	400	1	1.2	Cuboids	~20	<0.5	~70	85
S-2	TEAOH	CHA	400	1	1.2	Plate-like	0.5 × 0.3	6.5	~90	85
S-3	Methylamine (MA)	CHA	400	1	1.2	Cuboids	~10	1.5	~85	85
S-4	Dipropyl amine (DPR)	CHA	400	1	1.2	Typical cube-like rhombohedra	20 × 20 × 5	2.5	~80	85
S-5	DEA	CHA	400	1	1.2	Cuboids	~10	~1	~65	85
SAPO-34, SAPO-20, SAPO-35, SAPO-17	Piperidine	CHA formed at optimum piperidine/ $\text{Al}_2\text{O}_3 = 1.1$, otherwise impurities phases exist	300	2.5	N/A	Pseudo-cubic crystals	5–30	N/A	N/A	287
M1–M8	MOR/TEAOH/TEA	CHA over MOR, TEOAH or combination of the three templates	N/A	N/A	N/A	M1 (hexagonal and plate-like morphology), M2–M3 (cubic), M6–M7 (cube-like agglomerated to sphere), M8 (cubes)	M1: 10–20, M2: 20–45, M3: 3–8, M6: 5–7, M7: 5–7, M8: 3	N/A	N/A	288
M0	MOR : TEOAH (0 : 1)	SAPO-5 (92%) + SAPO-34 (8%)	250–500	1	1	Typical hexagonal of SAPO-5	<1	6.2	60.2 ^a	290
M5	MOR : TEOAH (0.25 : 0.75)	CHA	250–500	1	1	Spherical with nano-cubes aggregate	—	7.2	76.3 ^a	290
M10	MOR : TEOAH (0.5 : 0.5)	CHA	250–500	1	1	Irregular shape	Submicron	8.7	82 ^a	290
M15	MOR : TEOAH (0.75 : 0.25)	CHA	250–500	1	1	Spherical aggregates of nano-sized cubes	1	14	81.8 ^a	290
M20	MOR : TEOAH (1 : 0)	CHA	250–500	1	1	Cubes	5–20	2.7	81.9 ^a	290
SP34-X (X = 1–0)	MOR/TEAOH	For x > 0.4 CHA, otherwise impurities exist	425	1	2	Cubes	1–0.5	5.5	91.8	214
A,b,c,d,f,g	TEAOH	CHA at higher TEOAH concentration, otherwise SAPO-5	170–220	N/A	N/A	Cuboids	~1	N/A	N/A	216

^a Yield of C₂–C₃ olefins.

4.4.2 Influence of template. Alteration of the organic structure directing agent (*i.e.*, a vital component in zeolite synthesis) is one method that has been investigated with the intention of modifying the catalyst structure to ultimately enhance activity and overall performance (see Table 8 summary). This notion has also been investigated by research groups with the aim of reducing the costs associated with catalyst synthesis. This is because TEOAH, the primary template used in micron-sized pure SAPO-34 synthesis, imposes a high cost. Hence, cheaper templates such as morpholine (MOR), triethylamine (TEA), dipropylamine (DPA), piperidine, and diethylamine (DEA) have also been employed.^{85,287} However, the economic benefit of individually applying the latter three templates may be offset by the consequences imposed on the catalyst, such as larger particle size, decreased crystallinity, and generation of impurities or competing phases, which can ultimately worsen catalytic performance.^{288,289} To combat this, recently, research has been devoted to synthesizing SAPO-34 *via* a mixed template approach, which involves the idea of initial nucleation of pure SAPO-34 crystals *via* TEOAH followed by further crystal growth using one of the cheaper template alternatives.^{288,290–292} Thus, by implementation of this method, one can achieve more cost-effective SAPO-34 synthesis without having to compromise on phase purity and particle size. In fact, the use of the mixed template synthesis method has been shown to introduce new catalyst intergrowth structures that

makes internal active sites more accessible to reactants, reducing the rate of blockage of pore openings by coke species, thereby prolonging catalyst lifetime.⁷⁶

It is well known that single template usage of TEOAH can generate pure SAPO-34 phase void of other competing phases or irregularities. However, it is important to be attentive to the pH of the gel during synthesis, since, depending on the template's specific basicity strength, different quantities should be added to remain within an acceptable pH range for the commencing gel.⁷⁶ The consequences of this can be seen in one group that employed identical synthesis procedures for alternating single template samples and surprisingly observed the SAPO-5 impurity phase while using TEOAH.²⁹⁰ The use of a mixed template, specifically a combination of MOR and TEOAH, leads to a reduction in particle size and a change in morphology to a spherical type formed by the aggregation of nano-sized crystals. The mixed template approach allows for a better balance of acidic sites, which is crucial for MTO reaction. The presence of both templates in different concentrations can optimize the distribution and strength of the acidic sites, leading to improved catalytic performance in the MTO reaction as shown in Fig. 27.

With regards to the a cheaper templates (*e.g.*, DEA and TEA) as a substitute for TEOAH, these typically introduce impurity phases such as SAPO-5 and SAPO-11,^{288,289} as well as undesired amorphous phases, which can be a major factor (along with the

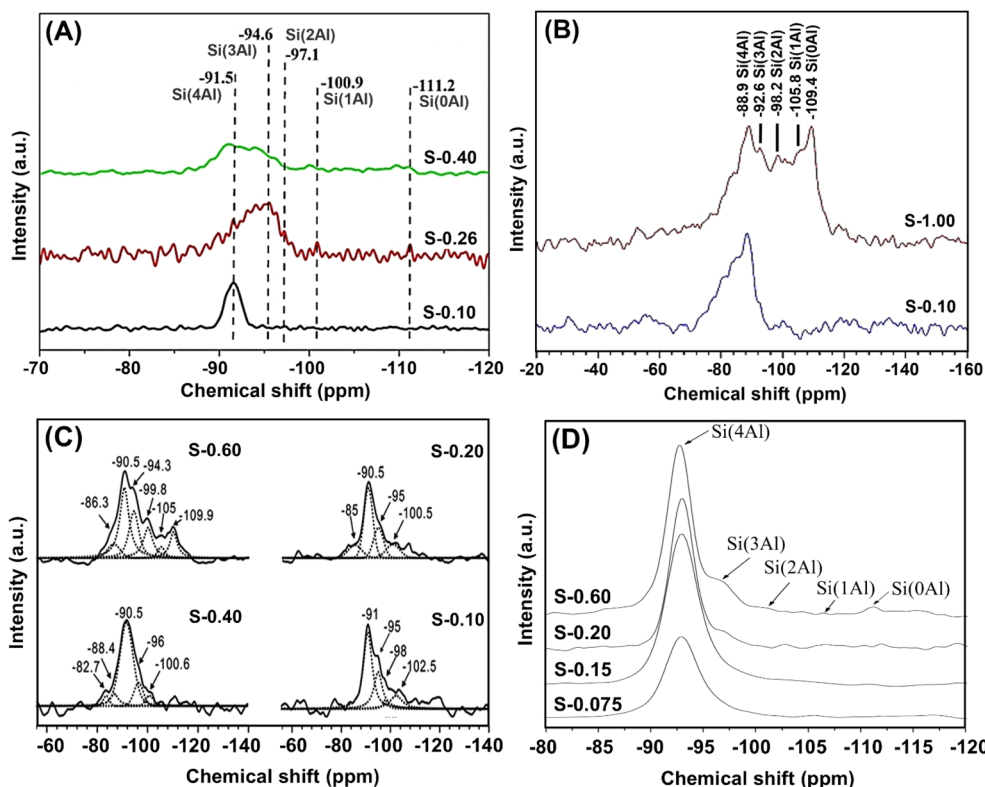


Fig. 30 ²⁹Si NMR spectrum for SAPO-34 (S-x) samples, where x stands for different SiO₂/Al₂O₃ ratios. (A) Reproduced from ref. 296 with permission from Elsevier, Copyright 2019. (B) Reproduced from ref. 280 with permission from Elsevier, Copyright 2008. (C) Reproduced from ref. 299 with permission from Royal Society of Chemistry, Copyright 2014. (D) Reproduced from ref. 300 with permission from Elsevier, Copyright 2008.



previously discussed catalyst particle size) in the downfall of catalytic activity of these samples, as can be seen in Fig. 29. For instance, the use of piperidine as an organic structure directing agent (OSDA) in catalyst synthesis generated a crystalline product that was a combination of different SAPO family structures (specifically SAPO-17, SAPO-20, and SAPO-35) in tandem with the desired SAPO-34 phase.²⁸⁷

Furthermore, based on the literature, alteration of the template undoubtedly influences the Si incorporation amount and mechanism, resulting in different acidities, despite having the same starting gel composition. Although one group²²⁴ reported that the higher the portion of TEOAH (using the MOR-TEAOH mixed-template catalyst) can result in lower catalyst acidity, in the majority of cases, no clear trend with regards to acidity could be established, with the results highly fluctuating between the different synthesized catalysts.^{293,294}

4.4.3 Influence of silicon content. Due to the strong correlation that exists between silicon content and its incorporation mechanism in SAPO-34's crystalline structure with the amount and strength of generated Brønsted acidic sites (BAS), a large number of studies have adjusted the Si quantity in the commencing catalyst synthesis gel to decipher the optimal acidity that can result in superior catalyst performance.^{131,295–297} According to ²⁹Si NMR analysis illustrated in Fig. 30, starting

with a low Si content in the gel (up to an SiO₂/Al₂O₃ ratio of roughly 0.2), the Si incorporation mechanism is mainly dominated by SM1 (P atom substitution for Si), resulting in so-called isolated Si (*i.e.*, Si(4Al)). However, when employing a higher amount of Si in the commencing gel, the Si incorporation mechanism is also comprised of SM2 (simultaneous adjacent Al-P pair substitution by two Si atoms), leading to Si-rich (island-like) environments (Si(*n*Al), *n* = 0–3).²⁹⁸

The general observation, according to literature,^{131,296} is that the larger the Si content incorporated into the bulk catalyst structure, the greater the total number of Brønsted acid sites generated – although there is the possibility that a portion of Si forms as an extra-framework amorphous silica phase. This trend occurs up to the point in which isolated Si environments begin to transform into island-like configurations. It is worth mentioning that the number of BAS per incorporated Si is lower for Si islands relative to isolated Si environments. As a result, the trend shifts to either stagnation¹³¹ or slight depletion²⁹⁶ in overall acidic site amount, as seen for the larger Si-containing samples in the NH₃ temperature programmed desorption (TPD) results of different research groups (Fig. 31).

By employing this Si alteration method, groups have found that the optimal starting SiO₂/Al₂O₃ molar ratio, with the aim of achieving superior catalyst lifetime, exists between 0.2 and 0.3,

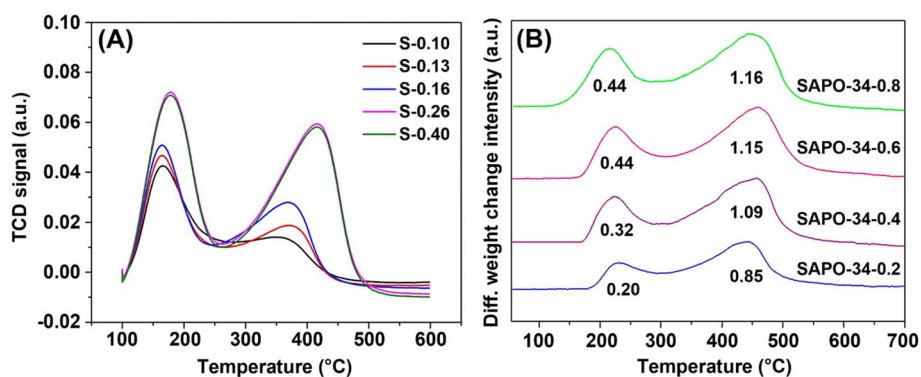


Fig. 31 NH₃ temperature programmed desorption of SAPO-34(-*x*) samples with different SiO₂/Al₂O₃ (*x*) ratios. (A) Reproduced from ref. 296 with permission from Elsevier, Copyright 2019 and (B) reproduced from ref. 131 with permission from American Chemical Society, Copyright 2018. The values beneath the peaks of (B) refer to acidic site density in mmolNH₃ g_{Cat}⁻¹.

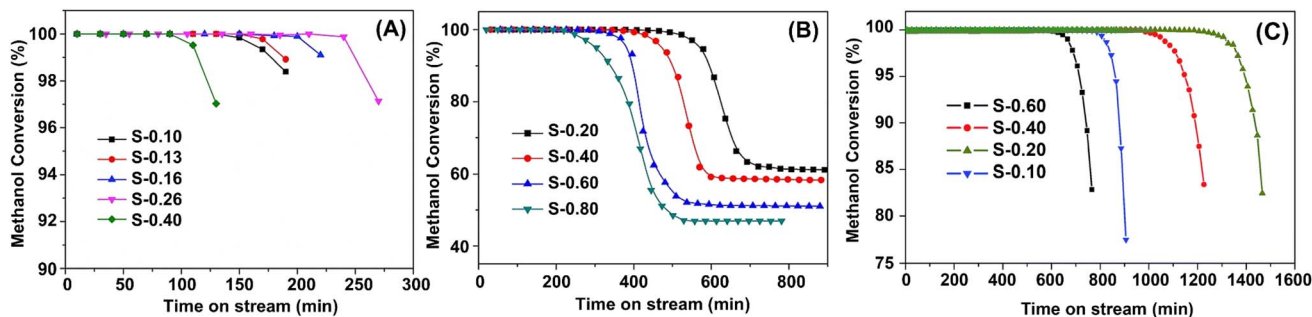


Fig. 32 Methanol conversion activity test results for SAPO-34 (S-*x*) samples, where *x* stands for different SiO₂/Al₂O₃ ratios. (A) At 450 °C and WHSV of 3 g_{MeOH} g_{Cat}⁻¹ h⁻¹. Reproduced from ref. 296 with permission from Elsevier, Copyright 2019. (B) At 425 °C and WHSV of 1 g_{MeOH} g_{Cat}⁻¹ h⁻¹. Reproduced from ref. 131 with permission from American Chemical Society, Copyright 2018. (C) At 400 °C and WHSV of 2 g_{MeOH} g_{Cat}⁻¹ h⁻¹. Reproduced from ref. 299 with permission from Royal Society of Chemistry, Copyright 2014.



Table 9 Summary of the influence of altering the Si content on SAPO-34 acidity and activity in the MTO reaction

Catalyst	Si/Al ratio	Acid density (mmol g ⁻¹)	Lifetime ^a (min)	Olefins (C ₂ -C ₃) S (%)	Note	Ref.
SAPO-34-0.2	0.4	1.05	351	~87	The Si content did not affect phase purity, morphology, particle size, or surface area	266
SAPO-34-0.4	0.8	1.41	287	~82	Acid density increased with the increase of Si/Al ratio	
SAPO-34-0.6	1.2	1.59	245	~79	Stability enhanced with lower Si/Al ratios	
SAPO-34-0.8	1.6	1.60	173	~79		
S-34(8)	0.8	—	120	~92 ^b	A lower Si/Al ratio require higher concentration of template to obtain a pure CHA phase	295
S-34(6)	0.6	—	120	~94 ^b	SEM micrographs showed plate-like morphology	
S-34(4)	0.4	—	<30	~87 ^b	The crystallite size decreased with the increase of silicon content	
S-0.10	0.2	0.92	160	~84	Samples with lower Si (S-0.10, S-0.13, S-0.16) contents formed AEI impurities beside the CHA phase	296
S-0.13	0.26	1.00	170	~84	Low Si content leads to thin-wall cube crystals with macropores	
S-0.16	0.32	1.16	220	~82	High Si content results in more solid crystals without macropores	
S-0.26	0.52	2.04	240	~84	Medium Si content produces hollow crystals with a more integrated surface	
S-0.40	0.80	2.04	110	~78	The increase in Si content also leads to a transformation from a hollow structure to a dense phase and a change in surface structure. The higher the Si content, the denser the crystal structure becomes, and the surface becomes more integrated without visible macroporous cracks	
S-0.50	1	2.13	—	—		
SAPO-34(0.3)	0.6	2.64	~1400	~90 ^b	Increasing the Si content generally leads to smoother and more cubic-like particles, while lower Si content can result in the formation of amorphous particles and rougher surfaces	297
LaAPSO-34(0.1)	0.2	—	~1500	~88 ^b	The Si/Al ratio affects the crystallinity of the catalysts	
LaAPSO-34(0.2)	0.4	1.92	~1000	~90 ^b		
LaAPSO-34(0.3)	0.6	1.29	~2100	~93 ^b		
LaAPSO-34(0.4)	0.8	—	~1900	~93 ^b		
LS	0.06	—	—	—	Si content influenced the crystallinity of SAPO-34	301
LMS	0.13	—	—	—	Si content affected the weak acid sites, which was linked to the degree of crystallinity	
MS	0.22	—	—	—	Weak acid sites had no role in the MTO reaction	
MHS	0.36	—	—	—	Hydrocarbon yield was maximum over the low medium silicon content (LMS) sample	
HS	0.50	—	—	—		

^a Lifetime was considered when the conversion dropped to lower than 100%. ^b C₂-C₄ olefins selectivity.

as seen in Fig. 32.^{131,296,301} With regards to light olefin selectivity, on the other hand, the greatest value was attained utilizing SAPO-34 of low acid site density (with an SiO₂/Al₂O₃ molar ratio between 0.1 and 0.2 in the gel),^{131,296,302} which possesses

a framework structure of primarily isolated silicon environments (see Fig. 30). It has been generally found that exceeding the SiO₂ to Al₂O₃ molar ratio of roughly 0.2 to 0.3 results in an excess of strong Brønsted active sites, which can facilitate both



aromatization of cycloalkanes, along with intermolecular hydrogen transfer reactions between polymethyl-benzenes/naphthalenes (PMBs and PMNs) and light olefins.²⁶⁶ In turn, this can amplify the formation rate of polycyclic aromatic compounds (as well as paraffinic species), ultimately resulting in much more rapid catalytic deactivation. The resulting decreased number of PMBs/PMNs, which are active components of the hydrocarbon pool mechanistic concept, can also result in a concurrent depletion of light olefin production. Furthermore, samples constructed with too large a silicon content in the gel have been found to possess dense cubic crystals with little to no macroporous cracks, relative to low Si-containing samples which seem to contain hollow, thin-walled SAPO crystals with a large range of pore sizes (meso- and macro-, in addition to micro-pores).²⁹⁶ Macro-pore presence in SAPO-34 can permit the diffusion of reactant and product species, despite the formation of coke (which initially blocks only microporous pathways). Hence, the elimination of such macro-cracks in high Si-containing samples can partly be blamed for the shortening of catalyst lifetime. On the other hand, moving below the optimal $\text{SiO}_2/\text{Al}_2\text{O}_3$ range of 0.2–0.3 can result in a catalyst that does not possess the required number of acidic/active sites to sustain long-term light olefin production. Once coke species begin to develop on this limited number of active sites, no further sites would be available for the adsorption and subsequent reactions involving further reactant methanol molecules. Although lower Si-containing samples (*i.e.*, a lower $\text{SiO}_2/\text{Al}_2\text{O}_3$ ratio such as 0.1–0.2) is beneficial for light olefin production, as it suppresses the hydrogen transfer and aromatization reactions leading to coke, it is still important to introduce enough silicon during the gel stage of synthesis to ensure an adequate active site density in the SAPO framework to achieve time-extended methanol conversion.²⁹⁶ Overall, the effects of Si content alteration on SAPO-34 acidity and activity in the MTO reaction have been detailed in Table 9.

The acidic strength of the proton attached to the bridging hydroxyl group (Si–OH–Al) can be highly affected by the electronegativity of the first and second shells of neighboring molecules.³⁰³ The higher the electronegativity of the surrounding atoms, the stronger the acidity of the Brønsted site (noting that the electronegativity of Al, Si, and P are 1.61, 1.90, and 2.19, respectively). Comparing the first and second shell of neighboring atoms for the isolated silica (Si (4Al)(8P)) and the silica island (Si (1Si 3Al)(3Si 5P)), the first neighboring shell of silica island is more electronegative (due to the presence of one Si instead of one Al). Taking this into account, it has been widely inferred that BAS associated with Si islands possess greater acidic strength than those with isolated environments,³⁰³ and consequently, catalysts possessing higher silica islands are prone to faster deactivation. However, it seems that the lower electronegativity of the second neighboring shell of the silica island has been overlooked (due to the presence of three Si instead of three P atoms). Thus, to correctly assess the strength of the acidic sites, an aggregate effect of the first and second shells should be considered.⁷⁶ Overall, there still exists the need for an exclusive study to properly address the matter.

4.4.4 Sub-micron SAPO-34 synthesis. Obtaining SAPO-34 morphology with reduced particle size is highly desired due to its contribution in prolonging catalyst lifetime.^{61,304} Groups have declared that smaller particles can increase the exposure and accessibility of active sites as well as enhance the mass transfer properties of the catalyst. This also implies a shortened diffusion pathway and subsequently lower residence time of reactant and product species in SAPO micro-pores and cavities, which therefore yields lower rates of coke formation. This notion seems rational and has been well accepted, so we speculate that one conflicting report,²⁹³ which introduced an optimum catalyst particle size (1.15 μm) and claimed that smaller ones have inferior catalytic performance, may have overlooked the other affecting parameters (*i.e.*, surface area, acidity, *etc.*). Accordingly, a large range of research groups have modified the conventional synthesis method to optimize this metric. Fig. 33 illustrates simplified schematics of these procedures. Furthermore, Table 10 provides an in-depth insight into how these different sub-micron techniques have affected SAPO-34 properties and activity in the MTO reaction.

Microwave-assisted hydrothermal synthesis is one such method that has been deemed capable in synthesizing nanocrystalline SAPO-34 catalysts relative to the conventional hydrothermal method *i.e.*, 0.075–0.15 μm (see Fig. 34(A)), compared to its parent of a few microns.^{304,305,311} This approach, involving microwave irradiation of the synthesis gel under static conditions, allowed for the reduction in crystallization time to only a few hours and more homogeneous nucleation of particles, yielding SAPO particle morphologies that were thin, smaller, more plate/sheet-like, and uniform. This ultimately resulted in roughly 50 to 60% enhanced catalyst lifetime (alongside maintained light olefin selectivity), justified by reductions in the diffusion pathway and residence time of the reactant and product species in the micro-pores of the catalyst.³¹²

Seed-assisted synthesis is an alternative technique that has been employed for the generation of higher surface area, nano-sized SAPO-34 (range of 0.2–0.7 μm as seen in Fig. 34(B)),^{222,306} which has displayed both a 4-fold enhanced catalytic lifetime and an improved average light olefin selectivity in the MTO reaction.^{306,313–315} In brief, this method operates *via* the addition of pre-constructed SAPO-34 seed crystals to the initial synthesis gel, followed by the normal hydrothermal synthesis approach.³¹⁶ The seed crystals, which partially dissolve in the gel, take on the role as surfaces or nuclei for newly growing crystals. As a result, one study³¹³ demonstrated that this method allowed for rapid nucleation, which effectively reduced the time for synthesis, as well as decreased the consumption of template (OSDA) by up to 52%, which was deemed to lower synthesis costs.

Alternatively, the generation of SAPO-34 nano-particles (0.05–0.35 μm in size, as seen in Fig. 34(C), relative to the parent of a few μm) has been achieved through the use of a top-down approach known as post-synthesis milling and recrystallization.^{307,317,318} In turn, implementation of this technique brought about improved MTO catalytic performance in terms of



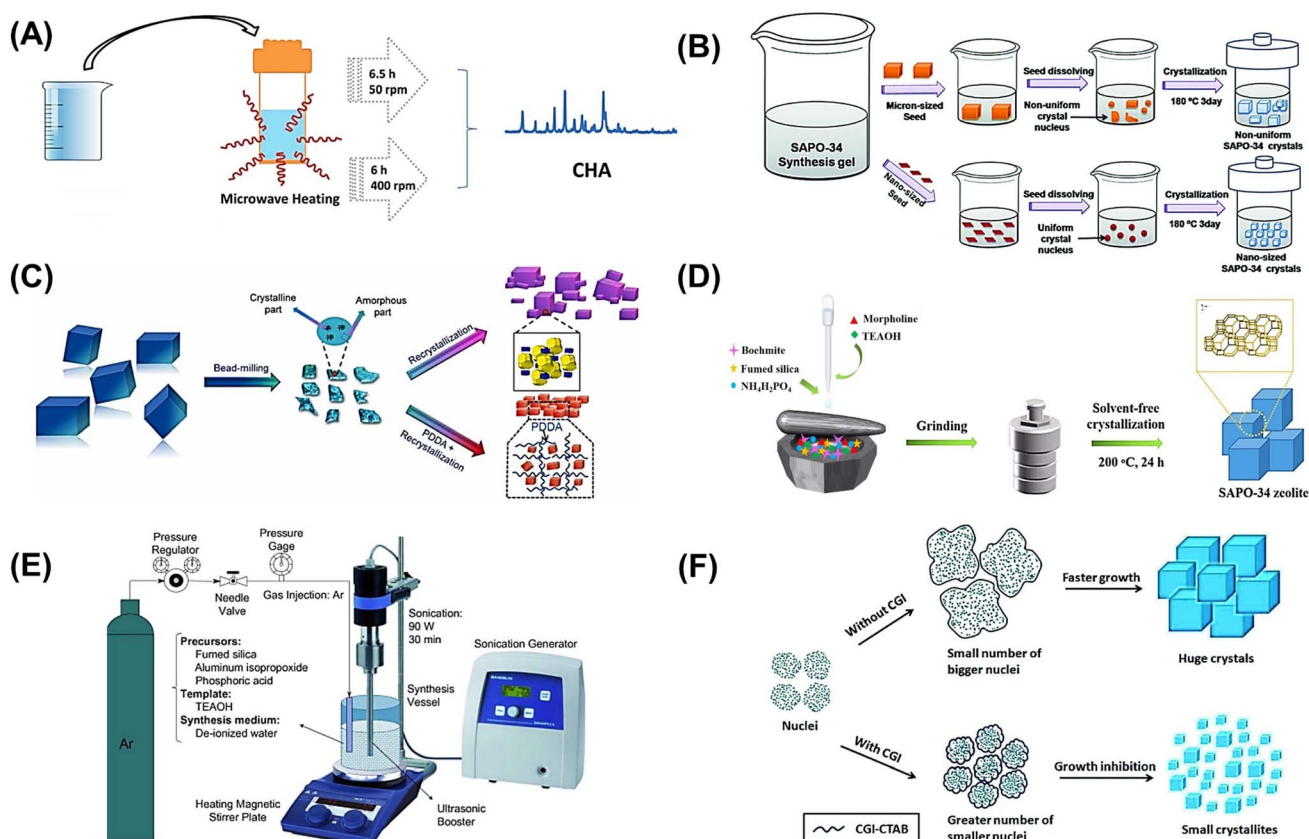


Fig. 33 Sub-micron SAPO-34 synthesis techniques: (A) microwave-assisted hydrothermal synthesis reproduced from ref. 305 with permission from American Chemical Society, Copyright 2019; (B) seed-assisted synthesis reproduced from ref. 306 with permission from Royal Society of Chemistry, Copyright 2017; (C) post-synthesis milling and recrystallization reproduced from ref. 307 with permission from Elsevier, Copyright 2022; (D) solvent-free hydrothermal synthesis reproduced from ref. 308 with permission from MDPI, Copyright 2021; (E) sonochemical-based synthesis reproduced from ref. 309 with permission from Elsevier, Copyright 2014; (F) crystal growth inhibitors (CGIs) involved synthesis reproduced from ref. 310 with permission from Royal Society of Chemistry, Copyright 2018.

both lifetime (a 5-fold enhancement) and LOS.³¹⁸ This method commences with synthesized SAPO crystals that are crushed *via* bead milling (such as using a planetary ball mill with water as a dispersing phase), followed by crystallinity recovery *via* hydrothermal recrystallization (utilizing an aluminophosphate solution). The use of X-ray photoelectron spectroscopy (XPS) in one study revealed that milling followed by post recrystallization resulted in decreased Si content on the surface of SAPO particles relative to the bulk interior.³¹⁸ Hence, other than the reduction in particle size, this decrease in particle surface acidity could suppress the hydrogen transfer and aromatization reactions responsible for coke formation, retaining the interior active sites as accessible.

Solvent-free hydrothermal synthesis, also termed dry gel conversion, exists as another method employed for the generation of nano-sized SAPO-34 crystals (in the range of 0.075 to 0.5 μm , as seen in Fig. 34(D)).^{286,308,319–321} This method can operate either by evaporating the solvents contained in the final catalyst gel prior to the hydrothermal step,³²² or alternatively, grinding and mixing the required precursors in the absence of a solvent entirely.³⁰⁸ In order to achieve a desirable $\text{H}_2\text{O}/\text{Al}_2\text{O}_3$ ratio, a certain quantity of water may potentially be introduced into the dry precursor, which is then followed by the typical hydrothermal

portion of the synthesis.³²² From an economic standpoint, an evident benefit of solvent-free hydrothermal synthesis is that it allows for significantly reduced template consumption. In addition, it is owed significantly reduced time for crystallization, with ~ 6 h of synthesis until fully crystalline SAPO-34 is successfully attained.^{286,320,321} Nano-sized catalysts constructed *via* this methodology possess a hierarchical structure containing mesopores (in addition to the traditional microporous pathways) due to the stacking of nanocrystals, which has been claimed to enable the mass transfer of reactant and product species.³²² Overall, the implementation of this method has resulted in prolonged catalyst lifetime (roughly two to four-fold increase) with equivalent to slightly greater light olefins selectivity, relative to that under the conventional hydrothermal method.^{320–322}

Sonochemical-based synthesis is introduced as another technique utilized in the construction of SAPO-34 nanocatalysts (in the range of 0.05 μm to 0.45 μm , as seen in Fig. 34(E)).^{57,227,323,324} This method involves the use of ultrasonic irradiation (rather than mechanical stirring) during the aging step, followed by a shortened hydrothermal step to minimize the overall growth of the crystal nuclei. Due to the chemical effects ultrasonic waves impose (as a result of acoustic cavitation phenomena in the solution), the rate of nucleation is



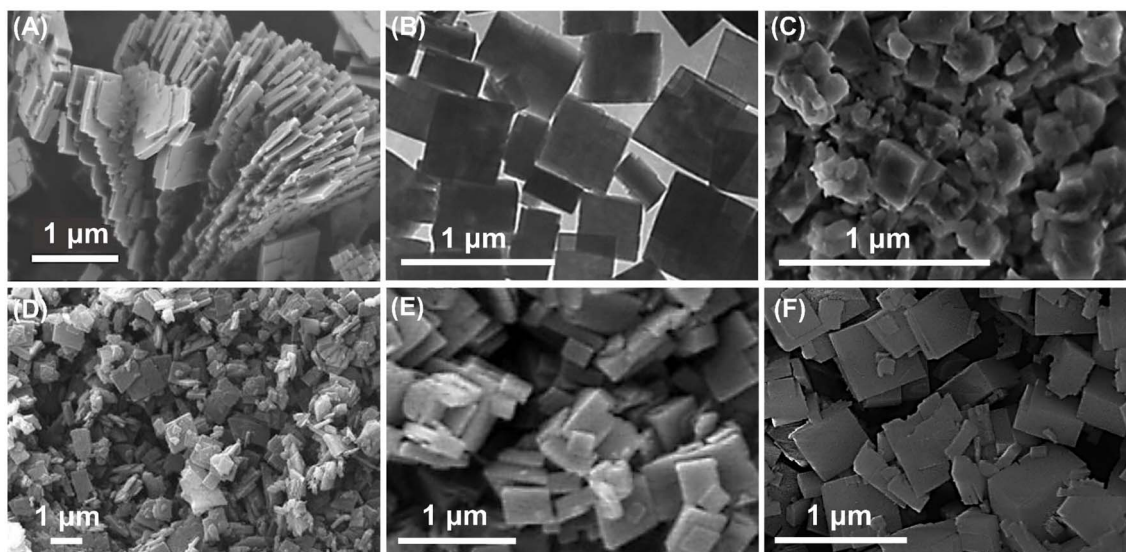


Fig. 34 SEM/TEM images of the catalysts prepared via different sub-micron SAPO-34 synthesis techniques: (A) microwave-assisted hydrothermal synthesis reproduced from ref. 312 with permission from Royal Society of Chemistry, Copyright 2014; (B) seed-assisted synthesis reproduced from ref. 314 with permission from Royal Society of Chemistry, Copyright 2015; (C) post-synthesis milling and recrystallization reproduced from ref. 318 with permission from Royal Society of Chemistry, Copyright 2014; (D) solvent-free hydrothermal synthesis reproduced from ref. 320 with permission from Springer Nature, Copyright 2016; (E) sonochemical-based synthesis reproduced from ref. 57 with permission from Elsevier, Copyright 2014; (F) crystal growth inhibitors (CGIs) involved synthesis reproduced from ref. 325 with permission from American Chemical Society, Copyright 2008.

enhanced while the agglomeration of particles is diminished, resulting in a catalyst with a larger total and external specific surface area than the traditionally prepared parent counterpart. Due to the reduced residence time of hydrocarbon species in nano-catalyst particles (*i.e.*, as a result of shorter intracrystalline diffusion length), sono-synthesized catalysts have displayed reduced coke generation rate, and thus prolonged activity (roughly two-fold), alongside a similar product distribution to the conventionally prepared parent SAPO-34 (displaying a 2 μm average particle size).³²³

Lastly, some groups have utilized crystal growth inhibitors (CGIs) during synthesis such as polyethylene glycol (PEG) (see Fig. 34(F)), polyoxyethylene lauryl ether (Brij-35), methylene blue (MB), and surfactant cetyltrimethylammonium bromide (CTAB)).^{310,325} This method consists of the addition of one or more of these compounds as either the final precursor(s) or prior to template addition during the gel stage of SAPO synthesis. The added inhibitors are generally removed during the final calcination step. On the other hand, the CGIs are suggested to interact with reactive sites of inorganic precursors in solution, such as to give rise to enhanced rates of nucleation. This ensues by the adsorption of CGIs onto the external surface of generated nuclei, allowing for their separation, and thus lowering the rate of crystal growth. Overall, this results in the formulation of a larger number of smaller-sized crystals, without imposing any significant structure-directing effects during the generation towards the final catalyst framework (*i.e.*, no clear major presence of competing phases). This method has resulted in catalysts with a much larger BET surface area than the conventionally prepared parent catalyst. A study that

investigated the use of CTAB during SAPO-34 synthesis found that the optimal resulting catalyst, sized between 0.8 and 1 μm (compared to the parent of nearly 7 μm), displayed three times longer catalyst lifetime than the parent counterpart alongside higher selectivity towards light olefins.³¹⁰

Although nano-sized SAPO-34 has proven useful in enhancing catalyst performance in the MTO reaction, the aforementioned synthesis methods generally still face issues involving low yield, higher production costs, and serious difficulties concerning scale-up. Hence, a comprehensive cost-benefit analysis of each technique seems to still be required to truly assess whether the benefits of improved catalyst performance outweigh the costs associated with these synthesis procedure adaptations. Furthermore, only a limited number of groups have investigated the effect of combined application of these methods, *e.g.*, the hydrothermal approach involving seed assistance followed by dry gel conversion.³²⁰ A detailed exploration of the possible synergistic effects of multiple technique integration (in a singular synthesis procedure) seems yet to be undertaken.

4.5 Hierarchical SAPO-34 synthesis by pre- and post-treatment methods

4.5.1 Overview. Another strategy that has been utilized by research groups to conquer fast catalyst deactivation by coking involves the introduction of secondary macro- and meso-pores to naturally microporous SAPO-34. This fundamental creation of a new level of porosity consequently results in a hierarchical structure.^{328–331} While micropores enable the molecular sieving process of SAPO-34, to maintain a high selectivity to light



Table 10 Summary of different approaches of sub-micron SAPO-34 synthesis and their effect on SAPO-34 properties and activity in the MTO reaction

Approach	Particle morphology		Activity	Note	Ref.
	Size (μm)	Shape			
Microwave-assisted approach	0.02–1	Flaky, nano-spherical, and cubic	Catalytic activity was linked to particle size. It was maximum over flake-like particles, followed by nano-sphere, submicron cube, and micron cube Coke formation was minimised with decreasing particle size	Particle size and shape vary with temperature, time, and source of Si precursor	304
Microwave-assisted approach	0.1–0.2	Sphere-like	Catalysts demonstrated 100% methanol conversion for 120 minutes and a total light olefins selectivity of ~85%	Spherical particles agglomerated to form a flake-like shape	269
Microwave-assisted approach	0.075–0.01	Varies between sphere- and cube-like	The study is limited to synthesis only, and the authors examined different parameters that affect crystallinity and morphology	Morphology and particle size were strongly influenced by microwave exposure time Microwave power delivery affected phase purity and morphology	311
Microwave-assisted approach	0.01–0.1	Plate-like	MW-assisted synthesized SAPO-34 exhibited a longer lifetime of up to 13 hours, compared to 8 hours with conventional heating. It also displayed higher initial olefins selectivity; however, after 3 h of TOS, all catalysts exhibited very similar olefins selectivity, reaching around 90% at all temperatures tested	Microwave irradiation favored lower levels of Si incorporation and thus lower acidity. It led to a significant variation in crystal shape and size compared to conventional heating, resulting in smaller size, shorter crystallization time, specific plate-like morphology, and conversely lower surface area	285
Seed-assisted synthesis	0.2–0.5	Cube-like	Methanol conversion was enhanced with the use of different-sized seeds. The smaller the crystal size of the seed, the smaller the SAPO-34 produced, leading to more effective catalytic activity & higher overall conversion rates and LOS compared to conventional SAPO-34	Seed-assisted synthesis of SAPO-34 reduced the crystal size to a range of 0.2–0.5 μm from 50–10 μm . This method also caused larger external surface areas than those obtained from conventional SAPO-34	222
Seed-assisted synthesis	0.035–0.1	Varies between sphere- and cube-like	The study analyzed the effect of the template and seed content on SAPO-34 synthesis. These factors significantly influenced the textural properties of SAPO-34. However, the analysis of the selected samples was inconclusive regarding the impact of seed content on catalytic activity due to simultaneous changes in more than one factor	Small particles (0.035–0.1 μm) agglomerated to form larger particles The introduction of seed in the synthesis reduced the use of OSDA by 50%	313
Seed-assisted synthesis	0.2	Nano-sheets	Since the seed-assisted method did not change the particle size and merely reduced the synthesis time to 20 min, there was no obvious effect on the catalytic activity	Using seeds accelerated crystallization of SAPO-34 significantly compared to methods without seeds. With seeds, the zeolite crystallized in just 10 minutes, while conventional methods could take over 40 minutes with low yields	314
Seed-assisted synthesis	0.3–0.8	Cube-like	The nano-sized seed -assisted hierarchical SAPO-34 exhibited a catalytic lifetime of up to 586 minutes, four times longer than that of the conventional micron-sized microporous catalyst. Its selectivity for ethylene and propylene reached 85.0%, a 10% improvement from conventional microporous SAPO-34 zeolites	The seed-assisted method is instrumental in achieving high yields, enhanced structural properties, and superior catalytic performance of SAPO-34 zeolites. It reduced the particle size by one-tenth of that of conventional SAPO-34 zeolite	315



Table 10 (Contd.)

Approach	Particle morphology		Activity	Note	Ref.
	Size (μm)	Shape			
Post-synthesis milling	0.05–0.15	Cube-like	Micro and nanoscale SAPO-34 catalysts achieved 100% methanol conversion. However, the lifetime of the bead-milled sample extended from around 65 to 140 minutes, increasing to approximately 290 minutes through porogen-assisted recrystallization. The catalysts had similar total olefin selectivity, but the bead-milled catalysts were more selective towards ethylene and less towards butylene	The bead-milling process effectively reduces the particle size from 20 μm to 0.05–0.15 μm , enhancing the material's properties for catalytic applications	307
Post-synthesis milling	0.05–0.35	Cube-like	After milling and recrystallization, the catalytic lifetime of the recrystallized SAPO-34 was significantly extended from 38 minutes to 224–258 minutes. The selectivity for ethylene and propylene increased from 71.6% to 82.6%. Additionally, the selectivity to propane was reduced from 6.3% to 0.6%	Milling and subsequent recrystallization not only improved particle size and shape, but also enhanced the catalytic performance of SAPO-34 in the MTO reaction, leading to longer catalyst lifetimes and higher selectivity for desired olefins	318
Solvent-free hydrothermal synthesis	10–30	Cubic	Conversion of methanol to olefins in SAPO-34 synthesised using the solvent-free method was found to be comparable to catalytic performance using the conventional hydrothermal method	Hierarchical macro- and meso-pores of 0.1–3 μm and 0.005–0.05 μm were also found in the cubic crystals, which was attributed to the presence of macro- and meso-structures in the starting materials	326
Sonochemical-based synthesis	0.05	Spherical	The study is limited to synthesis only, and the authors examined different parameters that affect crystallinity and morphology	Sonochemically prepared SAPO-34 was obtained in only 1.5 h, a significant improvement from the 24 h attributed to the conventional hydrothermal method. The former was found to be a much faster & efficient method to synthesise SAPO-34 nanocrystals	324
Crystal growth inhibitors	0.02–0.2	Cubic rhombohedral	Catalytic lifetime increased by 1.5 times when an inhibitor was used in comparison to conventionally synthesised SAPO-34. It was concluded that higher Si content in SAPO-34 gave rise to more acidic sites, thereby leading to higher conversion & superior selectivity towards light olefins	The addition of the inhibitor was deemed ineffective at suppressing intergrowth (given instances of unsuccessful modification and large crystallite size). It was found that the time of inhibitor addition to SAPO-34 is critical to attaining hierarchical structures	327

olefins, these secondary macro- and meso-pores can improve the mass transfer properties of the catalyst. They can keep interior active sites accessible to reactants despite progressive coke blockage of micropores over time, thereby prolonging catalyst lifetime. To achieve this, research groups have employed a variety of bottom-up (pre-treatment) and top-down (post-treatment) synthesis methods, which have been described in more detail in the subsequent sections. Pre-treatment methods generally involve the addition of secondary soft or hard templates into the catalyst gel during synthesis. Generally speaking, these will burn during calcination, leaving behind macro- and meso-sized pores within catalyst particles that depend on both the size and morphology of these secondary

templates. In contrast, the top-down method involves (generally chemically selective) post-treatment of the synthesized catalyst *via* the employment of acid or alkali-based etching agents that results in somewhat preferential extraction of framework cations.^{218,332,333}

In general, although bottom-up methods have generally been successful in preparing hierarchically structured catalysts, they may pose some drawbacks worth mentioning.³³² Firstly, since the process of combustion (through calcination) is generally required for template removal and simultaneous creation of desired meso-pores, this results in a sacrificial loss of potentially expensive template, in the case that no recovery-based methods are viable. Secondly, a significant quantity of



heat and water may be generated during the thermal decomposition of the mesopore agents, which could potentially contribute to the destabilization of the catalyst framework. Overall, while top-down methods may be substituted in place of bottom-up as they are more economically favorable and less complicated,³³⁴ it should be mentioned that they may yield their own consequences as well. For instance, the preferential extraction of framework cations (*i.e.*, dealumination or desilication), during chemically selective acid or alkali post-treatment, may trigger a loss in catalyst mass. This could subsequently hinder the catalyst's acidic (active) properties, and thus its potential regarding catalytic performance.^{332,335} It should also be noted that the etching post-treatment time and concentration (intensity) utilized are parameters that should be carefully considered and optimized to achieve superior catalytic performance. If chosen inadequately, its effects may be either too minimal (resulting in no significant introduction of meso-/macro-sized secondary pores) or over-bearing (yielding an unstable and demolished SAPO crystalline framework).³³³ Various instances of either approach will be discussed extensively in the following sections. Furthermore, Table 11 has been provided to summarize the different approaches of Hierarchical SAPO-34 synthesis and their effect on SAPO-34 properties and activity in the MTO reaction.

4.5.2 Soft template addition. With regards to soft templates, one that is commonly employed is the organosilane surfactant [3-(trimethoxysilyl)propyl]-octadecyldimethylammonium chloride (*i.e.*, TPOAC).^{272,336} This secondary template has been observed to adopt a dual role as a mesopore-generating agent and an alternative silica source precursor (alongside tetraethyl orthosilicate (TEOS)²⁷²) during synthesis. Due to the surfactant's adsorption behavior with regards to SAPO precursors, it has been also stated to possess crystal growth-inhibiting capabilities (see Fig. 38(A), displaying the spherical agglomeration of nanosheet-like crystals for the pre-treated catalyst). The resulting catalyst has displayed an improved catalyst lifetime (two to four-fold) as well as maintained comparable²⁷² to enhanced³³⁶ selectivity (by ~10%) towards light olefins (relative to its conventionally prepared parent counterpart). However, it can be argued that the use of organosilane surfactants as secondary templates is a relatively expensive approach.^{335,337} Therefore, another largely explored soft template includes the use of cost-effective polyethylene glycol (PEG) in a variety of molecular weights.^{338–342} This compound has demonstrated similar capability in formulating hierarchical SAPO-34 catalysts with intracrystalline micro-, meso-, and macro-porosity (see Fig. 38(B)). The resulting catalysts generated *via* this soft template have exhibited two to six times higher catalytic lifetime, alongside comparable to 10% improvement in light olefins selectivity. PEG employment has been shown to trigger a conflicting mix of either greater³⁴⁰ or lower³³⁹ concentration of strong acidic (primarily Brønsted) sites; however, groups have generally reported a depletion in average Brønsted acidic strength upon its use (with reference to PEG of higher molecular weight *i.e.*, 2000–4000 g mol⁻¹) (Fig. 35).

4.5.3 Hard template addition. Aside from the soft templates mentioned previously, the use of hard (solid) structure directing agents has been used as an alternative approach by groups to introduce mesoporosity in SAPO-34 crystals. Here, groups have primarily investigated the use of carbon nanoparticles and nanotubes as secondary (hard) templates added during the culmination of the gel stage of catalyst generation (prior to the crystallization step).^{335,337} Then, as previously touched upon, the combustion of these secondary templates during calcination is ultimately responsible for the mesopore generation seen in the catalyst structure. One group that investigated the use of carbon nanoparticles as a hard template demonstrated its ability in enhancing the catalyst external surface area and increasing the total pore volume, but the structures formulated were cavity-like rather than interconnected pores. This would have seriously hindered the catalyst mass transfer properties and internal diffusivity, and also limited the accessibility of reactant species to internal active sites.³³⁷ In turn, the resulting catalyst displayed no significant improvement in catalytic performance (*i.e.*, with regards to lifetime and light olefins selectivity) relative to the parent catalyst. In order to alleviate the previously mentioned issues associated with nanoparticle templating, carbon nanotubes have been utilized instead as an alternative secondary template.^{335,337} With respect to Fig. 38(C), its implementation has led to the generation of mesoporous channels (of varying diameters and orientation) which connect the inner section of the catalyst to the external surface. This has resulted in a three-dimensional multi-pore network, rather than simple cave-like and unconnected pores. This approach demonstrated prolonged catalytic lifetime in the MTO reaction with only slightly lower light olefins selectivity relative to the conventionally-prepared sample.³³⁵ However, the slight reduction in LOS could be argued to be related to the greater collapse of the SAPO crystalline framework structure, which could hinder its molecular sieving capabilities (Fig. 36).

4.5.4 Post-treatment desilication. Chemically selective etching (specifically desilication) of SAPO-34 catalysts has been undertaken under ultrasonic conditions *via* a fluoride medium post-treatment route involving an HF-NH₄F mixed aqueous solutions.^{218,332} The resulting post-treated catalysts (prepared individually with one of MOR, TEA or TEOH as micropore generating agents²¹⁸) displayed the desired hierarchical-like structure (see Fig. 38(D)) with slightly reduced average Brønsted acidic site strength, in turn performing better in terms of average light olefins selectivity (by ~2%). In terms of catalyst lifetime, one study displayed a range of worse (by 25–30%) to comparable values for post-treated samples compared to the parent counterparts.²¹⁸ This was largely attributed to the observed SAPO crystallinity depletion alongside insufficiency in the number of catalyst active sites due to reduced strong (primarily Brønsted) acidic site quantity. By contrast, another group³³² observed a slightly enhanced lifetime (less than 2-fold) of their treated samples, despite having a similarly large reduction in active site (BAS) concentration. Other than fluoride-based post-treatment, etching using TEOH aqueous



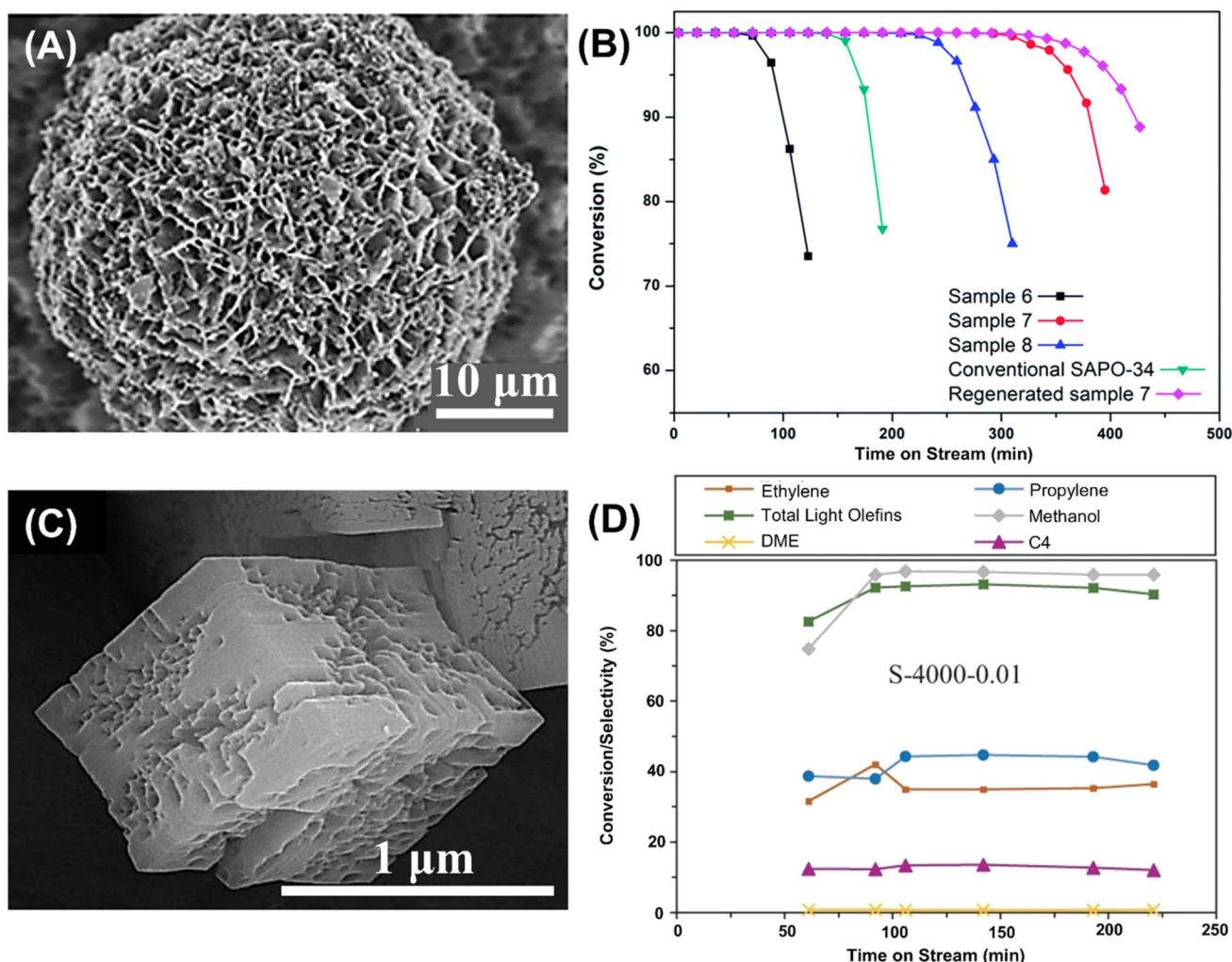


Fig. 35 SEM images and time on stream results of catalysts (fixed on one particle) produced via application of different soft templating treatment techniques: (A) TPOAC as a secondary (soft) template; (B) methanol conversion variation with time-on-stream over samples 6, 7, 8, conventional SAPO-34 and regenerated sample 7. Experimental conditions: $\text{WHSV} = 3 \text{ g}_{\text{MeOH}} \text{ g}_{\text{Cat}}^{-1} \text{ h}^{-1}$, $T = 450 \text{ }^\circ\text{C}$, catalyst weight = 300 mg. Reproduced from ref. 272 with permission from Royal Society of Chemistry, Copyright 2015; (C) PEG as a secondary (soft) template; (D) light olefin selectivities, methanol and DME conversions over the hierarchical SAPO-34 catalyst variation with time on stream for sample S-4000-0.01 with experimental conditions: $T = 425 \text{ }^\circ\text{C}$, $P = 1 \text{ atm}$, $\text{WHSV} = 2.3 \text{ g}_{\text{MeOH}} \text{ g}_{\text{Cat}}^{-1} \text{ h}^{-1}$, catalyst weight = 2 g. Reproduced from ref. 340 from Springer Nature, Copyright 2021.

solution is another procedure that has been employed for the preparation of hierarchical SAPO-34 crystals.³³³ The resultant catalysts displayed double the catalyst lifetime while maintaining the average LOS relative to the parent catalyst. However, the concentration of strong acidic sites (as well as strength) was reported to have increased over treated samples, which seems to conflict with the observed and expected preferential desilication upon treatment.¹⁶⁰ Generally, the choice of etching agent is crucial when it comes to controlling the preparation of hierarchical SAPO-34. For instance, the use of sodium hydroxide instead as a basic etching agent in the post-treatment of SAPO-34 has been observed to lead to a substantial reduction in crystallinity and selectivity remove phosphorus from the bulk structure, yielding amorphous aluminosilicate.³⁴³ Another important consideration is the time duration associated with etching post-treatment, which has also been partly investigated

by groups with regards to its effect on catalyst properties.^{218,332} It was uncovered that increased treatment time resulted in reduced SAPO-34 crystallinity, more deeply etched crystals, larger secondary pores, and greater desilication (which resulted in decreased Brønsted acidic site concentration) (Fig. 37).

4.6 Metal promotion

Metal modification of SAPO-34 has also been examined with regards to its effect on catalytic performance and activity. To the extent of our knowledge, the following metals have been utilized for SAPO-34 promotion in MTO centered studies: Co, Ni, Fe, La, Ce, Y, K, Ag, Ca, Cu, W, Mg, Mn, Ge, Zn, Ti, and Zr.^{231,233,239,241,344–360} These articles have attempted different methods of promoting parent SAPO-34 catalyst with metal, ranging from *in situ* metal incorporation (*i.e.*, isomorphous



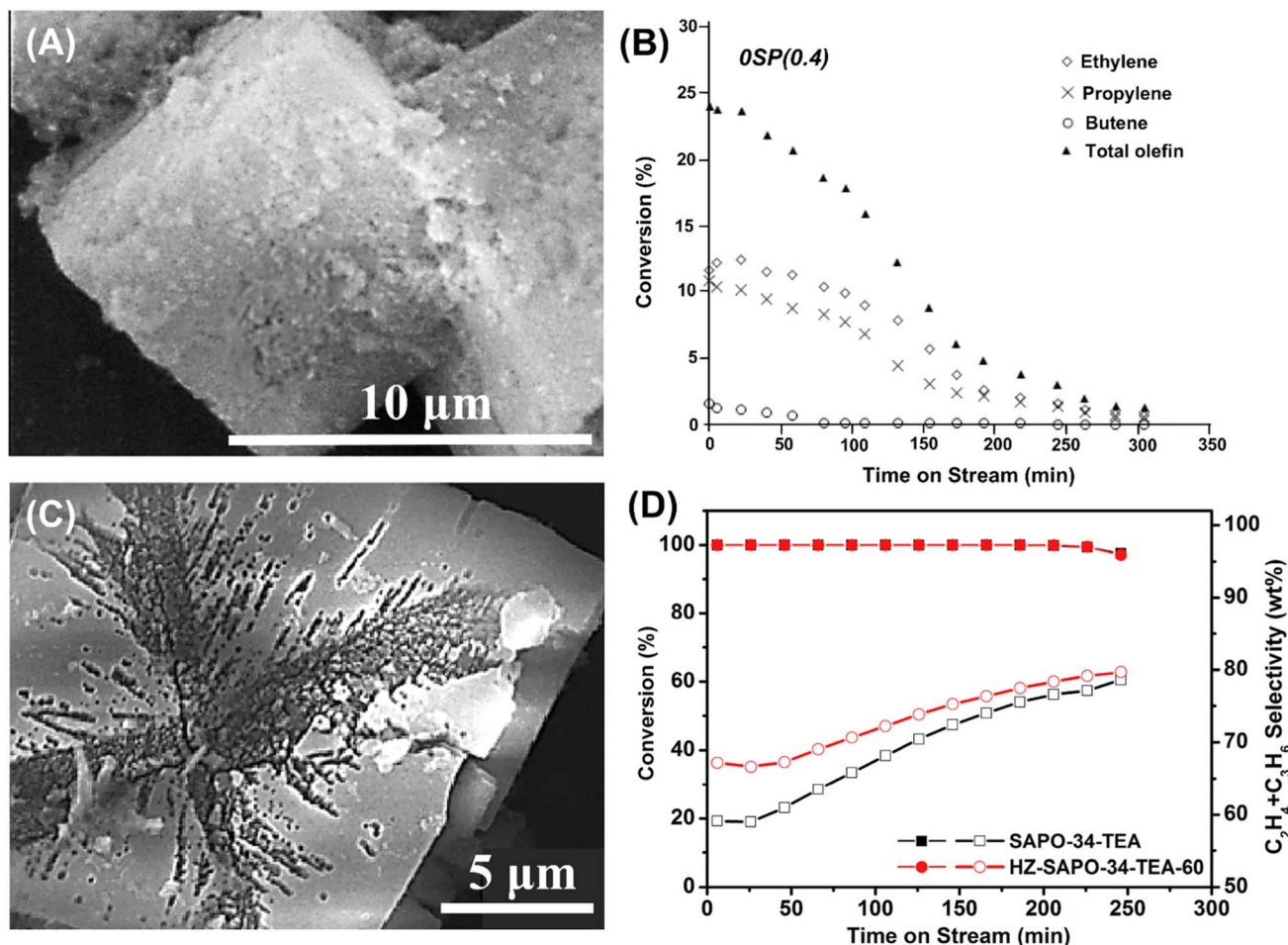


Fig. 36 SEM images of the catalysts (fixated on one particle) produced via application of different hard templating treatment techniques: (A) carbon nanotube as a secondary (hard) template; (B) yields of ethylene, propylene, butene, and total olefins (C_2-C_4) with time on stream over SAPO-34 catalyst OSP(0.4). Reproduced from ref. 335 with permission from Elsevier, Copyright 2016; (C) HF-NH₄F mixed aqueous solution as an acid-based etching agent; (D) methanol conversion (solid symbols) and selectivity of C₂H₄ plus C₃H₆ (hollow symbols) during the MTO reaction over the initial sample and the sample treated with a mixed HF-NH₄F solution for SAPO-34-TEA (WHSV = 2 g_{MeOH} g_{Cat}⁻¹ h⁻¹, T = 400 °C). Reproduced from ref. 218 with permission from Elsevier, Copyright 2016.

substitution), dry (incipient) and wet impregnation, as well as physical mixing.

The *in situ* metal incorporation approach typically involves adding a specific quantity of metal salts during the gel synthesis phase, prior to hydrothermal crystallization.^{231,233,241,344,346-349,351,352,354,355} This generally leads to the integration of metal atoms into the SAPO-34 framework. The impregnation method,^{239,344-346,350} on the other hand, involves the addition of metal solutions to already-prepared SAPO-34 catalyst. This results in metal atoms being largely deposited onto the catalyst surface (with only slight incorporation into the framework) *via* capillary force (dry approach) or diffusion (wet approach) from the metal salt solution. A minority of articles have also looked to a simple physical mixing/blending method for metal promotion.^{345,353,356} This typically comprises of the mixing of already-synthesized SAPO-34 sample with the metal salt, where these components are often mechanically ground together. Amongst these techniques, *in situ* metal incorporation is the most widespread method utilized by research groups for

metal promotion because of its superiority in uniform metal distribution (see Fig. 39(A) and (C)) and its capability of retaining SAPO-34's high surface area. Alternatively, impregnation usually leads to metal agglomerates (Fig. 39(B) and (D)) and SAPO pore blockage.³⁴⁴

Table 12 compares the application of different metals utilized for SAPO-34 promotion. The synthesis method involved, the type of metal used, the gel formula, along with the reaction conditions and activity performance for the promoted catalysts as well as the parent sample (to provide a fair comparison) has been presented. Generally, it has been claimed that metal promotion is able to both extend the catalyst lifetime and boost light olefins selectivity, relative to unmodified parent SAPO-34. With reference to the tabulated studies, there exists some clear trends with regards to the optimal type of metals utilized. For instance, several articles have demonstrated that non-transition metals (*i.e.*, rare earth metals such as La, Ce and Y, alkali metals, such as K, and alkaline earth metals, such as Ca) have displayed superior catalytic performance in relation to



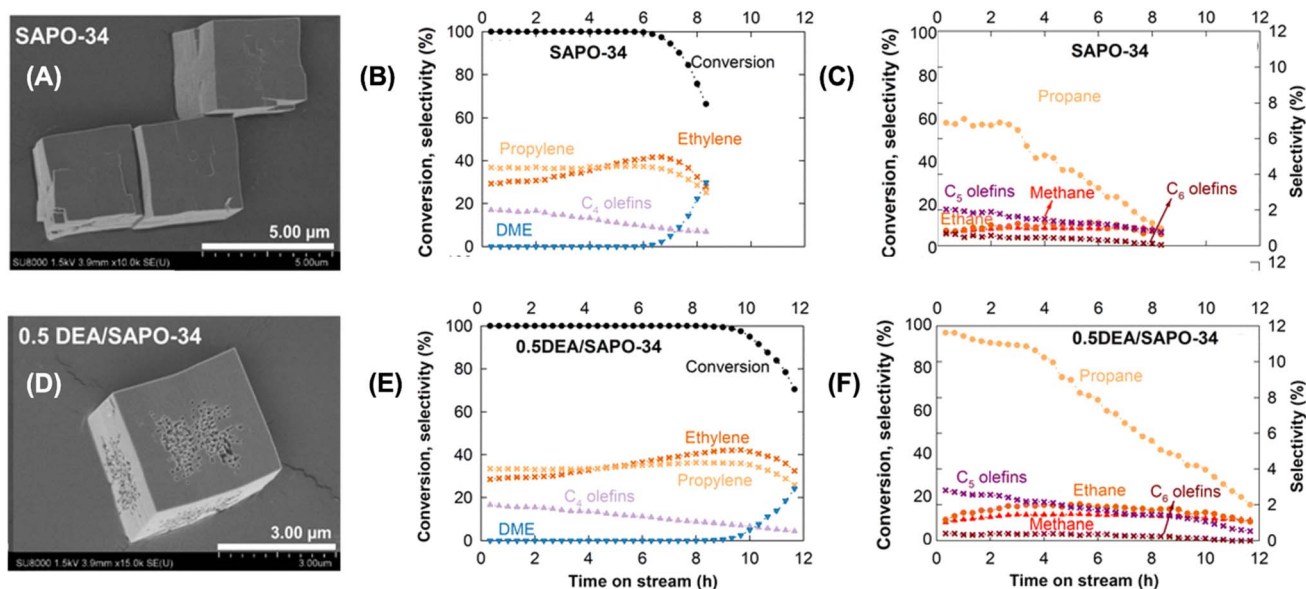


Fig. 37 (A) SEM image of the parent catalyst. (B) Methanol conversion and (C) products' selectivity versus time on stream for parent SAPO-34. (D) SEM image of the treated catalyst. (E) Methanol conversion and (F) selectivity versus time on stream for over 0.5 DEA/SAPO-34. Temperature 400 °C, atmospheric pressure, and at a WHSV of $1 \text{ g}_{\text{MeOH}} \text{ g}_{\text{Cat}}^{-1} \text{ h}^{-1}$. Reproduced from ref. 160 with permission from Royal Chemistry Society, Copyright 2023.

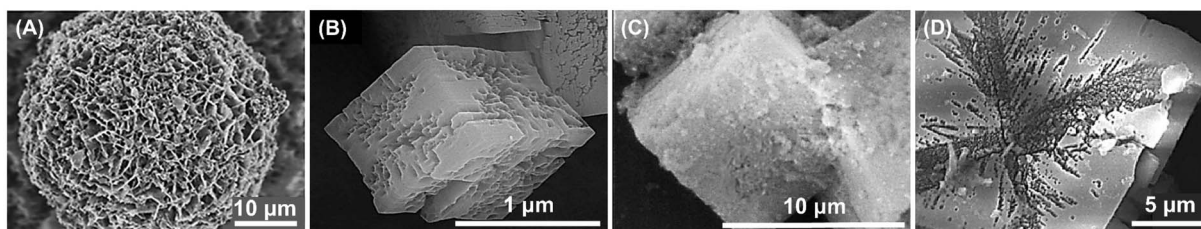


Fig. 38 SEM images of the catalysts (fixed on one particle) produced via application of the different pre- and post-treatment techniques: (A) TPOAC as a secondary (soft) template. Reproduced from ref. 272 with permission from Royal Society of Chemistry, Copyright 2015; (B) PEG as a secondary (soft) template. Reproduced from ref. 340 from Springer Nature, Copyright 2021; (C) carbon nanotube as a secondary (hard) template. Reproduced from ref. 335 with permission from Elsevier, Copyright 2016; (D) HF-NH₄F mixed aqueous solution as an acid-based etching agent. Reproduced from ref. 218 with permission from Elsevier, Copyright 2016.

the studied transition metals (*i.e.*, Fe, Co, Ni, Ag, Cu and W).^{231,345–347} Specifically, these non-transition metals have surpassed the other candidates in the domains of catalyst lifetime and total light olefins selectivity, as well as lower overall methane formation. Upon review, out of the wide range of metals studied, it has also been detected that certain metals were able to facilitate the selective formation of specific olefins over others. For instance, Ni-incorporated SAPO-34 demonstrated greater selectivity to ethylene as opposed to propylene.^{231,344,352} On the other hand, La-modified SAPO-34 displayed the contrary, favoring propylene production instead.^{231,345} However, it should be noted that the justifications for such selective formation of specific light olefins remain immature and are yet to be thoroughly explained.

A small portion of articles investigated the effect of metal loading on catalytic performance.^{241,350,351,354} A majority of the articles have tried a 0.05–0.08 Me/Al₂O₃ ratio in the initial

gel,^{231,346,348,355,357} while some groups have tested lower values (0.005–0.01),^{233,349,350} and some higher (around 0.1–0.12).^{351,353} Some studies^{350,351} have fixated on deciphering the optimum loading amount and have found that the metal content has a crucial effect on the promoted catalyst performance, but it should be noted that this optimum metal content cannot be generalized to all metals and could vary from one metal to another. However, it seems that exceptionally small loadings (relative to the other components in the gel) are required to achieve enhanced performance.³⁵⁰ Nevertheless, with such small loadings, one can question how much of a role the metal can truly play. It can even be argued as to whether the introduction of metals is truly the driving force behind any positive results seen, since it seems that the greater the loading, the poorer the performance.

Some articles have partially supported the idea of metal-promoted catalysts performing similarly to or even worse than



Table 11 Summary of different approaches of Hierarchical SAPO-34 synthesis and their effect on SAPO-34 properties and activity in the MTO reaction

Approach	Particle morphology	Activity	Reaction conditions	Note	Ref.
Soft template (TPOAC)	Spherical agglomeration of nanosheet-like crystals	2–4-fold increase in catalyst lifetime; comparable to ~10% improved light olefins selectivity (LOS)	WHSV = 3 g _{MeOH} g _{Cat} ⁻¹ h ⁻¹ , T = 450 °C, catalyst weight = 300 mg	Acts as mesopore-generating agent and silica source; relatively expensive	213 and 276
Soft template (PEG)	Intracrystalline micro-, meso-, and macro-porosity	2–6-fold increase in catalyst lifetime; comparable to ~10% improved LOS	WHSV = 2.3 g _{MeOH} g _{Cat} ⁻¹ h ⁻¹ , T = 425 °C, p = 1 atm, catalyst weight = 2 mg	Cost effective; variable Brønsted acid site concentration depending on molecular weight	338–342
Hard template (carbon nanoparticles)	Cave-like, unconnected pores	No significant improvement in lifetime or LOS	Not specified	Limited mass transfer due to the unconnected pores	275 and 277
Hard template (carbon nanotubes)	Interconnected mesoporous channels	Prolonged lifetime; slightly lower LOS	Not specified	Forms 3D multi-pore network; slight framework collapse may affect sieving	275 and 277
Post-treatment desilication (HF-NH ₄ F)	Hierarchical structure	Comparable to 25–30% worse lifetime; ~2% improved LOS	WHSV = 2 h ⁻¹ , T = 400 °C	Reduced Brønsted acid site strength; crystallinity depletion affects performance	218 and 332
Post-treatment desilication (TEAOH)	Hierarchical structure	2-Fold increase in lifetime; maintained LOS	WHSV = 1 g _{MeOH} g _{Cat} ⁻¹ h ⁻¹ , T = 400 °C, p = 1 atm	Increased acid site concentration; conflicts with expected desilication effects	160 and 333

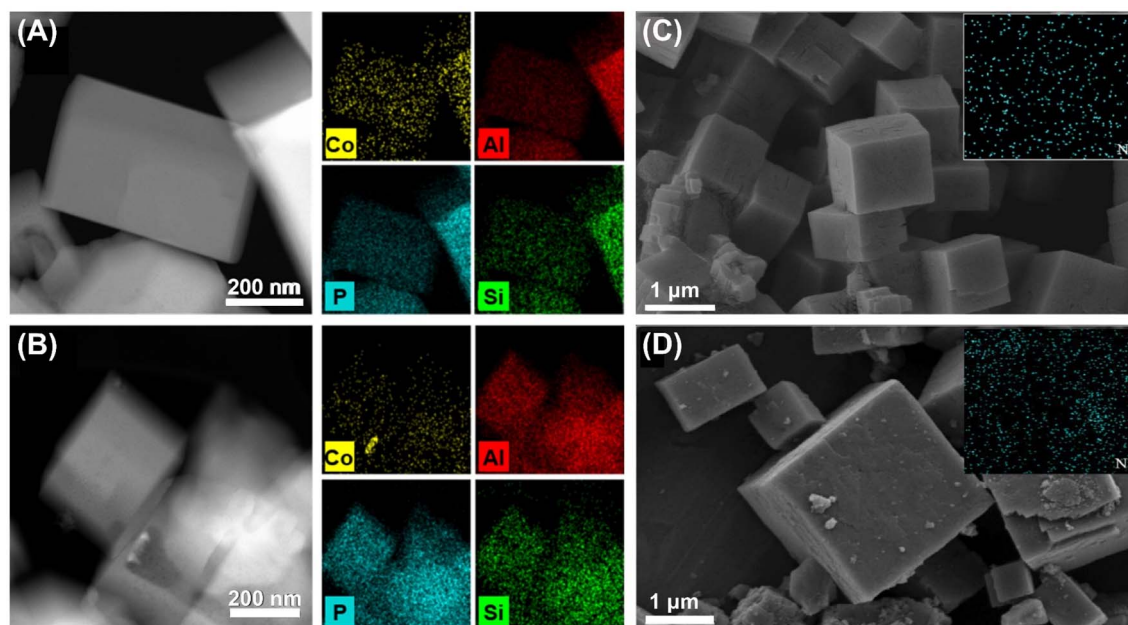


Fig. 39 (A) and (B) High-angle annular dark-field (HAADF) scanning TEM and energy-dispersive X-ray spectroscopy (EDS) elemental mapping images of (A) Co-incorporated SAPO-34 and (B) Co-impregnated SAPO-34. Reproduced from ref. 236 with permission from Elsevier, Copyright 2019. (C) and (D) SEM and EDS elemental mapping images of (C) Ni-incorporated SAPO-34 (using a template assisted approach) and (D) Ni-impregnated SAPO-34. Reproduced from ref. 239 from Springer Nature, Copyright 2021.

the parent SAPO-34. One study³⁴⁹ reported that the metal-modified catalysts possessed lifetimes shorter (for Co-SAPO-34) and similar (in the case of Ni-SAPO-34) to the parent catalyst, without considerable changes in light olefins selectivity. Another study³⁴⁶ explored the effect of Fe-incorporated SAPO-34

and found that the selectivity towards C₂ and C₃ olefins (under optimal conditions) was notably small due to substantial methane formation, particularly in comparison to the parent catalyst. Deterioration of catalytic performance by secondary metal addition has also been reported using the impregnation



Table 12 Summary of the key parameters and results from metal promoted SAPO-34 studies

Metal	Comment/ synthesis technique	Gel formula (molar) (Al ₂ O ₃ /P ₂ O ₅ /SiO ₂ / T(H ₂ O)/Me) ^a	Catalyst formula (molar) Al _a P _b Si _c M _d	Methanol processing ability ^d (g _{MeOH} g _{Cat} ⁻¹)	Average light olefins selectivity ^b (mol%)	Reaction conditions	Ref
—	Parent SAPO-34	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1TEAOH/1MOR/60H ₂ O	Al _{0.485} P _{0.405} Si _{0.108}	7.9	N/A	WHSV = 5.00 h ⁻¹ reaction T: 400 ° C MeOH: 19 mol% water: 81 mol%	344
—	Hierarchical SAPO-34	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1TEAOH/1MOR/60H ₂ O	Al _{0.474} P _{0.423} Si _{0.101}	10.4	N/A		
Co	Hierarchical + <i>in situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1TEAOH/1MOR/60H ₂ O/ 0.08Co	Al _{0.480} P _{0.426} Si _{0.079} Co _{0.014}	15.8	N/A		
	Hierarchical + impregnation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1TEAOH/1MOR/60H ₂ O	Al _{0.481} P _{0.409} Si _{0.093} Co _{0.016}	13.3	N/A		
Ni	Hierarchical + <i>in-situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1TEAOH/1MOR/60H ₂ O/ 0.08Ni	Al _{0.482} P _{0.421} Si _{0.087} Ni _{0.009}	14.2	N/A		
	Hierarchical + impregnation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1TEAOH/1MOR/60H ₂ O	Al _{0.489} P _{0.405} Si _{0.090} Ni _{0.015}	8.3	N/A		
—	Parent SAPO-34	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1.1 TEAOH/0.9 MOR/ 60H ₂ O	Al _{0.481} P _{0.407} Si _{0.112}	8.0	N/A	WHSV = 2.46 h ⁻¹ reaction T: 425 ° C MeOH: 19 mol% water: 81 mol%	231
Fe	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1.1TEAOH/0.9MOR/ 60H ₂ O/0.05Fe	Al _{0.467} P _{0.386} Si _{0.139} Fe _{0.008}	9.0	N/A		
Co	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1.1TEAOH/0.9MOR/ 60H ₂ O/0.05Co	Al _{0.468} P _{0.385} Si _{0.135} Co _{0.012}	11.0	N/A		
Ni	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1.1TEAOH/0.9MOR/ 60H ₂ O/0.05Ni	Al _{0.478} P _{0.381} Si _{0.138} Ni _{0.003}	10.0	N/A		
La	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1.1TEAOH/0.9MOR/ 60H ₂ O/0.05La	Al _{0.470} P _{0.378} Si _{0.137} La _{0.015}	15.0	N/A		
Ce	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.4SiO ₂ / 1.1TEAOH/0.9MOR/ 60H ₂ O/0.05Ce	Al _{0.472} P _{0.380} Si _{0.132} Ce _{0.016}	17.0	N/A		
—	Parent SAPO-34	N/A	Al _{0.448} P _{0.352} Si _{0.200}	4.9	81.0	WHSV = 2.10 h ⁻¹ reaction T: 450 ° C MeOH: 25 mol% water: 75 mol%	345
La	3 wt.% La loading, solid- state ion exchange	N/A	N/A	5.6	84.5		
	Liquid phase ion exchange	N/A	La _{0.002}	4.6	81.2		
Y	3 wt.% Y loading, solid- state ion exchange	N/A	N/A	5.8	85.0		
	Liquid phase ion exchange	N/A	Y _{0.003}	4.7	81.9		



Table 12 (Contd.)

Metal	Comment/ synthesis technique	Gel formula (molar) ($Al_2O_3/P_2O_5/SiO_2$ / $T/H_2O/Me$) ^a	Catalyst formula (molar) $Al_xP_ySi_zM_d$	Methanol processing ability ^d ($g_{MeOH} g_{Cat}^{-1}$)	Average light olefins selectivity ^b (mol%)	Reaction conditions	Ref
—	Parent SAPO-34	$1Al_2O_3/1P_2O_5/0.65SiO_2/$ 9DEA/110H ₂ O	$Al_{0.460}P_{0.364}Si_{0.176}^c$	12.0	85.0	WHSV = 6.00 h ⁻¹ reaction T: 450 ° C MeOH: 30 mol% water: 70 mol%	346
Fe	<i>In situ</i> metal incorporation	$1Al_2O_3/1P_2O_5/0.65SiO_2/$ 9DEA/110H ₂ O	$Al_{0.335}P_{0.325}Si_{0.114}Fe_{0.227}^c$	36.0	10.0		
K	Ion exchange -wet impregnation	N/A	N/A	60.0	90.0		
Ag	Ion exchange -wet impregnation	N/A	N/A	17.0	91.0		
—	Parent SAPO-34	$1Al_2O_3/1.15P_2O_5/$ $0.55SiO_2/2TEAOH/$ 110H ₂ O	$Al_{0.526}P_{0.357}Si_{0.117}^c$	7.3	40.3	WHSV = 4.85 h ⁻¹ reaction T: 350 ° C MeOH: 5 mol% helium: 95 mol%	347
Ca	<i>In situ</i> metal incorporation	$1Al_2O_3/1.15P_2O_5/$ $0.55SiO_2/2TEAOH/$ 110H ₂ O/0.007CaO _x	$Al_{0.544}P_{0.308}Si_{0.141}Ca_{0.007}^c$	9.7	58.7		
Cu	<i>In situ</i> metal incorporation	$1Al_2O_3/1.15P_2O_5/$ $0.55SiO_2/2TEAOH/$ 110H ₂ O/0.007CuO _x	$Al_{0.541}P_{0.341}Si_{0.116}Cu_{0.002}^c$	8.9	49.0		
W	<i>In situ</i> metal incorporation	$1Al_2O_3/1.15P_2O_5/$ $0.55SiO_2/2TEAOH/$ 110H ₂ O/0.007WO _x	$Al_{0.537}P_{0.326}Si_{0.130}W_{0.007}^c$	5.7	40.9		
—	Parent SAPO-34	$1Al_2O_3/1P_2O_5/0.65SiO_2/$ 1TEAOH/1MOR/60H ₂ O	$Al_{0.58}P_{0.29}Si_{0.12}^c$	N/A	1.8 ^d	WHSV = 8.70 h ⁻¹ reaction T: 400 ° C MeOH: 19 mol% water: 81 mol%	348
Ni	<i>In situ</i> metal incorporation	$1Al_2O_3/1P_2O_5/0.65SiO_2/$ 1TEAOH/1MOR/60H ₂ O/ 0.05NiO	$Al_{0.50}P_{0.31}Si_{0.19}Ni_{0.005}^c$	N/A	7.8 ^d		
Mg	<i>In situ</i> metal incorporation	$1Al_2O_3/1P_2O_5/0.65SiO_2/$ 1TEAOH/1MOR/60H ₂ O/ 0.05MgO	$Al_{0.36}P_{0.44}Si_{0.20}Mg_{0.006}^c$	N/A	4.0 ^d		
—	Parent SAPO-34	$1Al_2O_3/1P_2O_5/0.30SiO_2/$ 2.0TEAOH/50H ₂ O	N/A	15.8	85.0	WHSV = 0.48 h ⁻¹ reaction T: 400 ° C MeOH: 5 mol% nitrogen: 95 mol%	349
Co	<i>In situ</i> metal incorporation (method 1)	$1Al_2O_3/1P_2O_5/0.30SiO_2/$ 2.0TEAOH/50H ₂ O/ 0.007CoO	$Co_{0.003}$	11.0	87.0		
Mn	<i>In situ</i> metal incorporation (method 1)	$1Al_2O_3/1P_2O_5/0.30SiO_2/$ 2.0TEAOH/50H ₂ O/ 0.005MnO	$Mn_{0.003}$	25.0	86.0		
Ni	<i>In situ</i> metal incorporation (method 1)	$1Al_2O_3/1P_2O_5/0.30SiO_2/$ 2.0TEAOH/50H ₂ O/ 0.007NiO	Ni incorporation ~0	9.6	85.0		
—	<i>In situ</i> metal incorporation (method 2)	$1Al_2O_3/1P_2O_5/0.10SiO_2/$ 2.0TEAOH/50H ₂ O/0.01 NiO	$Ni_{0.003}$	17.3	85.0		



Table 12 (Contd.)

Metal	Comment/ synthesis technique	Gel formula (molar) ($Al_2O_3/P_2O_5/SiO_2/$ $T/H_2O/Me$) ^a	Catalyst formula (molar) $Al_xP_ySi_zM_d$	Methanol processing ability ^d ($g_{MeOH} g_{Cat}^{-1}$)	Average light olefins selectivity ^b (mol%)	Reaction conditions	Ref
—	Parent SAPO-34	$1Al_2O_3/1P_2O_5/0.65SiO_2/$ $2TEAOH/70H_2O$	$Al_{0.381}P_{0.477}Si_{0.142}^c$	119.1	90.0	WHSV = $14.3 h^{-1}$ reaction T: $400^\circ C$ C MeOH: 30 mol% water: 70 mol%	350
Ce	Impregnation (optimal: 0.5 wt.% CeO ₂ loading)	$1Al_2O_3/1P_2O_5/0.65SiO_2/$ $2TEAOH/70H_2O/$ $0.012CeO_2$	$Al_{0.380}P_{0.480}Si_{0.137}^c$ $Ce_{0.002}^c$	214.5	91.0		
—	Parent SAPO-34	$1Al_2O_3/1P_2O_5/0.25SiO_2/$ $0.7TEAOH/1.1DEA/$ $50H_2O$	$Al_{0.501}P_{0.425}Si_{0.074}^c$	N/A	82.0	WHSV = $3.5 h^{-1}$ reaction T: $450^\circ C$ MeOH: 27 mol% water: 73 mol%	351
Ge	<i>In situ</i> metal incorporation (optimal: 0.1 mol GeO ₂ loading)	$1Al_2O_3/1P_2O_5/0.15SiO_2/$ $0.7TEAOH/1.1DEA/$ $50H_2O/0.1GeO_2$	$Al_{0.503}P_{0.436}Si_{0.048}^c$ $Ge_{0.013}^c$	N/A	84.0		
—	Parent SAPO-34	$1Al_2O_3/1P_2O_5/0.45SiO_2/$ $2TEAOH/70H_2O$	N/A	15.3	80.1	WHSV = $4.0 h^{-1}$ reaction T: $450^\circ C$ MeOH: 36 mol% water: 64 mol%	233
AgPW	<i>In situ</i> metal incorporation	$1Al_2O_3/1P_2O_5/0.45SiO_2/$ $2TEAOH/70H_2O/$ $0.010AgPW$	AgPW _{0.009} ^f	32.7	92.3		
—	Parent SAPO-34	$1Al_2O_3/1P_2O_5/0.35SiO_2/$ $1TEAOH/50H_2O$	$Al_{0.503}P_{0.347}Si_{0.151}$	7.0 ^e	N/A	WHSV = $1.00 h^{-1}$ reaction T: $425^\circ C$ C MeOH: 15 mol% nitrogen: 85 mol%	352
Fe	<i>In situ</i> metal incorporation	$1Al_2O_3/1P_2O_5/0.35SiO_2/$ $2TEAOH/50H_2O/$ $0.075FeO$	$Al_{0.509}P_{0.351}Si_{0.137}^c$ $Fe_{0.002}$	4.5 ^e	N/A		
Co	<i>In situ</i> metal incorporation	$1Al_2O_3/1P_2O_5/0.35SiO_2/$ $2TEAOH/50H_2O/$ $0.075CoO$	$Al_{0.506}P_{0.329}Si_{0.162}^c$ $Co_{0.004}$	8.0 ^e	N/A		
Ni	<i>In situ</i> metal incorporation	$1Al_2O_3/1P_2O_5/0.35SiO_2/$ $2TEAOH/50H_2O/$ $0.075NiO$	$Al_{0.494}P_{0.361}Si_{0.143}^c$ $Ni_{0.001}$	5.0 ^e	N/A		
—	Hierarchical SAPO-34	$1Al_2O_3/0.8P_2O_5/0.45SiO_2/$ $2.0TEAOH/50H_2O$	N/A	16.7	82.3	WHSV = $10.0 h^{-1}$ reaction T: $500^\circ C$ C MeOH: 85 mol% nitrogen: 15 mol%	353
ZnO	Physical mixing (interpellet)	$1Al_2O_3/0.8P_2O_5/0.45SiO_2/$ $2.0TEAOH/50H_2O/$ $0.11ZnO$	N/A	15.0	82.4		
—	Physical mixing (intrapellet- AL <i>i.e.</i> , not calcined mixture)	$1Al_2O_3/0.8P_2O_5/0.45SiO_2/$ $2.0TEAOH/50H_2O/$ $0.11ZnO$	N/A	9.2	84.0		
—	Physical mixing (intrapellet-HY <i>i.e.</i> , calcined mixture)	$1Al_2O_3/0.8P_2O_5/0.45SiO_2/$ $2.0TEAOH/50H_2O/$ $0.11ZnO$	N/A	8.4	83.7		



Table 12 (Contd.)

Metal	Comment/ synthesis technique	Gel formula (molar) (Al ₂ O ₃ /P ₂ O ₅ /SiO ₂ / T/H ₂ O/Me) ^a	Catalyst formula (molar) Al _a P _b Si _c M _d	Methanol processing ability ^d (g _{MeOH} g _{Cat} ⁻¹)	Average light olefins selectivity ^b (mol%)	Reaction conditions	Ref
—	Parent H-SAPO-34 L (sample with relatively large crystals)	1Al ₂ O ₃ /1P ₂ O ₅ /0.3SiO ₂ / 2.0TEAOH/52H ₂ O	Al _{0.32} P _{0.34} Si _{0.14} ^c	11.7	65.0	WHSV = 10.0 h ⁻¹ reaction T: 400 °C C MeOH: 77 mol% nitrogen: 23 mol%	354
Ti	<i>In situ</i> metal incorporation (optimal: TiO ₂ loading in 0.006 relative molar ratio)	1Al ₂ O ₃ /1P ₂ O ₅ /0.3SiO ₂ / 2.0TEAOH/52H ₂ O/ 0.006TiO ₂	Al _{0.56} P _{0.31} Si _{0.13} Ti _{0.00034} ^c	30.0	78.0		
—	Parent SAPO-34	N/A	N/A	9.5 ^e	80.5	WHSV = 2.0 h ⁻¹ reaction T: 400 °C MeOH: 35.7 mol% nitrogen: 64.3 mol%	239
Co	Wet impregnation	N/A	Co _{0.0176}	7.0 ^e	76.7		
—	Template-assisted method	N/A	Co _{0.0024}	10.0 ^e	81.3		
Ni	Wet impregnation	N/A	Ni _{0.0175}	7.5 ^e	78.2		
—	Template-assisted method	N/A	Ni _{0.0028}	11.0 ^e	80.4		
Zn	Wet impregnation	N/A	Zn _{0.0178}	6.5 ^e	82.5		
—	Template-assisted method	N/A	Zn _{0.0050}	9.5 ^e	77.5		
Ni	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.6SiO ₂ / 1DEA/70H ₂ O/0.05NiO	Al _{0.470} P _{0.380} Si _{0.130} Ni _{0.02} ^c	2.5	88.0	WHSV = 0.4 h ⁻¹ reaction T: 400 °C MeOH: 30 mol% water: 70 mol%	355
Mn	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.6SiO ₂ / 1DEA/70H ₂ O/0.05NiO	Al _{0.480} P _{0.390} Si _{0.120} Mn _{0.01} ^c	4.2	94.0		
—	<i>In situ</i> metal incorporation (bimetallic)	1DEA/70H ₂ O/0.05MnO	Al _{0.380} P _{0.420} Si _{0.170} Ni _{0.02} Mn _{0.01} ^c	2.5	85.0		
—	Parent SAPO-34	N/A	Al _{0.411} P _{0.485} Si _{0.105}	8.3	60.0	WHSV = 1.0 h ⁻¹ reaction T: 350 °C MeOH: 83.5 mol% water: 16.5 mol%	241
Co	<i>In situ</i> metal incorporation, milling and recrystallization (optimal: 0.003 mol in relative molar ratio)	N/A	Al _{0.472} P _{0.415} Si _{0.086} Co _{0.027}	11.8	65.0		
—	Parent SAPO-34	1Al ₂ O ₃ /0.8P ₂ O ₅ /0.6SiO ₂ / 2.5MOR/80H ₂ O	Al _{0.43} P _{0.43} Si _{0.14}	12.3	79.0	WHSV = 2.0 h ⁻¹ reaction T: 380 °C MeOH: 50 mol% water: 50 mol%	356
t-ZrO ₂ (tetragonal- phase zirconia)	Physical blending method Hydro-thermal surface coating modification method	N/A	Al _{0.41} P _{0.41} Si _{0.18}	28.4	82.0		
—	Parent SAPO-34	N/A	Al _{0.42} P _{0.40} Si _{0.18}	37.3	82.0		
Zn	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.6SiO ₂ / 3TEA/50H ₂ O	Al _{0.470} P _{0.351} Si _{0.178}	5.5 ^e	77.0	WHSV = 2.0 h ⁻¹ reaction T: 425 °C MeOH: 50 mol% water: 50 mol%	357
—	<i>In situ</i> metal incorporation	1Al ₂ O ₃ /1P ₂ O ₅ /0.6SiO ₂ / 3TEA/50H ₂ O/ 0.04Zn(NO ₃) ₂ ·6H ₂ O	Al _{0.448} P _{0.339} Si _{0.198} Zn _{0.015}	8.5 ^e	80.0		

^a Methanol processing ability = $\text{WHSV}(\text{g}_{\text{MeOH}} \text{g}_{\text{Cat}}^{-1} \text{h}^{-1}) \times (\text{time taken to go to 80\% MeOH conversion})$ (h) ^b Average light olefin selectivity = amount of methanol

converted to C₂ $\frac{\text{amount of methanol converted on average (mol)}}{\text{C}_2 \text{ \& C}_3 \text{ on average (mol)}} \times 100\%$ ^c The surface composition was only measured/provided (*i.e.*, by XPS or SEM-EDS). ^d An alternative definition for light olefin selectivity was utilized: specified as the ratio of the sum of produced ethylene and propylene to undesirable products (others hydrocarbon). ^e For ref. 352, 239 and 357 none of the catalysts reached 80% conversion. The methanol processing ability was instead computed using the time taken to reach 95% conversion. ^f The numerical value presented depicts the molar ratio of AgPW to Al₂O₃ (*i.e.*, not with respect to all the precursors).



technique. Co-, Ni- and Zn-impregnated SAPO-34 catalysts in one study²³⁹ demonstrated significantly shortened lifetime relative to the parent catalyst with worse or comparable LOS. This inferior performance due to metal addition has also been observed in other circumstances, such as La- and Y- impregnated³⁴⁵ and physically-blended ZnO modified samples.³⁵³ A comprehensive study conducted by Ghavipour *et al.*,³⁶¹ in which 20 different metal promoted catalysts were tested, found that the majority of the catalysts suffered from a loss in crystallinity, which ultimately resulted in inferior catalytic performance.

Overall, by observing the results of the conducted research in metal modification of SAPO-34, clear contradictions can be spotted. Even while employing similar metals, opposing catalytic performances have been reported. For instance, in the case of Ni-incorporated SAPO-34, there have been reports of significantly improved,³⁴⁸ unchanged,³⁴⁹ and slightly worsened²³⁹ light olefins selectivity (relative to the parent catalyst). This demands a comprehensive discussion on the justifications provided by these articles in support of superior or inferior catalytic activity of the promoted catalysts. In this regard, some articles have claimed that the addition of secondary metal precursors enhances the nucleation, resulting in smaller crystals that have better mass transfer characteristics and can last longer within the reaction.³⁵¹ Alternatively, introduction of new mesopores into the structure due to secondary metal addition has also been claimed to be one of the underlying factors for superior catalytic performance,³⁴⁷ although the mechanism for the formation of these mesopores has not been clarified. One study³⁶⁰ has applied a distinct metal modification method termed hydrothermal surface coating, which resulted in elimination of the external active sites by tetragonal zirconia crystals addition. Subsequently, the group claimed that this may reduce the build-up of coke at the entry of SAPO-34 pores. In turn, the lack of pore blockage would lead to a greater diffusion of reactant methanol

molecules over time on stream, which has resulted in doubled catalyst lifetime. However, this statement does not seem to be completely valid, since the acidic sites that lie beneath the exterior neutralized layer are still active and can block the pores by coke build-up. Apart from that, the majority of metal promoted articles^{231,344,346,348,350} stated that lowering the quantity of Brønsted acidic sites *via* substitution of Si (as opposed to the other SAPO elements) by the added metal^{231,344,346–348} hinders the evolution of coke precursor formation reactions, while retaining an acidity level sufficient for the dehydration steps to light olefins. Thus, overall, this results in the decreased production of alkanes (specifically methane) and multi-ring aromatic species (the precursors of coke), leading to a delay in pore blockage in which the diffusion of species can continue to transpire (*i.e.*, prolonged catalytic activity).

Despite the consensus amongst these articles in support of this rationale, there still lies some ambiguities that need to be addressed. Firstly, these articles are quick to suggest that the decrease in acidity acts as the primary justification, when this alteration can be achieved by a more facile approach: reducing the silica content in the starting gel. This should in turn question the need to use secondary metals entirely, since they seem to increase costs unnecessarily when a more economic approach may be utilized instead.

Secondly, it should be noted that research groups have previously investigated the effect of silica content alterations on catalytic performance.^{131,296} They arrived at the conclusion that the optimal SiO₂/Al₂O₃ ratio, concerning catalyst lifetime, lies between 0.2 and 0.3. This silica content range is generally adopted by the aforementioned articles in the preparation of their parent catalyst. Hence, metal atom promotion (which causes acidity and silica content to fall below optimum levels) cannot appropriately explain why improved catalytic performance is achieved.

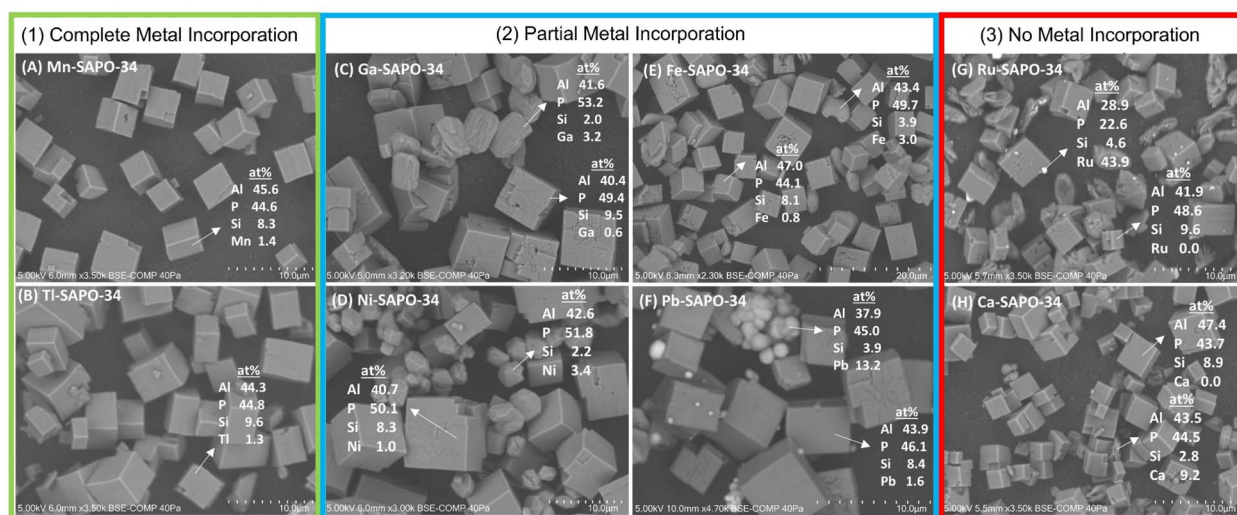


Fig. 40 SEM with SEM-EDX results overlaid with respect to different secondary metal-promoted SAPO-34 catalysts. The results have been categorized as follows: complete metal incorporation consisting of (A) Mn-SAPO-34 and (B) Ti-SAPO-34; partial metal incorporation consisting of (C) Ga-SAPO-34, (D) Ni-SAPO-34, (E) Fe-SAPO-34, and (F) Pb-SAPO-34; no incorporation consisting of (G) Ru-SAPO-34 and (H) Ca-SAPO-34. Reproduced from ref. 361 from John Wiley and Sons, Copyright 2024.



Finally, metal modification usually results in a worsening of physicochemical properties, such as a loss in both surface area and crystallinity; in turn, this finding further contradicts the idea that metal promotion can yield enhanced results. Nonetheless, assuming the legitimacy that improved performance may be achieved *via* secondary metal introduction, a fundamental justification for this may still need to be uncovered. It is possible that the promotion of metal with regards to the SAPO framework may in some way affect Brønsted acidic sites and/or influence the mechanistic route taken (by altering the activation energy barriers of elementary steps). There may even lie the possibility of metal atoms acting as new active sites themselves in addition to Brønsted acidic sites. Similarly, one article³⁵³ observed intermediate formaldehyde formation due to ZnO promotion and hypothesized that formaldehyde is able to facilitate ethylene production and simultaneously convert active hydrocarbon pool species into inactive carbon deposits. However, none of these articles, including the previously mentioned one, examined how the secondary metals could participate in the reaction mechanism. Although, utilizing SEM-EDX, one study³⁶¹ was able to separate different secondary metal modified catalysts into specific categories (see Fig. 40) *i.e.*, those that illustrated complete metal incorporation, partial metal incorporation, and no incorporation whatsoever. Such analysis may be useful for future work in determining if some metals that do incorporate into the crystalline framework play a role (a positive or negative one) in the reaction mechanistic scheme for light olefin production, or simply play no significant role in the case when incorporation did not occur (*i.e.*, when metals emerged as an amorphous phase instead). Hence, the role of secondary metals in the mechanistic scheme for the MTO reaction remains an area open to further exploration. For example, a recent study employed molecular dynamics (MD) simulations to provide mechanistic insights into interatomic interactions associated with coke suppression and enhanced olefin selectivity in mixed metal oxide-modified SAPO-34 catalysts.³⁶² However, to gain a more comprehensive understanding, complementary approaches such as density functional theory (DFT) calculations or other more exhaustive characterization techniques are still needed.

In summary, research groups have demonstrated that metal modification, considering the method of promotion, the metal loading, and the type of metal, can pose completely different effects on SAPO-34 catalytic performance (which can range from worsening, improving or unsubstantial effects). Nonetheless, the explanation as to why specific metals perform better than others still remain relatively immature.

4.7 Shaping of SAPO-34 catalyst

Since SAPO-34 suffers from rapid deactivation by inevitable coke generation, the only practical industrial-scale solution is the employment of a fluidized bed reactor accompanied by continuous regeneration of spent catalyst (see MTO process Section 2). That being said, catalyst particles entrained in the gas phase must be collected using multi-stage cyclones to then be directed to the regenerator. Hence, in order to ensure that

this catalyst-collecting procedure using cyclones is both feasible and efficient, the SAPO-34 catalyst should be present in the form of agglomerates of cubic crystals sized generally between 50 and 100 micron.³⁶³ Since the conventional (lab-scale) hydrothermal method generates particles in the size of a few micron, the larger desired SAPO-34 particle size for fluidized-bed application is traditionally achieved through the employment of an adhesive agent (binder) such as silica or alumina, followed by spray drying of the catalyst. However, the issue with this customary method is that the binder utilized has the capability to alter the physicochemical properties of the catalyst (potentially worsening catalytic performance), block catalyst pores, substantially reduce the total catalyst surface area, as well as make catalyst particles in the core of the agglomerates inaccessible to reactant species. Hence, groups have begun to investigate an alternate catalyst-shaping strategy by mounting SAPO-34 crystals over pre-synthesized micro-spherical supports made of alumina, silica, or aluminosilicates (*i.e.*, Kaolin, as seen in Fig. 41(A)).^{364–368} To achieve this, the microspheres are placed within the initial catalyst gel prior to the hydrothermal (*i.e.*, SAPO crystallization) phase, followed by all other conventional steps. The benefits of this technique include the fact that firstly, microspheres are inexpensive, and secondly, catalyst mounting on the outer surface of microspheres ensures that no part of the precious catalyst is wasted (unlike the binder-spray drying approach). Although this acts as a suitable alternate method, it still faces a crucial problem of not being able to achieve thorough adherence and high loading of SAPO particles onto microspheres. For instance, when α -Al₂O₃ was used as the microsphere support, the SAPO-34 formation was even on the outer surface but the content was low (6.5 wt%).³⁶⁶ In another case, the employment of silica microspheres resulted in an uneven distribution of SAPO-34 crystals (see Fig. 41(B)).³⁶⁶ This depicts the importance of attempting to choose a suitable support material and corresponding synthesis procedure to achieve successful catalyst mounting. Furthermore, to deal with these previously mentioned issues, one approach has involved the pre-treatment of microspheres. In one study, Kaolin microspheres were treated with NaOH aqueous solution of increasing concentration to increase its surface roughness, in pursuit of enhancing SAPO-34 loading on the external surface (see Fig. 41(C)–(F)).³⁶⁴ Another tactic involves supplying the silica and alumina required for SAPO-34 synthesis from the support itself (*i.e.*, *in situ* SAPO/Kaolin catalyst) as a means for successful SAPO-34 crystal adhesion, which resulted in up to 22 wt% SAPO-34 loading.³⁶⁵ Another group³⁶⁸ who grew SAPO-34 crystals on γ -Al₂O₃ microspheres introduced a new microwave-assisted synthesis approach, and were thus able to load up to even higher relative quantities, 31 and 44 wt%, when mounting SAPO-34 directed with TEOH and TEA templates, respectively. However, it is to be noted that procedures to introduce better adhesion between the microsphere support and catalyst still remains an area of study to be developed further. In terms of catalytic activity, it has been generally observed that SAPO-34 over microspheres are better performing relative to spray-dried samples (which deactivate much more rapidly) due to a more developed mesoporous structure.^{365,366,368} Relative to the



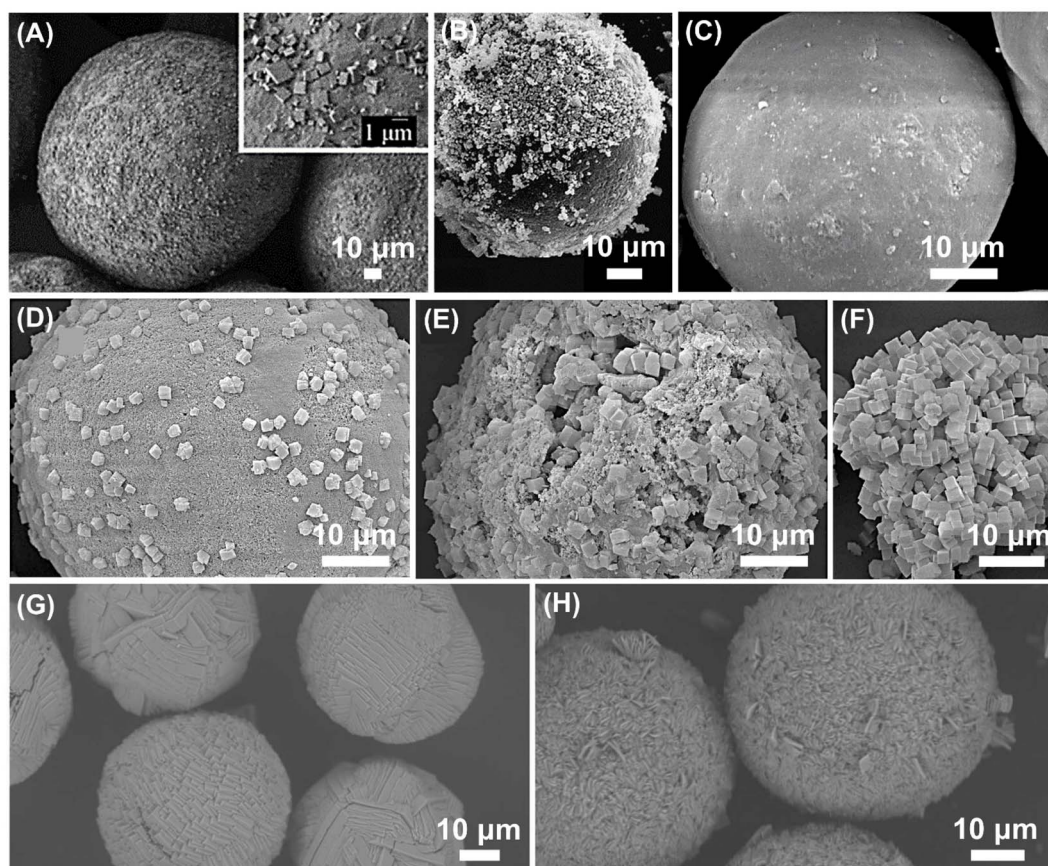


Fig. 41 SEM images depicting the morphology of SAPO-34 mounted on different microspheres: (A) SAPO-34/kaolin microsphere reproduced from ref. 365 with permission from Elsevier, Copyright 2010; (B) SAPO-34/silica microsphere reproduced from ref. 366 with permission from Elsevier, Copyright 2008. (C–F) SAPO-34 on calcined kaolin microspheres treated with NaOH solutions of (C) 2, (D) 4, (E) 10, (F) 14 wt% concentrations reproduced from ref. 364 with permission from American Chemical Society, Copyright 2011. (G) TEAOH-directed SAPO-34 crystals on γ - Al_2O_3 microspheres. (H) TEA-directed SAPO-34 crystals on γ - Al_2O_3 microspheres. Reproduced from ref. 368 with permission from Elsevier, Copyright 2024.

parent SAPO-34 catalyst, one group³⁶⁴ reported 9 times longer activity over their SAPO/Kaolin sample (treated with 4 wt% NaOH solution—as seen in Fig. 41(E)), but in the absence of a distinct and well-established justification. It could be argued that, in the best-case scenario (*i.e.*, a successful mounting of SAPO-34, with unaltered physicochemical properties, over the support), SAPO/microsphere catalyst should exhibit similar to slightly enhanced activity relative to the conventionally prepared SAPO-34 catalyst.³⁶⁸ Overall, Table 13 has been provided to summarize the different SAPO-34 shaping methods and their corresponding effects on structure, characteristics and MTO activity.

4.8 Catalyst design from an industrial perspective

The MTO process has been successfully commercialized in several large-scale plants around the world. For instance, Dalian Institute of Chemical Physics (DICP) and its partners developed the DMTO (Dalian MTO) technology with capacities exceeding 600 kt of olefins per year.^{369,370} UOP and Total also developed the UOP/Hydro MTO process, operating with a fluidized-bed reactor and regeneration system tailored for the SAPO-34 catalyst.

These commercial efforts demonstrate the maturity of the technology and emphasize the need for robust, scalable, and cost-effective catalyst designs to meet the demands of continuous high-throughput operations. Yet, much effort on the lab scale is still ongoing for the catalysts development.

The innovation in catalyst development should be translated into an industrial application; however, it requires a comprehensive evaluation of several factors beyond the catalytic activity. The lab-scale studies focus on optimizing SAPO-34 catalysts for activity and selectivity under controlled conditions, while industrial viability critically depends on the economic feasibility of raw materials, catalyst synthesis routes, mechanical robustness under reactor conditions, long-term regenerability, and alignment with sustainable and green chemistry principles. The different synthesis routes should be further examined from an industrial perspective. This should include, for instance, the green synthesis route, the mechanical and chemical stability of hierarchical and metal-promoted SAPO-34 catalysts, and trade-offs between activity and likelihood of scaling up the catalysts. The integration of these perspectives would assist in designing catalysts that are not only



Table 13 Summary of SAPO-34 shaping methods: structure, characteristics and MTO activity

Shaping method	Structure description	Characteristics	Activity	Additional remarks	Ref.
Spray-dried SAPO-34	Agglomerates of cubic crystals between 50 and 100 μm	Binder (silica or alumina) used to form the agglomerates of cubic crystals	Rapid deactivation; lower activity compared to microsphere-supported SAPO-34	Binder may block pores, reduce surface area, and limit core accessibility	363
Parent SAPO-34	Cubic crystals in the size of a few μm	Conventional hydrothermal synthesis product	Baseline activity; deactivates faster than microsphere-supported catalysts	Limited by small particle size for fluidized bed applications	363
SAPO-34 on $\alpha\text{-Al}_2\text{O}_3$	SAPO-34 crystals mounted on 50–100 μm microspheres	Even SAPO-34 distribution, low loading (6.5 wt%)	Not specified; expected similar to slightly enhanced activity vs. parent	Low loading limits catalytic efficiency	366
SAPO-34 on silica microspheres	SAPO-34 crystals mounted on 50–100 μm microspheres	Uneven SAPO-34 distribution	Not specified; expected similar to slightly enhanced activity vs. parent	Uneven distribution reduces effectiveness	366
SAPO-34 on Kaolin microspheres (NaOH-treated)	SAPO-34 crystals mounted on 50–100 μm microspheres	Enhanced loading due to NaOH-treated surface roughness	9 times longer activity than parent SAPO-34	NaOH treatment (4 wt%) improves adhesion; developed mesoporous structure	364 and 366
SAPO-34 on Kaolin microspheres (<i>in situ</i>)	SAPO-34 crystals mounted on 50–100 μm microspheres	<i>In situ</i> synthesis, up to 22 wt% SAPO-34 loading	Better performance than spray-dried	Supplied silica/alumina from support; improved adhesion	365
SAPO-34 on $\gamma\text{-Al}_2\text{O}_3$ microspheres (microwave-assisted)	SAPO-34 crystals mounted on 50–100 μm microspheres	High loading (31–44 wt%) with TEOH/TEA templates	Similar to slightly enhanced activity vs. parent; better than spray dried	Microwave-assisted synthesis enhances loading; mesoporous structure	368

highly active but also scalable, durable, and align with sustainability concerns, which facilitate bridging the gap between academic research and practical industrial deployment.

A major challenge in commercializing SAPO-34 is related to the catalyst production costs. The traditional hydrothermal synthesis, which we discussed earlier in detail, is widely adopted in industry; however, it utilizes the expensive templates, TEOH, which significantly increases catalyst cost. Fortunately, the mixed template method could mitigate this by combining cheaper amines (*e.g.*, MOR and DEA) with minimal TEOH. This has been studied and has effectively reduced template consumption and associated costs while still producing nanocrystalline and high-performance SAPO-34.^{371,372} Moreover, the expensive TEOH is reduced by introducing a pre-synthesized SAPO-34, known as seeds-assisted crystallization, into the synthesis gel to provide nucleation centers, allowing less OSDA to be used while still obtaining pure SAPO-34.^{373,374} Seeded syntheses not only reduce template cost but also accelerate crystallization and can yield smaller crystals with improved MTO performance. Therefore, the mixed-template and seed-assisted methods are cost-effective, which reduces the scalability cost.

The dry gel method is an eco-friendly method for synthesizing SAPO-34 in the absence of solvent with a minimal amount of water. This leads to high concentrations of ingredients, including the OSDA, which promotes rapid nucleation and the formation of smaller crystals.³⁷⁵ The dry gel method produced nano SAPO-34 crystals using the TEOH template,

which are significantly smaller than those from traditional hydrothermal methods.³⁷⁶ Similarly, the combination of TEOH with MOR yielded SAPO-34 nanoparticles (under 500 nm) in just 6 hours, much faster than the 1–2 days typically required.³⁷⁷ Moreover, the dry gel method enhances crystallization yield, which minimizes unconverted “waste gel,” reduces wastewater generation, and aligns with green chemistry principles by lowering the use of water and other ingredients, including OSDA.³⁷⁸ However, the industrial scalability remains a challenge to ensure uniform heating and hydration of a large amount of dry gel.

The ionothermal synthesis utilizes ionic liquids as solvents and templates, which eliminating the need for OSDA. For instance, SAPO-34 can be synthesized using 1-ethyl-3-methylimidazolium bromide (EMIM)Br in the absence of OSDA,^{379,380} leading to green chemistry advantages. This method reduces costs by avoiding expensive templates and eliminating the high-temperature calcination step needed to remove organics, thus saving energy and reducing CO₂ and NO_x emissions.³⁷⁴ We refer to this method as OSDA-Free, however, it worth mentioning that this method is different than the well-known OSD-Free CHA synthesis.^{381–384} Although ionic liquids are still not cheap (but less expensive than OSDA), however, they are recyclable, supporting waste minimization, with biodegradable solvents like choline chloride being explored for SAPO synthesis. This process generates minimal wastewater that aligns with green chemistry principles. However, challenges remain, including the high risk of impurities, which may necessitate mild processing steps.³⁸⁵ Scalability is another



concern, as handling viscous ionic liquids and crystal separation may require specialized equipment. Despite these challenges, advancements in ionothermal synthesis, including the production of hierarchical porous SAPO-34, indicate a promising future for green, sustainable scaling, provided that ionic liquid recycling and process efficiency are effectively managed.²⁸³

Microwave heating accelerates the SAPO-34 synthesis, improves crystal size uniformity, reduces the time from ~48 hours to about two hours, and producing SAPO-34 cube particles ~100 nm.^{375,386,387} This method enhances throughput and lowers costs due to its efficient energy use. Moreover, it allows for better morphology control in crystal formation. In regard to the activity test, microwave-synthesized SAPO-34 shows equal or

improved performance, as seen in longer catalyst lifetimes. However, scalability remains a challenge for larger reactors, but advances may enable rapid, energy-efficient SAPO-34 production at scale, justifying equipment costs.

In summary, each synthesis method involves trade-offs in cost, physicochemical properties, performance, and scalability. Hydrothermal is proven on an industrial scale, but uses expensive templates or yields larger crystals if cheap templates are used. Mixed-template and seed methods mitigate template cost and improve crystal size, with minimal changes needed to existing hydrothermal processes. Dry-gel conversion and concentrated gel methods significantly reduce waste and template use, supporting low-cost scale-up, but uniform scale crystallization must be controlled. Ionothermal synthesis aligns

Table 14 Comparison of different SAPO-34 synthesis methods for industrial scaling up feasibility

Synthesis method	Relative catalyst cost	Industrial feasibility	Effect on SAPO-34
Conventional hydrothermal	High cost if using expensive OSDA	Excellent scalability Well-established in the industry	Produces highly crystalline SAPO-34 Large crystals and stable SAPO-34. Minimum structure damage if properly calcined
Mixed-template hydrothermal method	Lower cost: a portion of the costly OSDA is replaced with a cheaper one	High scalability; uses the same autoclave setup as conventional	Reduced crystal size improving diffusional stability Stable catalyst compared to conventional SAPO-34
Seed-assisted method	Lower cost: it saves up to ~40% of OSDA Seeds require preparation but can be recycled	High scalability and ensure phase purity	Seeds yield smaller crystals and suppress impurities No loss of framework stability; it improves crystallinity at lower template usage
Dry-gel conversion method	Low cost: minimal solvent and high yield Reduces template and energy per unit product	Good scalability; eliminates handling of large liquids but requires uniform steam distribution. High flexibility to adopt simple synthesis equipment	Yields small, plate-like crystals with hierarchical porosity. ³⁹⁶ These have high structural integrity and improved coke resistance
Ionothermal (OSDA-free)	Potentially low cost: no OSDA and recyclable ion liquids	Moderate scalability; requires handling of viscous ion liquid and catalyst separation Recycling ion liquid is crucial for sustainability and waste minimization	Produces triclinic SAPO-34; slightly different crystal habit. ³⁸⁰ Higher framework stability; low calcination temperature ³⁸⁵
Microwave-assisted	Moderate cost with the advancement in microwave equipment Faster crystallization and lower energy consumption	Scale-up is challenging; limited by microwave penetration and uniformity in large volumes Dedicated microwave reactors are needed for industrial throughput	Produces uniform nanocrystalline SAPO-34, crystals may need binders for mechanical strength (see next section)
Ultrasonic-assisted method	Low-to-moderate cost: uses standard equipment plus ultrasonic transducers Reduce synthesis time and thus energy	Scalable in principle; large ultrasonic reactors exist (<i>e.g.</i> sonochemical tanks) Multi-point sonication may be required for uniform large batches	Produces ultra-small crystals (~50 nm) ²⁸⁴ enhanced catalyst life compared to conventional. As with microwave, nano-SAPO-34 needs to be formed into larger particles for use
Metal promotion of SAPO-34	A higher cost compared to the metal-free approach	Scalable and the dopant selection and loading method are crucial for mechanical stability ³⁹²	Tune the catalyst acidity, enhancing stability and selectivity ³⁸⁸⁻³⁹⁰ Metals sinter and lose dispersion High metal concentration may cause the formation of a secondary phase



with green chemistry by eliminating OSDAs and wastewater, though the economics rely on reusing ionic liquid. Microwave and ultrasonic techniques offer faster, energy-efficient synthesis of superior nano catalysts, but scaling the equipment is an obstacle. Table 14 provides a comparison of the cost and industrial feasibility of the different approaches of SAPO-34 synthesis and their effect on SAPO-34.

Metal promotion of SAPO-34, especially with rare earths like La³⁺ and Ce³⁺, can significantly extend catalyst lifetime by moderating acidity and suppressing coke formation. Comparative studies rank lifetime improvement as Ce > La > Co > Ni > Fe > undoped SAPO-34.³⁸⁸ La/Ce dopants typically double or triple cycle length and enhance light-olefin selectivity while reducing methane formation.^{388,389}

La and Ce form non-volatile oxides, maintaining stability under high-temperature regeneration (up to ~750 °C) and reinforcing framework integrity.³⁹⁰ The main risks are sintering and loss of dispersion, especially when metals are introduced *via* post-synthesis impregnation, whereas *in situ* substitution or ion exchange better anchors dopants and preserves crystallinity. Excessive loading may also introduce secondary phases or over-cracking, which can degrade lifetime if not controlled.³⁸⁸ Leaching is negligible under dry, gas-phase MTO conditions; rare-earth dopants remain stable during attrition-prone cycling.³⁹¹ While promoters can slightly shift product distribution, for example, La favoring propylene over ethylene, these changes are minor compared to the overall benefit of prolonged catalyst life.³⁸⁸ In industrial settings, dopant choice and incorporation methods must also preserve the mechanical integrity of catalyst particles.³⁹²

In industry, SAPO-34 powders must be shaped (*e.g.*, spray-dried with binders, please refer to Section 4.7) into robust, millimeter-scale particles to withstand thermal cycling and fluidized-bed abrasion. Attrition resistance depends on crystal size, binder material, and morphology. Optimal designs use ~0.5–5 μm crystals with alumina binders, which typically yield stronger particles than silica.^{393,394} Metal additives can influence mechanical integrity (*e.g.*, Ni/Cs incorporation may weaken particles) while hierarchical structuring or binderless pelletization can improve strength without heavy binder dilution.³⁹⁵

5 Sustainability of olefin production methods

5.1 Catalytic efficiency for emissions abatement

The negative environmental externalities arising from the production of base chemicals are significant, specifically with regards to petrochemicals such as olefins.³⁹⁷ In particular, the olefin industry comprises 30% of total direct CO₂ emissions produced from chemical plants,^{398,399} making this sector one of the most pertinent for consumption reduction. When considering the evolution of this industry, from 2000–2016, CO₂ emissions decreased by 29.4% per tonne of ethylene generated in China. This implies process improvements and efficiency advancements in a leading region of olefin production. However, it is coupled with ever-expanding industry growth and, as such,

increased overall emissions.⁴⁰⁰ Thus, it is crucial to account for emissions across Scope 1 (direct, on-site process and combustion), Scope 2 (indirect from purchased electricity/steam), and Scope 3 (all other value-chains upstream and downstream).⁴⁰¹ This helps ensure that olefin production can be assessed, quantified, and ultimately produced in a sustainable manner.³⁹⁷

Catalyst design, then, can inherently tackle reductions across Scopes 1, 2, and 3 respectively by abating emissions through enhanced process efficiency and reduced energy demand.⁴⁰² Catalysts can directly abate Scope 1 emissions by enhancing selectivity, which create pathways that bypass by-product formation and reduce CO₂ generation entirely.⁴⁰² Taking an example of the development of SAPO-34 catalysts for the methanol-to-olefin process,^{400,403} this on-site enhancement would lead to direct improvements across Scope 1 by enhancing methanol-to-olefin conversion and selectivity, thereby reducing the likelihood of GHG byproducts. Broadly speaking, however, an increase in catalytic efficiency could also allow for the integration of cleaner energy sources, thus decreasing the prospect of indirect CO₂ produced, which would then address Scope 2 emissions.⁴⁰² Finally, innovations in catalyst design could also propel significant improvements across Scope 3 emissions which are thought to be the most polluting in industry, and simultaneously the most difficult to track.⁴⁰⁴ By improving upon feedstock conversion and product yield, optimised catalysts can shave off upstream burdens, while also tackling embodied emissions downstream and enabling greener end-of-life practices.⁴⁰⁵

As such, the need for continuous improvement and process optimisation is further exacerbated by issues of commercial attractiveness and technical feasibility, which present hurdles to transitioning away from conventional, fossil-derived olefin production.³⁹⁷ One study by Reznichenko *et al.*³⁹⁸ conceptualises a colour scheme of olefin production technologies not unlike that of the “hydrogen rainbow”.⁴⁰⁶ They differentiate between feedstocks stemming from coal (brown), oil and gas (grey), biomass (green), and recycled plastic (pink), considering also the cases of carbon capture and storage (CCS, purple) and carbon capture and utilisation (blue).³⁹⁸ By coupling an analysis of both olefin production technology and its feedstock, the sustainability and circularity of the process can be more comprehensively understood.

5.2 LCA considerations in olefin production

Regarding the emissions drawn from olefin production processes, process feedstock is of paramount importance, from its sourcing, to its purification, to its eventual conversion.⁴⁰⁷ This choice of feedstock in olefin production is predominantly governed by resource availability and price, where Europe and Asia are more petroleum and coal based, whereas light hydrocarbons take the fore in North America and the Middle East.⁴⁰⁰ Irrespective of this regional variability, which inherently affects the preferred technology employed, steam cracking is regarded as the benchmark and most widely adopted production route in the olefin production industry, with coal-based processes largely explored due to their vast adoption in regions like China.^{399,408}



As with all life cycle assessments (LCAs), it is important to note that direct comparisons may either vary tremendously, or their results cannot be compared entirely. This can be attributed to 5 main LCA factors: functional unit, allocation method, system boundaries, region, and year of study.⁴⁰⁰ Evidently, the functional unit, system boundaries, and allocation method provide the reference frame against which LCA results can be compared.⁴⁰⁹ The year of study is directly related to the assessment method and database, given that new aggregates and benchmarks are established in tandem with their database sources.⁴¹⁰ Finally, given the availability of different technology and resources across the world, this creates inherent discrepancies in embodied emissions across the processes' value chains – particularly when considering energy use and provision sources (Scope 2), as well as overall supply chains and their counterparts like transportation (Scope 3).^{409,410}

A critical facet to LCAs available in literature is that of co-products and their allocation. When comparing the two key processes of steam cracking *vs.* MTO, steam cracking yields valuable co-products (such as propylene, but alongside other hydrocarbons and fuel gas/oil), whereby some environmental impacts are shared.^{411,412} Conversely, MTO is regarded as a more targeted production method in the context of LCAs, with olefins and water and/or heat as the primary byproducts;⁴¹¹ this concentrates more detriment per unit olefin, if there are no major by-products among which to distribute the process' impact. In addition, the allocation of functional unit can vary by entirely different scales (namely, mass, energy, or economic), which can skew results by tens of percentage points.³⁹⁸ For example, Zhao *et al.*⁴¹¹ allocated approximately 29% of steam cracking impact to propylene (which was the main product in their study), whereas in another study by Lv *et al.*⁴¹² only 15% was allocated, which justifies the subsequently reported CO₂ per ton disparities – even though both studies were conducted using a mass basis. Thus, caution must be heeded when drawing comparisons across LCAs from different sources, particularly when involving different functional unit bases and allocation methods.

5.3 Cradle-to-gate LCA findings in olefin production

Generally, studies have found that coal-based MTO has a significantly higher cradle-to-gate CO₂ footprint (*i.e.*, per amount of olefin product produced) than traditional steam (naphtha- or ethane-fed) cracking.^{398,400,413} Furthermore, when considering the MTO process specifically, natural gas-based production tends to fare better than its coal-fuelled counterpart.^{398,400,414–416} In particular, the product of interest that is typically examined for these light olefin applications is either ethylene or propylene, given their prevalence in industry.^{398,414} Various studies have considered offsetting upstream emissions by modifying the methanol carbon source to coke-oven gas^{417,418} or methane,⁴¹⁹ all of which have been reported to reduce CO₂ emissions. Another widely considered environmental improvement is the inclusion of carbon capture and storage (CCS) units,^{400,416} which have been stated to largely abate GHG emissions in olefin production applications.

For instance, comparing these routes, Chen *et al.*⁴¹⁶ found that the following olefin production method had the most environmental and commercial potential in China, albeit a historically coal-abundant and -dependent region: production and importing of methanol sourced from American natural gas/shale gas to obtain ethylene. This natural gas-to-ethylene (NTO) route was determined to have the highest “eco-efficiency” when compared against oil-to-ethylene (OTO, sourced from imported crude oil from the Middle East) and coal-to-ethylene (CTO, locally sourced coal within China).⁴¹⁶ The concept of eco-efficiency (E/E) – considering the product's added value against its environmental detriment,⁴²⁰ proved how NTO outperformed the other two methods in both economics and emissions.⁴¹⁶ As such, NTO was found to generate a larger net value than OTO (by 25%) and CTO (by 7.8%), as well as lower GHG emissions (4% less than OTO and 60% less than CTO) – making its eco-efficiency higher by 30.2% and 176.6% than OTO and CTO, respectively.⁴¹⁶ Even when pairing the CTO system with a CCS unit hosting a 50% capture rate – though its emissions did decrease by a significant 20% – the CTO-CCS system's emissions remained considerably higher than the OTO and NTO systems considered.⁴¹⁶

Corroborating these conclusions is a study by Xiang *et al.*,⁴⁰⁰ which highlighted the differences between ten production routes: steam cracking of mixed petroleum (PSC), steam cracking of conventional natural gas (NSC), steam cracking of shale gas (SSC), catalytic pyrolytic process (CPP), coal to olefins *via* methanol-to-olefins (CMTO), natural gas to olefins *via* methanol-to-olefins (NMTO), CO₂ to olefins *via* methanol-to-olefins (CO₂-MTO), biomass to olefins *via* methanol-to-olefins (BMTO), coal to olefins *via* Fischer-Tropsch to olefins (CFTO), and biomass to ethylene *via* ethanol-to-ethylene (BETE). In this study,⁴⁰⁰ PSC was taken as the emissions baseline (due to its industrial prevalence) at 1.7 tCO₂ per tonne of ethylene produced. Benchmarked against this, the other two steam cracking processes investigated (NSC and SSC) were found to modestly reduce emissions by 23.5% and 17.7%, respectively.⁴⁰⁰ Conversely, CPP released 94.1% more tCO₂ than PSC, making it ~2.4× more polluting than steam cracking.⁴⁰⁰ On the other hand, the MTO options varied drastically. NMTO (natural gas-based) emitted nearly 2× as much CO₂ as PSC, whereas CMTO (coal-based) was 5× more emitting than PSC, with CFTO (coal-based *via* Fischer-Tropsch) peaking at 12× more emissions than the baseline.⁴⁰⁰ By stark contrast, CO₂-MTO was almost carbon-neutral (101.2% decrease compared to benchmark), while BMTO (biomass-based) was carbon-negative at –1.3 tCO₂ per tonne of ethylene produced (176.5% decrease).⁴⁰⁰ When paired with CCS integration, BMTO had the lowest emissions contribution at –8.2 tCO₂, while CMTO and CFTO (coal-based) emissions demonstrated 55.2% and 36.9% reductions – though this marginal improvement was not drastic enough to make the coal-based MTO processes competitive, as they were still the most carbon-intensive options by far. With the “green” synthesis routes (biomass-based and/or coupled with CCS) making up the emissions rear, and the coal-based (“brown”) methods at the fore, it is evident that low-carbon technologies present the most sustainable MTO pathways,



whereas coal-fuelled production carries a substantial environmental premium.^{398,400} This reinforces the idea that the carbon burden of olefin production – especially for the MTO process – hinges on feedstock choice and the progression beyond and away from coal-based processes.

Another cradle-to-gate study by Zhao *et al.*⁴²¹ similarly compared major olefin production methods, analysing the following four: catalytic cracking (CC), steam cracking (SC), coal-to-olefins (CTO), and coal-to-propylene (CTP). Among these, CC was owed the lowest footprint, with 22.5% less GHG emissions than SC, whereas CTO and CTP produced emissions that were 5.8× and 7.6× higher (respectively) than CC. This was also reflected by the trend observed in the primary energy demand, where CTO consumed the most energy, followed by CTP and SC, while CC again performed the best. For the two top polluters (CTO and CTP), over 88% of emissions arose from the production processes themselves, which encompass many stages such as air separation, coal gasification, methanol synthesis, and MTO/MTP. Most significantly, coal gasification alone gave rise to 67.72% and 60.79% of the GHG emissions in CTO and CTP, respectively. However, when adopting CO₂ capture, GHG emissions were found to decrease by 32.4% and 26.1% – though energy usage spiked by 2.6% and 3.6% – for CTO and CTP, respectively. This further propels the notion that coal-based production processes are disproportionately carbon- and energy-intensive – and while advancements like CCS can certainly curb their environmental detriment, coal's unfavourable performance warrants phase-out.

A 2022 compilation by Reznichenko *et al.*³⁹⁸ presented one of the most comprehensive cradle-to-gate comparisons of olefin production methods, tabulating all the cases explored in previously mentioned studies and more. This analysis unsurprisingly revealed similar findings: the “brown” (coal-to-olefins MTO) baseline was the most emissions-intensive (even when compared to other coal conversion processes different than MTO), at a hefty maximum of 8.65 tCO₂ per t olefin.³⁹⁸ Yet again, coal-powered processes proved more detrimental than their “grey” counterparts' emissions peaks – with 2× more CO₂ generated than conventional naphtha steam (4.5 tCO₂) and 2.5× more than condensate-based cracking (3.4 tCO₂).³⁹⁸ Conversely, including CCS units (“purple”) into coal-based MTO lowered emissions by 66% (to 2.63 tCO₂); though this was a notable change, it still trailed behind several non-fossil alternatives.³⁹⁸ It is evident that – solely considering the MTO process alternatives – circular routes hold the most promise. With the low-value waste stream of black liquor sourced as a methanol feedstock, a “green” MTO route led to a 75% decrease in emissions (to 2.17 tCO₂), whereas a “blue” Power-to-X approach to olefins, starting from atmospheric CO₂ and using DAC, had carbon-negative performance (−1.5 tCO₂), 117% lower than the coal-MTO benchmark – albeit with ~2× higher energy demand.³⁹⁸ This study³⁹⁸ echoed the same findings as discussed earlier: coal-based MTO is currently the least sustainable in industry – though this can be abated through the incorporation of renewable or waste-derived methanol feed, making it a low-carbon or even carbon-negative route.

Other technologies like bioethanol to ethylene (“green”) have demonstrated high sensitivity to upstream intermediate production emissions, such as those related to farming biomass, fertilisers, and transportation.³⁹⁸ As such, the emissions associated with their olefin production vary drastically – from being carbon-negative⁴²² to net-positive and even highly net-positive.⁴²³ Some of these studies depict how these sources' intrinsically sustainable nature can be eroded by biogenic emissions in pre-processing during upstream stages. However, though drawing from various LCAs can relay highly variable results in the literature, most realistic scenarios depict a near 35–45% reduction compared to traditional naphtha-based cracking, thus performing better than both steam cracking (“grey”) and coal-MTO (“brown”).³⁹⁸

Similarly, different LCAs report mixed results for the recycled plastic (“pink”) route, depending on the accounting method employed. For instance, when considering a direct process basis to produce ethylene from waste, pyrolysis with steam cracking was carbon-intensive in nature, relaying 3 tCO₂ of reported emissions, which was not much improved from traditional naphtha cracking.³⁹⁸ However, common LCA practice is to adopt a “differential credit” system – accounting for the abated emissions of this route insofar as waste incineration is avoided (which would have been the fate for the plastics involved). Doing so alleviated the footprint across the process' life cycle drastically, amounting to 50% lower GHG than the original naphtha route.³⁹⁸ In other words, this “pink” recycled plastic feedstock paves the way for closing the carbon loop on polymers,⁴²⁴ where the CO₂ that would have been emitted by the disposal or burning of waste is harnessed for new value-added products. Comparable environmental benefits for this waste-to-methanol-to-olefin (WMO) process are also disclosed by Salah *et al.*,⁴²⁴ who express how this route can achieve meaningful GHG reductions in the so-called “linear” fossil economy. It must be noted, however, that this path was made viable by modelling a cleaner electricity mix (ideally renewable sources such as wind energy),⁴²⁵ necessarily also at the large-scale to avoid burden shifting to other impact categories (namely, eutrophication, particulate matter formation, acidification, and resource use).⁴²⁶

Overall, though the differences in regional energy sources and efficiencies across cradle-to-gate studies are vast, naphtha steam cracking is said to produce roughly 0.8–2.1 tons of CO₂ emissions per ton of olefin produced, an immense difference from the whopping 5.2–11.5 tons arising from coal-based MTO.⁴²⁷ Though both processes are very well-established in industry, they face imminent obsolescence without addressing their environmental implications. Steam cracking is anticipated to remain omnipresent in industry for another 10–20 years; to remain as such, however, mid-term improvements of optimised cracker efficiency and feedstocks from recycled (“pink”) and renewable (“green”) sources are crucial. Long-term advancements in cracker electrification propel this shift,³⁹⁸ as electrified crackers can achieve up to a 92% reduction in emissions relative to today's furnaces (assuming a carbon-free electricity supply).⁴²⁸ On the other hand, the best-case MTO scenarios outperform steam cracking when methanol is sourced from



low-carbon pathways like biomass or waste, resulting in nearly net-zero or even net carbon-negative GHG footprints (depending on the feed).⁴²⁹ Growth in this regard is predominantly staggered by technological or economic immaturity – especially for “green” and “blue” MTO solutions still at lab- or pilot-scale – when choosing to opt for less carbon-intensive feedstocks.⁴³⁰ Therefore, bio-based methanol sources cannot singlehandedly replace fossil feed at scale – at least in the near-short term.³⁹⁸ As for coal-powered processes, eventual phase-out or decarbonisation is unequivocally pertinent, though increased policy support and infrastructural changes are needed to accommodate for CCS (“purple”) solutions.³⁹⁸

5.4 Energy requirements in olefin production

A key environmental consideration – especially when benchmarking technologies and feedstocks across various studies – is that of energy requirement. When investigating olefin production, the conversion of methane or coal into methanol and then to olefins involves multiple reaction steps, where energy content of the feed is expended and efficiency losses are realised in stages such as syngas production, methanol synthesis, and the MTO reaction itself.⁴³¹ On the other hand, steam-cracking has been dubbed more energy-efficient due to its one-step nature and direct olefin yield – albeit endothermic⁴³¹ and energy-intensive, where crackers typically require furnace temperature between 750 and 1100 °C.⁴³² The disparity in energy requirement between these two conventional routes is equally as ingrained in literature as that in emissions – where multiple studies report notable differences between the energy demand of coal-based MTO (150–156 GJ per ton olefin) and steam cracking (54–66 GJ per ton olefin)^{421,433} again demonstrating the shortcomings in coal-powered MTO.

As a proposed fix to coal-based MTO, the caveat with CO₂-to-olefins is their immense energy requirement – and, as such, the high capital and operational costs. Power-to-X routes are energy-intensive, particularly due to electrolysis and CO₂ conversion steps. For example, Flores-Granobles *et al.*⁴⁰⁸ found that a two-step CO₂-to-methanol-to-olefin process would require ~59 GJ (16.3 MWh) of electricity per ton of olefin produced, which is roughly 5× higher than the conventional cracking benchmark used. Similarly, Reznichenko *et al.*³⁹⁸ noted that it is not viable to satisfy the global light olefin demand of 200 Mt per year solely through the CO₂-MTO production method, as it would require 644 GW of renewable electric power (roughly half of the United States' overall grid capacity of 1280 GW).⁴³⁴ This immense scale of renewable energy deployment poses a towering economic viability barrier. Thus, the only two scenarios in which these routes could compete with petrochemicals today is either through (1) strong carbon pricing, subsidies, or policies, or (2) the adoption of wide-scale renewable energy to the degree of abundance and affordability.³⁹⁸

Looking towards the prospect of a fully decarbonised olefin industry, a mixed-market solution may prove ideal to accommodate regional strengths, financing hurdles, and technological barriers. For instance, e-crackers can be employed in steam

cracking solutions, whereas other green fuels can be used to feed MTO units presently available in industry.³⁹⁸ In tandem, the proposed circular carbon storage (“purple”) and utilisation (“blue”) solutions can be further developed, while also accelerating the integration of renewable energy into local grids.³⁹⁷

5.5 Water usage & other impact categories in olefin production

Beyond GHG emissions and energy requirements, other environmental considerations are also considered to provide a more holistic overview of how different technologies and materials interact with both human life, the natural environment, and available resources. These are referred to as midpoint impact categories, all of which are related to an endpoint that is encompassed by an area of protection. As illustrated in Fig. 42 below,⁴³⁵ the selection of impact categories in most comprehensive LCAs spans across all three major nodes of protection – Human Health, Natural Environment, and Natural Resources.

To contextualise the results that will be disclosed across multiple LCAs, a definition for each impact category will be provided. An extensive discussion was provided on climate change in an earlier analysis – which is primarily measured in terms of GHG emissions (CO₂-equivalent units) – defined as a greenhouse gas-driven radiative forcing (or global warming), leading to sea level rise, extreme weather, and ecological disruptions, and impacting both human health and the natural environment.⁴³⁵ Ozone depletion involves chlorofluorocarbons (CFCs) and halocarbons, which reduce the ozone layer and increase UVB radiation; this direct causation of skin cancer, cataracts, and ecosystem loss also similarly affects both human health and the natural environment.⁴³⁵ Acidification involves sulfur dioxide and nitrogen dioxide emissions, which lead to acid rain and soil acidification, impacting both human health and the natural environment.⁴³⁵ Eutrophication involves phosphorus and nitrogen emissions, which lead to algal blooms and hypoxia, impacting both human health and the natural environment.⁴³⁵ Land use involves changes in land cover, which impact both human health and the natural environment.⁴³⁵ Water use involves the consumption of water resources, which impact both human health and the natural environment.⁴³⁵ Abiotic resource use involves the consumption of non-renewable resources, which impact both human health and the natural environment.⁴³⁵ Human toxicity involves the release of toxic substances, which impact human health.⁴³⁵ Particulate matter formation involves the release of particulate matter, which impact human health and the natural environment.⁴³⁵ Photochemical ozone formation involves the release of volatile organic compounds, which impact human health and the natural environment.⁴³⁵ Ecotoxicity involves the release of toxic substances, which impact the natural environment.⁴³⁵ Acidification involves the release of sulfur dioxide and nitrogen dioxide, which impact the natural environment.⁴³⁵ Eutrophication involves the release of phosphorus and nitrogen, which impact the natural environment.⁴³⁵ Land use involves changes in land cover, which impact the natural environment.⁴³⁵ Water use involves the consumption of water resources, which impact the natural environment.⁴³⁵ Abiotic resource use involves the consumption of non-renewable resources, which impact the natural environment.⁴³⁵ Human health involves the impact on human well-being, which is the primary area of protection.⁴³⁵ Natural environment involves the impact on the natural world, which is the primary area of protection.⁴³⁵ Natural resources involves the impact on the availability of natural resources, which is the primary area of protection.⁴³⁵

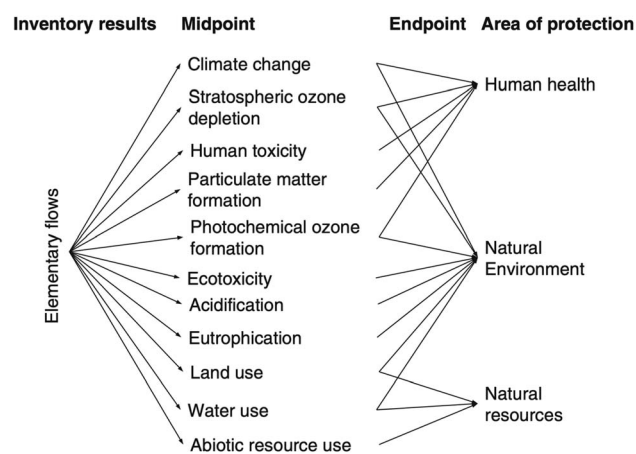


Fig. 42 Framework of environmental midpoint impact categories illustrating their relation to the areas of protection. Reproduced from ref. 435 from Springer Nature, Copyright 2015.



Photochemical ozone formation accounts for the ozone (or O₃) that is formed by NO_x and NMVOC emissions, causing both respiratory effects and crop (and greater ecosystem) damage, linking both to human health and the natural environment. Ecotoxicity assesses the damage from chemical exposure *via* effects on species across trophic levels, ultimately harming the natural environment.⁴³⁵ Acidification describes the ecosystem damage that arises specifically from emissions (namely – CO₂, N, and S compounds) causing acidification across terrestrial, freshwater, and marine ecosystems, which directly impacts the natural environment. Eutrophication refers to the ecosystem change that is driven by the production and release of excess nitrogen and phosphorus, leading to nutrient oversupply that disrupts natural pre-existing ecosystems and ultimately affects the natural environment.⁴³⁵ Land use reflects the disruptions from human land occupation and transformation, including the loss of biodiversity, which also affects the natural environment.⁴³⁵ Water use captures the impacts of water consumption, which can deprive ecosystems and human users of available water supplies, linking to both the natural environment and human health.⁴³⁵ Finally, abiotic resource use concerns the depletion of non-living resources such as metals, minerals, fossil and nuclear energy, and flow energy sources like wind, which directly links to natural resources.⁴³⁵

Among these categories, water depletion metrics are recurring figures of sustainability in the context of olefin production.⁴³⁶ The previously mentioned comparative study by Chen *et al.*⁴³⁷ revealed that the process water consumption of coal-to-olefins (CTO) was the worst of the three methods considered – 2.3× higher than NTO (natural gas-to-olefin) and 3.3× higher than OTO (oil-to-olefin). Revisiting the study by Lv *et al.*,⁴³³ the authors also emphasised a similar trend across four propylene production routes: coal-to-olefins (CTO), propane dehydrogenation (PDH), steam cracking (SC), and catalytic cracking (CC). As expected, water consumption was highest in CTO, averaging 35.4 tons of water per ton of propylene produced – attributed to its intensive cooling and gasification operations.⁴³³ Furthermore, the three other impact categories of eutrophication, acidification, and respiratory inorganics also peaked with CTO – the latter two arising predominantly from electricity and steam input.⁴³³ On the other hand, the reason why eutrophication was heavily skewed towards CTO was because of its direct nitrogen emissions⁴³³ – which has the most immediate effect on eutrophication potential⁴³⁵ – reaching levels approximately 2800×, 1900×, and 900× as high as CC, CC, and PDH (respectively).⁴³³ However, PDH exhibited both severe abiotic depletion and ozone depletion, which was propelled by the process' propane feedstock contributions to these categories (94% and 98%, respectively).⁴³³ Furthermore, although SC and CC fared better than CTO in terms of water use, their raw material impacts were considerable; for instance, when looking at the same metrics of abiotic and ozone depletion potential, naphtha and refinery dry gas drove 66–77% of SC's contributions to these categories, whereas heavy oil accounted for 73% of CC's contributions.⁴³³ Nonetheless, an important sustainability aspect to the CC technology was its ability to recover steam from coke combustion, which alone offset 160% of respiratory inorganics, 13% of

its water use, and 12% of its acidification impact – though this favourable performance was not reflected across all categories (*e.g.*, ozone depletion, where it is the second most detrimental after PDH).⁴³³ Although this analysis depicts how coal-based MTO is undeniably the most water- and pollution-intensive, propane dehydrogenation shifts the burden towards resource and ozone depletion-related categories, making it challenging to identify which process singlehandedly had the worst overall footprint.⁴³³

In a study covered earlier by Salah *et al.*,⁴²⁴ a prospective LCA defined a linear olefin production route (or LPR) based on traditional naphtha steam cracking to compare against a waste-to-methanol-to-olefins (WMO) route. Beyond the climate change effects covered earlier, 15 other environmental impact categories were considered – which depicted not only circular improvements, but also burden-shifting, measured in terms of probability (percentages).⁴²⁴ For one, WMO illustrated high likelihood of reducing impacts across multiple categories: freshwater ecotoxicity (99%), resource use (100%), land use (87%), and ozone depletion (100%), largely because of the exclusion of naphtha extraction and refining, which are credited for land disturbance, toxic effluents, and fossil resource depletion.⁴²⁴ However, WMO was not without its shortcomings; it increased the likelihood of burden-shifting across several categories due to its electricity-intensive nature: acidification (94%), freshwater eutrophication (100%), terrestrial eutrophication (87%), metal/mineral resource use (100%), and particulate matter formation (97%).⁴²⁴ These burdens stem from the global grid mix (2020) – dominated by coal (36% make-up) – which emit NO_x, SO_x, and particulates, in addition to requiring extensive mineral inputs for generation and transmission.⁴²⁴ For the remaining six categories – namely, marine eutrophication, human toxicity (carcinogenic and non-carcinogenic), ionising radiation, photochemical oxidant formation, and water use – the results were unclear, with probabilities of burden-shifting between 25 and 75%, making a firm conclusion across these metrics impossible. Despite this, a low-carbon electricity mix by 2050 would likely omit this notion of burden-shifting, which would make the WMO route a clear sustainable fix across almost all the indicators investigated. Altogether, when aggregating all these performance indicators into one final environmental footprint score, WMO exceeded LPR by 16.8% in 2020 (time of study) and 45.1% by 2050 (future renewable energy grid case), demonstrating that while this circular solution presented many environmental benefits, these advantages could realistically not be realised without grid decarbonisation.⁴²⁴

Centred on comparing prospects of oil-based steam cracking against coal-fuelled MTO, Xiang *et al.*⁴³⁸ reinforce the striking differences between feedstocks illustrated previously in the context of emissions. Analogous to what was found earlier (for emissions), coal-to-methanol-to-olefins (CMTO) and coke oven gas-based methanol-to-olefins (CGMTO) were the most water-intensive, consuming 44.2 and 55.0 tons of water per ton of olefin, respectively.⁴³⁸ By contrast, natural gas-based MTO (NGMTO) consumed only 27.2 t/t, whereas oil-to-olefins (OTO) had the lowest usage at 22.1 t/t.⁴³⁸ Looking more closely at the



indirect water use (associated with energy and feedstock preparation), the demand distribution expressed as a proportion of total water usage among the top three processes was 32.1%, 25.5%, and 21.3%, for CMTO, CGMTO, and NGMTO, respectively – which echoes the earlier idea of accounting for and improving upon upstream process savings.⁴³⁸ In the context of this study, geographic misalignment of resources like coal reserves and water exacerbates this impact: if CMTO plants are concentrated in arid regions that are disconnected from a direct water supply, it manifests into a structural water-resource conflict. Thus, steering away from coal-based operations and instead pivoting towards a more diversified olefin production portfolio (*e.g.*, with co-feeding routes or better geographic positioning) could cut average water intensity by 15.8% compared to business as usual. These findings demonstrate how coal-based olefins remain excessive from a consumption standpoint, propelling the need for gas-based alternatives to mitigate water scarcity and regional imbalances.⁴³⁸

Another study discussed by Xiang *et al.*⁴³⁹ focused on the comparison of feedstocks, depicting the stark contrast in environmental burdens as a byproduct. Yet again, coal-to-olefins (CTO) shouldered a markedly heavier burden in both water and resource consumption in comparison to oil-to-olefins (OTO). For example, the CTO route necessitates 30 t of fresh water per 1 t of olefins produced – 4.2× greater than the water demand of OTO. Furthermore, CTO draws far more heavily on utilities, with 6.0 GJ per t of electricity and 8.8 GJ per t of steam, compared to a mere 0.27 GJ per t and 1.16 GJ per t demanded by OTO, respectively – which points to auxiliary resource requirements beyond direct feedstock burden. As expressed previously, this could pose resource conflicts in water-scarce regions, suggesting that coal-based MTO could worsen resource scarcity unless mitigated in the near-term. To address these coal-backed drawbacks, process-level optimisations (*e.g.*, cooling through air instead of water) and policy intervention could address sustainability bottlenecks in industrial MTO applications.⁴³⁹

6 Future opportunities

Although the MTO process over SAPO-34 offers a promising route for producing light olefins, challenges such as rapid catalyst deactivation *via* coke formation, complex reaction mechanisms, and demanding process conditions require innovative solutions to enhance efficiency, selectivity, and scalability. This section outlines key opportunities for future research and development to optimise catalyst longevity, process efficiency, and environmental sustainability, paving the way for next-generation MTO technologies that can realistically cater to the global olefin demand.

One critical area for improvement is that of SAPO-34 catalyst design; given that it is the benchmark MTO catalyst in industry,⁴⁴⁰ its synthesis can be further optimised. As covered extensively in this review, one major improvement is that of its crystallite size and morphology – where reductions of crystal size can alleviate the key limitations of diffusion and coke build-up, increasing selectivity and lifetime.⁴⁴¹ Given that SAPO-34's cage structure can impose shape-selectivity for light olefins,⁴⁴²

engineering hierarchical or nano-sized porosity can alleviate the diffusion limitations of SAPO-34's small 8-ring channels. This allows for the suppression of coke deposition and prolonged catalyst lifetime compared to conventional large crystals, since coke molecules can diffuse out or get confined to smaller domains.^{443,444} For instance, Guo *et al.*⁴⁴⁴ prepared SAPO-34 with a novel aluminium source (pseudoboehmite) with lower Si content and a hierarchical pore structure, resulting in a longer catalytic lifetime than standard SAPO-34 in MTO tests. This approach was paired with the strategy of tuning Si content by optimising the Si/(Al + P) ratio. Balancing activity and stability is key in this regard – as lower Si can improve longevity and boost selectivity, but at the expense of higher temperature.⁴⁴⁴ Altogether, enhanced coke resistance is a practical solution as it directly translates into longer operational cycles in industrial reactors.⁴⁴⁴ As such, future work should focus on precisely controlling morphology and acid site distribution, given their direct ties to olefin selectivity and catalyst efficiency.

In fluidised bed reactor applications, catalyst regeneration is a potential avenue to mitigate the accumulation of coke (as highlighted in Section 2.2),⁴⁴² typically through high-temperature air combustion. However, steam-assisted regeneration offers a milder alternative (or even a supplement to traditional air/oxygen burning), operating at 400–500 °C to gasify coke into CO, CO₂, and H₂, thereby preserving the catalyst's chabazite framework.^{445,446} Unlike air-based methods at 600–700 °C, which risk hydrothermal dealumination, Zhao *et al.* showed that steam regeneration can extend catalyst lifetime by 15–20% by reducing polyaromatic coke formation.¹⁴⁰ Another study by Zhou *et al.*⁴⁴⁷ proved how partial regeneration of coked SAPO-34 using steam gasification boosted both activity and ethylene selectivity (from 23% for fresh SAPO-34 catalyst to 53% after steam treatment). The occurrence of the latter was attributed to the formation of residual “hydrocarbon pool” coke (mostly methylated benzenes) on the catalyst, which promoted ethylene formation and slowed deactivation, whereas even partial air regeneration left less desirable coke (*e.g.* oxygenated compounds and naphthalene) that accelerated deactivation.⁴⁴⁷ In addition, Liang *et al.*⁴⁴⁵ suggest steam regeneration as a means to both increasing revenue and abating emissions – especially in comparison to conventional air combustion. The authors found that the integration of steam-assisted regeneration improved the catalyst's carbon utilisation rate by ~10% and energy efficiency by ~6%, in addition to exhibiting near net-zero emissions potential.⁴⁴⁵ Future research could focus on optimising this potential regeneration for the MTO process.⁴⁴⁷ Furthermore, it is important to recall how industrial MTO catalysts are typically spray-dried into millimetre-sized particles (often with a binder like kaolin or alumina) to circulate in fluidised beds.⁴⁴⁸ Thus, forthcoming research should also link morphology and binder content of the SAPO-34 particles to their performance and durability since stronger, attrition-resistant particles can reduce catalyst losses over time.⁴⁴⁸ What all these insights illuminate is that the deliberate control of the extent of coke removal – rather than catalyst regeneration to a pristine and unadulterated state – may prove most applicable in prolonging catalyst longevity in commercial MTO



operations. In addition, integrating dilute oxygen streams in fluidised bed applications could further reduce CO₂ emissions, enhancing environmental sustainability in a largely polluting chemical sector.^{398,431} On the other hand, recent *in situ* nuclear magnetic resonance (NMR) studies also demonstrate how maintaining a modest level of coke on the catalyst (not eliminating it entirely) and co-feeding water during the reaction can significantly preserve catalyst framework and suppress dealumination.⁴⁴² For this purpose, employing *in situ* spectroscopic techniques (such as *operando* infrared or NMR spectroscopy) would provide real-time insights into coke evolution and dynamics, enabling precise process control.⁴⁴⁹ These advancements could significantly reduce operational downtime and improve economic viability of MTO plants, particularly for large-scale fluidised-bed applications.

Another promising direction is hydrogen co-feeding to suppress coke formation,⁹⁸ as highlighted in Section 2.3. By hydrogenating polyaromatic coke precursors, hydrogen can extend SAPO-34's active lifetime by up to 80 hours; though it effectively removes most coke-prone intermediates,⁹⁸ it can often reduce light olefin selectivity due to this alkene hydrogenation.⁹⁸ Corroborating this is the study by Ghavipour *et al.*,⁴⁵⁰ which captures how the addition of hydrogen extended the time on-stream by ~27%, with only a minor increase in alkane (paraffin) byproducts. Future efforts should focus on designing such catalysts to optimise hydrogenation specificity. To achieve this, kinetic modelling could identify ideal H₂/CH₃OH ratios to balance lifetime extension with selectivity, while density functional theory (DFT) studies could explain hydrogen's interactions with hydrocarbon pool intermediates. By enabling stable operation in fixed-bed reactors – where continuous regeneration is impractical – this approach could enhance MTO's process flexibility and adaptability to diverse feedstock compositions.

With regards to the findings from the earlier sustainability assessment, much of the readily available LCA and techno-economic data on the industrial application of MTO technology is centred on that of coal feedstock.³⁹⁸ From this sustainability analysis, it becomes evident that the four MTO technologies that are presently or will prospectively be employed in industry are the following: “brown” (coal-based), “green” (bio- or waste-based), “blue” (CO₂ point source-based, carbon capture and utilisation), and “purple” (incorporation of carbon capture and storage).³⁹⁸ With the innumerable cases of coal-powered MTO leading to grave environmental concerns,^{421,433,437–439,451} alternatives to “business as usual” operations must expediently be developed – particularly when considering how sensitive the coal-to-olefin technology is to national regulatory changes and carbon policies.⁴³⁸ In terms of the deployment of CCS solutions to pre-existing coal-MTO technology, this technology is regarded as relatively mature, though it needs to be complemented by systemic infrastructural changes to be made feasible.⁴³⁸ In addition, a study by Xu *et al.*⁴⁵² emphasises how “brown” solutions may pose a competitive advantage over other methods (like the extensively explored steam cracking, which is largely petroleum-based) in regions that are coal-abundant, since this would

reduce reliance on crude oil imports and bolster local economic activity. As such, CCS becomes a key lever in ensuring the sustainable advancement of this technology, since coal's long-term phase-out will need to be rectified by short-to medium-term emissions interventions.⁴³⁸ Consequently, “purple” technologies also provide a more inexpensive route to lowering CO₂ emissions than imposing carbon taxes.⁴⁵² As for solutions in both CCU (obtaining methanol from CO₂-based sources) and renewable routes (obtaining methanol from biomass-based sources), major leaps need to be made to address technological pitfalls.⁴³⁸ Though “green” solutions are expected to remain a marginal segment of the olefin production portfolio, great strides in economic viability could render it competitive with traditional “grey” methods (like naphtha cracking).⁴³⁸ Improvements in gasification and synthesis concepts⁴⁵³ could manifest in a first-of-its-kind demonstration plant within the next decade.⁴³⁸ Finally, “blue” (CCU) technologies will be largely dependent on the provision and price of low-carbon electricity – where, if enhanced, could help actualise the several proposed industrial demonstration cases in reality.⁴³⁸

On another note, some researchers have postulated that it is possible to directly convert syngas or even CO₂ into light olefins using bi-functional catalysts^{454–457} (see Fig. 43) – bypassing the intermediate methanol stage altogether in a single reactor. These catalysts combine a metal-based component with a zeolite such as SAPO-34 to achieve relatively high olefin selectivities. This approach is said to eliminate the energy-intensive methanol synthesis step and reduce capital costs – and researchers have made rapid progress in this area.^{458–463} Accelerating this development would present immense advantages to the olefin industry at large – since direct syngas-to-olefins would accommodate for a wide variety of feedstocks (namely, coal, natural gas, and biomass).⁴⁵⁸ However, challenges such as low single-pass conversion (due to thermodynamic limits and the production of CO₂ in the water-gas shift equilibrium) and catalyst stability under operating conditions still remain.⁴⁵⁷ Future efforts could focus on developing catalysts with abundant oxygen vacancies or optimised promoters to improve conversion but decrease CO₂ formation.⁴⁵⁸ Furthermore, tailoring metal-zeolite interactions or lowering acidity would optimise active site distribution and suppress secondary hydrogenation reactions, such as the formation of methane as opposed to the olefin desired product.⁴⁵⁸ Tackling these drawbacks would prove essential to propelling this innovative approach into viable industrial applications.

Overall, real-world applications of MTO technology have confirmed many of the bottlenecks identified herewith, illustrating which challenges persist in industry and what strategies can be pursued to overcome them. For instance, China's DMTO (Dalian Institute's methanol-to-olefins technology) exemplified how the highly exothermic nature of methanol conversion and rapid SAPO-34 coking necessitate a circulating fluidised-bed reactor with continuous regeneration.^{465,466} This principle was previously emphasised, as fluidised bed technology is a major node of industrial viability for the MTO process at scale. The first commercialised DMTO unit achieved >99% methanol conversion and ~80% light olefin selectivity by using this fixed



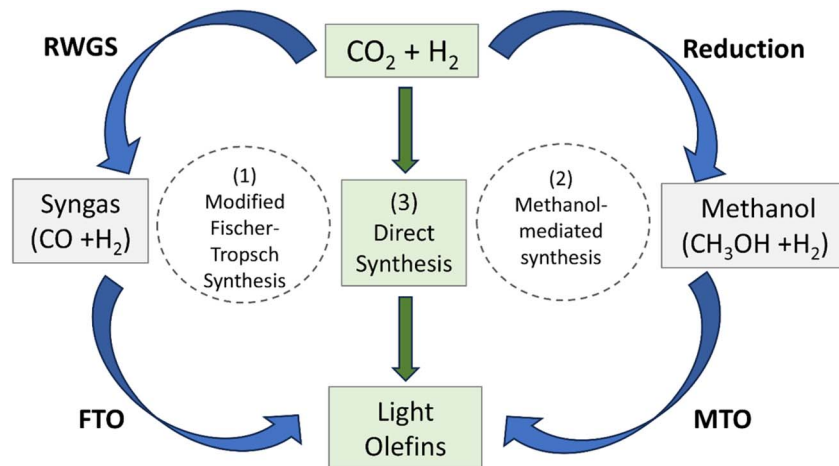


Fig. 43 General scheme of the three routes of olefins formation from CO_2 : (1) modified Fischer–Tropsch to Olefins (FTO) via Reverse Water–Gas Shift (RWGS); (2) methanol to olefins (MTO); (3) direct olefin formation from CO_2 . Reproduced from ref. 464 with permission from MDPI, Copyright 2021.

catalytic cracking style reactor-regenerator design.^{465,466} This configuration also allowed for coke burn-off on SAPO-34 *in situ* and maintained near-isothermal operation, which proved superior to the alternative of fixed beds in handling heat removal and deactivation.⁴⁶⁵ Industrial implementation also showcased how partial regeneration can occur: rather than burning off the catalyst until it was completely purified, DMTO units typically leave a small hydrocarbon residual on SAPO-34 during regeneration to act as active “hydrocarbon pools”. This approach, deployed in commercial units,⁴⁶⁵ was found to boost ethylene and total olefin selectivity in the following cycle, thereby validating the laboratory-scale notion of the benefits of retaining minor quantities of coke on the catalyst surface.

General industrial applications of MTO point towards its energy-^{421,428} and water-intensive^{436,467} nature – particularly when considering downstream effects. One key factor contributing to this metric (and, in turn, to operating costs) is that of the refrigeration duty for cryogenic separation.⁴⁶⁸ However, advances are being made in prefractionation and absorption techniques to handle the very light components in MTO's product mix (H_2/CH_4) separation – which typically mirrors that of a steam cracker (in that it requires many of the same steps).⁴⁶⁸ One such design study proposed an oil absorption process to remove this residual methane and carbon monoxide, allowing the demethaniser temperature to be raised and bypassing some deep refrigeration steps.^{468,469} Other technologies like membranes and adsorption are also being considered; to this end, zeolite membranes could selectively permeate olefins over paraffins, potentially de-bottlenecking the C_2/C_3 split and saving energy, such as was the case with the use of metal-organic framework (MOF) adsorbents in the context of MTO purification.⁴⁷⁰ As for water management, given the need for resource savings,^{436,467} researchers are beginning to investigate reuse technologies.⁴⁷¹ This would prove instrumental in resource-challenged, dryer regions⁴³⁸ – whereby wastewater streams can be avoided, and water recovery can simultaneously be attained.⁴⁷¹

7 Conclusion and outlook

Growing from concept to commercial reality, at present, the influx of research in the topic of the methanol-to-olefin process is catalysed by a need for more sustainable production routes. By enabling the manufacturing of essential petrochemicals, MTO supports energy diversification by opting for alternative feedstocks, thereby reducing reliance on cruel oil (upon which conventional steam cracking relies). Olefins' significance is evidenced by a rise in global demand of around 20% from 2015–2022 for both ethylene and propylene. This steady increase is anticipated to only persist over the next decade, particularly due to the potential of the MTO process as a gateway to a closed-loop carbon cycles, by sourcing methanol into the process from low-carbon or waste-derived feedstocks. Thus, MTO's unique value offering is its ability to convert feedstock flexibility into chemical output that is highly sought after in an increasingly decarbonised sector.

In this literature review, alternative methods to produce light olefins have been presented. Two longstanding frontrunners of light olefin production are thermal cracking and fluid catalytic cracking, though they are widely recognised as being emissions-intensive and low-yielding, respectively. In contrast, the prospects of MTO as a feasible route to sustainable olefin production are quite favourable, those of which are only heightened when paired with the supply of renewable feedstock at the onset. To actualise these, critical parameters and major reactor operating factors such as temperature, WHSV, and co-feeding of water and hydrogen are paramount, as illustrated in the main process flow diagram presented in this article.

The development and implementation of the MTO process have taken great leaps in the past few decades. In this article, this was substantiated and chronologically documented in the discussion of MTO reaction mechanisms. Among the numerous proposed schemes, the hydrocarbon pool concept is the most generally agreed-upon, with two key mechanisms which light olefin formation can be attributed to: the side chain mechanism



and the expansion–contraction mechanism. One major finding is that preferential behaviour towards one or the other is largely dependent on the acidity of the zeolite catalyst used. Nevertheless, by considering the sheer volume of proposed reaction routes and their corresponding justifications, there remains the need for studies to be conducted to ascertain which schemes are of primary relevance. A similar consensus arises when examining studies of the process's kinetics. After assessing various proposed kinetic models, it can readily be inferred that some studies disregard the effect of coke formation on the catalyst altogether, which is a primary consideration in the context of performance optimisation. On the other hand, other experiments utilise reaction conditions that are not representative of the industrial application of this process and are therefore not applicable to the scope of this review. Therefore, further documentation and testing must be conducted for a more conclusive assessment.

On the note of catalyst considerations, this review proceeded with the introduction of various materials employed in the MTO process, with SAPO-34 standing out as one of the most viable. To actualise its suitability for this application, various strategies to improve its performance were discussed. Though SAPO-34 is most commonly synthesised by means of the hydrothermal method, other approaches like the dry-gel conversion, ionothermal, ultrasonic, or microwave synthesis methods were explored, offering advantages such as reduced solvent usage, controlled crystallisation, and rapid nucleation. In terms of the template employed during synthesis – though TEOAH is the most prevalent alternative – more inexpensive materials have also been investigated with varying degrees of success, since one opportunity cost of doing so is overall poorer catalytic performance. Therefore, a mixed template approach (with TEOAH followed by a cheaper alternative) offers a competitive edge to capitalise on both benefits: performance and cost-efficiency. On the other hand, silicon content was determined to play a crucial role for selective light olefin production, underscored by a delicate balance of the SiO_2 to Al_2O_3 molar ratio in SAPO-34. Overall, the prevailing consensus is that catalyst morphology of reduced particle size offers enhanced activity and mass transfer properties, thereby proving sub-micron synthesis to be a viable objective for SAPO-34 development. Various means of attaining hierarchical SAPO-34 synthesis by use of soft or hard templates or post-treatment desilication were also found to be noteworthy paths to overcoming deactivation by coking. In contrast, metal promotion seems to lack reliable analysis on the contribution of secondary metals and their influence on the reaction mechanism, despite numerous claims of its potential to improve the catalyst. This contradictory finding was exacerbated by the antithetical results obtained from different research groups' studies, casting further doubt on the effectiveness of metal promotion overall.

In terms of catalyst use, the spray-dried technique has demonstrated inextricable drawbacks in the fluidised bed application. Catalyst shaping, therefore, paves the way for an open area of research that could substantially improve the economic viability and efficiency of the process. However, instances of *in situ* SAPO-34 synthesis over micro-spherical supports still encounter challenges of low loading and proper

adhesion. With regards to the nature of the process itself, current MTO applications have been limited to thermo-catalytic implementation. This presents a pathway for research groups that can try to implement MTO in the context of other viable methods such as electrocatalysis, photocatalysis, homogeneous catalysis, or plasma-based catalysis.

Realising the full potential of MTO in practical applications will require continued innovation focused on both technical viability and environmental sustainability. One core challenge of doing so is the trade-off between catalyst longevity and olefin selectivity – which is where synthesis procedures play an instrumental role in balancing these facets. Alongside catalyst design improvements, optimised process regenerations and co-feeds are on the horizon to address coking issues in industry. In particular, research shows how there is an inextricable link between catalyst structure and performance requirements; these materials must be engineered for reactant conversion and product selectivity, but they must also be robust and regenerable under real-world commercial operating conditions. As a result, recent studies demonstrate the benefit of tuning catalyst morphology, porosity, and acidity to influence reaction pathway and coke dynamics, which can help predict system efficiency and identify any shortcomings. The implication of this is designing catalysts for the MTO process with intention – and with the priority of heightening key performance indicators at the plant-scale.

On the process side, practical bottlenecks will be solved through the continued translation of lab-scale insights to industrial cases. For instance, further development in the context of catalyst regeneration and downstream process innovations must be made. In terms of the former, regeneration can simultaneously reduce energy demand and waste by-products, whereas the latter could improve the overall carbon efficiency of less immediate concerns in the MTO industry. Altogether, auxiliary improvements are critical in terms of making MTO a more competitive route for olefin production.

Finally, the key pillar to overcome current MTO challenges and unlock its full potential as an alternative to traditional olefin production is sustainability. With the basic chemical industry striving for reduced GHG emissions, the ability of MTO to harness renewable feedstocks (and even serve as a carbon sink by integrating CCU-derived sources) uniquely positions this technology as a circular solution frontrunner in its own right. Further assessments will need to be made to both quantify and analyse the current life-cycle environmental detriments and future benefits of transitioning to a low-carbon electricity grid, of improving water and energy use, or of sourcing greener feedstocks. In any of these cases, leveraging SAPO-34 catalyst design in tandem with efficient industry practices will ensure that MTO remains commercially viable and becomes environmentally reliable.

Author contributions

Ralph Al Hussami: conceptualization; investigation; methodology; data curation; formal analysis; visualization; writing – original draft; validation. Mohammad Ghavipour: conceptualization; investigation; methodology; data curation; formal analysis; visualization; writing – original draft; validation; Galal



Nasser: conceptualization; investigation; methodology; data curation; formal analysis; visualization; writing – original draft; validation. Chasty Duah: conceptualization; data curation; formal analysis; visualization; writing – final draft; validation. Shaza Yousef: conceptualization; investigation; data curation; formal analysis; visualisation; writing – original draft; validation; permissions & licensing rights acquisition. Jan Kopyscinski: funding acquisition; project administration; supervision; writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

Abbreviations

AES	Atomic emission spectroscopy
ANN	Artificial neural network
BAS	Brønsted acidic sites
BETE	Biomass to ethylene <i>via</i> ethanol-to-ethylene
BMTO	Biomass to olefins <i>via</i> methanol-to-olefins
BSE	Back scattered electrons
CC	Catalytic cracking
CCS	Carbon capture and storage
CCU	Carbon capture and utilisation
CFCs	Chlorofluorocarbons
CFTO	Coal to olefins <i>via</i> fischer-tropsch to olefins
CGIs	Crystal growth inhibitors
CGMTO	Coke oven gas based methanol to olefins
CHA	Chabazite
CMTO	Coal to methanol to olefins
CPP	Catalytic pyrolytic process
CTAB	Cetyltrimethylammonium bromide
CTO	Coal to olefins
D6R	Double six rings
DAC	Direct air capture
DEA	Diethylamine
DICP	Dalian institute of chemical physics
DME	Dimethyl ether
DMTO	Dalian methanol to olefins
DPA	Dipropylamine
DFT	Density functional theory
EDS	Energy dispersive X-ray spectroscopy
EMIM	1-Ethyl 3-methylimidazolium
ESR	Electron spin resonance
FCC	Fluid catalytic cracking
FTIR	Fourier transform infrared spectroscopy
FTO	Fischer-tropsch to olefins
GC-MS	Gas chromatography mass spectrometry
GHG	Greenhouse gas
HAADF	High angle annular dark field
HCP	Hydrocarbon pool
heptaMB ⁺	Heptamethylbenzenium cation
HMB	Hexamethylbenzene
HMMC	Hexamethyl methylene cyclohexadiene
ICP	Inductively coupled plasma
LAS	Lewis acidic sites

LCA	Life cycle assessment
LH	Langmuir–Hinshelwood
LMS	Low medium silicon content
LOS	Light olefin selectivity
LPG	Liquified petroleum gas
LPR	Linear olefin production route
MA	Methylamine
MAS	Magic angle spinning
MB	Methylene blue
MD	Molecular dynamics
MDOH	Methanol and dimethyl ether
ML	Machine learning
MOF	Metal organic framework
MOR	Morpholine
MS	Mass spectrometry
MTO	Methanol to olefins
MTG	Methanol to gasoline
NDE	Nondestructive evaluation
NGMTO	Natural gas based methanol to olefins
NMR	Nuclear magnetic resonance
NMTO	Natural gas to olefins <i>via</i> methanol to olefins
NMVOC	Non-methane volatile organic compounds
NSC	Steam cracking of conventional natural gas
NTO	Natural gas to ethylene
OSDA	Organic structure directing agent
OTO	Oil to ethylene
PDH	Propane dehydrogenation
PEG	Polyethylene glycol
PMBs	Polymethylbenzenes
PMNs	Polymethylnaphthalenes
poly(MN)	Polymethylnaphthalene
polyMBs	Polymethylbenzenes
polyMNs	Polymethylnaphthalenes
PSC	Steam cracking of mixed petroleum
PX	<i>p</i> -Xylene
RVM	Relevance vector machine
RWGS	Reverse water–gas shift
SAPO	Silicoaluminophosphate
SC	Steam cracking
SCR	Selective catalytic reduction
SEM	Scanning electron microscopy
SEM-EDXS	Scanning electron microscopy with energy dispersive X-ray spectroscopy
SSC	Steam cracking of shale gas
TEA	Triethylamine
TEAOH	Tetraethylammonium hydroxide
TeMB	1,2,4,6-Tetramethylbenzene
TEM	Transmission electron microscopy
TGA	Thermogravimetric analysis
TMB	1,2,4-Trimethylbenzene
TOS	Time on stream
TPD	Temperature programmed desorption
TPO	Temperature programmed oxidation
UCC	Union carbide corporation
UOP	Honeywell universal oil products
UV	Ultraviolet
UVB	Ultraviolet B radiation
VGO	Vacuum gas oil
WHSV	Weight hourly space velocity



WMO	Waste to methanol to olefin
XPS	X-ray photoelectron spectroscopy
XRD	X-ray diffraction
XRF	X-ray fluorescence
ZSM	Zeolite socony mobil

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.

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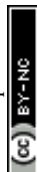
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