

Cite this: *Catal. Sci. Technol.*, 2023,
13, 4020

Zeolite-based catalysts for oxidative upgrading of methane: design and control of active sites

Mizuho Yabushita, ^{*a} Ryota Osuga, ^{*b}
Toshiyuki Yokoi ^c and Atsushi Muramatsu^{*de}

Oxidative upgrading of methane into carbon monoxide, methanol, lower hydrocarbons (such as ethane and ethylene), and other oxygenates has been achieved at relatively low temperatures over zeolite-based catalysts. The high tunability afforded by the framework structures of zeolites offers a unique and precious opportunity for the design of active species, resulting in outstanding catalytic performance. Thus, we categorize the various reported active species in zeolite-based catalysts into three groups in terms of their origin, viz., active species originated from intra-framework heteroatoms, active species introduced by ion-exchange, and active species prepared by impregnation. We present an overview of the performance, structure, and catalysis of each category and comment on the general outlook.

Received 8th April 2023,
Accepted 5th June 2023

DOI: 10.1039/d3cy00482a

rsc.li/catalysis

1. Introduction

Methane, the simplest hydrocarbon, is the main component of natural gas and shale gas. This abundant gaseous compound is currently used as a fuel, while its transformation into platform chemicals in industry has been central to C1 chemistry. The exemplified reactions for methane upgrading are dry/steam reforming into a gaseous mixture of CO and H₂ (*i.e.*, synthesis gas, also called syngas),^{1–3} partial oxidation into syngas,⁴ partial oxidation to methanol and other oxygenates,^{5,6} oxidative/non-oxidative coupling of methane into lower hydrocarbons such as ethane and ethylene,⁷ and dehydroaromatization.^{8,9} Among these reactions, oxidative upgrading is an attractive approach to synthesizing useful compounds directly from methane by skipping the reforming steps, which are endothermic reactions and thus require very high reaction temperatures to achieve high conversion of methane. The extremely high chemical stability of methane, as evidenced by the C–H dissociation energy of *ca.* 440 kJ mol^{–1}, is a common obstacle

and typically necessitates high reaction temperatures for the activation of methane.¹⁰ In addition to the requirement of high energy input, such harsh conditions in the presence of oxidant(s) have another drawback, namely, low selectivity for the desired product(s) because the products are more reactive than methane and readily undergo complete oxidation to CO₂. Thus, the development of catalysts that can greatly activate methane has been desired and investigated extensively for these decades. Furthermore, the shale gas boom (also called the shale gas revolution) that commenced in the USA in the late 2000s has enhanced the research activity for catalytic upgrading of methane worldwide.^{10,11} Because of the ease of continuous operation by flow reactors and separation from unreacted methane and products as well as the high thermal stability, heterogeneous catalysts have attracted much attention.

Among the variety of heterogeneous catalysts reported, such as metal-oxide-based ones,^{10,11} zeolite-based materials, which consist of crystalline and porous frameworks, have great potential to be active, selective, and durable catalysts for methane upgrading. Indeed, the number of publications and citations on methane conversion over zeolite-based catalysts has gradually increased since the 1990s, and as described above, the shale gas boom has boosted this trend (Fig. 1).¹² The unique tunability of zeolites arises from the replacement of intra-framework Si⁴⁺ by heteroatoms (*i.e.*, isomorphous substitution). In particular, the isomorphous substitution by trivalent heteroatoms such as Al³⁺ generates negatively charged frameworks, which require charge-compensating cations. Various cations such as proton and metal cations can be introduced as active sites onto the thus-generated ion-exchange sites. For supported metal

^a Department of Applied Chemistry, School of Engineering, Tohoku University, 6-6-07 Aoba, Aramaki, Aoba-ku, Sendai, Miyagi 980-8579, Japan.

E-mail: m.yabushita@tohoku.ac.jp

^b Institute for Catalysis, Hokkaido University, Kita 21 Nishi 10, Kita-ku, Sapporo, Hokkaido 001-0021, Japan. E-mail: osuga@cat.hokudai.ac.jp

^c Institute of Innovative Research, Tokyo Institute of Technology, 4259 Nagatsuta-cho, Midori-ku, Yokohama, Kanagawa 226-8503, Japan

^d International Center for Synchrotron Radiation Innovation Smart, Tohoku University, 2-1-1 Katahira, Aoba-ku, Sendai, Miyagi 980-8577, Japan.

E-mail: mura@tohoku.ac.jp

^e Institute of Multidisciplinary Research for Advanced Materials, Tohoku University, 2-1-1 Katahira, Aoba-ku, Sendai, Miyagi 980-8577, Japan

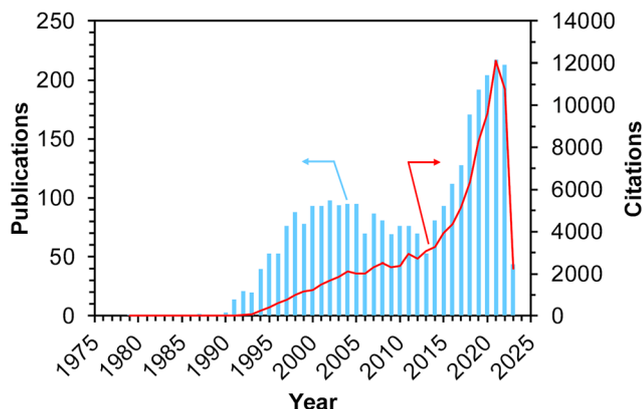


Fig. 1 Number of publications and citations searched with the keywords of “methane + zeolite + catalyst” on Web of Science™ (accessed and searched on April 1st, 2023).¹²

nanoparticles, negatively charged frameworks are beneficial for fixation and isolation of the supported species through the anchoring effect.^{13,14} Likewise, isomorphously introduced species themselves are able to function as active sites. Another important and attractive feature imparting tunability to zeolites is their well-defined porous structures. In some cases, relatively wide nanopores such as cages and channel intersections are also present. Such nanopores of zeolites offer not only high surface area but also the confinement effect,^{13–15} both of which lead to the high dispersion and anti-aggregation nature of supported metal (or metal oxide) species that act as active sites for methane conversion.

In light of these insights, we have categorized the zeolite-based catalysts reported for oxidative upgrading of methane into three groups (Fig. 2): active sites originated from intra-framework heteroatoms (section 2); active sites created in zeolite micropores *via* ion-exchange (section 3); and active sites prepared *via* impregnation on zeolite surfaces (section 4). For each category, we have summarized the performance, structure, and catalysis of representative materials.

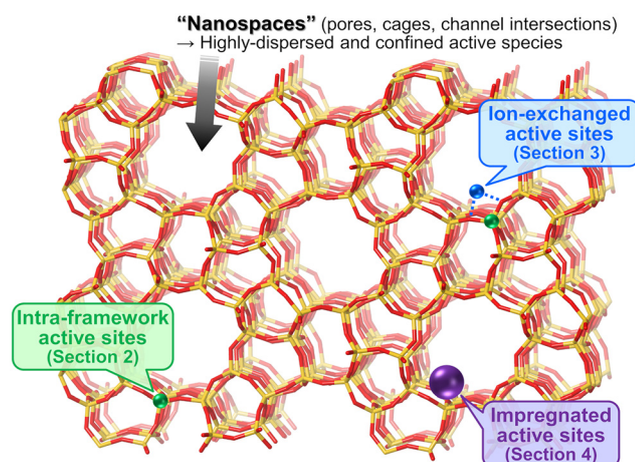


Fig. 2 Approaches to designing zeolite-based catalysts.

2. Active sites originated from intra-framework heteroatoms

A variety of heteroatoms such as Al^{3+} , Ga^{3+} , Fe^{3+} , and Ti^{4+} are known to be introduced into zeolite frameworks *via* isomorphous substitution of Si^{4+} , as mentioned above. Such isomorphously substituting heteroatoms function as catalytic active sites themselves, and cationic species introduced *via* ion-exchange sites also contribute to catalytic reactions (details are described in section 3, except for the case of H^+). In this section, we discuss active sites originated from intra-framework heteroatoms for oxidative upgrading of methane. The exemplified catalysts introduced in this section are summarized in Table 1.

Upon the isomorphous substitution of Si^{4+} by trivalent Al^{3+} and the introduction of H^+ as a charge-compensating cation, Brønsted acid sites are generated in zeolites. Although the activation of methane *via* typical acids is extremely difficult owing to its low proton affinity (544 kJ mol^{-1}) and weak acidity ($\text{p}K_{\text{a}} \sim 40$),¹¹ superacids, which exhibit stronger acidity than 100% sulfuric acid (the Hammett acidity function H_0 of sulfuric acid = -12), have been reported to activate methane to a pentacoordinated carbocation, *i.e.*, $\text{CH}_4 + \text{H}^+$ (superacid) $\rightarrow \text{CH}_5^+ \rightarrow \text{CH}_3^+ + \text{H}_2$.¹⁶ The typical heterogeneous superacid catalyst for methane conversion is sulfated zirconia,¹⁷ but zeolite-based superacid catalysts have also been investigated. In the presence of N_2O as an oxidant, a sample of commercially available H-MOR provided a methane conversion of at most 0.2% at 698 K, possibly owing to its insufficient Brønsted acidity for methane activation ($H_0 = -10.6$), while the fluorination of H-MOR enhanced its acidity ($H_0 = -13$) and more than doubled the methane conversion.¹⁸ In this reaction, the major products at 623–673 K were C2–C4 compounds without the formation of CO or CO_2 (entry 1 in Table 1), probably as a result of the coupling of methane with the surface-generated carbocations containing a pentacoordinated carbon atom as suggested for homogeneous catalytic systems.¹⁶ In contrast, at 698 K, the oxidation of methane and produced hydrocarbons proceeded readily to form CO and CO_2 . In a related study by Spivey *et al.*, the surface of H-MFI (*i.e.*, H-ZSM-5) was modified by $\text{HBr} + \text{AlBr}_3$, a combination known to be a superacid,¹⁹ and the resulting catalyst converted methane into C2, C3, C4, and aromatic compounds in the absence of an oxidant at 573 K.²⁰ However, the methane conversion was only $\sim 1\%$. These reports suggest that the activation of methane remains a great challenge even with state-of-the-art zeolite-based superacid catalysts, owing to the almost inert nature of methane against acid–base reactions. Thus, different activation approaches are much preferred.

We found that the Ga species introduced in the MFI zeolites (Si/Ga molar ratio = 41) by a two-step synthetic procedure called the mechanochemical method, which consists of the mechanochemical treatment of SiO_2 and a heteroatom source to obtain a mixed metal oxide and

Table 1 Catalysts for oxidative upgrading of methane relying on active sites originated from intra-framework heteroatoms, exemplified in section 2^a

Entry	Catalyst	Framework topology	Plausible active site	Oxidant	Reactor type	Solvent	T^b/K	t^c/h	$SV^d/mL h^{-1}$	P^e/MPa	Conv./% product(s)	Target product		Ref.		
												Yield/%	SeL/%			
1	Fluorinated H-MOR	MOR	H ⁺ (superacid site)	N ₂ O	Flow	—	623	—	4.2 × 10 ⁴	n.d.	0.18	C2 compounds	0.17	93	MeOH	18
2	[Ga]-MFI _{MC} ^f	MFI	Ga species	O ₂	Flow	—	923	—	7.5 × 10 ³	0.1	10	CO	8.0	80	CO ₂	22
3	[Ga]-MFI _{Hr} ^f	MFI	Ga species	O ₂	Flow	—	923	—	7.5 × 10 ³	0.1	6.7	CO ₂	6.3	94	CO	22
4	Fe/ZSM-5	MFI	Fe dinuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.05	0.7	MeOH	0.084	12	HCOOH, MeOOH, CO ₂	30
5	Cu-Fe/ZSM-5	MFI	Fe dinuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.05	0.7	MeOH	0.68	85	HCOOH, MeOOH, CO ₂	30
6	Fe-silicalite-1	MFI	Fe dinuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.05	0.3	MeOH	0.051	17	HCOOH, MeOOH, CO ₂	30
7	1.5% Cu-1.5% Fe/ZSM-5	MFI	Fe dinuclear complex	H ₂ O ₂	Flow	H ₂ O	323	—	4.0 × 10 ²	2.0	0.5	MeOH	0.46	92	CO ₂	32
8	Fe-MFI	MFI	Fe species	H ₂ O ₂	Batch	Sulfolane/H ₂ O	323	2	—	3	n.d.	MeOH	n.d.	85	HCOOH, HCHO	35
9	Mn-Na ₂ WO ₄ /Ti-MWW	MWW	W ⁶⁺ species ^g	O ₂	Flow	—	993	—	8.0 × 10 ³	0.1	26	C2 + C3 compounds	20 ^h	76 ^h	CO, CO ₂	36

^a Abbreviations in this table: MeOH = methanol; MeOOH = methyl hydroperoxide. ^b Reaction temperature. ^c Reaction time (only for batch-type reaction). ^d Space velocity (only for flow-type reaction). ^e Reaction pressure. ^f The catalytic performance at various temperatures is shown in Fig. 3. ^g Proposed catalytic cycle consisting of W-, Mn-, and Ti-including species is shown in Fig. 6. ^h These values include both C2 and C3 compounds.

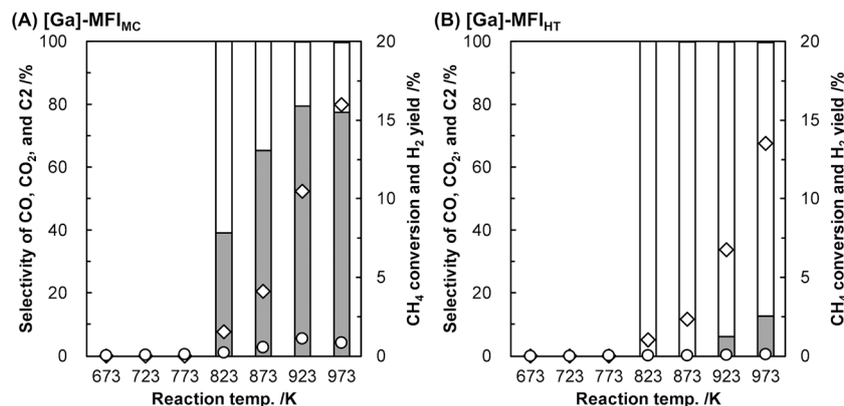


Fig. 3 Oxidative conversion of methane over (A) [Ga]-MFI_{MC} (Si/Ga = 41) and (B) [Ga]-MFI_{HT} (Si/Ga = 37).²² Reaction conditions: CH₄/O₂/Ar = 8.0/2.0/2.5 mL min⁻¹; SV = 7.5 × 10³ mL h⁻¹ g_{cat}⁻¹; 673–973 K; 0.1 MPa. Legends: grey bars = CO selectivity; white bars = CO₂ selectivity; black bars = C₂ selectivity; diamonds = CH₄ conversion; circles = H₂ yield. Although C₂ hydrocarbons were detected, their selectivity was <1%.

subsequent hydrothermal treatment of the resulting mixed metal oxide to synthesize heteroatom-containing zeolites,²¹ exhibited a high selectivity for CO in the oxidative conversion of methane.²² This catalyst (hereafter, [Ga]-MFI_{MC}) produced CO as the main product (80% selectivity) with CO₂ (20%) at 10% conversion of methane at 923 K in the presence of O₂ as an oxidant (Fig. 3A and entry 2 in Table 1). Given the lack of stoichiometry between CO and H₂ (the H₂/CO molar ratio was at most 1.1), this partial oxidation was assumed to proceed *via* an indirect pathway, *i.e.*, the complete oxidation of methane followed by steam/dry reforming reactions and the reverse water-gas shift reaction, rather than the direct partial oxidation pathway. The fact that the Brønsted acidity of Ga-incorporated zeolites is weaker than that of Al-incorporated zeolites^{23,24} indicates that the methane activation over [Ga]-MFI_{MC} occurs in a completely different manner from the cases of superacid catalysts (*vide supra*). Interestingly, the product distribution given by [Ga]-MFI_{MC} was in stark contrast to that obtained using the control specimen prepared by the typical one-step hydrothermal treatment for tetraethyl orthosilicate (TEOS) and Ga(NO₃)₃ ([Ga]-MFI_{HT}), which yielded CO₂ as the main product (Fig. 3B and entry 3 in Table 1). This could arise from the difference in the nature of the Ga species in [Ga]-MFI_{MC} and [Ga]-MFI_{HT}. Our related studies on zeolites synthesized *via* the mechanochemical method^{25,26} suggest that the Ga species in [Ga]-MFI_{MC} are distributed in the particles more uniformly than those in [Ga]-MFI_{HT}, and these well-distributed Ga species result in the unique catalytic activity. For the aromatization of propane, Giannetto *et al.* demonstrated that extra-framework Ga species and Brønsted acid sites in steam-treated [Al,Ga]-MFI zeolites cooperatively functioned as active sites by altering the ratio of extra-framework Ga species to acid sites.²⁷ Stepanov *et al.* revealed *via* isotope tracer analysis relying on solid-state ¹³C cross-polarization/magic angle spinning nuclear magnetic resonance (¹³C CP/MAS NMR) spectroscopy that the dissociative adsorption of methane to form Ga-methyl species (upper route in Fig. 4)

was the major pathway over Ga₂O₃ supported on H-beta, and another activation pathway to generate Ga-methoxy species (lower route in Fig. 4) was also possible but minor.²⁸ On the basis of these reports, the highly dispersed Ga species in [Ga]-MFI_{MC} achieved by the mechanochemical method²² could enhance the formation of binary active sites, extra-framework Ga species and acid sites, leading to the unique catalytic performance of [Ga]-MFI_{MC} for the oxidative conversion of methane.

Fe-containing MFI zeolites were reported as catalysts for the methane-to-methanol (MTM) reaction with the oxidant H₂O₂ under mild conditions.²⁹ As in the case of Ga-incorporated zeolites (*vide supra*), the weak acidity of Fe-incorporated zeolites^{23,24} rules out the possibility of acid-base-type activation of methane. Hutchings *et al.* found that Fe-containing ZSM-5 (Si/Al molar ratio = 15, denoted as Fe/ZSM-5) prepared *via* solid-state ion exchange (SSIE) produced methanol in 12% selectivity along with formic acid, methyl hydroperoxide, and CO₂ in the presence of H₂O₂ at 323 K (entry 4 in Table 1).³⁰ The addition of Cu species to this catalyst suppressed the over-oxidation of methanol to formic acid and CO₂ and enabled the selective production of methanol (85% selectivity, entry 5 in Table 1) because Cu species suppress the formation of 'OH radicals, which cause such undesired over-oxidation.³¹ From *ex situ* X-ray absorption spectroscopy (XAS), ultraviolet-visible (UV-vis) spectroscopy, and density functional theory (DFT) calculations, Hutchings *et al.* identified the dinuclear Fe complex containing antiferromagnetically coupled high-spin

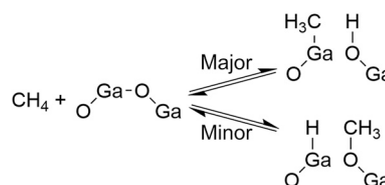


Fig. 4 Methane activation over Ga₂O₃ species supported on H-beta.

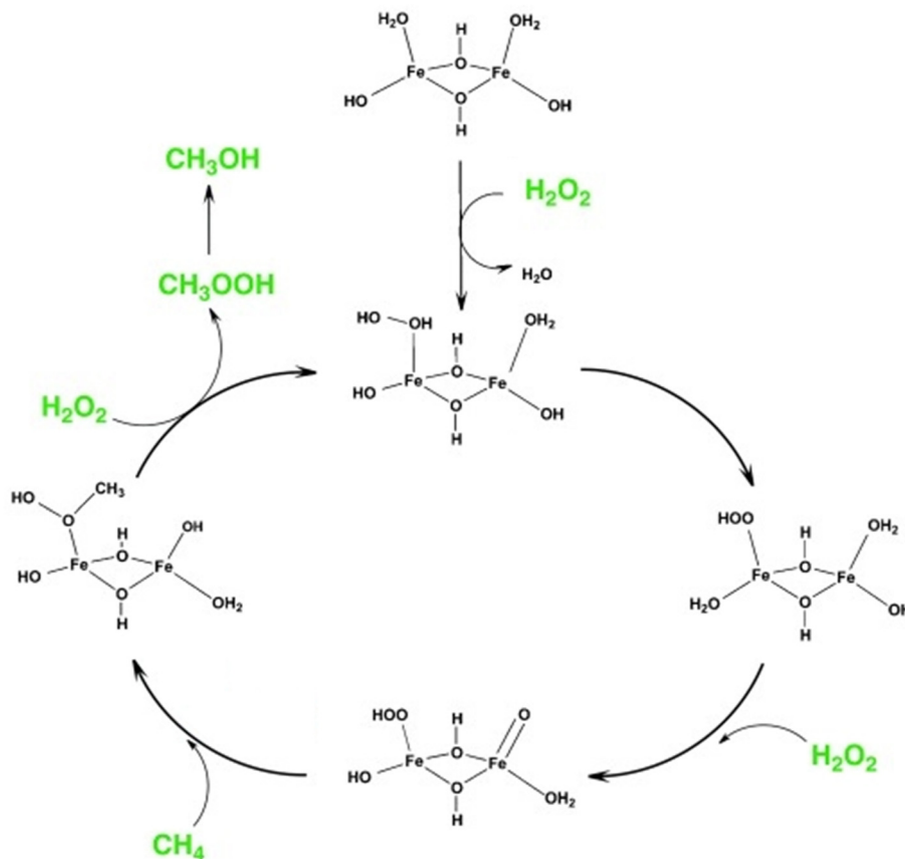


Fig. 5 Proposed reaction mechanism of MTM reaction over the active site of $[\text{Fe}_2(\mu_2\text{-OH})_2(\text{OH})_2(\text{H}_2\text{O})_2]^{2+}$ in Fe-containing ZSM-5. Reprinted from ref. 30 with permission from John Wiley and Sons, copyright 2012.

octahedral Fe^{3+} centers, $[\text{Fe}_2(\mu_2\text{-OH})_2(\text{OH})_2(\text{H}_2\text{O})_2]^{2+}$, as the most likely active site and proposed the catalysis mechanism shown in Fig. 5. This active site seemed to be anchored on the divalent ion-exchange sites (so-called pair sites) of ZSM-5, which allowed it to perform constantly in a continuous operation inside a fixed-bed flow reactor at 323 K (entry 7 in Table 1).³² In the same report, the hydrothermally-synthesized Fe-containing silicalite-1 (silicalite-1 is an Al-free (*i.e.*, pure-silica) MFI-type zeolite) was found to exhibit a catalytic activity similar to Fe/ZSM-5 (entry 6 in Table 1).³⁰ Although the Fe atoms were present in the framework just after hydrothermal synthesis, a part of the Fe species were extracted to form the octahedral Fe species located outside the framework upon the calcination performed to remove the molecules used as an organic structure-directing agent (OSDA). This phenomenon could generate both ion-exchange sites on the remaining intra-framework Fe species and the dinuclear Fe complex $[\text{Fe}_2(\mu_2\text{-OH})_2(\text{OH})_2(\text{H}_2\text{O})_2]^{2+}$. However, spectroscopic discrimination of these two Fe species is quite difficult and the local structure of actual active sites has remained unclear. Previous studies have reported intra-framework Fe species and Fe oligomers as active sites for Fenton-type oxidation of hydrocarbons with H_2O_2 over Fe-containing zeolites.^{33,34} These reports suggest that such sites also are responsible

for the oxidative conversion of methane with H_2O_2 . For the same reaction, one member of our research team investigated solvent effects with H_2O_2 and the hydrothermally synthesized Fe-containing MFI zeolite catalyst in a batch reactor.³⁵ A mixed solvent of 50 vol% sulfolane/ H_2O yielded the highest selectivity for methanol (85%) (entry 8 in Table 1). Given that the formation of CO_2 was negligible in the presence of sulfolane in the catalytic system, this solvent seems to have a similar role to the Cu additive described above (*i.e.*, the suppression of the formation of $\cdot\text{OH}$ radicals).

As mentioned above, intra-framework heteroatoms themselves can be used as catalytic active sites or as precursors to active sites. For the latter scenario, Lu *et al.* employed a Ti-incorporated MWW zeolite as a support for Mn- Na_2WO_4 species,³⁶ which have been known to be outstanding active sites for the oxidative coupling of methane (OCM) since their discovery.³⁷⁻³⁹ The Mn- $\text{Na}_2\text{WO}_4/\text{Ti-MWW}$ catalyst exhibited a lower activation energy (80–100 kJ mol^{-1}) than the typical Mn- $\text{Na}_2\text{WO}_4/\text{SiO}_2$ (180–200 kJ mol^{-1}), enabling the production of C2 and C3 compounds at relatively low reaction temperatures of 993–1073 K (entry 9 in Table 1). Ti-MWW behaved as a source of both SiO_2 and Ti, the former being a typical support for the Mn- Na_2WO_4 catalyst.³⁷⁻³⁹ The Ti species derived from Ti-MWW and the

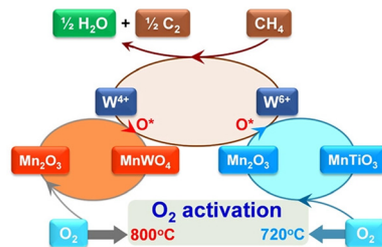


Fig. 6 Proposed catalytic cycle for OCM reaction mediated by the redox cycle $\text{Mn}^{2+} \leftrightarrow \text{Mn}^{3+}$ over $\text{Mn-Na}_2\text{WO}_4/\text{Ti-MWW}$. Reprinted from ref. 36 with permission from The American Association for the Advancement of Science, copyright 2017.

Mn species supplied *via* the incipient wetness impregnation (IWI) method formed the MnTiO_3 phase. In the presence of both CH_4 and O_2 , this new phase triggered the redox cycle of $\text{Mn}^{2+} \leftrightarrow \text{Mn}^{3+}$ and resulted in a high O_2 activation rate (Fig. 6), leading to the low-temperature OCM activity of $\text{Mn-Na}_2\text{WO}_4/\text{Ti-MWW}$.

3. Active sites created in zeolite micropores *via* ion exchange

Ion exchange can offer various types of chemical species in the zeolite micropores, including isolated and multi-nucleus cations, sub-nano clusters, and nano-sized metals (metal oxides). Such metal species exhibit unique and excellent catalytic performance in various catalytic reactions. Here, the active sites on the zeolite surfaces prepared *via* the ion exchange method are introduced and examined with some exemplified studies summarized in Tables 2 and 3.

Among the numerous studies, Cu ion-exchanged zeolites are the most widely investigated zeolite catalyst for methane conversion, where Cu-oxo nanoclusters have emerged as promising active sites for the partial oxidation of methane into methanol.^{40,41} The design of active sites in these catalysts is based on enzymes. Methane monooxygenase (MMO), a naturally occurring enzyme, produces methanol from methane even under ambient conditions.⁴² The active sites of MMOs are the ionic binuclear metal centers Cu and Fe, which possess metal-oxo structures with highly active oxygens (Fig. 7).^{43–45} Industrial application of MMOs for methanol production, however, is not realistic from the viewpoint of cost and reaction efficiency. To overcome these issues, specific nanospaces within the zeolite micropores have attracted much attention because the confinement effect afforded by such nanospaces can stabilize metal and metal oxide clusters. Thus, the MMO-mimetic catalyst design, relying mainly on the ion exchange of Cu species on the zeolite surface, has been one of the mainstreams for the development of zeolite-based catalysts.

In 2005, Groothaert *et al.* reported that Cu ion-exchanged ZSM-5 (Cu-ZSM-5) enabled the selective oxidation of methane to methanol, where $8.2 \mu\text{mol g}^{-1}$ of methanol was obtained with 98% of selectivity (entry 1 in Table 2).⁴⁶ The zeolite, which was pretreated with O_2 at 573 K, converted methane to

methanol at 398 K under flowing methane. In the UV-vis spectra of the O_2 -activated catalyst, a band attributed to the bis(μ -oxo)dicopper species was observed at 22700 cm^{-1} , and the integrated intensity of the band decreased with methane flowing. This result suggests that the bis(μ -oxo)dicopper species are active sites in this reaction. On the other hand, the extraction of methoxy species generated on the zeolite surface as methanol using water is essential and inevitable for completing the reaction cycle, owing to the high adsorption enthalpy of methanol on the active sites (Fig. 8), resulting in low reaction efficiency. Such disadvantages of the chemical looping process are serious and need to be overcome for industrial applications.

Dicopper oxygen clusters have been regarded as primary active species for some time now, but trinuclear copper oxygen clusters in MOR-type zeolites have also been found to behave as active species for the partial oxidation of methane to methanol.⁴⁷ The study cited as ref. 47 employed the chemical looping process, which is almost identical to the process used in ref. 46. In the case of ref. 47, methanol and dimethyl ether were detected as reaction products, and the total yield of methanol and dimethyl ether reached $160 \mu\text{mol g}^{-1}$ (entry 2 in Table 2), an order of magnitude higher than that of related studies reported at the time.^{46,48} Despite the different Si/Al ratios, the total yield of methanol and dimethyl ether was proportional to the Cu concentration (Fig. 9A), suggesting that only one type of active site was formed regardless of the different Cu concentrations. Some spectroscopic and computational approaches proposed that $[\text{Cu}_3(\mu\text{-O})_3]^{2+}$ clusters functioned as active sites and were located at the 8-ring (8R) side pocket of the MOR-type structure (Fig. 9B and C). The authors inferred that the hydrophobic environment provided by the 8R side pockets enhanced the activity because of the presence of a similar hydrophobic environment in MMOs. In this respect, highly active zeolites are thought to require a high local concentration of framework Al atoms at the 8R side pockets. Thus, these results suggest the importance of the location of the framework Al atoms in zeolite frameworks. In 2017, a breakthrough study in the partial oxidation of methane to methanol over Cu ion-exchanged zeolites with a chemical looping process was reported by Sushkevich and van Bokhoven *et al.*⁴⁹ Notably, their study used water molecules as an oxidant for methane. The chemical looping process used for the partial oxidation of methane to methanol consisted of three steps as mentioned above (see also Fig. 8): (i) activation with O_2 , (ii) reaction of methane, and (iii) extraction of methoxy species as methanol from the catalyst surface with water. In contrast, the direct stepwise method was successfully used for converting methane into methanol with $204 \mu\text{mol g}^{-1}$ of productivity and 97% of selectivity in the study cited as ref. 49 (entry 3 in Table 2). The active site was determined to be the “mono”(μ -oxo)dicopper species by using XAS and Fourier transform infrared (FT-IR) measurements combined with DFT calculations. In the first step of the reaction, methane reacted with the mono(μ -oxo)

Table 2 Catalysts for oxidative upgrading of methane to methanol relying on active sites created by ion exchange, exemplified in section 3 (catalyst-weight-based yield or space time yield)^{a,c}

Entry	Catalyst	Framework topology	Plausible active site	Oxidant type	Reactor type	Solvent	T^b/K	t^c/h	$SV^d/mL h^{-1} g_{cat}^{-1}$	P^e/MPa	Conv./%	Target C-based product(s)	Target product			Ref.	
													Yield/ $\mu mol g_{cat}^{-1} h^{-1}$	Yield/ $mmol g_{cat}^{-1} h^{-1}$	Sel./%		By-product(s)
1	Cu-MFI	MFI	Cu dinuclear complex	O ₂	Chemical looping	—	398	—	—	0.1	—	MeOH	8.2	—	98	CO ₂	46
2	Cu-MOR	MOR	Cu trinuclear complex	O ₂	Chemical looping	—	473	—	—	0.1	—	MeOH, DME	160	—	80	CO, CO ₂	47
3	Cu-MOR	MOR	Cu dinuclear complex	H ₂ O	Chemical looping	—	473	—	—	0.7	—	MeOH	204	—	97	CO ₂	49
4	Cu-H-ZSM-5 ^f	MFI	Cu _x O _y	O ₂	Flow	—	483	—	2.4 × 10 ³	0.1	—	MeOH	—	1.81 × 10 ⁻³	70	CO ₂	50
5	Cu-BEA ^g	Beta	Cu dinuclear complex	N ₂ O	Flow	—	593	—	1.2 × 10 ⁴	0.1	1.0	MeOH	—	0.24	72	CO ₂ , C ₂ H ₄ , DME	52
6	5Cu/AEI(Na)-850 ^h	AEI	Cu dinuclear complex	N ₂ O	Flow	—	623	—	1.5 × 10 ⁴	0.1	1.0	MeOH	—	1.64	50	CO, CO ₂ , DME, hydrocarbons	62
7	Cu-H/CHA-FAU ⁱ	CHA	Cu dinuclear complex	N ₂ O	Flow	—	623	—	1.5 × 10 ⁴	0.1	0.6	MeOH, lower olefins	—	0.90	66	CO, CO ₂ , DME	63
8	Cu-H/CHA-Am ⁱ	CHA	Cu dinuclear complex	N ₂ O	Flow	—	623	—	1.5 × 10 ⁴	0.1	0.2	MeOH, lower olefins	—	0.30	57	CO, CO ₂ , DME	63
9	Fe/ZSM-5 ^{SSIE}	MFI	Fe mononuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.0	—	C1 oxygenates	—	21.5	96	CO ₂	65
10	Fe/ZSM-5 ^{IWI}	MFI	Fe mononuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.0	—	C1 oxygenates	—	9.42	97	CO ₂	65
11	Fe/ZSM-5 ^{IE}	MFI	Fe mononuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.0	—	C1 oxygenates	—	12.5	98	CO ₂	65
12	Fe/ZSM-5(TPA)	MFI	Fe mononuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.0	—	C1 oxygenates	—	25.9	—	CO ₂	66
13	Fe/ZSM-5(TPA + Na)	MFI	Fe mononuclear complex	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.0	—	C1 oxygenates	—	200	n.d.	CO ₂	66

^a Abbreviations in this table: MeOH = methanol; DME = dimethyl ether. ^b Reaction temperature. ^c Reaction time (only for batch-type reaction). ^d Space velocity (only for flow-type reaction). ^e Reaction pressure. ^f Reaction data under various conditions are shown in Fig. 11. ^g Water was co-flowed. The reaction data including product selectivity are shown in Fig. 12. ^h Reaction data for TOS of 30 h are shown in Fig. 13G. ⁱ Reaction data for TOS of 190 min are shown in Fig. 14.

Table 3 Catalysts for oxidative upgrading of methane to syngas relying on active sites created by ion exchange, exemplified in section 3 (mole-based yield)

Entry	Catalyst	Framework topology	Plausible active site	Oxidant	Reactor type	T^c/K	$SV^b/mL\ h^{-1}$	P^d/MPa	Conv./%	Target product		Ref.
										Yield/%	Sel./%	
1	Rh-MOR	MOR	Rh sub-nano cluster	O ₂	Flow	873	1.2×10^6	0.1	84	CO	CO	71
2	Rh-MFI[TPA] ^d	MFI	Rh sub-nano cluster	O ₂	Flow	873	1.2×10^6	0.1	86	CO	CO ₂	72
3	Rh-MFI[TPA,Na] ^d	MFI	Rh sub-nano cluster	O ₂	Flow	873	1.2×10^6	0.1	62	CO	CO ₂	72
4	Rh-Y	FAU	Rh cation	O ₂	Flow	873	1.2×10^6	0.1	78	CO	CO ₂	73
5	Rh-ASA	—	Rh oxide	O ₂	Flow	873	1.2×10^6	0.1	86	CO	CO ₂	73

^a Reaction temperature. ^b Space velocity. ^c Reaction pressure. ^d Reaction data at different SV are shown in Fig. 16.

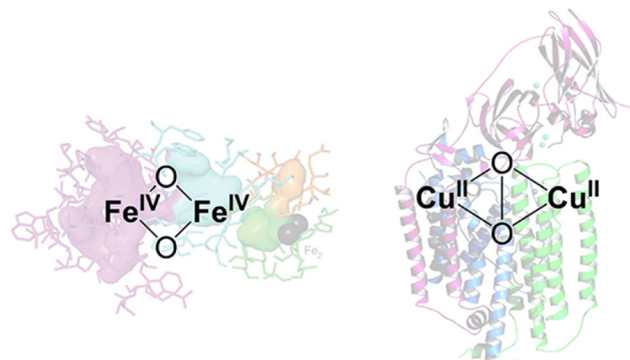


Fig. 7 Proposed Fe- and Cu-based binuclear active sites in MMOs. Reprinted from ref. 45 with permission from American Chemical Society, copyright 2018.

dicopper species at 473 K, resulting in the formation of methoxy species and Brønsted acid sites, accompanied by the reduction of Cu²⁺ to Cu⁺ (Fig. 10). After the reaction, methanol and hydrogen were formed with the regeneration of mono(μ -oxo)dicopper species by flowing water vapor at the same temperature as the flowing methane. In addition, the zeolite was reusable with high activity for at least five cycles.

As mentioned above, the chemical looping process is a powerful method for the partial oxidation of methane to methanol. However, the establishment of a continuous-flow system has been highly desired to realize the MTM reaction in the chemical industry. Román-Leshkov and his co-workers reported the first demonstration of the partial oxidation of methane to methanol using O₂ as an oxidant over Cu ion-exchanged zeolites at low reaction temperatures (483–498 K).⁵⁰ This continuous system was achieved by co-feeding methane, O₂, and water, where Cu ion-exchanged ZSM-5 (Cu-H-ZSM-5) showed high catalytic activity (entry 4 in Table 2). The catalytic activity depended on the framework types of zeolites, and the catalytic yield for methanol reached 491

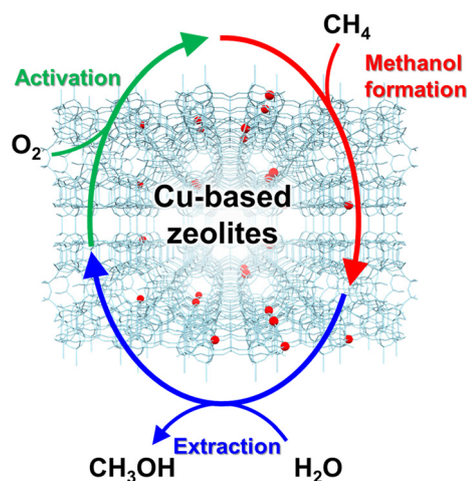


Fig. 8 Schematics of chemical looping process for partial oxidation of methane to methanol over Cu-based zeolites.

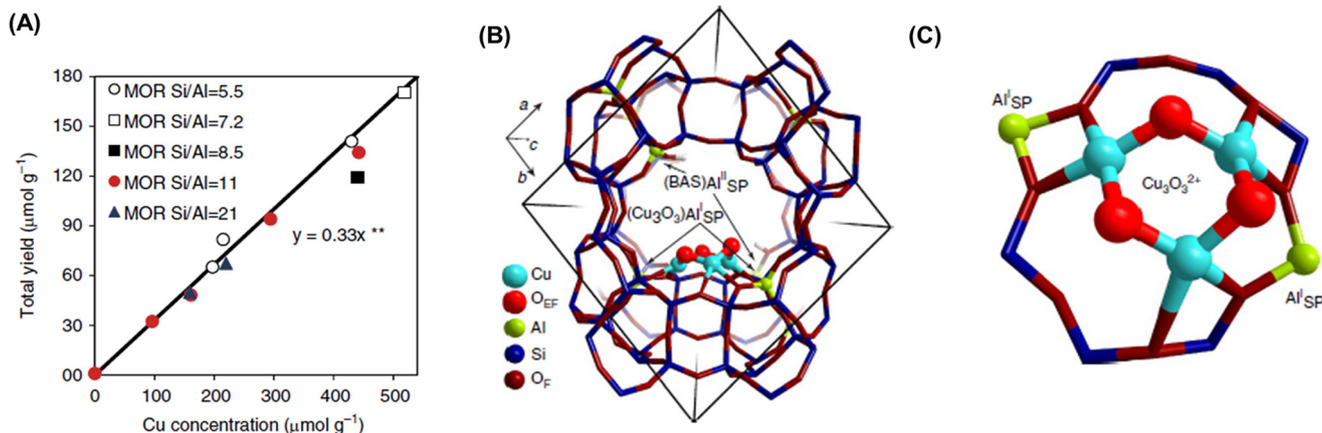


Fig. 9 (A) Total yield of methanol over Cu-MOR with different Si/Al ratios and Cu concentrations. (B) Location and (C) structure of $[\text{Cu}_3(\mu\text{-O})_3]^{2+}$ cluster in MOR determined by DFT calculations. Reproduced from ref. 47 with permission from Springer Nature, copyright 2015.

$\mu\text{mol g}^{-1}$ (Fig. 11). This was the highest value reported by any study at the time, owing to the continuous-flow system. In the study by Román-Leshkov *et al.*, UV-vis spectra revealed the generation of copper oxide species during the reaction; however, the catalytically active sites were not identified. Starting with this report, the partial oxidation of methane to methanol *via* the continuous-flow system has been actively investigated by several research groups.^{41,51,52} O₂ and water have been widely accepted as effective oxidants for the partial oxidation of methane in both the chemical looping and the continuous-flow processes. In addition, stronger oxidants such as N₂O were used for the continuous-flow system. Liu *et al.* reported the continuous methane oxidation to methanol over Cu ion-exchanged beta zeolite (Cu-BEA).⁵² This reaction was boosted by the presence of water; methanol productivity was much higher in the presence of water than in the absence of water (Fig. 12A and B and entry 5 in Table 2). The authors demonstrated that water molecules contributed to the reaction *via* a high-speed proton transfer bridge between the generated CH₃⁻ and OH⁻ moieties (Fig. 12C). The catalytically active sites for this reaction were identified by XAS as the mono(μ -oxo)dicopper species. The

methanol productivity was recorded as $242.9 \mu\text{mol g}_{\text{cat}}^{-1} \text{h}^{-1}$ with 71.6% selectivity for methanol. Moreover, this catalyst showed a long catalyst lifetime (70 h). This excellent catalytic performance achieved in ref. 52 can be considered as one of the benchmarks for analogous systems.

Enormous efforts have been dedicated to achieving a high productivity for the partial oxidation of methane to methanol. The general consensus is that the catalytic active sites are cationic species such as $[\text{Cu}_2(\mu\text{-O})]^{2+}$, $[\text{Cu}_2(\mu\text{-O})_2]^{2+}$, and $[\text{Cu}_3(\mu\text{-O})_3]^{2+}$. To selectively form these clusters in the zeolite micropores, a precise materials design, in particular

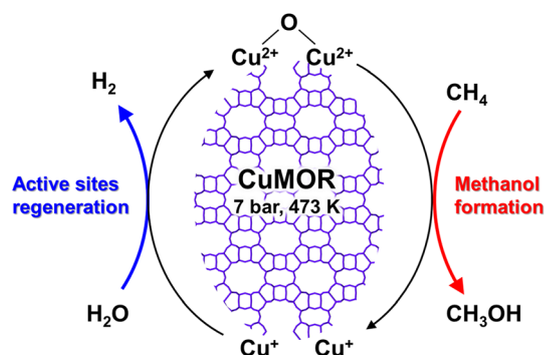


Fig. 10 Plausible reaction scheme for partial oxidation of methane to methanol over Cu-MOR *via* the chemical looping process, using water as an oxidant.

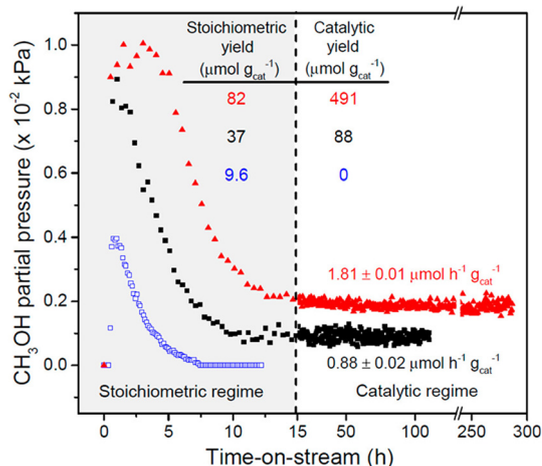


Fig. 11 Catalytic performance of Cu-ZSM-5 in the partial oxidation of methane. Pretreatment conditions: O₂ flow at 823 K for 5 h. Reaction conditions (chemical looping) for Cu-Na-ZSM-5 (blue open squares, Cu/Al = 0.37, Na/Al = 0.26): reaction with methane, WHSV = 2400 mL h⁻¹ g_{cat}⁻¹, 483 K, 0.5 h; extraction of methanol, WHSV = 2400 mL h⁻¹ g_{cat}⁻¹, 483 K, P_{He} = 98.1 kPa, P_{H₂O} = 3.2 kPa, P_{O₂} = 0.0025 kPa. Reaction conditions (continuous-flow) for Cu-Na-ZSM-5 (black filled squares) and Cu-H-ZSM-5 (red filled triangles, Cu/Al = 0.31): WHSV = 2400 mL h⁻¹ g_{cat}⁻¹, 483 K, P_{CH₄} = 98.1 kPa, P_{H₂O} = 3.2 kPa, P_{O₂} = 0.0025 kPa. Reprinted from ref. 50 with permission from American Chemical Society, copyright 2016.

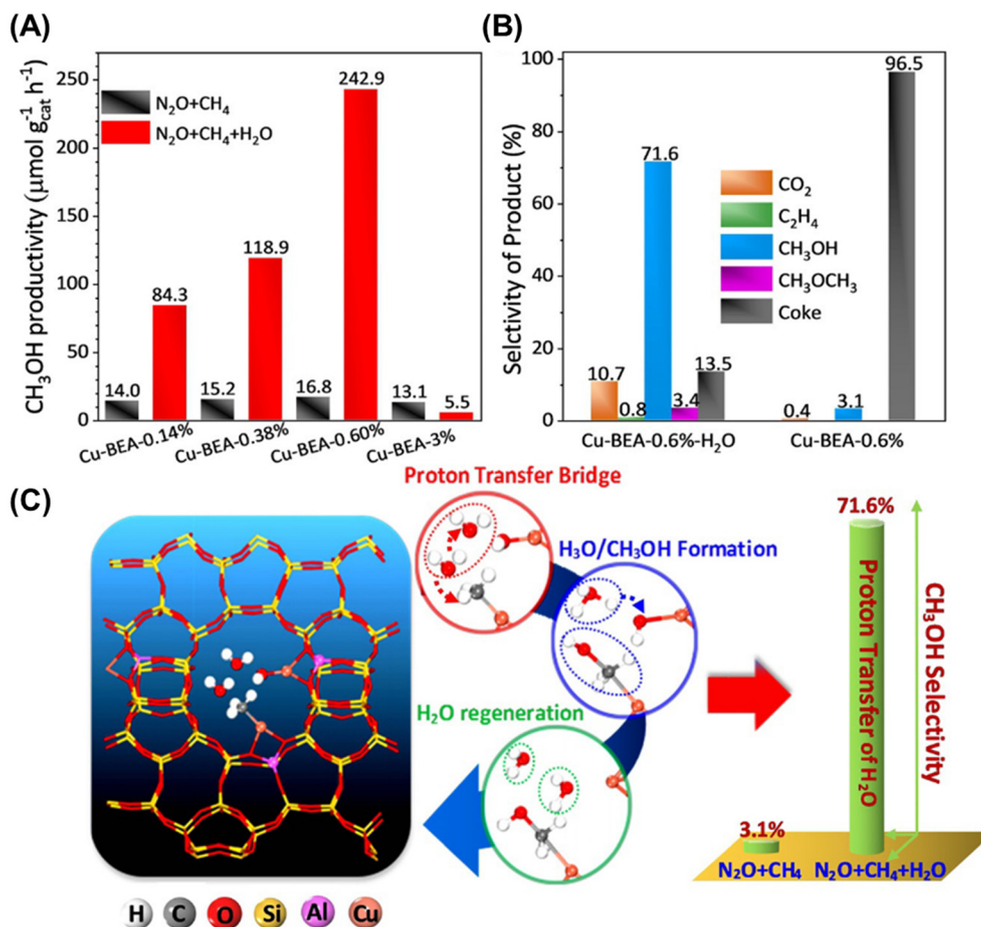


Fig. 12 (A) Methanol productivity of Cu-BEAs for methane to methanol reaction in the presence (red) and absence (black) of H_2O . Reaction conditions: $\text{N}_2\text{O} : \text{CH}_4 : \text{H}_2\text{O} : \text{He} = 30 : 15 : 10(0) : 45(55)$; $\text{GHSV} = 12\,000 \text{ h}^{-1}$; 593 K; TOS = 6 h. (B) Selectivity of products over Cu-BEA under the same reaction conditions as figure A, TOS = 70 h. (C) Schematic of high-speed proton transfer bridge between the generated CH_3^- and OH^- moieties. Reprinted from ref. 52 with permission from John Wiley and Sons, copyright 2021.

the location of the framework Al atoms and the framework structure of the zeolites, is important. Therefore, control of the location of framework Al atoms has been a hot topic in zeolite chemistry over the past several decades.^{53–56} Because the capture of divalent Cu clusters is crucial for the preparation of MMO-mimetic active sites in zeolite micropores, it is essential to construct the framework Al atoms in close proximity to each other, which has been the goal of many studies including some of our own.^{57–61} Recently, we developed a highly active catalyst for the partial oxidation of methane into methanol using N_2O as an oxidant, which was achieved by controlling the position of the framework Al atoms.⁶² OSDAs with/without Na cations strongly influence the position of the framework Al atoms in AEI-type structures; the AEI-type zeolite synthesized with Na cations (denoted as AEI(Na)) had a greater number of paired Al sites compared to that synthesized without Na cations (AEI(Na-free)). Such a paired Al-rich environment generated a relatively larger number of dicopper species ($[\text{Cu}_2(\mu\text{-O})]^{2+}$) *via* ion exchange, resulting in higher catalytic performance (Fig. 13G). Furthermore, the activity and stability of Cu ion-exchanged AEI-type zeolites (5Cu/AEI(Na) and 5Cu/AEI(Na-

free)) were controlled by calcination accompanied by partial dealumination. As a result, 5Cu/AEI(Na) calcined at 1123 K (5Cu/AEI(Na)-850) exhibited excellent catalytic performance, a methanol formation rate of $1638 \mu\text{mol g}_{\text{cat}}^{-1} \text{h}^{-1}$ and a selectivity for methanol of 50% at 623 K (entry 6 in Table 2), with a long catalyst lifetime. After this finding, our group further improved these catalytic systems with a novel approach very recently.⁶³ As already mentioned, methanol is easily sequentially oxidized, owing to its higher reactivity compared to methane. In order to overcome this issue, we attempted to develop a novel catalytic system for the rapid conversion of the produced methanol to lower olefins *via* the methanol-to-olefins (MTO) reaction. Here again, control of the location of the framework Al atoms is crucial because active copper species and Brønsted acid sites need to be placed in juxtaposition for efficient consecutive reactions consisting of MTM and MTO reactions. In this case, CHA-type zeolites were synthesized using amorphous silica (CHA-Am) or FAU-type zeolite (CHA-FAU) as starting materials under OSDA-free conditions. For these samples, different Al distributions were observed: CHA-FAU showed a higher proportion of paired Al sites than CHA-Am. The unique

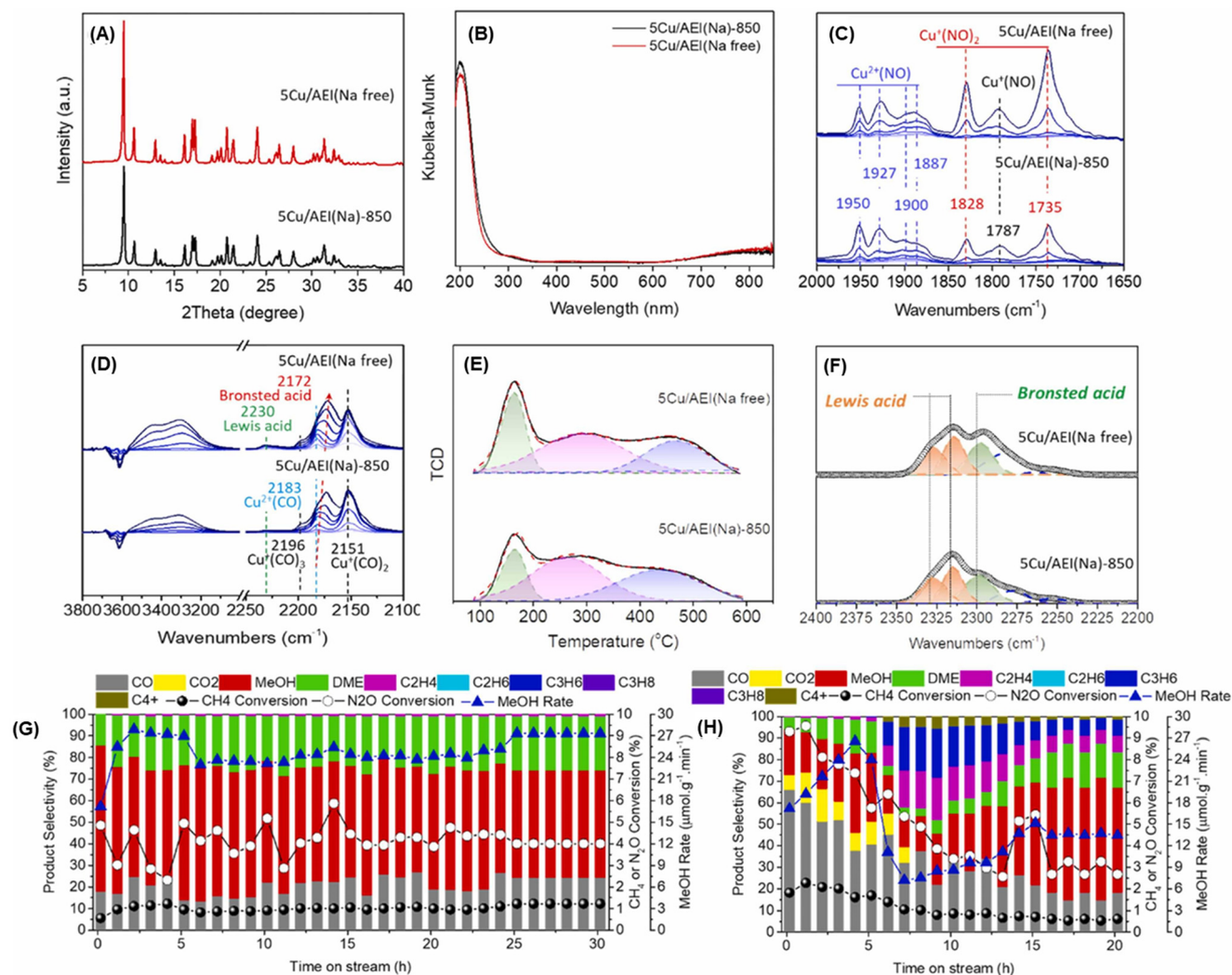


Fig. 13 (A) XRD patterns and (B) UV-vis spectra of 5Cu/AEI(Na)-850 and 5Cu/AEI(Na free). *In situ* FT-IR spectra of adsorbed (C) NO (5–120 Pa) and (D) CO (5–1000 Pa) on Cu ion-exchanged AEI-type zeolites at 153 K. (E) NH_3 -TPD profiles. (F) *In situ* FT-IR spectra of adsorbed CD_3CN species on Cu ion-exchanged AEI-type zeolites at room temperature. Catalytic performance of (G) 5Cu/AEI(Na)-850 and (H) 5Cu/AEI(Na free) in the MTM reaction. Reaction conditions: 100 mg catalysts; $\text{CH}_4/\text{N}_2\text{O}/\text{H}_2\text{O}/\text{Ar} = 10/10/2/3$ mL min^{-1} ; WHSV = 15 000 mL h^{-1} $\text{g}_{\text{cat}}^{-1}$; 623 K. Reprinted from ref. 62 with permission from Elsevier, copyright 2023.

distribution of the framework Al atoms in CHA-FAU facilitated the pairing of active Cu and Brønsted acid sites, which led to a higher selectivity for lower olefins than in the case of CHA-Am (Fig. 14 and entries 7 and 8 in Table 2). This approach based on the distribution of framework Al atoms would open a new door for the development of zeolite-based catalysts for the oxidative upgrading of methane.

Cu ion-exchanged zeolites are widely recognized as promising catalysts for the MTM reaction, while Fe species, which are also found in MMOs,^{43,45} are also promising candidates as catalytically active sites. Yoshizawa *et al.* proposed that Fe, Co, and Ni species are candidates for the MTM reaction on the basis of DFT calculations.⁶⁴ According to these calculation results, the orders of increasing reactivity to C–H bond dissociation and of the selectivity for methanol were estimated to be $\text{CoO}^+-\text{ZSM}-5 < \text{NiO}^+-\text{ZSM}-5 < \text{FeO}^+-\text{ZSM}-5 < \text{CuO}^+-\text{ZSM}-5$ and $\text{FeO}^+-\text{ZSM}-5 < \text{CoO}^+-\text{ZSM}-5 <$

$\text{NiO}^+-\text{ZSM}-5 < \text{CuO}^+-\text{ZSM}-5$, respectively. Weckhuysen and Luo *et al.* reported Fe ion-exchanged ZSM-5 to be an efficient catalyst for the low-temperature oxidation of methane.⁶⁵ In that study, Fe-containing ZSM-5 samples were prepared by one of three methods: IWI, liquid-phase ion exchange (IE), and SSIE. In the cases of $\text{Fe}/\text{ZSM}-5_{\text{IWI}}$ and $\text{Fe}/\text{ZSM}-5_{\text{IE}}$, Fe-derived particles were observed in the scanning transmission electron microscope (STEM) images, while $\text{Fe}/\text{ZSM}-5_{\text{SSIE}}$ did not contain such visible Fe-derived particles (Fig. 15A). In addition, UV-vis and Mössbauer spectroscopies revealed that a larger number of monomeric Fe^{3+} species were present in $\text{Fe}/\text{ZSM}-5_{\text{SSIE}}$ compared to the others (Fig. 15B and C). $\text{Fe}/\text{ZSM}-5_{\text{SSIE}}$ exhibited a remarkably high total yield for the target products (CH_3OH , CH_3OOH , HOCH_2OOH , and HCOOH), as compared with $\text{Fe}/\text{ZSM}-5_{\text{IWI}}$ and $\text{Fe}/\text{ZSM}-5_{\text{IE}}$ (entries 9, 10, and 11 in Table 2). Recently, more highly active Fe ion-exchanged MFI-type zeolites ($\text{Fe}/\text{ZSM}-5$) have been

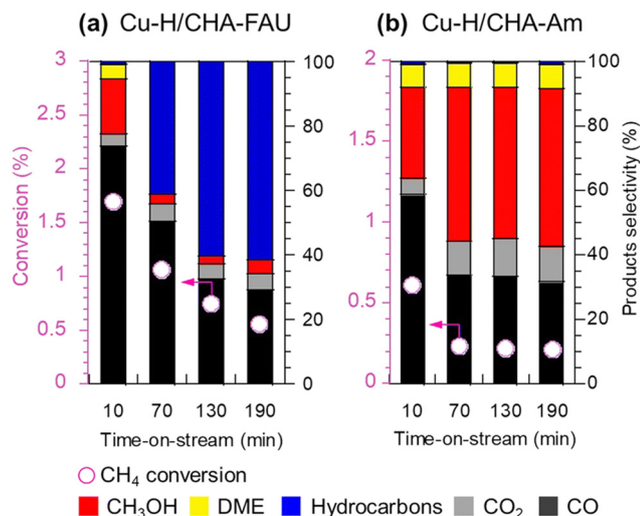


Fig. 14 Catalytic performance of (a) Cu-H/CHA-FAU and (b) Cu-H/CHA-Am in the partial oxidation of methane. Reaction conditions: 100 mg catalyst; $P_{\text{CH}_4}:P_{\text{N}_2\text{O}}:P_{\text{H}_2\text{O}}:P_{\text{Ar}} = 0.4:0.4:0.08:\text{balance}$; total pressure 0.1 MPa; $\text{SV} = 15\,000\text{ mL h}^{-1}\text{ g}_{\text{cat}}^{-1}$; $W/F = 3.7\text{ g}_{\text{cat}}\text{ h mol}_{\text{CH}_4}^{-1}$; 623 K. Reprinted from ref. 63 with permission from Royal Society of Chemistry, copyright 2023.

reported. In these catalysts, the local environment of the framework Al atoms is the key to creating active single Fe

species and suppressing the oligomerization of Fe species.⁶⁶ The MFI-type structure has straight and sinusoidal 10R channels, and the widths of both channels are *ca.* 5.5 Å. In contrast to these channels, a relatively larger spherical space (*ca.* 10 Å) is located at the intersection at these channels. In general, tetrapropylammonium (TPA) hydroxide is used as an OSDA for the synthesis of MFI-type zeolites, and TPA^+ can sit only at the intersection in the MFI-type framework since TPA^+ is too large for the 10R channels. Thus, owing to the electrostatic interaction between TPA^+ and negatively charged building blocks for the zeolite framework, the framework Al atoms are selectively located at the channel intersections when only TPA cations are used for zeolite synthesis.⁶⁷ The catalyst prepared *via* Fe-ion exchange adopted to the ZSM-5 synthesized by using TPA only (denoted as Fe/ZSM-5(TPA)) showed higher catalytic activity than the catalyst prepared *via* Fe-ion exchange for ZSM-5 synthesized by using both TPA and Na cations (Fe/ZSM-5(TPA + Na)) (entries 12 and 13 in Table 2). This clearly indicates the importance of the location of the framework Al atoms (*i.e.*, ion-exchange sites). In addition, the reactive Fe center was confirmed by Fe K-edge XAS measurement, where no oligomeric Fe species were detected. Thus, it was concluded that the active sites of Fe/ZSM-5(TPA) were single Fe species anchored on the ion-exchange sites.

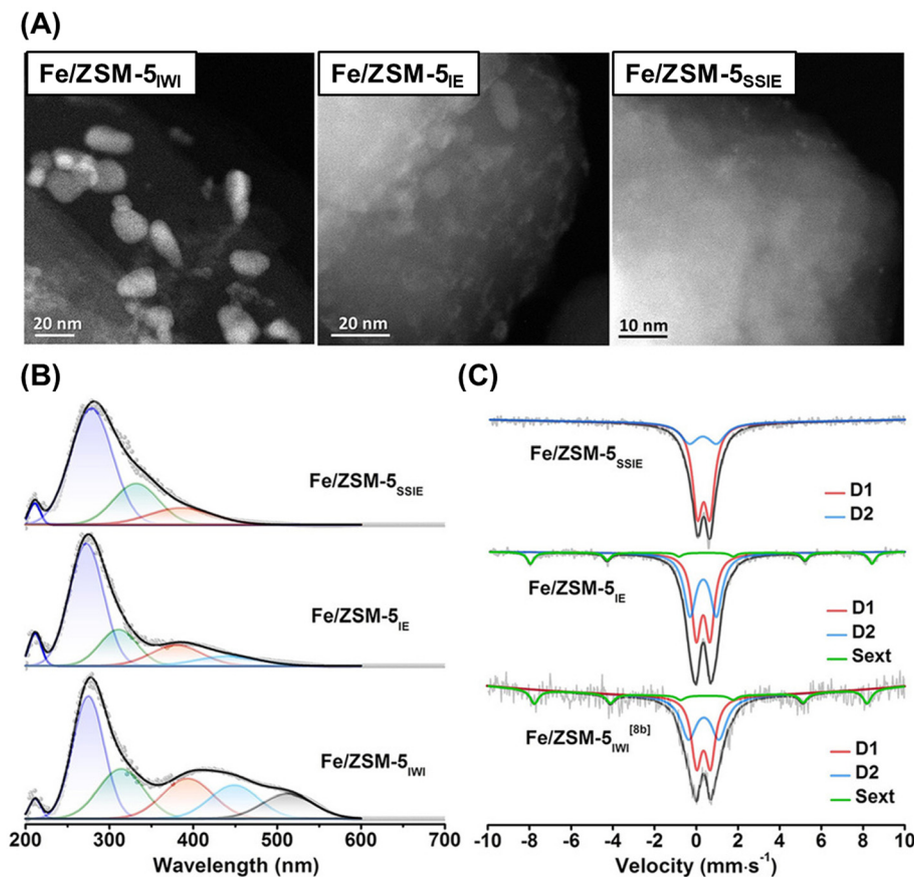


Fig. 15 (A) STEM images, (B) UV-vis spectra, and (C) ^{57}Fe Mössbauer spectra of Fe/ZSM-5s. Reprinted from ref. 65 with permission from John Wiley and Sons, copyright 2021.

Ion-exchanged zeolites are applicable to the oxidation of methane into not only methanol but also other chemicals.^{68–70} Here, we introduce syngas production over Rh ion-exchanged zeolites. Direct steam reforming of methane is a common means of syngas production but requires a high reaction temperature because the reaction is endothermic. By contrast, the partial oxidation of methane into syngas has great potential in terms of achieving a catalytic system that can be operated at low reaction temperatures since the partial oxidation of methane is exothermic. Hou *et al.* prepared a MOR-type zeolite-supported Rh sub-nano cluster (Rh-MOR) *via* ion exchange as a highly active catalyst for the low-temperature selective methane oxidation to syngas.⁷¹ This catalyst lowered the reaction temperature, generally 1073 K or higher, to 873 K. Moreover, almost the equilibrium CO yield was obtained even under high space velocity (SV) conditions (entry 1 in Table 3). In this reaction, the Rh sub-nano clusters with a size of *ca.* 0.6 nm, formed *via* ion exchange, served as active sites. Inspired by this research, our team has tried to selectively form such active Rh sub-nano clusters in the zeolite micropore on the basis of the location of the framework Al atoms.⁷² In the case of MFI[TPA], which is analogous to ZSM-5(TPA), the

framework Al atoms are located selectively at the channel intersection. The ion exchange sites, derived from such site-controlled Al species, provide enough space for the formation of Rh sub-nano clusters, because the sizes of the channel intersections and the Rh sub-nano clusters are *ca.* 1.0 and 0.6 nm, respectively, as mentioned above. In addition, no clusters were expected to form in the channels (*ca.* 0.55 nm) owing to the size limitation. In the study cited as ref. 72, the chemical state of the Rh species was characterized by UV-vis and *in situ* FT-IR spectroscopies. The isolated Rh cations were observed for Rh-MFI[TPA,Na], in which the ion-exchange sites were present both in the channels and at the intersections, while Rh oxide clusters were found in Rh-MFI[TPA] (Fig. 16A and B). A significant difference in catalytic activity was found between Rh-MFI[TPA] and Rh-MFI[TPA,Na] even at almost the same Rh loading (*ca.* 0.2 wt%) (Fig. 16C and D, entries 2 and 3 in Table 3): Rh-MFI[TPA] showed greater catalytic performance and higher stability compared to Rh-MFI[TPA,Na], indicating that active Rh clusters of a suitable size were successfully created in the channel intersections owing to the selective sitting of the framework Al atoms in MFI[TPA]. Furthermore, we investigated the role of the zeolite framework in this reaction by using model catalysts.⁷³ To this

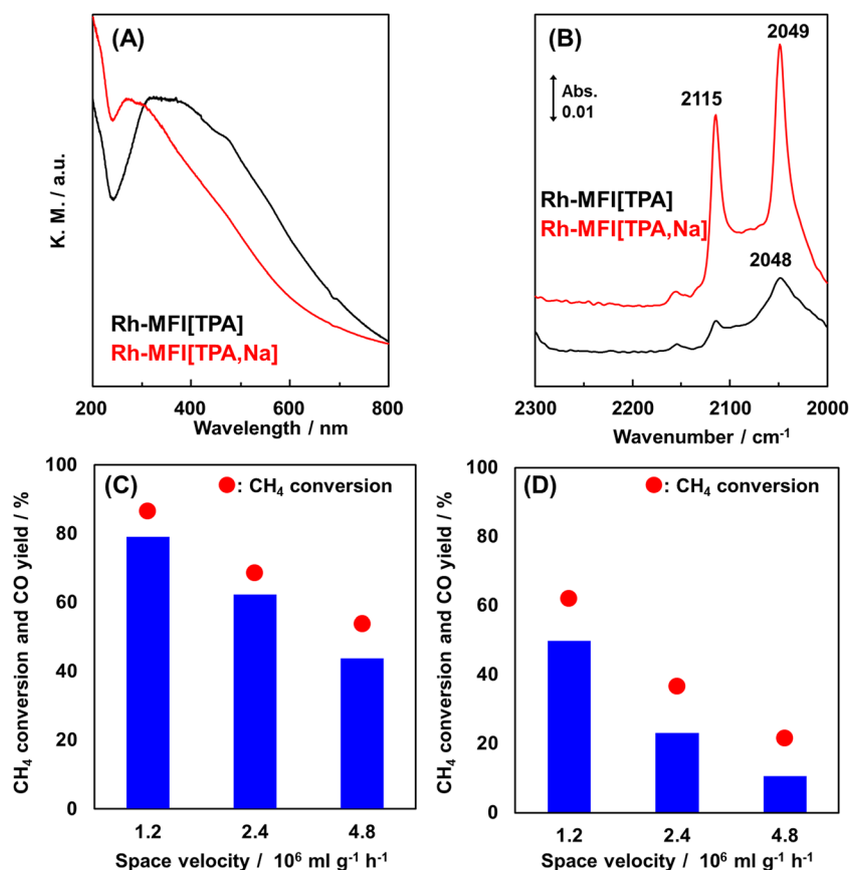


Fig. 16 (A) UV-vis spectra of Rh ion-exchanged MFI-type zeolites. (B) *In situ* FT-IR spectra of adsorbed CO species on Rh-MFI[TPA] and Rh-MFI[TPA,Na] at 298 K. Catalytic performance of (C) Rh-MFI[TPA] and (D) Rh-MFI[TPA,Na] in the partial oxidation of methane at 873 K with different space velocities. Reaction conditions: 10–25 mg catalyst; $P_{\text{CH}_4}:P_{\text{O}_2}:P_{\text{Ar}} = 3.33:1.67:95$; total pressure 0.1 MPa. Reprinted from ref. 72 with permission from Royal Society of Chemistry, copyright 2020.

end, we prepared two kinds of model catalysts: Rh ion-exchanged Y zeolite (Rh-Y) and amorphous silica-alumina (Rh-ASA). The Rh species were characterized by UV-vis and *in situ* FT-IR spectroscopies, transmission electron microscopy (TEM), and H₂ temperature-programmed reduction (H₂-TPR) measurements. Owing to the abundance of ion-exchange sites of Y zeolites, isolated Rh cations were preferentially formed on Rh-Y, while Rh oxide particles (or clusters) were generated on Rh-ASA by the reaction. In the comparison of the catalytic performance of these model catalysts, Rh-ASA showed a slightly higher CO yield than Rh-Y (entries 4 and 5 in Table 3), which is consistent with the results for the Rh-MFI zeolites described above. On the other hand, the stability and catalytic performance of Rh-Y were higher than those of Rh-ASA. In TEM images of the spent catalysts, some aggregated Rh species were observed for Rh-ASA, indicating that the zeolite framework suppressed the aggregation of metal species, owing to the electrostatic interaction between the cations and the anionic field of the zeolite framework (Fig. 17). These results suggest that the active Rh clusters in Rh-MFI[TPA] were preserved by the zeolite framework. In other words, the zeolite framework has two main roles: the creation and preservation of catalytic active sites during the reaction.

4. Active sites prepared *via* impregnation on zeolite surfaces

The impregnation technique, used commonly to prepare various supported metal (or metal oxide) catalysts, has also been widely employed in the field of zeolite-based catalysts. This section discusses zeolite-based catalysts that possess active sites prepared *via* impregnation technique. The zeolite-based catalysts introduced in this section and their catalytic performance for methane upgrading are summarized in Table 4.

The Pb species impregnated over NaX zeolite were reported to be active for the oxidative coupling of methane, producing the C₂ compounds ethane and ethene (entry 1 in

Table 4).⁷⁴ The conversion of methane and C₂ selectivity exhibited a volcano-type dependence on the loading amount of Pb species, the optimum loading amount being 6.4 wt%. At 1073 K, the catalyst with this Pb loading produced a 5.9% yield of C₂ compounds with 69% selectivity. The X-ray diffraction (XRD) patterns of the catalyst before/after the reaction indicated that the redox cycle $\text{Pb}^{2+} \leftrightarrow \text{Pb}^0$ was the key to methane coupling. Control experiments with acidic zeolites were missing in the paper cited as ref. 74, but the lattice oxygen of the basic zeolite, NaX, was proposed to act as a hydrogen acceptor in the reaction. For metal oxide catalysts employed for methane activation, namely oxidative coupling of methane, the importance of their basicity due to lattice oxygens is widely recognized.¹⁰ A part of the metal sources (typically metal cations) can be introduced onto the ion-exchange sites of zeolites even by impregnation, and these species lead to the formation of metal (or metal oxide) nanoparticles anchored on the negatively charged framework during the calcination treatment. In other words, active metal (or metal oxide) species are in proximity to basic lattice oxygens, which is beneficial for methane activation. Thus, the combination of a metal oxide that causes a redox reaction and a zeolite framework with appropriate surface properties is a promising approach to designing catalysts for methane activation.

As for the case of Rh clusters supported on MFI-type zeolite frameworks (*vide supra*), an anchoring effect afforded by negatively charged zeolite frameworks is also useful for isolating small metal clusters/nanoparticles and suppressing their aggregation.^{13,14} NiO particles were prepared on the surface of a CHA-type framework substituted by B, Al, and Ga *via* impregnation and subsequent calcination.⁷⁵ At the same Ni loading amount of 2 wt%, the size of NiO particles was found from XRD to be dependent on a class of framework-substituting atoms, and the NiO particle size increased in the order [Al]-CHA < [Ga]-CHA < [B]-CHA. This order might be connected to the deprotonation energy (*i.e.*, basicity) of these frameworks, given the consistent trend;⁷⁶ thus, the consideration of such parameters could help researchers design and prepare zeolite-based catalysts with supported metal species of a desirable size. The smaller and more highly dispersed NiO particles on [Al]-CHA produced CO and H₂ along with CO₂ in the oxidative conversion of methane at lower reaction temperatures (entry 2 in Table 4), compared to those on [Ga]-CHA and [B]-CHA.⁷⁵

Although it is unclear whether an anchoring effect was at play, ultra-small metal nanoparticles deposited on the MOR-type zeolite were reported to maintain their isolated and highly dispersed state during the partial oxidation of methane and show a long catalyst lifetime. In the first example introduced here, ultra-small NiO particles with a mean diameter of 1.6 nm were successfully deposited on the MOR surface *via* an IWI method; the TEM image for 5 wt% Ni/MOR is shown in Fig. 18A.⁷⁷ This catalyst produced CO (91–92% selectivity) and H₂ with a molar ratio of 2.0 at 97–98% methane conversion at 973 K for a time on stream (TOS)

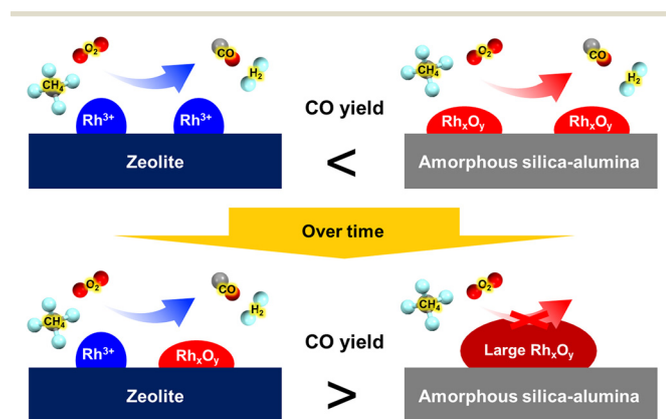


Fig. 17 Schematics for changing behavior of both model catalysts during the reaction. Reprinted from ref. 73 with permission from American Chemical Society, copyright 2021.

Table 4 Catalysts for oxidative upgrading of methane relying on active sites prepared via impregnation on zeolite surfaces, exemplified in section 4^a

Entry	Catalyst	Framework topology	Plausible active site	Oxidant	Reactor type	Solvent	T^b/K	t^c/h	SV ^d /mL h ⁻¹	P^e/MPa	Conv./%	Target C-based product(s)	Target product		Ref.	
													Yield ^d /%	Sel. ^d /%		
1	6.4 wt% Pb/NaX	FAU	PbO species	O ₂	Flow	—	1023	—	3.6×10^3	0.1	8.6	C ₂ compounds	5.9	69	CO ₂	74
2	2 wt% Ni/[Al]-CHA	CHA	Small NiO	O ₂	Flow	—	773	—	1.5×10^4	0.1	22	CO	5.7	26	CO ₂	75
3	5 wt% Ni/MOR ^f	MOR	Ultra-small NiO	O ₂	Flow	—	973	—	3.0×10^4	0.1	98	CO	90	92	CO ₂	77
4	0.005 wt% Rh-3 wt% Co/MOR ^g	MOR	Ultra-small Co	O ₂	Flow	—	923	—	1.2×10^6	0.1	86	CO	78	91	CO ₂	79
5	0.3 wt% Rh/ZSM-5-2D	MFI	Ultra-small Rh	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.0	0.2	MeOOH	0.15	73.2	MeOH, HCOOH	80
6	Pd ₁ O ₄ @ZSM-5 + 2 wt% CuO ^h	MFI	Pd ₁ O ₄	H ₂ O ₂	Batch	H ₂ O	368	0.5	—	3.0	n.d.	MeOH	n.d.	86	MeOOH, CO ₂	81
7 ⁱ	Rh ₁ O ₃ @ZSM-5 ^j	MFI	Rh ₁ O ₃	O ₂	Batch	H ₂ O	423	12	—	6.8	n.d.	AcOH	n.d.	70.1	MeOH, HCOOH	82
8	3 wt% Pd/Ce-MFI _{MC} ^k	MFI	Small Ce species with Pd	O ₂	Flow	—	523	—	7.5×10^3	0.1	3.8	C ₂ H ₆	0.19	4.9	CO ₂	85
9	1 wt% Pd/Ce-MFI _{MC}	MFI	Small Ce species with Pd	O ₂	Flow	—	573	—	7.5×10^3	0.1	3.0	C ₂ H ₆	0.11	3.7	CO ₂	88
10	1 wt% Pd-1 wt% Co/Ce-MFI _{MC}	MFI	Small Ce species with Pd and Co	O ₂	Flow	—	573	—	7.5×10^3	0.1	4.7	C ₂ H ₆	0.16	3.4	CO ₂	88
11	AuPd@ZSM-5-C ₁₆ ^l	MFI	Au-Pd alloy	H ₂ O ₂ ^m	Batch	H ₂ O	343	0.5	—	3.0	17.3	MeOH	16	92	HCOOH, CO ₂	91
12	AuPd/HZSM-5 ⁿ	MFI	Au-Pd alloy	H ₂ O ₂	Batch	H ₂ O	323	0.5	—	3.05	n.d.	MeOH	n.d.	16.9	HCOOH, MeOOH, CO ₂	92

^a Abbreviations in this table: MeOH = methanol; MeOOH = methyl hydroperoxide; AcOH = acetic acid. ^b Reaction temperature. ^c Reaction time (only for batch-type reaction). ^d Space velocity (only for flow-type reaction). ^e Reaction pressure. ^f Reaction data for TOS of 8000 min are shown in Fig. 18B. ^g Reaction data for TOS of 50 h are shown in Fig. 19B. ^h Pd loading was 0.01 wt%. ⁱ CO was also used as a reactant along with methane. ^j Rh loading was 0.10 wt%. ^k Reaction data at different temperatures are shown in Fig. 22B. ^l Au and Pd loading amounts were 3.24 wt% and 1.76 wt%, respectively. ^m *In situ* formed H₂O₂ from H₂ and O₂ over Au-Pd alloy nanoparticles. ⁿ Au and Pd loading amounts were both 0.5 wt%.

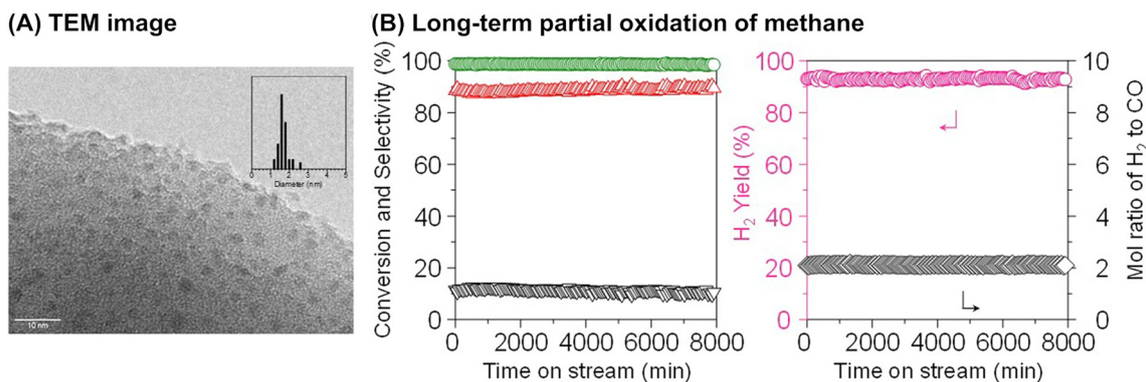


Fig. 18 (A) TEM image of 5 wt% Ni/MOR catalyst and (B) long-term partial oxidation of methane over the Ni/MOR catalyst. Reaction conditions: $\text{CH}_4/\text{O}_2/\text{Ar} = 0.06/0.03/0.91$ atm; $\text{SV} = 3.0 \times 10^4 \text{ mL h}^{-1} \text{ g}_{\text{cat}}^{-1}$; 973 K. Reproduced from ref. 77 with permission from Springer Nature, copyright 2020.

as long as 8000 min (entry 3 in Table 4 and Fig. 18B), in which the lattice oxygen atoms of NiO were suggested to participate (*i.e.*, Mars-van Krevelen mechanism). The high selectivity and long lifetime of this catalyst arose from the ultra-small sized NiO particles. Particle size was

demonstrated to greatly affect catalytic activity, *viz.*, both methane conversion and selectivity for CO; NiO particles larger than 15 nm exhibited at least a 2.4-fold lower methane conversion as compared to NiO particles smaller than 3 nm and accelerated the complete oxidation of methane, yielding

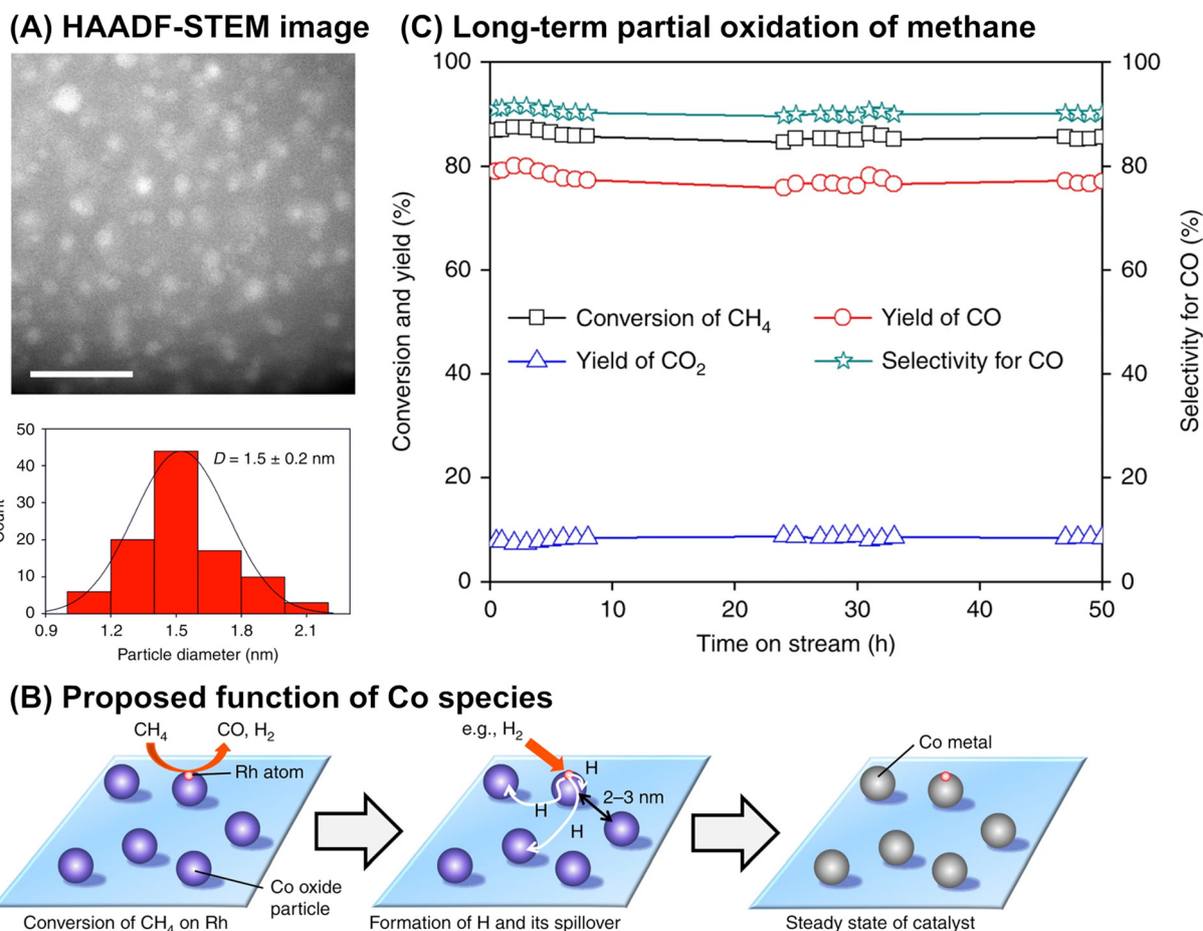


Fig. 19 (A) HAADF-STEM image of 0.005 wt% Rh-3 wt% Co/MOR catalyst, (B) proposed function of Co species for reducing Rh species via hydrogen spillover, and (C) long-term partial oxidation of methane over the Rh-Co/MOR catalyst. Reaction conditions: $\text{CH}_4/\text{O}_2/\text{N}_2 = 3.33/1.67/95$; $\text{SV} = 1.2 \times 10^6 \text{ mL h}^{-1} \text{ g}_{\text{cat}}^{-1}$; 923 K; 0.1 MPa. Reproduced from ref. 79 with permission from Springer Nature, copyright 2018.

CO₂. In addition, methane decomposition into H₂ and carbon nanofibers is known to occur readily on large Ni particles in the absence of O₂.⁷⁸ In contrast to large Ni particles, ultra-small NiO nanoparticles on MOR did not form carbonaceous species, which resulted in a long catalyst lifetime. In the second example, Co nanoparticles with a mean diameter of 1.5 nm were loaded on the MOR surface *via* the standard impregnation method, as seen in the image obtained by a high-angle annular dark-field STEM (HAADF-STEM; Fig. 19A).⁷⁹ This 3 wt% Co/MOR catalyst pre-reduced with H₂ initially produced CO in 91% selectivity at a methane conversion of 7.6% at 873 K in the presence of O₂ but lost this activity for CO production in the steady state, where CO₂ was the main product. This observed alteration of product selectivity resulted from the transformation of Co⁰ into Co²⁺ species, the former of which is necessary for the partial oxidation of methane to CO. In stark contrast, the surface modification of the 3 wt% Co/MOR catalyst with a tiny amount of Rh (0.005 wt%) drastically improved the catalytic performance to yield

CO with 90% selectivity at a methane conversion of 15% even in the steady state under the same reaction conditions. The Rh species were revealed by extended X-ray absorption fine structure (EXAFS) to be mono-atomically dispersed and attached on some of the Co nanoparticles. Such Rh species were proposed to cause the spillover of hydrogen atoms onto the catalyst surface, which allowed Co species to be in the zero-valence state (Fig. 19B). As a result, this 0.005 wt% Rh–3 wt% Co/MOR catalyst exhibited an outstanding lifetime and produced a mixture of CO and H₂ with a molar ratio of 2.0 at an SV as high as 1.2×10^6 mL h⁻¹ g_{cat}⁻¹, as seen in Fig. 19C (see also entry 4 in Table 4).

Along with this anchoring effect, confinement of supported particles in nanospaces consisting of micropores, cages, and/or channel intersections is another attractive approach to keeping supported metal particles isolated and highly dispersed.^{13–15} Subnanometric Rh particles were successfully deposited in the micropores of two-dimensional ZSM-5 nanosheets (ZSM-5-2D).⁸⁰

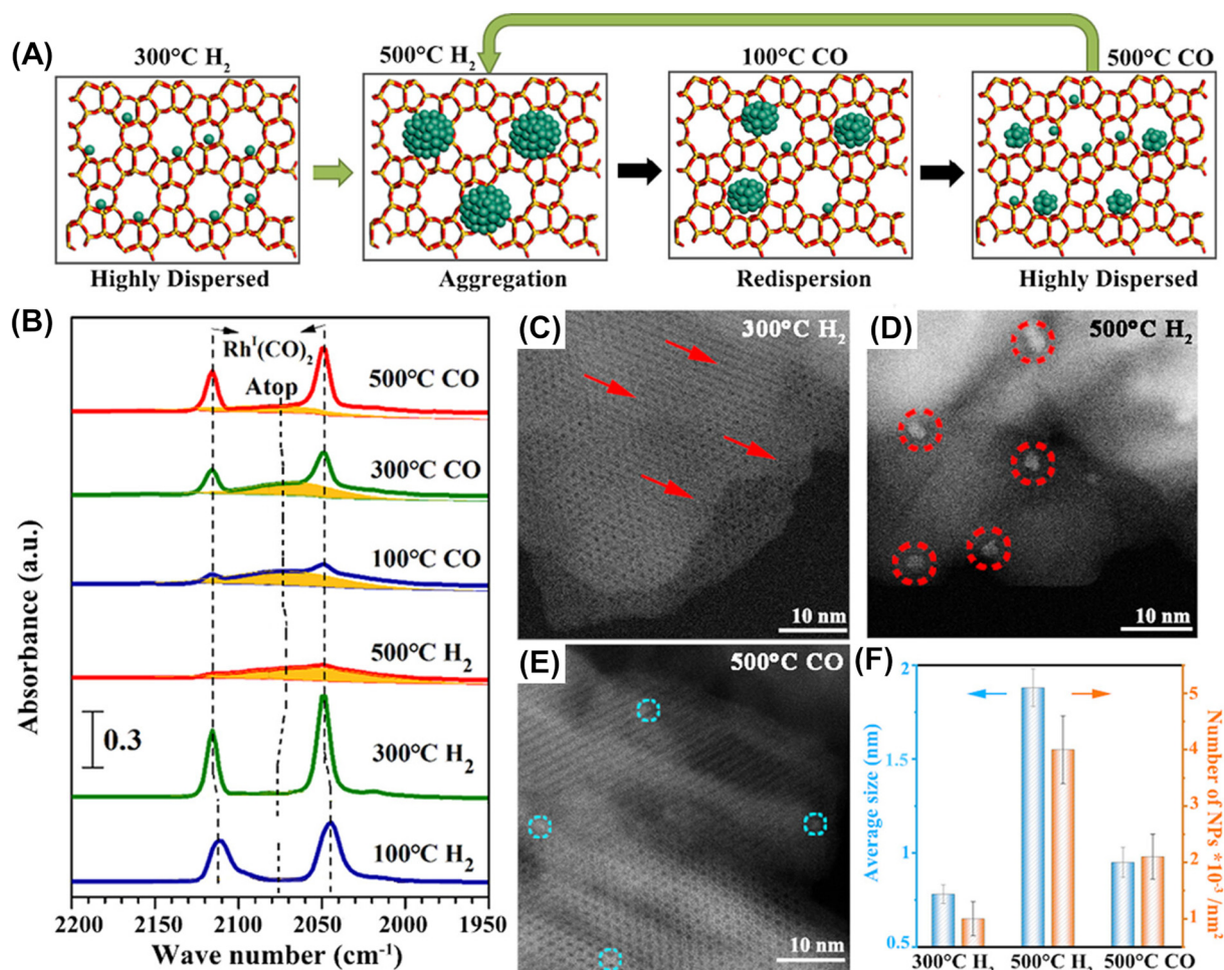


Fig. 20 Reversible size control of Rh nanoparticles ZSM-5-2D. (A) Schematic of reversible size control of Rh particles upon heat treatment in H₂ or CO. (B) CO FT-IR spectra of Rh/ZSM-5-2D heat-treated in H₂ and CO step-by-step. (C–E) HAADF-STEM images of Rh/ZSM-5-2D heat-treated under different conditions. (F) Average particle size and density of Rh nanoparticles on ZSM-5-2D heat-treated under different conditions. Reprinted from ref. 80 with permission from American Chemical Society, copyright 2023.

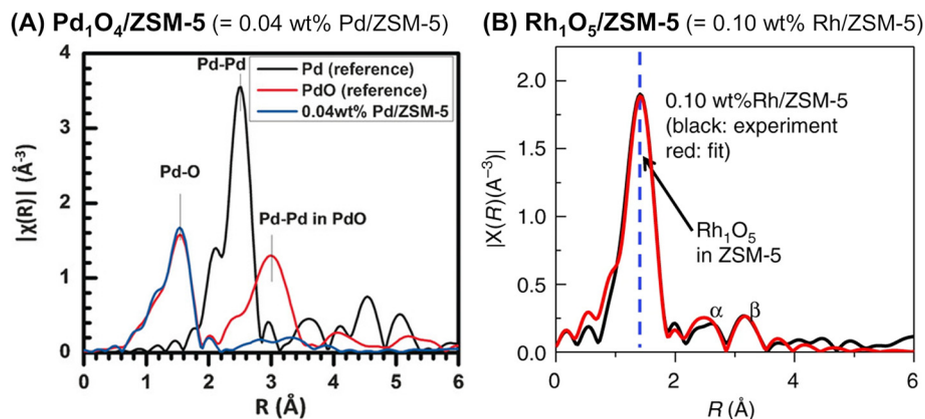


Fig. 21 Fourier transform of EXAFS spectra of (A) Pd₁O₄@ZSM-5 (*k*²-weighted Pd K-edge) and (B) Rh₁O₅@ZSM-5 (*k*²-weighted Rh K-edge). Reprinted from (A) ref. 81 with permission from John Wiley and Sons, copyright 2016, and (B) ref. 82 with permission from Springer Nature, copyright 2018.

Interestingly, their size could be reversibly controlled by heat-treatment in H₂ or CO, which was confirmed by FT-IR after CO adsorption and HAADF-STEM (Fig. 20). The catalytic performance for methane oxidation was dependent on the size of Rh nanoparticles; the subnanometer-sized Rh particles (~0.9 nm) prepared *via* two-step heat treatment in H₂ at 773 K and in CO at 773 K exhibited the highest productivity for methyl hydroperoxide (CH₃OOH) (entry 5 in Table 4).

Ultimately, so-called single-site catalysts can be developed by an appropriate technique for introducing metal species. Tao *et al.* developed two different zeolite-supported single-site catalysts: Pd₁O₄ anchored on the internal surface of ZSM-5 (Pd₁O₄@ZSM-5)⁸¹ and Rh₁O₅ on ZSM-5 (Rh₁O₅@ZSM-5).⁸² In both cases, the pores of the ZSM-5 support were evacuated by a vacuum pump, and then an aqueous solution containing a metal nitrate was

introduced. As in the case of a variety of single-site catalysts,^{83,84} the absence of the second shell of Pd-Pd (or Rh-Rh) in all EXAFS spectra revealed the local structure of the supported Pd and Rh species (Fig. 21). Pd₁O₄@ZSM-5 with CuO and H₂O₂, which were the co-catalyst and oxidant, respectively, converted methane to methanol with 86% selectivity (entry 6 in Table 4). Meanwhile, Rh₁O₅@ZSM-5 accelerated the coupling of methane, CO, and O₂ to produce acetic acid (entry 7 in Table 4).

As mentioned above, zeolite frameworks offer a unique opportunity to form highly dispersed and small species that can activate methane. We recently demonstrated that the combination of the Ce-containing MFI-type zeolite prepared *via* the two-step mechanochemical method and Pd nanoparticles (denoted as Pd/Ce-MFI_{MC}) showed low-temperature activity for oxidative coupling of methane.⁸⁵ The 3 wt% Pd/Ce-MFI_{MC} produced ethane in 0.19% yield at 523 K

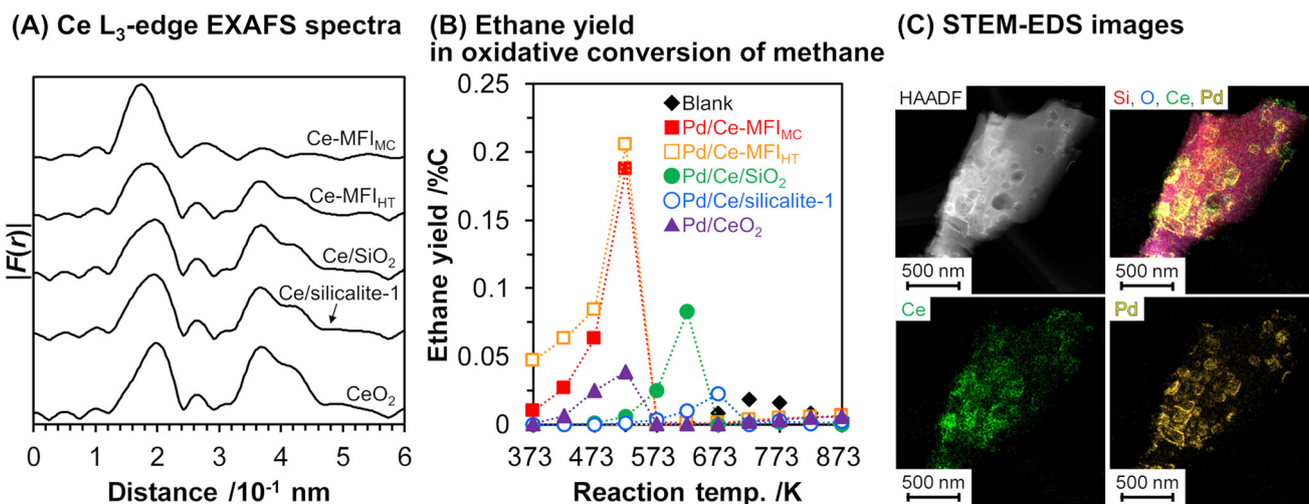
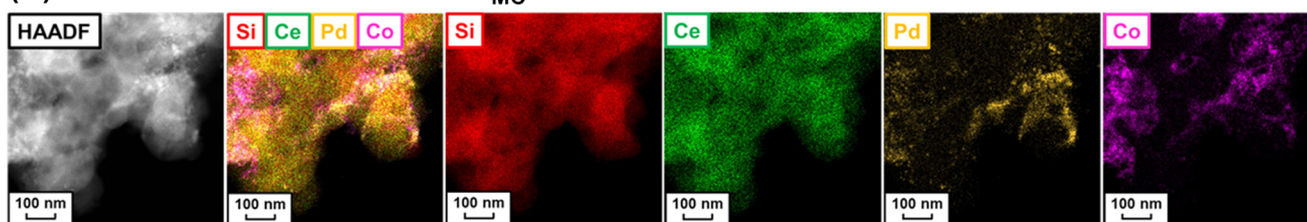


Fig. 22 (A) Fourier transform of *k*³-weighted Ce L₃-edge EXAFS spectra for Ce-containing supports, (B) ethane yield in oxidative conversion of methane over 3 wt% Pd/support, and (C) STEM-EDS images of 3 wt% Pd/Ce-MFI_{MC}. Reaction conditions: CH₄/O₂/Ar₂ = 8.0/2.0/2.5 mL min⁻¹; SV = 7.5 × 10³ mL g_{cat}⁻¹ h⁻¹; 373–873 K; 0.1 MPa. Reproduced from ref. 85 with permission from American Chemical Society, copyright 2021.

(entry 8 in Table 4). The ethane production was probably induced by the redox cycle of CeO_x species mediated by supported Pd, as suggested by H_2 -TPR measurements as well as a previous report where Pd/ZSM-5 without Ce species accelerated the complete oxidation of methane to CO_2 and H_2O .⁸⁶ As seen in the EXAFS spectra in Fig. 22A, Ce-MFI_{MC} did not provide the second shell corresponding to Ce–Ce, while the other Ce-containing supports—Ce-MFI_{HT} (Ce-containing zeolite synthesized *via* typical one-step hydrothermal synthesis), Ce/SiO₂ (SiO₂-supported Ce, prepared by impregnation), Ce/silicalite-1 (silicalite-1-supported Ce, prepared by impregnation), and bulk CeO₂—showed a peak to some extent. These spectra indicated that the size of Ce species increased in the order Ce-MFI_{MC} < Ce-MFI_{HT} < Ce/SiO₂ < Ce/silicalite-1 < CeO₂. Among these supports, Ce-MFI_{MC} and Ce-MFI_{HT} combined with Pd yielded ethane at low reaction temperatures below 573 K (Fig. 22B), indicating the requirement of highly dispersed and small Ce species for the low-temperature oxidative coupling of methane. Owing to the high affinity between Ce and Pd,⁸⁷ Pd species were in proximity to Ce species (STEM images with elemental mapping by energy-dispersive X-ray spectroscopy (EDS) are shown in Fig. 22C) and facilitated the completion of the redox cycle of Ce species for activating methane. Such highly dispersed Ce species with Pd, *i.e.* Pd/Ce-MFI_{MC}, improved catalytic activity through further surface modification by third species. When Co was loaded on Pd/Ce-MFI_{MC}, new active sites, *i.e.*, concerted Ce, Pd, and Co species, were formed (Fig. 23A) and gave a 1.5-fold higher ethane yield than Pd/Ce-MFI_{MC} (entries 9 and 10 in Table 4).⁸⁸ Owing to the poorer dispersion of Ce species in Ce-MFI_{HT} than in Ce-MFI_{MC}, such triad sites were difficult to form (Fig. 23B) and the catalytic activity was not improved.

Encapsulation of active nanoparticles is effective in controlling and maintaining their size and performance.^{89,90} Xiao *et al.* fixed AuPd alloy nanoparticles in ZSM-5 crystals *via* a two-step synthetic method.⁹¹ A mixture of AuPd colloids and TEOS was hydrothermally treated to prepare AuPd nanoparticles coated with amorphous silica. The resulting material was further treated in the presence of TPA hydroxide and boehmite to obtain AuPd nanoparticles encapsulated in ZSM-5 crystals, whose external surface was further modified with organosilanes (AuPd@ZSM-5-C₁₆). The structure of this catalyst differed completely from the control materials prepared *via* the conventional impregnation (AuPd/ZSM-5). This well-designed AuPd@ZSM-5-C₁₆ catalyst converted methane to methanol in the presence of *in situ* generated H₂O₂ from H₂ and O₂ (entry 11 in Table 4). This catalyst was the analogue of the catalyst reported by Hutchings *et al.*,⁹² but the latter was prepared by a conventional co-impregnation method and used only for methane conversion with H₂O₂, not with H₂ and O₂ (entry 12 in Table 4). AuPd alloy nanoparticles had a dual function in this reaction system: the *in situ* formation of H₂O₂ from H₂ and O₂ and methane activation.⁹² Meanwhile, the organosilane modification of ZSM-5 provides hydrophobic external surfaces, which impede the outward diffusion of *in situ* formed H₂O₂ from inside the ZSM-5 crystals. This greatly enhanced the local concentration of H₂O₂ within the crystals, promoting methane conversion. Indeed, methanol productivity with AuPd@ZSM-5 (without surface modification) became 2.8-fold lower than that with AuPd@ZSM-5-C₁₆ (with modification). Metal nanoparticles encapsulated in zeolites have also been employed for other classes of methane conversion such as combustion and dry reforming.^{93,94}

(A) 1 wt% Pd-1 wt% Co/Ce-MFI_{MC}



(B) 1 wt% Pd-1 wt% Co/Ce-MFI_{HT}

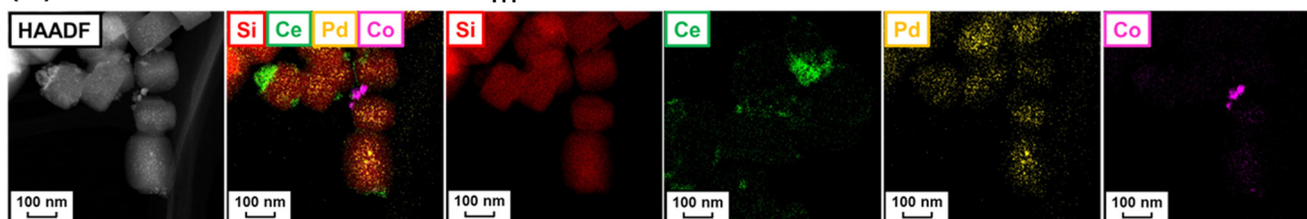


Fig. 23 STEM-EDS images of (A) 1 wt% Pd-1 wt% Co/Ce-MFI_{MC} and (B) 1 wt% Pd-1 wt% Co/Ce-MFI_{HT}. Reproduced from ref. 88 with permission from American Chemical Society, copyright 2022.

5. Summary of advantages and disadvantages of each preparation method

Based on the exemplified catalysts and discussion in the former sections, the advantages and disadvantages of the three preparation methods—incorporation of heteroatoms into frameworks, ion-exchange, and impregnation—are overviewed in this section. These insights are expected to be guidelines for preparation of desired structures and active sites in zeolite-based catalysts for not only oxidative upgrading of methane but also other reactions.

The incorporation of heteroatoms into zeolite frameworks as active sites easily prepares single-site catalysts due to the complete isolation of each introduced heteroatom by inert siliceous frameworks. Meanwhile, metal elements that can be employed in this method are severely limited. Trivalent cations consisting of Al^{3+} and Ga^{3+} are known to be introduced into zeolite frameworks relatively easily, while the complete incorporation of other metal cations into frameworks is quite challenging even in the case of widely investigated Fe^{3+} regardless of the same valence as the trivalent cations described above (typically, intra- and extra-framework Fe species are generated).²⁶ The limited element scope makes the freedom of catalyst design narrower, compared to the other two preparation methods. Besides, one-pot hydrothermal synthesis of heteroatom-incorporated zeolites is sometimes impossible depending on heteroatoms, framework topologies, and molar ratios of Si to heteroatoms. In such situations, post-synthetic incorporation is necessary, making the overall catalyst preparation relatively complicated. Furthermore, the formation of binuclear heteroatom species in the framework-incorporation method is impossible due to the violation of the widely-accepted Loewenstein's rule, which omits the possibility of the formation of M–O–M species (M = heteroatom) in zeolite frameworks due to their thermodynamic instability.⁹⁵ In this respect, this approach is not appropriate for creating binuclear active sites that mimic MMOs.

In contrast to the framework incorporation method, the ion-exchange method is a powerful means of preparing MMO-mimetic binuclear active sites, as introduced in section 3. This method is also useful for preparing highly dispersed metal species since each cationic metal ions are introduced *via* electrostatic interaction with negatively-charged sites originated from isomorphously substituting trivalent heteroatoms. Such electrostatic interaction is beneficial to anchoring introduced ions tightly, leading to the suppression of aggregation of active metal species. Yet, for the introduction of multivalent cationic species *via* the ion-exchange method, the presence of paired heteroatom sites in zeolite frameworks, which can compensate two positive charges, is the prerequisite; otherwise, undesired oxygen-containing ligands inevitably attach cationic metal ions for charge compensation, hampering the formation of well-

designed active sites or rather triggering undesired side reactions.

The impregnation method has the broadest scope of metal elements among the three approaches introduced here; in principle, all metal elements are applicable. However, the introduction of highly-dispersed active metal species precisely into zeolite micropores requires a carefully-handled impregnation method combined with a vacuum technique. Otherwise, metal species are present on outer surfaces of zeolites and the confinement effect by zeolite micropores cannot be expected in such cases.

6. Conclusions and outlook

Zeolite-based materials have a great potential to be highly active and selective catalysts for the oxidative conversion of methane into platform chemicals such as carbon monoxide, methanol, lower hydrocarbons, and other oxygenates, owing to their high tunability. Active sites of zeolite-based catalysts consist of intra-framework heteroatoms, ion-exchanged species (H^+ and metal ions), supported metal or metal oxide nanoparticles, single sites, and combinations thereof. For the preparation and stabilization of these species that function as catalytic active sites, their environment, typically nanospaces (*i.e.*, pores, cages, and channel intersections) of siliceous frameworks, plays a significant role and needs to be tuned properly. In most cases, catalyst regeneration is quite difficult once active sites are aggregated. Therefore, isolation and fixation by both anchoring and confinement effects of nanospaces, which are inaccessible to typical catalyst supports, are among the characteristic and attractive features of zeolite-based materials. To this end, a thorough understanding of the precise design and preparation of zeolite-based catalysts is required.

The conventional *a posteriori* approach relying on screening of each parameter (*e.g.*, catalyst component, content, and preparation technique) is undoubtedly still important and necessary for the development of effective catalysts; however, recent advances in computational chemistry, including DFT calculations and machine learning, provide useful guidelines for catalyst design.^{96–99} For example, previous MMO-mimetic zeolites rely on the preparation of binuclear (or trinuclear) Cu and Fe species on the zeolite surfaces *via* the ion-exchange method as described in section 3. The resulting Cu- and Fe-based active sites successfully convert methane to methoxy species, which is an important intermediate of MTM reaction but also exhibit too high affinity with methoxy species, leading to the requirement of the extraction step of methoxy species with water (Fig. 8). In other words, the chemical looping system or co-feeding of water is inevitably required to complete the reaction cycle over these active sites; namely, in the case of chemical looping, the impossibility of exceeding the turnover number of one is a drawback (*i.e.*, reactions with chemical looping are not catalytic). To establish more simplified and efficient catalytic system, suitable active sites that enable the

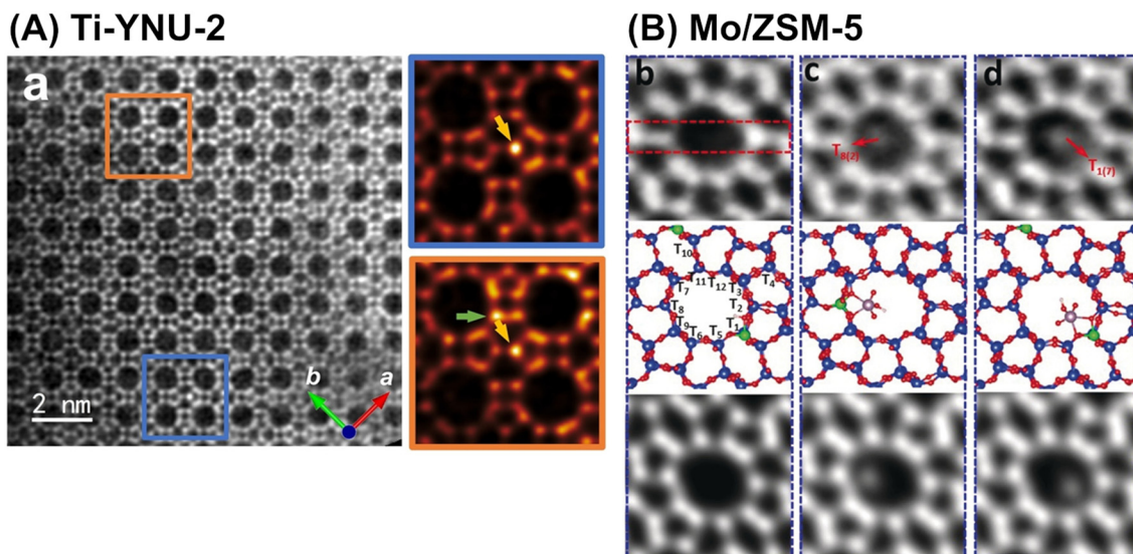


Fig. 24 (A) C_s -corrected STEM LAADF image along (001) incidence of Ti-YNU-2. Two regions colored by blue and orange rectangles with distinct contrast differences are enlarged on the right. Green and golden arrows in the expanded figures represent Ti atoms at the T1/T2 and T7/T8 sites, respectively. Reprinted from ref. 102 with permission from John Wiley and Sons, copyright 2022. (B) iDPC-STEM images of Mo/ZSM-5, where (b–d) are different fields of view for the same sample: top = raw images; middle = calculated structural models; and bottom = simulated projected electrostatic potential. Reprinted from ref. 106 with permission from John Wiley and Sons, copyright 2019.

spontaneous desorption of methoxy species need to be devised. According to computational calculations by Mahyuddin and Yoshizawa, Ni-based binuclear and trinuclear sites (*i.e.*, $[\text{Ni}_2(\mu\text{-O})_2]^{2+}$ and $[\text{Ni}_3(\mu\text{-O})_3]^{2+}$) on ZSM-5 have such potential along with the methane activation;^{64,100} however, such sites have not yet been developed. Such computational-chemistry-driven catalyst design is a promising approach but has remained a grand challenge.

In addition to advanced computational chemistry, state-of-the-art atomic-level characterization techniques help researchers to gain a deeper understanding of the structure–activity relationship for each synthesized catalyst. One of the techniques introduced in this manuscript is high-resolution microscopy. Spherical-aberration-corrected STEM using annular dark field and low angle annular dark field (abbreviated as C_s -corrected STEM ADF and C_s -corrected STEM LAADF, respectively) have been employed to identify the locations (*i.e.*, tetrahedral site (T-site)) of framework-substituting heteroatoms, *e.g.*, Fe atoms in MFI¹⁰¹ and Ti atoms in YNU-2 (Fig. 24A).¹⁰² High-resolution NMR spectroscopy is another powerful tool used for the same objective, as described in many review papers.^{103–105} Integrated differential phase contrast STEM (iDPC-STEM) also enables the imaging of metal species anchored on the inner surfaces of zeolites, as exemplified by Mo species in a ZSM-5 pore (Fig. 24B).¹⁰⁶ Furthermore, dynamic atomic-level imaging by iDPC-STEM directly visualizes organic molecules confined in nanopores; for example, the dynamic adsorption and desorption of pyridine in a ZSM-5 pore have been directly observed.¹⁰⁷ These “seeing-is-believing” techniques are powerful tools for understanding both catalytic active sites and the adsorption states of substrate molecules.

Both microscopy and NMR spectroscopy can provide atomic-level structural information for zeolites as described above, yet these methods are relatively difficult to apply to *in situ/operando* modes, as compared to other characterization techniques. The static structural information of each catalyst is surely important, but dynamic structural analyses offer more direct and detailed insight into catalysis. To reveal the working states of active metal or metal oxide species over zeolites, UV-vis spectroscopy, FT-IR spectroscopy, Raman spectroscopy, and XAS have been employed in *in situ/operando* modes.¹⁰⁸ FT-IR spectroscopy is also useful in terms of characterizing surface-adsorbed species. Among these techniques, *in situ/operando* XAS is the most popular and attractive since it provides both the valence state and local structure of active metal or metal oxide species. For example, Beato *et al.* successfully tracked the gradual changes in Cu active sites over SSZ-13 (*i.e.*, Cu-SSZ-13), a typical catalyst for the MTM reaction (see section 3), by *operando* XAS, monitoring the step-by-step alteration of the environment and temperature necessary to complete the entire reaction process, *i.e.*, activation of Cu species by O_2 , reaction with methane, and subsequent water-mediated extraction of the produced methoxy species as methanol (see Fig. 25).¹⁰⁹ They found that the Cu^{2+} species, after activation by O_2 and flushing with He, partially underwent reduction to Cu^+ , which was successively re-oxidized to Cu^{2+} during the water-assisted extraction of methoxy species. We should note that in the oxidative coupling of methane to lower hydrocarbons, the selectivity for target product(s) is typically low, except when the methane conversion is carefully controlled to be low,^{7,10} and CO and CO_2 are major products. In such situations, *in situ/operando* spectroscopy should provide

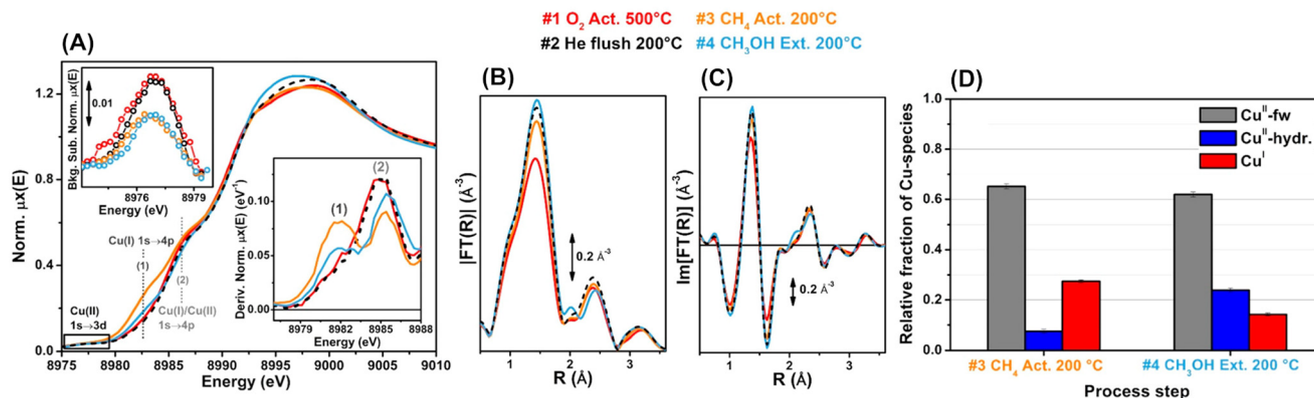


Fig. 25 Operando Cu K-edge XAS spectra of Cu-SSZ-13 after sequential treatments: step 1 (red) = O₂ activation at 773 K; step 2 (black) = He flushing after cooling to 473 K in O₂; step 3 (orange) = CH₄ treatment at 473 K; step 4 (blue) = H₂O-assisted extraction of methoxy species produced on the catalyst. (A) X-ray absorption near edge structure (XANES) spectra, where labels (1) and (2) represent the rising-edge from 1s to 4p peaks that are characteristic of Cu⁺ and Cu⁺/Cu²⁺. (B) Magnitude part of Fourier transform of EXAFS spectra. (C) Imaginary part of Fourier transform of EXAFS spectra. (D) Relative fractions of Cu species estimated via linear combination fitting (LCF) of XANES spectra: Cu^{II}-fw = Cu²⁺ species that interact with the framework; Cu^{II}-hydr. = mobile Cu²⁺ aquo complexes. Reproduced from ref. 109 with permission from American Chemical Society, copyright 2017.

catalytic insights into partial/complete oxidation of methane, not oxidative coupling. Similar situation of low selectivity to target product(s) is also possible in other types of oxidative upgrading of methane due to the ease of over-oxidation of product(s). Therefore, the operation conditions need to be fine-tuned to gain precise and reliable data for each catalytic reaction as well as *in situ/operando* analysis.

Zeolitic materials such as metal–organic frameworks (MOFs) also offer opportunities to tune active sites precisely *via* design of nodes and linkers as well as modifiers.¹¹⁰ Similar to the case of zeolite-based catalysts introduced in section 3, the creation of MMO-mimetic active sites has also been investigated for MOF-based catalysts.¹¹¹ Higher surface area of MOFs compared to zeolites¹¹² enables the preparation of highly dispersed metal nanoparticles even by using typical impregnation technique, which could lead to better productivity per loading amount of metals. Meanwhile, due to the presence of organic linkers, thermal stability of MOFs is inferior to that of zeolites. This limitation of MOFs cannot allow reaction operation at high temperatures, and thus, reaction rates are apt to be low. This fact also points out that fine-tuned sites that are capable of highly activating C–H bond(s) of methane even at low reaction temperatures are always necessary and also that careful selection of reaction conditions is required to suppress thermal degradation of MOFs completely. With respect to the latter issue, isotope tracer analysis using ¹³C-labeled methane as a substrate could be helpful for clearly confirming and distinguishing methane-derived and MOF-derived products. Besides, high selectivity toward target product(s) is mandatory for MOF-based catalysts since even if carbonaceous materials are deposited onto the catalyst surfaces, these catalysts cannot be regenerated *via* simple calcination. In this respect, zeolites and zeolitic materials need to be carefully chosen for one's objective.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

A part of this work was supported financially by the following grants: Core Research for Evolutional Science and Technology of the Japan Science and Technology Agency (JST CREST, JPMJCR16P3); a Grant-in-Aid for Scientific Research (S) (KAKENHI, 21H05011), a Grant-in-Aid for Scientific Research (C) (23K04494), and a Grant-in-Aid for Early-Career Scientists (21K14459) from the Japan Society for the Promotion of Science (JSPS); a research grant from The Japan Petroleum Institute; Kurita Water and Environment Foundation (21A023 and 22K002); Nippon Life Insurance Foundation; and the Ministry of Education, Culture, Sports, Science, and Technology (MEXT) Dynamic Alliance for Open Innovation Bridging Human, Environment and Materials in Network Joint Research Center for Materials and Devices.

References

- D. Li, Y. Nakagawa and K. Tomishige, *Appl. Catal., A*, 2011, **408**, 1–24.
- D. Pakhare and J. Spivey, *Chem. Soc. Rev.*, 2014, **43**, 7813–7837.
- M. Usman, W. M. A. Wan Daud and H. F. Abbas, *Renewable Sustainable Energy Rev.*, 2015, **45**, 710–744.
- B. C. Enger, R. Lødeng and A. Holmen, *Appl. Catal., A*, 2008, **346**, 1–27.
- P. Tomkins, M. Ranocchiari and J. A. van Bokhoven, *Acc. Chem. Res.*, 2017, **50**, 418–425.
- M. Ravi, M. Ranocchiari and J. A. van Bokhoven, *Angew. Chem., Int. Ed.*, 2017, **56**, 16464–16483.

- 7 A. Galadima and O. Muraza, *J. Ind. Eng. Chem.*, 2016, **37**, 1–13.
- 8 J. J. Spivey and G. Hutchings, *Chem. Soc. Rev.*, 2014, **43**, 792–803.
- 9 T. Zhang, *Chem. Sci.*, 2021, **12**, 12529–12545.
- 10 P. Schwach, X. Pan and X. Bao, *Chem. Rev.*, 2017, **117**, 8497–8520.
- 11 R. Horn and R. Schlögl, *Catal. Lett.*, 2015, **145**, 23–39.
- 12 Web of Science, <https://www.webofscience.com/>, (accessed April 2023).
- 13 Y. Wang, C. Wang, L. Wang, L. Wang and F.-S. Xiao, *Acc. Chem. Res.*, 2021, **54**, 2579–2590.
- 14 Q. Zhang, S. Gao and J. Yu, *Chem. Rev.*, 2023, **123**, 6039–6106.
- 15 S.-M. Wu, X.-Y. Yang and C. Janiak, *Angew. Chem., Int. Ed.*, 2019, **58**, 12340–12354.
- 16 N. J. Gunsalus, A. Koppaka, S. H. Park, S. M. Bischof, B. G. Hashiguchi and R. A. Periana, *Chem. Rev.*, 2017, **117**, 8521–8573.
- 17 P. Batamack, I. Bucsí, Á. Molnár and G. A. Olah, *Catal. Lett.*, 1994, **25**, 11–19.
- 18 S. Kowalak and J. B. Moffat, *Appl. Catal.*, 1988, **36**, 139–145.
- 19 D. Farcasiu, S. L. Fisk, M. T. Melchior and K. D. Rose, *J. Org. Chem.*, 1982, **47**, 453–457.
- 20 S. Kanitkar, J. H. Carter, G. J. Hutchings, K. Ding and J. J. Spivey, *ChemCatChem*, 2018, **10**, 5019–5024.
- 21 K. Yamamoto, G. S. E. Borjas, F. Saito and A. Muramatsu, *Chem. Lett.*, 2006, **35**, 570–571.
- 22 M. Yabushita, M. Yoshida, F. Muto, M. Horie, Y. Kunitake, T. Nishitoba, S. Maki, K. Kanie, T. Yokoi and A. Muramatsu, *Mol. Catal.*, 2019, **478**, 110579.
- 23 C. T. W. Chu and C. D. Chang, *J. Phys. Chem.*, 1985, **89**, 1569–1571.
- 24 M. S. Stave and J. B. Nicholas, *J. Phys. Chem.*, 1995, **99**, 15046–15061.
- 25 M. Yabushita, H. Kobayashi, R. Osuga, M. Nakaya, M. Matsubara, S. Maki, K. Kanie and A. Muramatsu, *Ind. Eng. Chem. Res.*, 2021, **60**, 2079–2088.
- 26 R. Osuga, G. Tanaka, M. Yabushita, K. Ninomiya, S. Maki, M. Nishibori, K. Kanie and A. Muramatsu, *Sekiyu Gakkaishi*, 2022, **65**, 67–77.
- 27 A. Montes and G. Giannetto, *Appl. Catal., A*, 2000, **197**, 31–39.
- 28 M. V. Luzgin, A. A. Gabrienko, V. A. Rogov, A. V. Toktarev, V. N. Parmon and A. G. Stepanov, *J. Phys. Chem. C*, 2010, **114**, 21555–21561.
- 29 Y. Liu, L. Wang and F.-S. Xiao, *Chem. Res. Chin. Univ.*, 2022, **38**, 671–676.
- 30 C. Hammond, M. M. Forde, M. H. Ab Rahim, A. Thetford, Q. He, R. L. Jenkins, N. Dimitratos, J. A. Lopez-Sanchez, N. F. Dummer, D. M. Murphy, A. F. Carley, S. H. Taylor, D. J. Willock, E. E. Stangland, J. Kang, H. Hagen, C. J. Kiely and G. J. Hutchings, *Angew. Chem., Int. Ed.*, 2012, **51**, 5129–5133.
- 31 C. Hammond, R. L. Jenkins, N. Dimitratos, J. A. Lopez-Sanchez, M. H. ab Rahim, M. M. Forde, A. Thetford, D. M. Murphy, H. Hagen, E. E. Stangland, J. M. Moulijn, S. H. Taylor, D. J. Willock and G. J. Hutchings, *Chem. – Eur. J.*, 2012, **18**, 15735–15745.
- 32 J. Xu, R. D. Armstrong, G. Shaw, N. F. Dummer, S. J. Freakley, S. H. Taylor and G. J. Hutchings, *Catal. Today*, 2016, **270**, 93–100.
- 33 S. Navalon, M. Alvaro and H. Garcia, *Appl. Catal., B*, 2010, **99**, 1–26.
- 34 P. Xiao, Y. Wang, J. N. Kondo and T. Yokoi, *Appl. Catal., A*, 2019, **579**, 159–167.
- 35 P. Xiao, Y. Wang, T. Nishitoba, J. N. Kondo and T. Yokoi, *Chem. Commun.*, 2019, **55**, 2896–2899.
- 36 P. Wang, G. Zhao, Y. Wang and Y. Lu, *Sci. Adv.*, 2017, **3**, e1603180.
- 37 S. L. X. Fang, J. Lin, J. Gu and D. Yan, *J. Mol. Catal.*, 1992, **6**, 255–262.
- 38 S. L. X. Fang, J. Lin and Y. Chu, *J. Mol. Catal.*, 1992, **6**, 427–433.
- 39 S. Arndt, T. Otremba, U. Simon, M. Yildiz, H. Schubert and R. Schomäcker, *Appl. Catal., A*, 2012, **425–426**, 53–61.
- 40 M. Ravi, V. L. Sushkevich, A. J. Knorpp, M. A. Newton, D. Palagin, A. B. Pinar, M. Ranocchiari and J. A. van Bokhoven, *Nat. Catal.*, 2019, **2**, 485–494.
- 41 L. Tao, I. Lee and M. Sanchez-Sanchez, *Catal. Sci. Technol.*, 2020, **10**, 7124–7141.
- 42 R. S. Hanson and T. E. Hanson, *Microbiol. Rev.*, 1996, **60**, 439–471.
- 43 S.-P. Huang, Y. Shiota and K. Yoshizawa, *Dalton Trans.*, 2013, **42**, 1011–1023.
- 44 J. C. S. Da Silva, R. C. R. Penniford, J. N. Harvey and W. R. Rocha, *Dalton Trans.*, 2016, **45**, 2492–2504.
- 45 K. T. Dinh, M. M. Sullivan, P. Serna, R. J. Meyer, M. Dincă and Y. Román-Leshkov, *ACS Catal.*, 2018, **8**, 8306–8313.
- 46 M. H. Groothaert, P. J. Smeets, B. F. Sels, P. A. Jacobs and R. A. Schoonheydt, *J. Am. Chem. Soc.*, 2005, **127**, 1394–1395.
- 47 S. Grundner, M. A. C. Markovits, G. Li, M. Tromp, E. A. Pidko, E. J. M. Hensen, A. Jentys, M. Sanchez-Sanchez and J. A. Lercher, *Nat. Commun.*, 2015, **6**, 7546.
- 48 E. M. C. Alayon, M. Nachtegaal, E. Kleymentov and J. A. van Bokhoven, *Microporous Mesoporous Mater.*, 2013, **166**, 131–136.
- 49 V. L. Sushkevich, D. Palagin, M. Ranocchiari and J. A. van Bokhoven, *Science*, 2017, **356**, 523–527.
- 50 K. Narsimhan, K. Iyoki, K. Dinh and Y. Román-Leshkov, *ACS Cent. Sci.*, 2016, **2**, 424–429.
- 51 K. T. Dinh, M. M. Sullivan, K. Narsimhan, P. Serna, R. J. Meyer, M. Dincă and Y. Román-Leshkov, *J. Am. Chem. Soc.*, 2019, **141**, 11641–11650.
- 52 R. Xu, N. Liu, C. Dai, Y. Li, J. Zhang, B. Wu, G. Yu and B. Chen, *Angew. Chem., Int. Ed.*, 2021, **60**, 16634–16640.
- 53 M. Yabushita, R. Osuga and A. Muramatsu, *CrystEngComm*, 2021, **23**, 6226–6233.
- 54 R. Osuga and T. Yokoi, in *Crystalline Metal Oxide Catalysts*, ed. W. Ueda, Springer Nature Singapore, Singapore, 2022, pp. 167–196.
- 55 J. Bae and M. Dusselier, *Chem. Commun.*, 2023, **59**, 852–867.

- 56 J. Li, M. Gao, W. Yan and J. Yu, *Chem. Sci.*, 2023, **14**, 1935–1959.
- 57 J. Dědeček, Z. Sobalík and B. Wichterlová, *Catal. Rev.: Sci. Eng.*, 2012, **54**, 135–223.
- 58 J. R. Di Iorio and R. Gounder, *Chem. Mater.*, 2016, **28**, 2236–2247.
- 59 T. Nishitoba, N. Yoshida, J. N. Kondo and T. Yokoi, *Ind. Eng. Chem. Res.*, 2018, **57**, 3914–3922.
- 60 J. R. Di Iorio, S. Li, C. B. Jones, C. T. Nimlos, Y. Wang, E. Kunkes, V. Vattipalli, S. Prasad, A. Moini, W. F. Schneider and R. Gounder, *J. Am. Chem. Soc.*, 2020, **142**, 4807–4819.
- 61 M. Yabushita, Y. Imanishi, T. Xiao, R. Osuga, T. Nishitoba, S. Maki, K. Kanie, W. Cao, T. Yokoi and A. Muramatsu, *Chem. Commun.*, 2021, **57**, 13301–13304.
- 62 P. Xiao, Y. Wang, Y. Lu, T. De Baerdemaeker, A.-N. Parvulescu, U. Müller, D. De Vos, X. Meng, F.-S. Xiao, W. Zhang, B. Marler, U. Kolb, H. Gies and T. Yokoi, *Appl. Catal., B*, 2023, **325**, 122395.
- 63 K. Nakamura, P. Xiao, R. Osuga, Y. Wang, S. Yasuda, T. Matsumoto, J. N. Kondo, M. Yabushita, A. Muramatsu, H. Gies and T. Yokoi, *Catal. Sci. Technol.*, 2023, **13**, 2648–2651.
- 64 M. H. Mahyuddin, A. Staykov, Y. Shiota and K. Yoshizawa, *ACS Catal.*, 2016, **6**, 8321–8331.
- 65 T. Yu, Y. Su, A. Wang, B. M. Weckhuysen and W. Luo, *ChemCatChem*, 2021, **13**, 2766–2770.
- 66 A. Oda, K. Aono, N. Murata, K. Murata, M. Yasumoto, N. Tsunoji, K. Sawabe and A. Satsuma, *Catal. Sci. Technol.*, 2022, **12**, 542–550.
- 67 T. Yokoi, H. Mochizuki, S. Namba, J. N. Kondo and T. Tatsumi, *J. Phys. Chem. C*, 2015, **119**, 15303–15315.
- 68 K. Narsimhan, V. K. Michaelis, G. Mathies, W. R. Gunther, R. G. Griffin and Y. Román-Leshkov, *J. Am. Chem. Soc.*, 2015, **137**, 1825–1832.
- 69 A. M. Rabie, M. A. Betiha and S.-E. Park, *Appl. Catal., B*, 2017, **215**, 50–59.
- 70 Q. Zhang, J. Yu and A. Corma, *Adv. Mater.*, 2020, **32**, 2002927.
- 71 Y. Hou, S. Ogasawara, A. Fukuoka and H. Kobayashi, *Catal. Sci. Technol.*, 2017, **7**, 6132–6139.
- 72 R. Osuga, S. Bayarsaikhan, S. Yasuda, R. Manabe, H. Shima, S. Tsutsuminai, A. Fukuoka, H. Kobayashi and T. Yokoi, *Chem. Commun.*, 2020, **56**, 5913–5916.
- 73 R. Osuga, S. Yasuda, M. Sawada, R. Manabe, H. Shima, S. Tsutsuminai, A. Fukuoka, H. Kobayashi, A. Muramatsu and T. Yokoi, *Ind. Eng. Chem. Res.*, 2021, **60**, 8696–8704.
- 74 N. Davidova, P. Kovacheva and A. H. Weiss, *Catal. Today*, 1992, **13**, 625–626.
- 75 S. Yasuda, Y. Kunitake, R. Osuga, K. Nakamura, T. Matsumoto, K. Sago, J. N. Kondo, M. Yabushita, A. Muramatsu and T. Yokoi, *Chem. Lett.*, 2022, **51**, 46–49.
- 76 R. A. Schoonheydt, P. Geerlings, E. A. Pidko and R. A. van Santen, *J. Mater. Chem.*, 2012, **22**, 18705–18717.
- 77 S. Yasuda, R. Osuga, Y. Kunitake, K. Kato, A. Fukuoka, H. Kobayashi, M. Gao, J. Hasegawa, R. Manabe, H. Shima, S. Tsutsuminai and T. Yokoi, *Commun. Chem.*, 2020, **3**, 129.
- 78 S. Takenaka, S. Kobayashi, H. Ogihara and K. Otsuka, *J. Catal.*, 2003, **217**, 79–87.
- 79 Y. Hou, S. Nagamatsu, K. Asakura, A. Fukuoka and H. Kobayashi, *Commun. Chem.*, 2018, **1**, 41.
- 80 H. Li, Y. Shen, X. Xiao, H. Jiang, Q. Gu, Y. Zhang, L. Lin, W. Luo, S. Zhou, J. Zhao, A. Wang, T. Zhang and B. Yang, *ACS Catal.*, 2023, **13**, 1197–1206.
- 81 W. Huang, S. Zhang, Y. Tang, Y. Li, L. Nguyen, Y. Li, J. Shan, D. Xiao, R. Gagne, A. I. Frenkel and F. Tao, *Angew. Chem., Int. Ed.*, 2016, **55**, 13441–13445.
- 82 Y. Tang, Y. Li, V. Fung, D.-e. Jiang, W. Huang, S. Zhang, Y. Iwasawa, T. Sakata, L. Nguyen, X. Zhang, A. I. Frenkel and F. Tao, *Nat. Commun.*, 2018, **9**, 1231.
- 83 T. Zhang, Z. Chen, A. G. Walsh, Y. Li and P. Zhang, *Adv. Mater.*, 2020, **32**, 2002910.
- 84 L. Fang, S. Seifert, R. E. Winans and T. Li, *Small Methods*, 2021, **5**, 2001194.
- 85 M. Yabushita, M. Yoshida, R. Osuga, F. Muto, S. Iguchi, S. Yasuda, A. Neya, M. Horie, S. Maki, K. Kanie, I. Yamanaka, T. Yokoi and A. Muramatsu, *Ind. Eng. Chem. Res.*, 2021, **60**, 10101–10111.
- 86 A. W. Petrov, D. Ferri, M. Tarik, O. Kröcher and J. A. van Bokhoven, *Top. Catal.*, 2017, **60**, 123–130.
- 87 S. Hosokawa, M. Taniguchi, K. Utani, H. Kanai and S. Imamura, *Appl. Catal., A*, 2005, **289**, 115–120.
- 88 R. Osuga, A. Neya, M. Yoshida, M. Yabushita, S. Yasuda, S. Maki, K. Kanie, T. Yokoi and A. Muramatsu, *Ind. Eng. Chem. Res.*, 2022, **61**, 9686–9694.
- 89 H. Wang, L. Wang and F.-S. Xiao, *ACS Cent. Sci.*, 2020, **6**, 1685–1697.
- 90 C. Gao, F. Lyu and Y. Yin, *Chem. Rev.*, 2021, **121**, 834–881.
- 91 Z. Jin, L. Wang, E. Zuidema, K. Mondal, M. Zhang, J. Zhang, C. Wang, X. Meng, H. Yang, C. Mesters and F.-S. Xiao, *Science*, 2020, **367**, 193–197.
- 92 R. J. Lewis, A. Bara-Estaun, N. Agarwal, S. J. Freakley, D. J. Morgan and G. J. Hutchings, *Catal. Lett.*, 2019, **149**, 3066–3075.
- 93 R. L. Mortensen, H.-D. Noack, K. Pedersen, S. Mossin and J. Mielby, *ChemCatChem*, 2022, **14**, e202101924.
- 94 S. Ma, X. Li, Z. Yang and H. Li, *J. Catal.*, 2023, **417**, 368–378.
- 95 W. Loewenstein, *Am. Mineral.*, 1954, **39**, 92–96.
- 96 T. Toyao, K. Suzuki, S. Kikuchi, S. Takakusagi, K. Shimizu and I. Takigawa, *J. Phys. Chem. C*, 2018, **122**, 8315–8326.
- 97 J. Ohyama, A. Hirayama, N. Kondou, H. Yoshida, M. Machida, S. Nishimura, K. Hirai, I. Miyazato and K. Takahashi, *Sci. Rep.*, 2021, **11**, 2067.
- 98 S. Ma and Z.-P. Liu, *Top. Catal.*, 2022, **65**, 59–68.
- 99 S. Ma and Z.-P. Liu, *Chem. Sci.*, 2022, **13**, 5055–5068.
- 100 M. H. Mahyuddin and K. Yoshizawa, *Catal. Sci. Technol.*, 2018, **8**, 5875–5885.
- 101 A. Mayoral, Q. Zhang, Y. Zhou, P. Chen, Y. Ma, T. Monji, P. Losch, W. Schmidt, F. Schüth, H. Hirao, J. Yu and O. Terasaki, *Angew. Chem., Int. Ed.*, 2020, **59**, 19510–19517.
- 102 J. Li, A. Mayoral, Y. Kubota, S. Inagaki, J. Yu and O. Terasaki, *Angew. Chem., Int. Ed.*, 2022, **61**, e202211196.

- 103 Y. Jiang, J. Huang, W. Dai and M. Hunger, *Solid State Nucl. Magn. Reson.*, 2011, **39**, 116–141.
- 104 S. Li, O. Lafon, W. Wang, Q. Wang, X. Wang, Y. Li, J. Xu and F. Deng, *Adv. Mater.*, 2020, **32**, 2002879.
- 105 X. Zhao, J. Xu and F. Deng, *Front. Chem. Sci. Eng.*, 2020, **14**, 159–187.
- 106 L. Liu, N. Wang, C. Zhu, X. Liu, Y. Zhu, P. Guo, L. Alfilfil, X. Dong, D. Zhang and Y. Han, *Angew. Chem., Int. Ed.*, 2020, **59**, 819–825.
- 107 B. Shen, H. Wang, H. Xiong, X. Chen, E. G. T. Bosch, I. Lazić, W. Qian and F. Wei, *Nature*, 2022, **607**, 703–707.
- 108 D. Kiani, S. Sourav, Y. Tang, J. Baltrusaitis and I. E. Wachs, *Chem. Soc. Rev.*, 2021, **50**, 1251–1268.
- 109 D. K. Pappas, E. Borfecchia, M. Dyballa, I. A. Pankin, K. A. Lomachenko, A. Martini, M. Signorile, S. Teketel, B. Arstad, G. Berlier, C. Lamberti, S. Bordiga, U. Olsbye, K. P. Lillerud, S. Svelle and P. Beato, *J. Am. Chem. Soc.*, 2017, **139**, 14961–14975.
- 110 W.-G. Cui, G.-Y. Zhang, T.-L. Hu and X.-H. Bu, *Coord. Chem. Rev.*, 2019, **387**, 79–120.
- 111 L. S. Andrade, H. H. L. B. Lima, C. T. P. Silva, W. L. N. Amorim, J. G. R. Poço, A. López-Castillo, M. V. Kirillova, W. A. Carvalho, A. M. Kirillov and D. Mandelli, *Coord. Chem. Rev.*, 2023, **481**, 215042.
- 112 J. Liang, Z. Liang, R. Zou and Y. Zhao, *Adv. Mater.*, 2017, **29**, 1701139.