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Catalytic halogenation of methane: a dream reaction with practical scope?

Vladimir Paunović * and Javier Pérez-Ramírez *

Development of catalysts that could surpass the activity and selectivity constraints of the non-catalytic radical-mediated halogenation of methane constitutes a long-standing challenge, which exhibits great potential to valorise this readily available resource for the production of commodities. This study presents comprehensive performance maps of a large library of materials, comprising carriers (quartz, SiO₂, SiC, α-Al₂O₃, γ-Al₂O₃ and carbon), noble metals (Pt, Pd and Ru), metal oxides (Fe₂O₃ and CeO₂), chlorides (PdCl₂ and CuCl₂) and oxyfluorides (TaOF₃) supported on SiO₂, γ-Al₂O₃, carbon or H-ZSM-5 carriers, sulfated systems (S-ZrO₂, S-ZrO₂-SBA-15, S-TiO₂, S-Nb₂O₅, S-Ta₂O₅ and Nafion) and zeolites (3A, H-USY, H-MOR, H-SAPO-34, H-BETA and H-ZSM-5), in the chlorination and bromination of methane under practically relevant conditions and gains insights into the nature of the catalytic effects as a function of the catalyst and halogen of choice. The chlorination activity of different catalyst beds at low temperatures (473–523 K) was 2–5.5 times higher compared to that of the empty reactor of identical volume, while the bromination rate was almost unaffected by the solids in the whole temperature range (643–723 K). Except for zeolites and Pt/carbon, which promoted polyhalogenation, selectivities to halomethanes over most of the catalysts were similar to those in the non-catalytic reactions and were higher in bromination ($S_{\text{CH}_3\text{Br}} = 80\text{--}95\%$ versus $S_{\text{CH}_3\text{Cl}} = 52\text{--}90\%$ at $X_{\text{CH}_4} = 5\text{--}18\%$). The formation of carbon oxides ($S_{\text{CO}_x} = 2\text{--}28\%$) over several materials in chlorination and virtually all systems in bromination implied the decomposition of halomethanes, which at higher temperatures led to coking, particularly in the latter reaction. The kinetic fingerprints along with the marginal impact of the Si:Al ratio, counter ions and extraframework aluminium species on the performance of the most active H-ZSM-5 catalyst indicated that methane chlorination over various materials is governed by the radical-chain mechanism, which limits the scope for breaking the selectivity–conversion relationships by tailoring the catalyst acidity. Nonetheless, the enhancement of chlorination activity over zeolites that followed a volcano-like dependence on their micropore size coupled with a more significant impact of the intracrystalline mesoporosity and crystallite size on the product distribution revealed the important role of confinement effects in this reaction, which may pave the way for advancements in the production of chloromethanes.

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Introduction

The reactions of methane with molecular halogens (X₂, X = F, Cl, Br or I, eqn (1)–(4)) constitute the earliest known routes to functionalize this astonishingly inert hydrocarbon, the history of which dates back to 1840 when the French chemist Jean Baptist Dumas reported the conversion of marsh gas by the action of chlorine (Fig. 1).^{1,2} The extensive kinetic studies of these thermally or photo-activated halogenation processes performed between the 1910s and the 1940s led to the postulation of the radical-chain mechanism that was verified for all halogens and paraffins (Fig. 1).^{3–11} The halogen radicals (X[•])

initially formed *via* the homolytic bond dissociation of X₂ and regenerated through the reaction of alkyl radicals with X₂ play a central role in this reaction network. The differences in stability between the X[•], which increase in the order F[•] < Cl[•] < Br[•] < I[•], cause methane fluorination to be extremely exothermic and often explosive, and chlorination and bromination to display medium and mild exothermicity, respectively, and moderate activity, while iodination is endothermic and equilibrium limited (eqn (1)–(4)).¹ The differences in radicals' reactivity induce substantial variations in the product distribution as a function of the halogen, since mono- (CH₃X), di- (CH₂X₂) and trihalomethanes (CHX₃) could react in an analogous fashion to methane leading to polyhalogenation. In contrast to fluorination which inevitably leads to a complete hydrogen substitution, chlorination could provide a high selectivity to monohalomethanes if operated at a relatively high methane-

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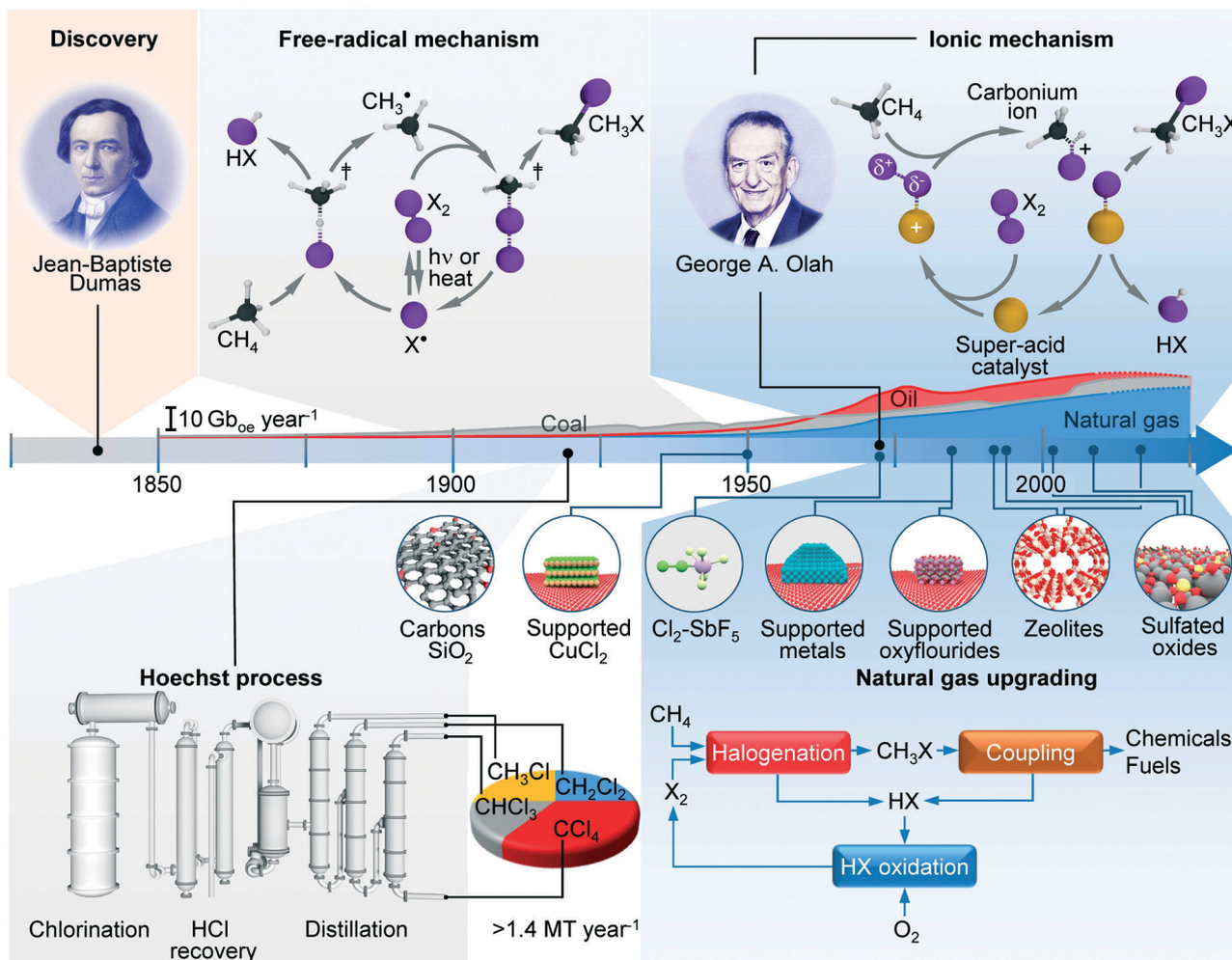
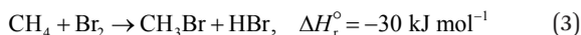


Fig. 1 Chronological overview of the major developments in methane halogenation presented along with the evolution in production of different carbon resources (X = Cl, Br).

to-chlorine ratio (typically molar $\text{CH}_4:\text{Cl}_2$ ratio > 10), although CH_2Cl_2 and CHCl_3 are formed in significant amounts at close to the stoichiometric feed composition.^{12,13} In contrast, methane bromination yields a high selectivity to CH_3Br ($>90\%$) at smaller methane excess ($\text{CH}_4:\text{Cl}_2 > 5$), while poly-bromination is generally limited to the formation of CH_2Br_2 .¹

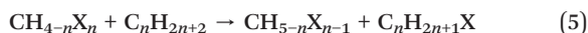


The industrial interest in methane halogenation already ignited in the early 20th century, when following a series of

patents, the first industrial process based on non-catalytic thermal chlorination of methane was established in 1923 by Hoechst.² After incorporating several refinements during its lifespan, this remains the state-of-the-art technology for the manufacture of chloromethanes (mostly CH_2Cl_2 and CHCl_3), the world-wide production of which is estimated to exceed 1.4 Mt per year, with a market size of *ca.* 1000 M\$ (Fig. 1).² Although the latter chemicals are mostly used as solvents and precursors in synthesis of polymers and fine chemicals, as early as the 1920s CH_3Cl was recognized as an attractive intermediate to produce methanol and ethers by reacting it with oxides such as CaO .¹⁴ This concept was advanced in the 1980s, when George A. Olah proposed a disruptive halogen-mediated route to valorise methane, the main constituent of highly abundant and inexpensive natural gas, by mitigating the costly and energy-intensive syngas-based processes.¹⁵ The concept is based on a selective C–H bond activation using Cl_2 or Br_2 to yield CH_3X , which are versatile platform molecules that can be readily coupled into value-added chemicals and liquid fuels *via* catalytic pathways that are similar to those of methanol (Fig. 1).^{15–19} The implementation of this technology



is contingent on the full recycling of the hydrogen halide (HX) formed in the halogenation and coupling steps and the ability to maximize the productivity of CH₃X over polyhalomethanes and its coupling efficiency.^{1,20} The recent decade demonstrated significant progress in HX recovery *via* cataloreactant systems,^{21,22} electrochemical and thermochemical HX oxidation,^{23–26} and methane oxyhalogenation,^{26–28} as well as CH₃X coupling,^{16–19} which brought this technology closer than ever to practical implementation. In contrast, the selective monohalogenation was tackled in a much less depth.



One approach to enhance the selectivity to CH₃X is based on the use of a substantial methane excess in the non-catalytic halogenation, which, however results in a large CH₄ recycle stream.^{1,12,13} Alternatively, polyhalomethanes can be reformed by reproporationation with another alkane (eqn (5)),^{28–30} or hydrodehalogenation (eqn (6)).³¹ Still, the key to selective monohalogenation relies on mitigating the radical steps *via* an alternative reaction pathway that is facilitated by the catalyst. Although the early reports on halogenation describe the enhancement of the reaction rate in the presence of different “inert” solids (*e.g.*, silica and active carbons) as well as CuCl₂-based melts and fluidized beds,^{11,32} the fundamental landmark to this goal was set by Olah, who devised the concept of electrophilic C–H bond activation enabled through the polarization of the X₂ molecule by the action of super-acid catalysts (Fig. 1).^{15,33} This pathway displays an inherent propensity to CH₃X as the addition of strong electron withdrawing halogen substituents destabilizes the intermediate carbonium ion complex and hinders further substitution. The proof of concept was demonstrated using the homogeneous SbF₅ catalyst, but aiming to achieve more practical space–time yields, Olah *et al.* studied supported noble metals as well as solid-acid catalysts based on oxyfluorides (Fig. 1).^{15,33} The former systems were found active at low temperatures (≤ 543 K), providing unprecedented selectivity to CH₃Cl (80–99%) and CH₃Br (99%) even when a substantial excess of halogen was applied. The same group also reported the catalytic effect of sulfated zirconia and several zeolites in methane chlorination.^{13,34} The high selectivity to CH₃Cl (99%) can be obtained over mordenite using a molar ratio of CH₄:Cl₂ ≥ 4 at >623 K, while lower temperatures and small methane excess favour the radical pathways and consequent polyhalogenation. Nevertheless, the zeolite catalysts displayed a fast deactivation that was associated with excessive dealumination. The interest in selective halogenations has been rekindled in the early 2000s as a response to a shale gas surge. GRT Inc. described sulfated Nb₂O₅ and ZrO₂ as selective bromination catalysts ($\leq 95\%$ CH₃Br),²¹ while Degirmenci *et al.*³⁵ reported high

CH₃Br selectivity ($>99\%$) over sulfated ZrO₂ included in SBA-15 structures (S-ZrO₂-SBA-15) at large bromine excess. Joo *et al.*³⁶ showed that methane chlorination over H–Y, Pt/H–Y and Pt/Na–Y leads to a slightly higher activity compared to the non-catalytic reaction, although the selectivity patterns remain almost unaffected. Batamack *et al.*³⁷ reported the one-pot conversion of methane to valuable hydrocarbons by integrating methane halogenation and CH₃X coupling over a SAPO-34 catalyst under a high CH₄:X₂ molar ratio of *ca.* 10.

In spite of these encouraging results, the previous reports were typically limited to the reaction of one halogen (*i.e.*, Cl₂ or Br₂) and few catalytic systems, which were investigated under very different regimes, precluding thus the assessment of the catalytic over non-catalytic effects and derivation of the generalized performance trends as a function of the catalyst family and type of halogen, which are essential for catalyst development. Besides, the stability of halomethanes over different materials and the propensity of the latter towards coking or reactions with halogens in a corrosive halogenation environment were rarely discussed.

Herein, aiming to set the basis that will revitalize the research in this field, we conducted the first comprehensive evaluation of a large library of catalytic materials comprising inorganic and carbon carriers, supported noble metals, metal oxides, chlorides and oxyfluorides, as well as sulfated systems and zeolites in both methane chlorination and bromination under identical reaction conditions and provided a general overview on their potential to promote these reactions. In addition, the impact of the framework structure and topology on the performance of zeolite catalysts in methane chlorination was investigated, which along with the kinetics analysis uncovered the significant role of confinement effects on C–H bond activation with chlorine.

Experimental

Catalyst preparation

Supports. Quartz (Thommen-Furler, $\approx 99\%$), SiO₂ (Evonik, AEROPERL 300/30, $\geq 99.0\%$), SiC (Alfa Aesar, $\geq 98.8\%$), α -Al₂O₃ (Alfa Aesar, 99.98%) and γ -Al₂O₃ (Sasol, PURALOX SCFa 140, $\geq 98\%$) were calcined prior to their use as catalyst carriers or in the catalytic tests. Activated carbon (CABOT NORIT RX 1.5 EXTRA, $\approx 100\%$) was applied as received.

Supported metal-based catalysts. Pt/Al₂O₃ (Aldrich, 5 wt% Pt) was purchased commercially and reduced just prior to the catalytic tests in flowing H₂ (20 mol% in He, gas hourly space velocity, GHSV = 680 h⁻¹) at 623 K for 2 h (5 K min⁻¹). Pt/SiO₂ (1 wt% Pt), Ru/SiO₂ (1 wt% Ru), Pd/SiO₂ (1 wt% Pd), PdCl₂/SiO₂ (2 wt% PdCl₂), CuCl₂/SiO₂ (10 wt% CuCl₂), Fe₂O₃/SiO₂ (5 wt% Fe₂O₃) and CeO₂/ZSM-5-140 (10 wt% CeO₂) were synthesised by incipient wetness impregnation. The respective precursors, [Pt(NH₃)₄]Cl₂·nH₂O (Aldrich, 98%), RuCl₃·nH₂O (ABCR, 39–42% Ru), Pd(NO₃)₂·nH₂O (Aldrich, $\approx 40\%$ Pd), CuCl₂ (Aldrich, 99%), FeCl₃·6H₂O (Sigma-Aldrich, 99%) and Ce(NO₃)₃·6H₂O (Aldrich, 99%), were dissolved in a



volume of deionized water that is equal to the pore volume of the carrier in appropriate amounts to achieve the desired loading of the active phase. The precursor solution was gradually added to the support and the wet solid was kneaded for 1 h at room temperature and then dried. Pt/SiO₂, Ru/SiO₂ and Pd/SiO₂ catalysts were calcined at 573 K and reduced just prior to the catalytic tests using the same protocol as detailed for the commercial Pt/Al₂O₃ catalyst. PdCl₂/SiO₂ and CuCl₂/SiO₂ catalysts were thermally treated in a N₂ atmosphere at 523 K and 573 K, respectively, for 5 h (5 K min⁻¹). Fe₂O₃/SiO₂ and CeO₂/ZSM-5-140 were calcined at 543 K and 823 K, respectively. Pt/carbon (1 wt% Pt) and TaOF₃/Al₂O₃ (20 wt% TaOF₃) were obtained by wet impregnation using previously reported protocols.^{15,38} In brief, the Pt/carbon catalyst was synthesized by dropwise addition of 0.025 M H₂PtCl₆·6H₂O (ABCR, 99.9%) solution in *aqua regia* to the carbon support (2.5 cm³ g_{carbon}⁻¹) for 30 min under continuous stirring. The material was dried at ambient pressure and 333 K, and then thermally treated in flowing N₂ (GHSV = 10 000 h⁻¹) at 673 K for 5 h (5 K min⁻¹). TaOF₃/Al₂O₃ was prepared by gradually dissolving TaF₅ (ABCR, 99%) in cold (195 K) methanol (Acros, 99.8%) followed by the addition of the resulting solution (0.48 M) to γ-Al₂O₃ (2.0 cm³ g_{Al₂O₃}⁻¹). Thereafter, the material was calcined at 523 K (1.5 K min⁻¹).

Sulfated oxides. S-ZrO₂ (Alfa Aesar, 2–4% active SO₃) and Nafion (Aldrich, SAC-13, 10–20% Nafion on SiO₂) were purchased commercially. S-TiO₂ was synthesized by stirring TiO₂ (Aldrich, nanopowder, 99.5%) in 1 M H₂SO₄ (Aldrich, 99.999%) aqueous solution (30 cm³ g_{TiO₂}⁻¹) for 30 min. The solid was recovered by filtration, dried and finally calcined at 773 K (2 K min⁻¹). S-Ta₂O₅ and S-Nb₂O₅ were prepared by adjusting a previously reported protocol.²¹ Herein, H₂SO₄ was added to Ta₂O₅ (Acros, 99.99%) or Nb₂O₅ (Aldrich, 99.9%) powder (1.7 mol_{H₂SO₄} mol_{oxide}⁻¹) and the mixture was stirred for 30 min, dried and calcined using the same procedure as that described for S-TiO₂. S-Zr-SBA-15 was obtained as reported by Degirmenci *et al.*³⁵ In brief, tetraethyl orthosilicate (TEOS, Aldrich, ≥99%) and ZrOCl₂·8H₂O (Fluka, ≥99%) were added (0.3 mol_{ZrOCl₂} mol_{TEOS}⁻¹) to a solution of Pluronic P123 polymer (Aldrich, Mn = 5800) in 1.6 M HCl (Acros, 37%) aqueous solution (0.027 g_{polymer} cm⁻³, 0.058 g_{TEOS} cm⁻³) and stirred at 313 K for 24 h. The mixture was transferred to a Teflon-lined autoclave and heated at 373 K for 12 h under static conditions. The solid was recovered by filtration, washed with deionized water, dried and calcined at 773 K for 5 h. Finally, the material was stirred with 0.5 M H₂SO₄ aqueous solution (100 cm³ g_{Zr-SBA-15}⁻¹) for 1 h, separated by filtration, dried and calcined at 573 K. S-Nb₂O₅-SiO₂ (20 wt% Nb₂O₅) was synthesized by wet impregnation. A 0.28 M solution of NH₄Nb(C₂O₄)₃·nH₂O (Aldrich, 99.99%) in 1 M H₂C₂O₄ (Acros, 98%) was added to a SiO₂ carrier (6.7 cm³ g_{SiO₂}⁻¹), followed by drying and calcination at 773 K. The obtained material was finally sulfated using the same protocol as that detailed for the S-Nb₂O₅ catalyst.

Zeolites. Commercial zeolites, 3A (Metrohm), H-USY-6 (Zeolyst International, CBV712), H-MOR-15 (Tosoh Corp.),

H-SAPO-34-6 (Tosoh Corp.), H-BETA-15 (Clariant, HCZB 30), NH₄-ZSM-5-15 (Zeolyst International, CBV 3024E), NH₄-ZSM-5-25 (Zeolyst International, CBV 5524G), NH₄-ZSM-5-40 (Zeolyst International, CBV 8014), NH₄-ZSM-5-140 (Zeolyst International, CBV 28014), H-ZSM-5-1000 (Tosoh Corp., HSZ-8090H0A) and TS-1-25 (ACS Material), the ionic form and Si:Al ratio of which are denoted by the prefix and suffix in the respective codes, were calcined prior to their use in the catalytic tests or as precursors in the further modification of the parent material. The zeolites containing ammonium ions were thus transformed into the protonic (H) form.

Li-ZSM-5-25, Na-ZSM-5-25, Cs-ZSM-5-25, Ca-ZSM-5-25 and Fe-ZSM-25 were prepared by triple ion exchange of H-ZSM-25 zeolite using a 0.1 M aqueous solution of the respective salts, LiNO₃ (FLUKA, 98%), NaNO₃ (Acros, ≥99%), CsNO₃ (Acros, 99.99%), Ca(NO₃)₂·4H₂O (Sigma-Aldrich, ≥99%) and FeCl₂·4H₂O (Sigma-Aldrich, 98%), at room temperature (100 cm³ g_{zeolyte}⁻¹) for 12 h, followed by drying and calcination. Pt-ZSM-5-25 (1 wt% Pt) was obtained by incipient wetness impregnation of the H-ZSM-5-25 material with a solution of [Pt(NH₃)₄]Cl₂·nH₂O using the analogous procedure detailed for supported metal catalysts. Mg-ZSM-5-140 (2 wt% MgO) was prepared following the method of Liu *et al.*¹⁸ Herein, a 0.16 M solution of Mg(NO₃)₂·6H₂O (Sigma-Aldrich, 99%) was mixed with H-ZSM-5-140 zeolite for 2 h (3.2 cm³ g_{zeolite}⁻¹). The suspension was dehydrated at 353 K for 4 h, dried and calcined at 723 K for 8 h.

H-ZSM-25-ox was prepared by refluxing the parent H-ZSM-5-25 sample with 1 M H₂C₂O₄ aqueous solution (50 cm³ g_{zeolyte}⁻¹) for 13 h. Thereafter, the material was recovered by filtration, washed with deionized water, dried and calcined. H-ZSM-5-25-st was derived by steaming (100 mol% H₂O) the parent H-ZSM-5-25 material in a fluidized bed reactor (GHSV = 5350 h⁻¹) for 12 h. H-ZSM-5-40-at zeolite was prepared using the well-established alkaline treatment protocol.³⁹ In brief, the parent H-ZSM-5-40 zeolite was exposed to a 0.2 M NaOH (Acros, 98.5%) aqueous solution (3.75 cm³ g_{zeolyte}⁻¹) and the resulting suspension was stirred at 338 K for 30 min in a Mettler Toledo Easymax 102 reactor system. The treatment was quenched by diluting the mixture with a twenty-fold excess of deionized water, followed by solid separation by filtration and washing until neutral pH. The material was ion exchanged three times with 0.1 M NH₄NO₃ (Acros, 99%) aqueous solution (100 cm³ g_{zeolyte}⁻¹) at 338 K for 12 h, and finally dried and calcined. Nanocrystalline H-ZSM-5-40-nc zeolite was synthesized according to the procedure of Van Grieken *et al.*⁴⁰ Aluminium isopropoxide (Al-iPr, Aldrich, ≥98%), NaOH, TEOS and ethanol (Acros, 99.8%) were added in the as-listed order to a 1 M TPAOH aqueous solution (molar ratio of TEOS:Al-iPr:NaOH:ethanol:TPAOH = 40:1:0.16:100:25) and stirred at room temperature for 12 h. The clear solution was transferred into a Teflon-lined autoclave and heated under static conditions at 438 K for 4 days. The final product was collected by filtration, washed with deionized water, dried and calcined. H-SnBETA-220 was synthesized using alkaline-assisted stannation,⁴¹ which is



equivalent to the above detailed alkaline-treatment in all aspects, except that the parent BETA-220 zeolite (Tosoh Corp.) was exposed to an aqueous solution comprising 0.3 M NaOH, 0.04 M SnSO₄ (ABCR, 95%) and 0.2 M tetrapropylammonium hydroxide (TPAOH, Alfa Aesar, 20 wt%).

Unless stated otherwise, the standard drying conditions involve a pressure of 5 kPa (vacuum), a temperature of 373 K and a treatment time of 12 h, while the standard calcination conditions involve a static air atmosphere, a temperature of 823 K, a time of treatment of 5 h and a heating rate of 5 K min⁻¹.

Catalyst characterization. The structure of the crystalline materials was confirmed by X-ray diffraction analysis performed using a PANalytical X'Pert PRO-MPD diffractometer with a Bragg-Brentano geometry using CuK α radiation ($\lambda = 1.54060 \text{ \AA}$). The presence of the sulfate groups in sulfated oxides was evidenced by diffuse-reflectance infrared Fourier-transformed spectroscopy (DRIFTS) performed using a Bruker Optics Vertex 70 spectrometer equipped with a high-temperature DRIFT cell (Harrick) and an MCT detector. N₂ isotherms of selected catalysts were recorded at 77 K using a Micromeritics TriStar II analyzer. Samples (*ca.* 0.15 g) were evacuated to 5 kPa at 573 K for 12 h prior to the measurement.

Catalyst testing. Chlorination and bromination of methane as well as the reactions of CH₂Cl₂ and CH₂Br₂ with different solids were performed at ambient pressure in a continuous-flow fixed-bed reactor setup described in Fig. 2. Gases, CH₄ (PanGas, purity 5.0), Cl₂ (PanGas, purity 2.8), O₂ (PanGas, purity 5.0), Ar (PanGas, purity 5.0) and He (PanGas, purity 5.0), were fed by a set of digital mass-flow controllers (Bronkhorst). Br₂ was supplied by passing a CH₄ flow through a bubbler containing liquid Br₂ (ABCR, 99%) which was placed in a water bath, the temperature of which was maintained at 302 ± 0.5 K. Water, CH₂Cl₂ (Sigma-Aldrich, 99.5%) and CH₂Br₂ (ABCR, 99%) were supplied by a syringe pump (Nexus 6000, Chemyx). The catalyst (catalyst volume, $V_{\text{cat}} = 2.7$ or 15 cm³, particle size, $d_p = 0.4$ –0.6 mm) was loaded in the central wide section of the glass reactor (characteristic dimensions, $d_r = 10.5$ mm, $d_t = 4$ mm, $h_r = 55$ mm and $h_c = 5.5$ or $d_r = 18.5$ mm, $d_t = 4.8$ mm, $h_r = 80$ mm and $h_c = 10.2$ mm, Fig. 2 inset) between two plugs of quartz wool. The reactor was heated by a home-built electric oven. The temperature was controlled by a thermocouple inserted in a glass thermowell (outside diameter, $d_{\text{tw}} = 3$ mm), which was positioned along the vertical reactor axis with its tip aligned with the bottom of the bed, while the pressure was monitored by the pressure sensor installed prior to the reactor. The thermowell and the bed composed of glass beads were used to additionally reduce the volume of the narrow top and bottom reactor sections, respectively. The aluminium foil shielding the reactor, opaque tubing, connections and valves were used to preclude the contact of light with the reaction mixtures and hence photo-induced halogenation. The gas lines used to feed Br₂, water, CH₂Cl₂ and CH₂Br₂, as well as the downstream lining were maintained at 393 K to prevent

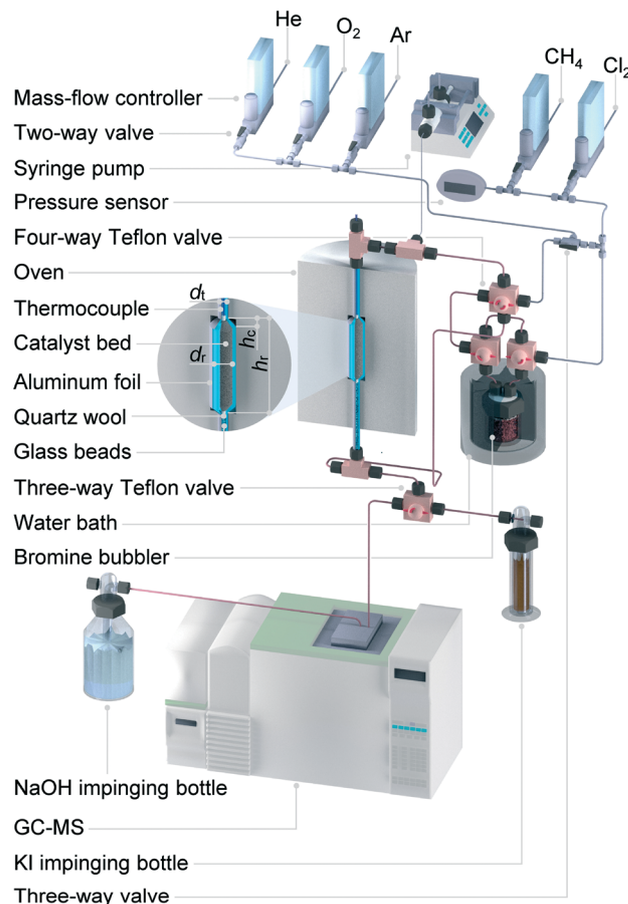


Fig. 2 Flowsheet of the continuous-flow fixed-bed reactor setup for methane halogenation. The red-coloured sections of the linings were heated at 393 K to prevent condensation.

the condensation of these reactants and reaction products such as CH₂Cl₂, CH₂Br₂ and CHCl₃.

Prior to testing, the solids were heated to 723 K for 2 h in flowing He (GHSV = 13 700 h⁻¹) to eliminate the traces of moisture, O₂ and other possible gas impurities. The only exceptions were PdCl₂/SiO₂, CuCl₂/SiO₂, TaOF₃/Al₂O₃ and Nafion catalysts, which were treated at 573 K to avoid their potential thermal decomposition or volatilization. Thereafter, the bed was cooled down to the desired temperature and left to stabilize for at least 30 min before the catalytic test. Unless stated otherwise, the reactant mixture comprising CH₄:Cl₂ in a molar ratio of 8.5:3.5 was fed at GHSV = 267 h⁻¹ or 50 h⁻¹ in methane chlorination and the mixture containing CH₄:Br₂ in a molar ratio of 8.5:3.2 ± 0.1 was introduced at GHSV = 260 h⁻¹ or 48.5 h⁻¹ in methane bromination. The reactor effluent was diluted with a He–Ar mixture (He:Ar = 71.5:4.5) and directed towards the analysis section. Carbon-containing compounds (CH₄, CH₃Br, CH₂Br₂, CH₃Cl, CH₂Cl₂, CO and CO₂) and Ar were quantified on-line *via* a gas chromatograph equipped with a GS-carbon PLOT column coupled to a mass spectrometer (GC-MS, Agilent GC 6890, Agilent MSD 5973N). Quantification of Cl₂ and Br₂ at the reactor outlet was performed by their absorption in an impinging bottle filled



with 0.1 M KI aqueous solution ($X_2 + 3I^- \rightarrow I_3^- + 2X^-$) followed by iodometric titration (Mettler Toledo G20 Compact Titrator) of the formed triiodide ($I_3^- + 2S_2O_3^{2-} \rightarrow 3I^- + S_4O_6^{2-}$) with 0.01 M $Na_2S_2O_3$ (Aldrich, 99.99%).

The conversion of methane, X_{CH_4} , the conversion of halogen, X_{X_2} , the reaction rate expressed with respect to methane, r_{CH_4} , the selectivity to product j , S_j , the space-time yield of carbon oxides, STY_{CO_x} , and the error of the carbon mass balance, ε_C , were calculated using eqn (7)–(12), respectively, in which n_i^{inlet} and n_i^{outlet} are the molar flows of reactant i at the reactor inlet and outlet, respectively, n_j^{outlet} is the molar flow of product j at the reactor outlet and W_{cat} is the catalyst weight. To minimize the errors, parameter k in eqn (7) was set to 0, except for those measurements for which the error of the carbon mass balance systematically showed negative deviation indicative of coke formation, where it was set to 1/2. The absence of mass and heat transfer limitations in the catalytic tests was confirmed by the evaluation of the dimensionless moduli based on the criteria of Carberry, Mears and Weisz-Prater.⁴² The reported values of conversion, selectivities and reaction rates were determined as an average of 3–5 consecutive measurements performed during 3 h of operation under specified reaction conditions, with the first sample taken after 40 min of stabilization, while the standard deviation of the data set is indicated by the error-bars where appropriate. Unless stated otherwise, the error of the carbon mass balance was less than 3% in all experiments.

$$X_{CH_4} = \left[\left(\frac{1}{2} + k \right) \times \frac{n_{CH_4}^{inlet} - n_{CH_4}^{outlet}}{n_{CH_4}^{inlet}} + \left(\frac{1}{2} - k \right) \times \frac{\sum_{outlet} n_j^{outlet}}{n_{CH_4}^{inlet}} \right] \times 100\% \quad (7)$$

$$X_{X_2} = \frac{n_{X_2}^{inlet} - n_{X_2}^{outlet}}{n_{X_2}^{inlet}} \times 100\% \quad (8)$$

$$r_{CH_4} = \frac{n_{CH_4}^{inlet} \times X_{CH_4}}{100 \times W_{cat}}, \text{ mol}_{CH_4} \text{ s}^{-1} \text{ g}_{cat}^{-1} \quad (9)$$

$$S_j = \frac{n_j^{outlet}}{\sum_{outlet} n_j^{outlet}} \times 100\% \quad (10)$$

$$STY_{CO_x} = \frac{n_{CO}^{outlet} + n_{CO_2}^{outlet}}{V_{cat}}, \text{ mol}_{CO_x} \text{ s}^{-1} \text{ cm}_{cat}^{-3} \quad (11)$$

$$\varepsilon_C = \frac{n_{CH_4}^{inlet} - \left(n_{CH_4}^{outlet} + \sum_{outlet} n_j^{outlet} \right)}{n_{CH_4}^{inlet}} \times 100\% \quad (12)$$

Results and discussion

Catalyst performance in methane halogenation

The preliminary insights into the catalytic effects in methane halogenation were acquired by analysing both the chlorination and bromination performances of Pt/SiO₂, TaOF₃/Al₂O₃, S-ZrO₂ and H-SAPO-34 (Fig. 3), which are the main representatives of the previously reported catalyst families (Fig. 1).^{13,15,34,37} The activity and selectivity profiles of the materials were recorded by using the undiluted reactant mixtures comprising methane and halogen in a molar ratio of CH₄:X₂ = 3.5 in the chlorination and CH₄:X₂ = 3.2 ± 0.1 in the bromination reaction. Moderate methane excess simulates the practically relevant operating regime, which strives at full halogen consumption that facilitates the downstream product separation while minimizing the size of the hydrocarbon recycle. Moreover, the selected ratios were smaller than those at which free-radical halogenations display an inherently high selectivity to CH₃X, enabling thus the resolution of the catalytic from the non-catalytic effects.^{15,34} With regard to the latter problems, the catalyst activities were compared at constant GHSV and constant temperature (GHSV = 267 h⁻¹ and $T = 523$ K in methane chlorination, GHSV = 260 h⁻¹ and $T = 643$ K in methane bromination) and benchmarked to the activity patterns of the empty glass reactor and reactor accommodating the bed of quartz particles, whereby the values of the latter two reaction parameters are consistent with the typical conditions applied in previous studies.^{13,15,34,35,37} This approach, which was practiced in the work of Olah *et al.*,^{13,15,34} enables the decoupling of the catalytic from the non-catalytic gas-phase contributions, since the volume of the interparticle voids does not differ significantly between the different catalyst beds. Besides, it also has practical relevance as it shows the level of productivity enhancement of a given reactor in the presence of catalysts compared to the inherent activity of its void volume or bed composed of inert particles.

The assessment of both methane and halogen conversion in methane chlorination at 523 K indicated that the bed of quartz particles displays an almost identical performance to an empty reactor of the same volume (Fig. 3a). Considering the fact that the effective void volume of the bed is *ca.* 2 times smaller than that of the empty reactor (bed porosity, $\varepsilon \approx 0.4$),⁴³ this result suggests that the chlorination reaction is promoted even in the presence of inert and low-surface area solids. The conversion of methane and Cl₂ over Pt/SiO₂, TaOF₃/Al₂O₃, S-ZrO₂ and H-SAPO-34 catalysts was *ca.* 2–2.5 times higher compared to that over quartz, suggesting the further activity promotion over these solids. The onset of methane bromination was shifted to *ca.* 120 K higher temperature with respect to chlorination. The activity of quartz particles was comparable to that of an empty reactor, but in contrast to chlorination, methane conversion in bromination was almost unaffected by the catalytic materials. Still, the conversion of bromine was higher over the catalysts, especially H-SAPO-34, which can be associated with a more pronounced polybromination, oxidation and reactions with a catalyst (*vide infra*).



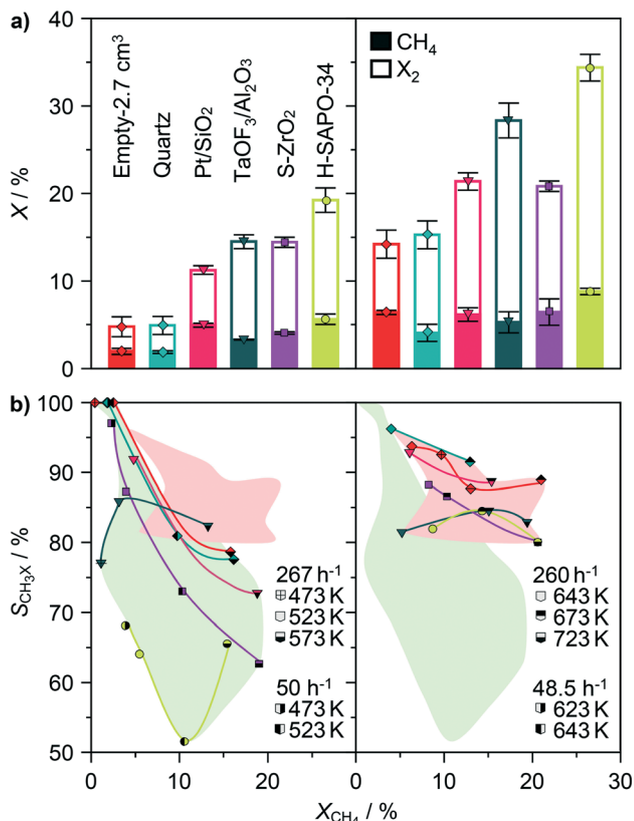


Fig. 3 a) Conversion of reactants in methane chlorination at $T = 523$ K and GHSV = 267 h^{-1} (left panel) and bromination at $T = 643$ K and GHSV = 260 h^{-1} (right panel) over representative catalysts. b) Selectivity to CH_3X in methane chlorination (left panel) and bromination (right panel) over representative catalysts as a function of methane conversion, which was adjusted by varying the temperature or GHSV. The catalysts are identified according to the shape and colour code presented in a), while the pattern of the symbol interior defined in b) provides the information on the reaction temperature and GHSV at which the data point was recorded. The green- and pink-coloured areas in b) outline the range of CH_3X selectivities in methane chlorination and bromination, respectively, achieved in the void reactor and over various materials. Other conditions: $\text{CH}_4 : \text{Cl}_2 = 8.5 : 3.5$, $\text{CH}_4 : \text{Br}_2 = 8.5 : 3.2 \pm 0.1$ and $P = 100$ kPa.

In addition to the conversion profiles, the differences in the selectivity to CH_3X were evaluated as another indicator of the potential catalytic effects (Fig. 3b). In general, the operation under comparable $\text{CH}_4 : \text{X}_2$ molar ratios and GHSVs led to a higher fraction of CH_3X in bromination compared to that in chlorination ($S_{\text{CH}_3\text{Br}} = 80\text{--}95\%$ versus $S_{\text{CH}_3\text{Cl}} = 52\text{--}90\%$ at $X_{\text{CH}_4} = 5\text{--}18\%$). Besides, the selectivity to CH_3X declined less steeply upon increasing the methane conversion in bromination, corroborating the higher inherent propensity to selective monohalogenation in this reaction. Nevertheless, the selectivity to CH_3X in both reactions decreased in a similar order, empty reactor \approx quartz \approx Pt/SiO₂ > TaOF₃/Al₂O₃ \approx S-ZrO₂ > H-SAPO-34, indicating the prominent impact of the catalyst on the product distribution.

In order to generalize the performance trends, the analysis was extended to a significantly broader range of materials

which can be classified into four distinct groups, supports, metal-based materials, sulphated oxides and zeolites. Aiming to provide a comprehensive overview of their catalytic behaviours and the facile benchmarking of the activity and selectivity patterns of the materials with respect to the void reactors, supports, other material families and their individual members, the circular plots are devised, as presented in Fig. 4 and 5. These are composed of the top semicircle, which provides the information of the catalyst activity, and the bottom semicircle, which outlines the product distribution patterns. The semicircles are further divided into four circular sectors, a–d (top) and e–h (bottom), which present the performances of the materials within a specific family. The activity sectors provide the information on the halogen and methane conversion at constant temperature and GHSV, which both increase in the direction from the circle centre. The bottom sectors present the selectivities to CH_3X , CH_2X_2 , CHX_3 and CO_x as a function of methane conversion, which increases in a clockwise direction, and indicate the formation of coke deposits. The red- and dark-coloured background areas of the subsections denote the performance regions of empty reactors and inorganic carriers, respectively, serving thus as the reference for the evaluation of the catalytic contributions.

The conversion levels attained over various inorganic supports, *i.e.*, SiO₂, SiC, α -Al₂O₃ and γ -Al₂O₃ are similar to or higher than that of the empty reactor, even when its volume exceeds the volume of the carrier bed by a factor of 5.5 (Fig. 4, sector a), corroborating thus the beneficial effect of solids on the low-temperature methane chlorination. Thereby, mesoporous SiO₂ and γ -Al₂O₃ carriers displayed *ca.* 2 times higher conversion of reactants as compared to their nonporous counterparts, quartz, SiC and α -Al₂O₃, suggesting the positive impact of the surface area on methane chlorination. Nonetheless, the selectivity profiles of the former carriers proved the lower fraction of CH_3Cl as compared to those of the latter group, which exhibited a similar product distribution pattern to the empty reactor (Fig. 4, sector e). Notably, while CH_2Cl_2 and to a small extent CHCl_3 constituted the only by-products over quartz, SiO₂ and SiC, the evolution of CO_x along with CH_2Cl_2 was very prominent over alumina supports. Given the oxygen-free inlet feed and the absence of reaction between methane and α -Al₂O₃ and γ -Al₂O₃ in the temperature range of interest, the formation of CO_x could only arise from the decomposition of chloromethanes over these carriers (*vide infra*, Fig. 7b).⁴⁴ Another peculiar behaviour was observed for the carbon support, which although exhibiting *ca.* 1.5–2 times higher methane conversion than SiO₂ and γ -Al₂O₃ and a selectivity profile that closely matched that of quartz and the empty reactor led to an immense Cl_2 conversion. The latter significantly exceeded the stoichiometric level calculated on the basis of chloromethane production (*ca.* 30%) as well as the Cl_2 conversion levels achieved over the materials displaying similar methane consumption (*e.g.*, H-ZSM-5-40, *vide infra*), indicating that a part of Cl_2 might react with the carbon support. This hypothesis was verified by exposing the carbon carrier



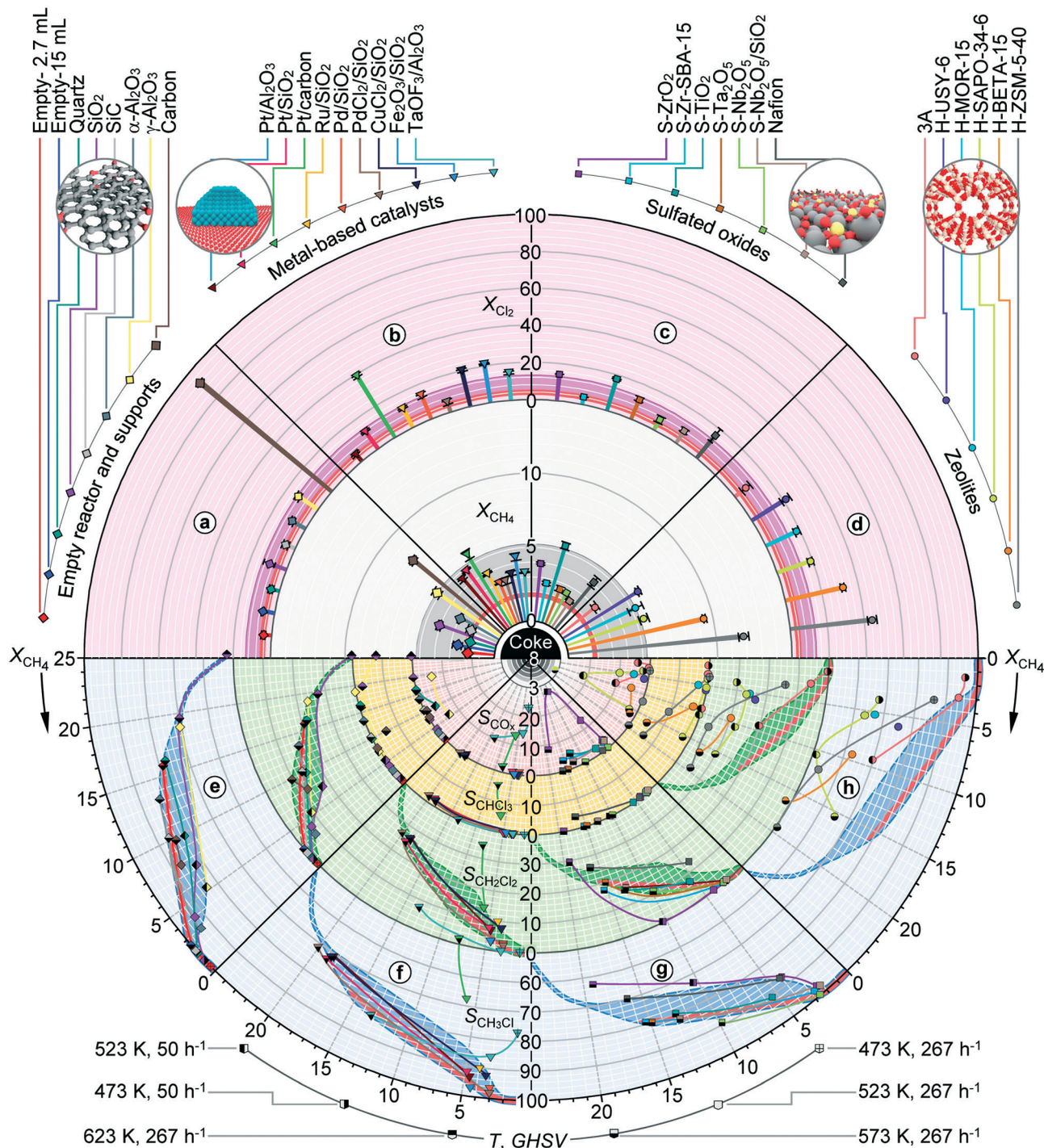


Fig. 4 Conversion of reactants in methane chlorination at $T = 523 \text{ K}$ and $\text{GHSV} = 267 \text{ h}^{-1}$ a) in empty reactors and over supports, b) supported metal-based catalysts, c) sulfated oxides and d) zeolites (top semicircle). Selectivity to different products in methane chlorination e) in empty reactors and over supports, f) supported metal-based catalysts, g) sulfated oxides and h) zeolites as a function of methane conversion which was adjusted by varying the temperature or GHSV (bottom semicircle). Conversions and selectivities are expressed in %. The catalysts are identified according to the shape and colour code presented above the top semicircle, while the pattern of the symbol interior defined below the bottom semicircle provides the information on the reaction temperature and GHSV at which the data point was recorded. The central part of the bottom semicircle displays the errors of the carbon mass balance in % for those catalytic points for which the consistent negative deviation of this parameter was observed, indicative of coke formation. The red- and darker-coloured areas outline the benchmark range of conversions and selectivities displayed by the empty reactor and different supports, respectively. More information regarding the plot is provided in the main text. Other conditions: $\text{CH}_4 : \text{Cl}_2 = 8.5 : 3.5$ and $P = 100 \text{ kPa}$.



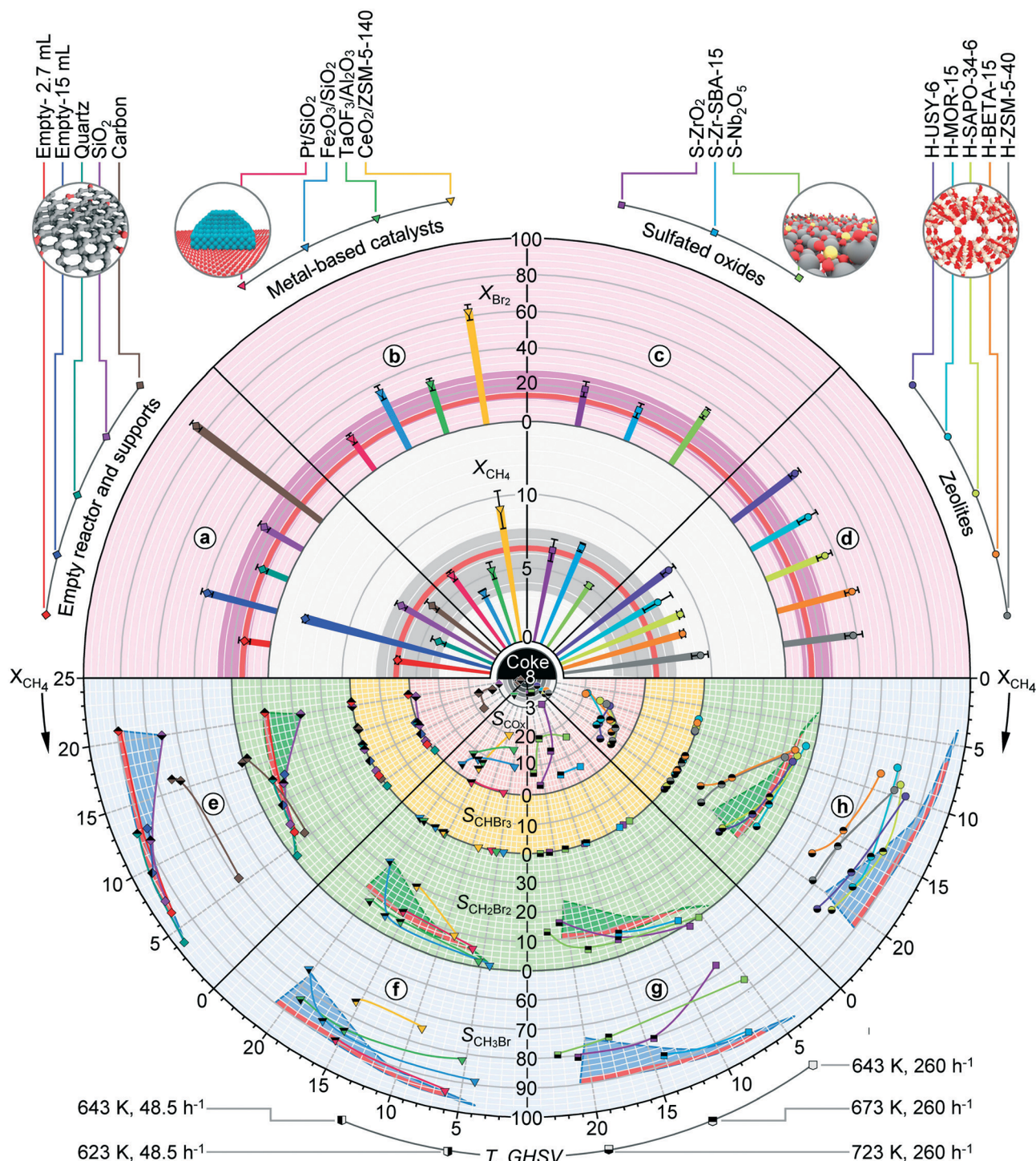


Fig. 5 Conversion of reactants in methane bromination at $T = 643\text{ K}$ and $GHSV = 260\text{ h}^{-1}$ a) in empty reactors and over supports, b) supported metal-based catalysts, c) sulfated oxides and d) zeolites (top semicircle). Selectivity to different products in methane bromination e) in empty reactors and over supports, f) supported metal-based catalysts, g) sulfated oxides and h) zeolites as a function of methane conversion, which was adjusted by varying the temperature or GHSV (bottom semicircle). Conversions and selectivities are expressed in %. The catalysts are identified according to the shape and colour code presented above the top semicircle, while the pattern of the symbol interior defined below the bottom semicircle provides the information on the reaction temperature and GHSV at which the data point was recorded. The central part of the bottom semicircle displays the errors of the carbon mass balance in % for those catalytic points for which the consistent negative deviation of this parameter was observed, indicative of coke formation. The red- and darker-coloured areas outline the benchmark range of conversions and selectivities displayed by the empty reactor and different supports, respectively. More information regarding the plot is provided in the main text. Other conditions: $\text{CH}_4:\text{Br}_2 = 8.5:3.2 \pm 0.1$ and $P = 100\text{ kPa}$.



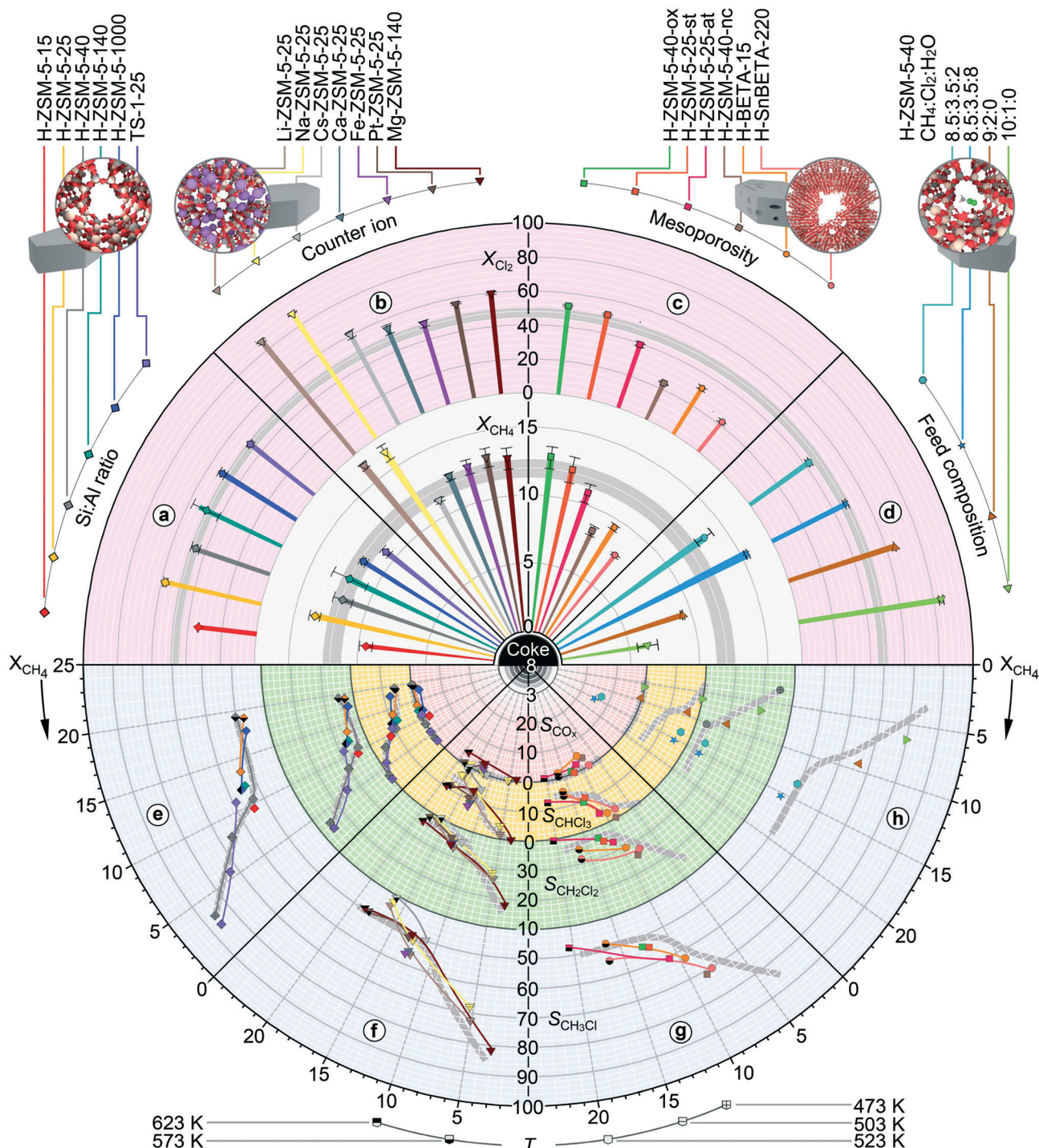


Fig. 6 Conversion of reactants in methane chlorination at $T = 523$ K and GHSV = 267 h^{-1} over ZSM-5 and BETA zeolites with different a) Si : Al ratios, b) counter ions, c) mesoporosities and crystallite sizes and d) at various feed compositions (top semicircle). Selectivity to different products in methane bromination over ZSM-5 and BETA zeolites with different e) Si : Al ratios, f) counter ions, g) mesoporosities and crystallite sizes and h) at various feed compositions as a function of methane conversion which was adjusted by varying the temperature (bottom semicircle). Conversions and selectivities are expressed in %. The catalysts are identified according to the shape and colour code presented above the top semicircle, while the pattern of the symbol interior defined below the bottom semicircle provides the information on the reaction temperature at which the data point was recorded. The central part of the bottom semicircle displays the errors of the carbon mass balance in % for those catalytic points for which the consistent negative deviation of this parameter was observed, indicative of coke formation. The gray-coloured areas outline the benchmark range of conversions and selectivities displayed by the H-ZSM-5-40 catalyst. More information regarding the plot is provided in the main text. Other conditions: $\text{CH}_4 : \text{Cl}_2 = 8.5 : 3.5$ and $P = 100$ kPa.



to a pure chlorine stream which was fed at a similar temperature and GHSV to those applied in chlorination, whereby a halogen uptake of *ca.* 50% was measured.

The recorded performance profiles of various carriers define the benchmark regions (denoted as darker-coloured areas in Fig. 4) which enabled the potential catalytic contributions of the supported metal-based active phases to be decoupled. Specifically, the conversion of methane over all the studied systems was of similar magnitude to that of the respective carriers (Fig. 4, sector b). Pt/carbon, CuCl₂/SiO₂ and Fe₂O₃/SiO₂ displayed a higher chlorine conversion, which can be associated with the carrier halogenation for the first (*vide supra*) and active phase chlorination for the latter two catalysts. In addition, CuCl₂/SiO₂ and Fe₂O₃/SiO₂ catalysts exhibited the formation of deposits on the reactor walls and glass-beads located below the catalyst bed, which are indicative of the volatilization of the metal-halide species. The

product distribution patterns of most of the metal, metal halide and metal oxide catalysts paralleled those of the respective supports and empty reactors, with PdCl₂/SiO₂ and Fe₂O₃/SiO₂ exhibiting the highest selectivity to CH₃Cl (Fig. 4, sector f). The most prominent deviations from these trends were displayed by Pt/carbon and TaOF₃/Al₂O₃, which favoured CO_x production, while the first system also displayed a marked polyhalogenation activity.

The conversion levels of methane and Cl₂ over sulfated catalysts were comparable to those of SiO₂ and γ -Al₂O₃, with S-TiO₂ displaying the highest activity (Fig. 4, sector c), while the selectivity to CH₃Cl for most of these systems followed almost identical patterns to those recorded over quartz and in an empty reactor (Fig. 4, sector g). More peculiar deviations from the general selectivity trends were displayed by S-ZrO₂ and S-Nb₂O₅ catalysts. Specifically, in the case of S-ZrO₂, CH₃Cl was the only chlorinated product at low-to-moderate

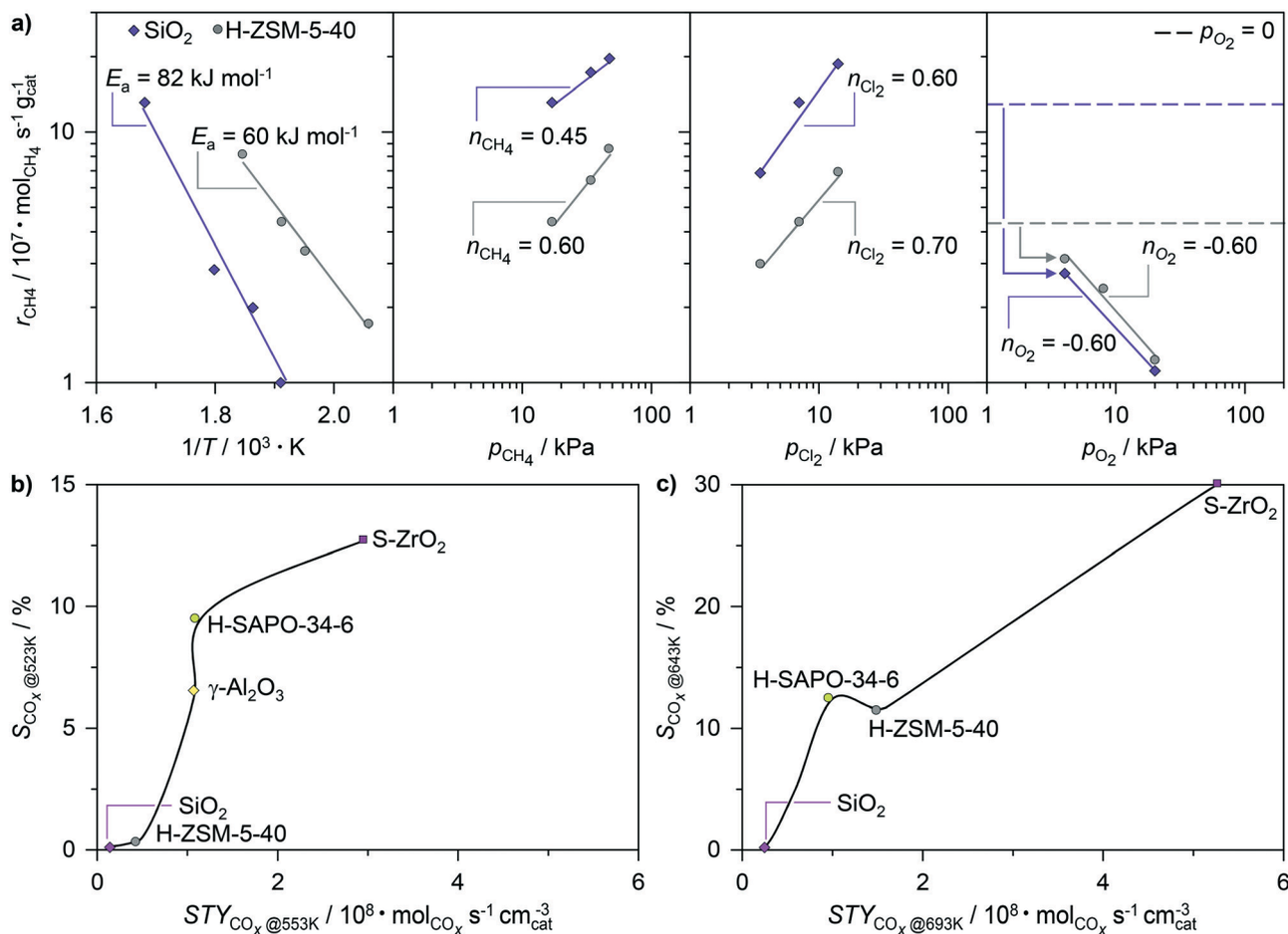


Fig. 7 a) Rate of methane chlorination as a function of reciprocal temperature and partial pressure of methane, chlorine and oxygen over SiO₂ and H-ZSM-5-40. b) Selectivity to CO_x in methane chlorination at 523 K versus space-time yield of CO_x in the reaction of CH₂Cl₂ with different solids at 553 K. c) Selectivity to CO_x in methane bromination at 643 K versus space-time yield of CO_x in the reaction of CH₂Br₂ with different solids at 693 K. Other conditions: a) CH₄:Cl₂:O₂:Ar:He = 17:7:0:4.5:71.5 (temperature variation), 17–47.5:7:0:4.5:71.5–41 (methane variation), 17:3.5–14:0:4.5:75–64.5 (chlorine variation) or 17:7:0–20:4.5:71.5–51.5 (oxygen variation). The impact of temperature on the reaction rate was studied at $T = 485$ –523 K (H-ZSM-5-40) or 518–594 K (SiO₂) and the impact of the partial pressures of reactants and oxygen was evaluated at $T = 523$ K (H-ZSM-5-40) or 594 K (SiO₂) using GHSV = 2220 h⁻¹ (H-ZSM-5-40) or 4440 h⁻¹ (SiO₂). The conversions of reactants were in the range of $X_{\text{CH}_4} = 1$ –9% and $X_{\text{Cl}_2} = 1$ –37%. b and c) Methane halogenation: CH₄:Cl₂ = 8.5:3.5, CH₄:Br₂ = 8.5:3.2 ± 0.1 and GHSV = 267 h⁻¹ (chlorination) or 260 h⁻¹ (bromination). CH₂X₂ reaction with solids: CH₂X₂:Ar:He = 2:4.5:93.5, GHSV = 1110 h⁻¹. All tests were performed at $P = 100$ kPa.



methane conversions ($\leq 11\%$). However, this behaviour was linked with high selectivity to CO_x ($S_{\text{CO}_x} = 2\text{--}27\%$, mostly CO), likely produced through the decomposition of polychlorinated products (*vide infra*, Fig. 7b). The evolution of CO_x was also pronounced over the S-Zr-SBA-15 catalyst. In turn, S-Nb₂O₅ provided the highest selectivity to CH_3Cl , which was *ca.* 2% higher than that of the empty reactor. The supporting of the latter active phase on SiO_2 did not enhance the activity, while the CH_3Cl selectivity curve was shifted to slightly lower values that are still above those of SiO_2 . Nevertheless, a small fraction of CO_x ($S_{\text{CO}_x} = 2\%$) was also formed over the supported system.

More significant performance differences with respect to the previously discussed catalyst families were observed when the chlorination reaction was performed over zeolites. In particular, while the methane and Cl_2 conversions achieved over zeolite 3A, H-USY-6, H-MOR-15 and H-SAPO-34 fall in the similar range to those of SiO_2 and $\gamma\text{-Al}_2\text{O}_3$ supports, H-BETA-15 and H-ZSM-5-40 provided *ca.* 2–2.5 times higher conversion levels, enabling thus up to 5.5 times higher activity with respect to the benchmark quartz and empty reactor systems. Another peculiar feature of zeolites is related to the increased fraction of CH_2Cl_2 and CHCl_3 with respect to CH_3Cl . In addition, similar to the alumina carriers, all zeolite catalysts except H-ZSM-5-40 favoured the production of CO_x . This was especially pronounced for H-SAPO-34, which was also the only system leading to coke deposition at 573 K, as inferred both from more significant deviations of the carbon balance and visual inspection of the used catalyst. The evaluation of the H-ZSM-5-40 zeolite in a 10 h test using a $\text{CH}_4:\text{X}_2$ ratio of 8.5:3.5 and GHSW = 267 h^{-1} at 573 K indicated no performance deterioration, whereby the X-ray diffraction pattern and N_2 -isotherm of the used catalyst do not show significant variations with respect to the fresh material.

In analogy to methane chlorination, the performance analysis was further extended to additional supports and catalytic systems in methane bromination (Fig. 5). The conversions of methane and Br_2 over the porous SiO_2 support were *ca.* 2 times higher as compared to those over quartz, but still comparable to those of the empty reactor of identical volume. This suggests that the external surface has a smaller impact on the bromination activity, which appears to be primarily governed by the available void volume. This is also corroborated by the significantly higher conversion increase upon expanding the void reactor volume as compared to methane chlorination. In view of the product distribution, the SiO_2 carrier displayed a similar behaviour to quartz and the empty reactor in the low temperature regime. However, a prominent drop in selectivity to CH_3Br (*ca.* 15%) coupled with the evolution of small amounts of CO_x and coking occurred at higher temperatures (*ca.* 723 K), suggesting the interaction of brominated products with the SiO_2 carrier. The fraction of CO_x as well as the propensity to coke was particularly high for the carbon support, which also displayed a marked bromine conversion in analogy to its behaviour in methane chlorination.

Compared to the empty reactor and inorganic carriers, supported metal-based catalysts (Fig. 5, sector b) and sulfated oxides (Fig. 5, sector c) generally led to comparable conversion levels, while their selectivity patterns were characterized by a lower fraction of CH_3Br coupled with pronounced CO_x production. The exceptions to these trends were Pt/ SiO_2 and to some extent S-ZrO₂-SBA-15, whose selectivity curves did not deteriorate substantially from the benchmark systems. Similar to methane chlorination, the volatilization of $\text{Fe}_2\text{O}_3/\text{SiO}_2$ was also observed. In the case of sulfated oxides, SO_2 was detected in the outlet feed, indicating their limited stability in a bromination environment. In addition, coke deposits were formed on S-ZrO₂ at 723 K. Zeolite catalysts with the exception of H-MOR yielded slightly higher conversions of methane (*ca.* 1–2%) compared to SiO_2 . However, the product pool of all the zeolite catalysts was negatively affected by the increased fraction of CO_x as well as coking, which was prominent at temperatures ≥ 723 K.

Impact of zeolite structure on methane chlorination

The noticeable activity promotion of the H-ZSM-5-40 catalyst with respect to the empty reactor and various inorganic carriers in methane chlorination coupled with the considerable compositional and modification flexibility of the MFI framework makes this catalytic system highly suitable to study the structure–performance relationships that could cast light on the origins of its catalytic action (Fig. 6). In this respect, the activity of H-ZSM-5 materials was first analysed as a function of their Si:Al ratio, which is the central parameter controlling the acid properties and polarity of zeolites (Fig. 6, sectors a and e).⁴⁵ The conversion of reactants was slightly lower over H-ZSM-5-15 as compared to the other H-ZSM-5 catalysts. Nonetheless, the overall variations in activity and product distribution were of limited extent in a broad range of Si:Al ratios (15–1000), which along with the similar behaviour of the TS-1-25 catalyst implies a weak correlation between the zeolite performance and the concentration and strength of Brønsted acidic sites. In addition, the introduction of various cations in the ZSM-5-25 framework had almost no influence on the activity and selectivity patterns (Fig. 6, sectors b and f). The only exception to the latter trend were Li-ZSM-5-25 and Na-ZSM-5-25, which led to a more significant conversion enhancement compared to the benchmark H-ZSM-5-40 and H-ZSM-5-25. This result might be the sign of the additional catalytic effect that could arise from the strong Lewis acidic (electrophilic) character of the relatively small and solvent-free Li^+ and Na^+ cations that has been found beneficial for the chlorination of aromatic compounds.⁴⁶ Nevertheless, the selectivity–conversion profiles of both of these catalysts did not deflect significantly from those recorded over other ZSM-5 systems, except that similar to Cs-ZSM-5-25 it led to a slightly lower fraction of CH_3Cl compared to CH_2Cl_2 and CHCl_3 (Fig. 6, sector f). Consistent with the above observations, the treatments of the H-ZSM-5-40 catalyst with steam or oxalic acid, which cause the formation of extraframework aluminium species, loss of acidity and moderate increase in mesoporosity,^{47,48}



did not affect the performance (Fig. 6, sectors c and g). Still, the alkaline-assisted desilication of this zeolite, which introduces a more significant fraction of intracrystalline mesopores that are also more accessible from the outside surface,³⁹ led to a slight increase in the fraction of monochlorinated over polychlorinated products at essentially unaltered activity. A similar change in the selectivity profile at almost unaltered activity was observed for H-SnBETA-220 (prepared by alkaline-assisted stannation) with respect to the reference H-BETA-15 catalyst (Fig. 6, sectors c and g). Given the weak sensitivity of the chlorination reaction to the nature and strength of acid sites, the shift in the selectivity pattern observed in the samples with more developed mesoporosity can be associated with a decrease in the characteristic length of the microporous domains that could reduce the extent of polyhalogenation by facilitating the escape of CH₃Cl. This hypothesis is further corroborated by the behaviour of H-ZSM-5-40-ns comprising crystals of smaller size with respect to H-ZSM-5-40, which also induced a slight decrease in the fraction of CH₂Cl₂ and CHCl₃ with respect to CH₃Cl (Fig. 6, sector g).

In addition to the variations in the framework structure, the impact of the reaction conditions on the chlorination performance of ZSM-5 zeolites was investigated (Fig. 6, sectors d and h). In particular, the impact of water co-feeding was evaluated as it was previously devised as a strategy to boost the selectivity to CH₃Cl and minimize coking over H-SAPO-34.³⁷ In fact, the addition of water to the chlorination feed decreased the fraction of CH₂Cl₂ and almost eliminated the formation of CHCl₃ at virtually unaltered conversion levels. Nevertheless, it provoked a significant CO_x (mostly CO) evolution, indicating the preferential hydrolysis of the polychlorinated products, which was not considered before. This result also corroborates that the evolution of the CO_x that was observed over other zeolite frameworks in chlorination and was especially pronounced in bromination stems from the interaction of halomethanes with bridging oxygen sites, which could be replenished by water. Finally, the CH₄:Cl₂ ratio was increased to explore the scope for the enhancement of selectivity to CH₃Cl. Notably, the operation with ten-fold methane excess caused only a small increase in selectivity to CH₃Cl (*ca.* 6%) with respect to the use of the reference CH₄:Cl₂ ratio of 8.5:3.5 (Fig. 6, sectors d and h). Comparatively, the use of 10-fold methane excess in the case of SiO₂ led to *ca.* 15% increase in selectivity to CH₃Cl with respect to the above stated reference feed.

Mechanistic insights into methane halogenation over solid surfaces

The evaluation of a large number of materials in methane halogenations revealed their positive impact on the chlorination activity, but limited effect on the bromination rate as compared to the respective non-catalysed reactions (Fig. 3–5). Nonetheless, metal-based and sulfated catalysts displayed similar conversion levels to the supports in methane chlorination, while their CH₃Cl selectivity profiles were comparable to those achieved in the empty reactor and over the bare car-

riers. Notably, the variations of the Si:Al ratio and counter cation as well as the formation of the extraframework aluminium species, which are expected to alter the polar intermediates and transition states, had almost no effect on the performance over the most active ZSM-5 zeolite. These results provide strong indication that the radical-chain mechanism dominates in the overall process, since the alternative activation pathways would induce more significant deviations in activity and selectivity patterns. To further support these findings, the apparent kinetic fingerprints of methane chlorination acquired over the H-ZSM-5-40 catalyst were compared to those recorded over SiO₂ (Fig. 7a). The apparent activation energy of the zeolite catalyst was *ca.* 20 kJ mol⁻¹ lower as compared to that of SiO₂, in agreement with the observed activity enhancement. Still, the apparent reaction orders with respect to methane and Cl₂ have very similar values for the two materials, suggesting the similarity in the mechanism of reactant activation. Additional experimental evidence supporting the involvement of radical intermediates in the chlorination over zeolite catalysts stems from a prominent decrease of the chlorination rate over H-ZSM-5-40 upon the addition of O₂, which is a well-known inhibitor of radical processes.^{3,6,11} Moreover, the O₂ inhibition constants over both H-ZSM-5-40 and SiO₂ are essentially identical (*ca.* -0.6), whereby their absolute value closely matches the apparent reaction order measured with respect to Cl₂. This is consistent with the mechanism of O₂ inhibition involving its reaction with methyl radicals, which hampers the regeneration of chlorine radicals from Cl₂ and hence chain propagation (Fig. 1).⁶

The promotional effect of the solid surfaces in the catalytic chlorination may stem from the evolution of methyl radicals, as postulated in other methane functionalization processes such as oxidative coupling.⁴⁹ Nonetheless, considering the significantly lower temperature window of methane chlorination (<623 K) with respect to oxidative coupling (>873 K), the surface-mediated formation of methyl radicals is expected to be less favored in the former reaction. This is corroborated by the negligible rate of methane oxidation over the most active H-ZSM-5 catalyst. Hence, the higher rates of methane chlorination over solid catalysts are likely associated with the enhancement of chlorine dissociation by the surfaces, which has the major contribution to the overall reaction barrier in this reaction at low temperatures (Fig. 8a).^{1,29} Namely, the unimolecular dissociation of Cl₂ necessitates the interaction with the second molecular entity, referred to as the mediator (M), which excites the halogen molecule.⁵⁰ In this context, one part of the promotional effect could come from the non-specific chlorine–surface collisions, generally known as “wall effects”.^{11,50} In addition, theoretical studies pointed out that Cl₂ interaction with the defect sites on metal and metal-oxide surfaces (Fig. 8a, inset) may lead to the transfer of the electron density to the antibonding orbitals of the adsorbate.^{51,52} This weakens the Cl–Cl bond and might induce practically barrierless dissociation of Cl₂ into atomic Cl'. The dominant impact of the solid surfaces on the chlorine dissociation step is also corroborated by the observation



that the rate difference between methane chlorination over the solids and methane chlorination in the empty reactor diminishes at higher reaction temperatures, since the latter reaction parameter increases the effectiveness of $\text{Cl}_2\text{-M}$ collisions and shifts the equilibrium towards dissociation, yielding thus sufficiently high concentration of Cl^\cdot radicals that can initiate the reaction.

The radical nature of halogenation reactions over zeolite catalysts raises the question about the origin of activity enhancement over the H-ZSM-5 and BETA materials with respect to the other zeolites as well as other catalytic systems. The results discussed so far suggest that these are not correlated with their electronic, but rather geometric properties. The latter could have an impact on the confinement of the reactants, radical intermediates or transition states, facilitating thus the reaction. In fact, in addition to their acidity, the ability of zeolites to stabilize the reactive intermediates and transition states by the action of the dispersive van der Waals interactions within the well-defined system of micropores, generally referred to as shape selectivity, represents another important aspect of their catalytic action, particularly in the reactions involving hydrocarbons.⁵³ Notably, if the activities of various zeolites attained in the low-temperature regime are plotted against their characteristic pore openings,⁵⁴ a volcano-like dependence is obtained, which exhibits a maximum for micropore diameters of 0.56–0.6 nm (Fig. 8b). Strikingly, the latter micropore size range is comparable to the sum of the kinetic diameters of methane (0.38 nm) and chlo-

rine radicals (0.16 nm),⁵⁵ which are the intermediates participating in the second step that has a smaller contribution to the overall reaction barrier (Fig. 8a). These results indicate that although the chlorination of methane over zeolite catalysts preserves its radical character, the appropriate size of BETA and particularly ZSM-5 structures enables the optimal confinement of the reactive species, enabling thus a higher activity (Fig. 8b).

The above reasoning might also explain the lack of a significant promoting effect of the catalysts in methane bromination. Owing to weaker Br–Br and C–Br bonds as compared to the respective Cl–Cl and C–Cl bonds, the major part of the activation barrier in bromination is determined by C–H bond scission with the Br^\cdot radical (Fig. 8a).²⁹ This leads to a higher reaction temperature of methane bromination as compared to chlorination, at which the concentration of Br^\cdot radicals is high enough to initiate the reaction, so that the promoting effect of the surface on the generation of these species becomes practically irrelevant. As the barrier for CH_4 activation is not substantially affected by the solid, the catalytic and non-catalytic reactions display a comparable rate.

Another peculiar aspect of the catalytic halogenation relates to the formation of CO_x , which was not discussed in previous studies in this field. As shown by the experiments in which CH_2X_2 was taken as a model halogenated product and fed over the representative catalysts and carriers under conditions comparable to those applied in methane halogenation

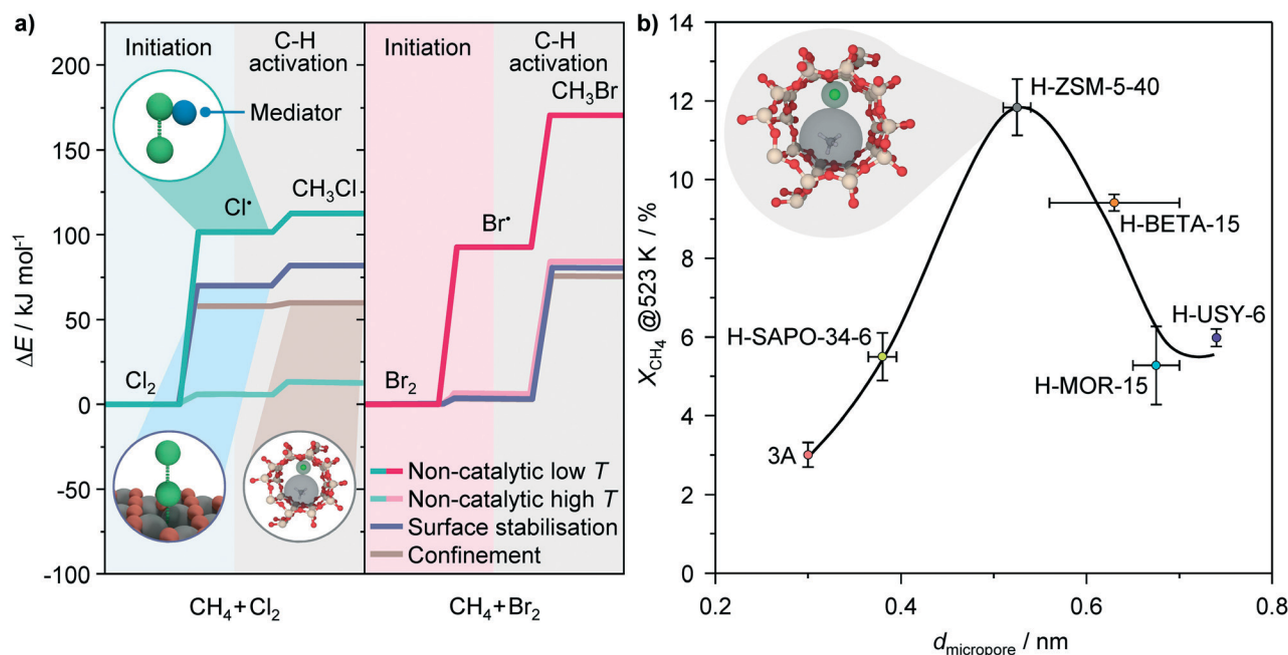


Fig. 8 a) Rate determining steps involved in non-catalytic free-radical methane chlorination (left) and bromination (right) at low and high temperatures, along with the proposed impact of the catalysts on their energy barriers. The energy profiles of the initiation and C–H activation in non-catalytic processes are taken from literature.^{29,56} The insets depict the interaction of halogens with a mediator and defect sites on catalyst surfaces facilitating halogen dissociation, and confinement within the zeolite pores favoring C–X bond formation. b) Conversion of methane in methane chlorination as a function of the micropore diameter of various zeolites. Conditions: $\text{CH}_4:\text{Cl}_2 = 8.5:3.5$, $T = 523 \text{ K}$, $\text{GHSV} = 267 \text{ h}^{-1}$ and $P = 100 \text{ kPa}$.



(Fig. 7b and c), the formation of CO_x primarily originates from the interaction of the halomethanes with the active oxygen sites of supports. In particular, the rate of carbon oxide evolution was high over S-ZrO₂ for both CH₂X₂ and CH₂Br₂ and was generally higher for CH₂Br₂, consistent with the selectivity patterns observed in methane halogenations (Fig. 4 and 5).

In general, our results provide strong hints that radical pathways play the major role in methane halogenation over solid surfaces. In this regard, they are coherent with our previous studies on methane oxyhalogenation, which showed that the catalysts primarily act as the source of reactive halogen radicals that activate methane in the gas phase.^{28,56}

Conclusions

The evaluation of a large number of materials comprising supports, noble metals, halides, oxyhalides, and oxides supported on different carriers, sulfated oxides and zeolites in methane halogenations revealed their positive impact on the chlorination activity, but limited effect on the bromination rate as compared to the respective non-catalysed reactions. Particularly high conversion levels in methane chlorination were attained over the beds of ZSM-5 and BETA zeolites, displaying ≥ 5 times higher productivity than the void reactor of equivalent volume. Still, the fraction of CH₃X over most of the studied catalysts was similar to that achieved in the void reactor or over the bare carriers and was generally higher in methane bromination than in methane chlorination. The major deviations from these trends were displayed by Pt/carbon and zeolite catalysts in methane chlorination, and BETA and ZSM-5 in bromination, which favoured polyhalogenation. Nonetheless, the formation of halomethanes over alumina carriers, alumina-supported catalysts, sulfated oxides, Pt/carbon and most of the zeolites in methane chlorination and virtually all catalytic systems in methane bromination was negatively affected by the evolution of CO_x. In addition, higher operating temperatures led to coke deposition over multiple catalysts in methane bromination as well as H-SAPO-34 in methane chlorination, while the carbon support displayed substantial halogen uptake in both reactions. These side processes stemming from the interaction of halomethanes and halogens with catalysts were typically not considered in previous studies, which in turn could lead to false conclusions on the catalytic performances. The variations of the Si:Al ratio and counter cation as well as the formation of the extraframework aluminium had almost no effect on the activity and selectivity patterns in methane chlorination over the most active ZSM-5 zeolite, which along with significant kinetic similarity between this material and SiO₂ as well as the pronounced O₂ inhibition effect indicates that the reaction primarily proceeds *via* a non-polar radical-chain mechanism. The promotion of activity is likely caused by the confinement of the radical intermediates and/or transition states, which is strongly supported by the volcano-like dependence of the chlorination activity on the size of zeolite

micropores. Thereby, the maximum is achieved for the ZSM-5 channels, the diameter of which corresponds to the sum of kinetic diameters of methane and chlorine radicals. Overall, our results suggest that the radical mechanism prevails in the halogenation over a broad range of catalysts, limiting thus the scope for improving the selectivity to CH₃X. Nonetheless, marked enhancement of the chlorination activity over ZSM-5 and BETA catalysts along with an increased fraction of CH₂Cl₂ and CHCl₃ might exhibit interesting potential for advancements in the production of these important chemicals.

Conflicts of interest

There are no conflicts to declare.

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