Structure and performance of zeolite supported Pd for complete methane oxidation

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\textbf{ABSTRACT}

The influence of zeolite support materials and their impact on CH\textsubscript{4} oxidation activity was studied utilizing Pd supported on H-beta and H-SSZ-13. A correlation between CH\textsubscript{4} oxidation activity, Si/Al ratio (SAR), the type of zeolite framework, reduction-oxidation behaviour, and Pd species present was found by combining catalytic activity measurements with a variety of characterization methods (operando XAS, NH\textsubscript{3}-TPD, SAXS, STEM and NaCl titration). Operando XAS analysis indicated that catalysts with high CH\textsubscript{4} oxidation activity experienced rapid transitions between metallic- and oxidized-Pd states when switching between rich and lean conditions. This behaviour was exhibited by catalysts with dispersed Pd particles. By contrast, the formation of ion-exchanged Pd\textsuperscript{2+} and large Pd particles appeared to have a detrimental effect on the oxidation-reduction behaviour and the conversion of CH\textsubscript{4}. The formation of ion-exchanged Pd\textsuperscript{2+} and large Pd particles was limited by using a highly siliceous beta zeolite support with a low capacity for cation exchange. The same effect was also found using a small-pore SSZ-13 zeolite due to the lower mobility of Pd species. It was found that the zeolite support material should be carefully selected so that the well-dispersed Pd particles remain, and the formation of ion-exchanged Pd\textsuperscript{2+} is minimized.

1. Introduction

Methane is a highly valued fuel and the main component in natural and biogas, however, it is also a greenhouse gas with a global warming potential 28 times higher than that of CO\textsubscript{2} over a 100-year period \cite{1}. The emission of unburnt CH\textsubscript{4} from natural gas and biogas engines can be decreased through the oxidation of the CH\textsubscript{4} remaining in exhaust gas to CO\textsubscript{2}. However, an efficient oxidation catalyst is required to operate under the typical temperature conditions found in an exhaust gas stream, as CH\textsubscript{4} is the most stable hydrocarbon. Pd-based catalysts have been shown to be the most efficient catalysts for the CH\textsubscript{4} oxidation reaction \cite{2}. The kinetic rate limiting step under dry conditions is considered to be an abstraction of the first hydrogen atom, often referred to as the activation of the C-H bond \cite{3,4}.

The efficiency of the CH\textsubscript{4} oxidation of Pd-based catalysts is highly dependent on the oxidation state of Pd. The active phase for CH\textsubscript{4} oxidation is considered to be PdO \cite{5-7}. However, some studies have proposed that a low amount of metallic Pd\textsuperscript{0} can promote CH\textsubscript{4} oxidation, i.e. a mixed phase of PdO/Pd\textsuperscript{0} is more active than PdO \cite{8-11}. It has also been reported that the formation of oxidized Pd is accompanied with an increase in the level of CH\textsubscript{4} oxidation until a certain amount of PdO has been formed. At that point, any further formation of PdO has no discernible influence on the catalytic activity \cite{7,12}. Murata et al. \cite{13} have observed the highest level of CH\textsubscript{4} oxidation using support materials with moderate oxide formation enthalpy resulting in particles consisting of a metallic Pd core surrounded by a PdO shell. Consequently, the relationship between the oxidation state of Pd and CH\textsubscript{4} oxidation is complex. The oxidation process of Pd\textsuperscript{0} is initiated by the quick chemisorption of oxygen onto the Pd\textsuperscript{0} surface \cite{6,7,14}. It is agreed that metallic Pd particles covered with chemisorbed oxygen exhibit some, but low, CH\textsubscript{4} oxidation activity \cite{5,7,15}. The oxidation process of Pd proceeds by the dissolution of oxygen atoms into the Pd\textsuperscript{0} core \cite{14},

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resulting in the slow formation of PdO with a high level of CH4 oxidation \[5,7,11,15\]. PdO reduction can be accomplished at relatively low temperatures under CH4-rich conditions. This makes it possible to study the relation between the oxidation state of Pd and low-temperature CH4 oxidation activity using alternated rich (CH4) and lean (O2/CH4) pulses \[11,16,17\]. The oxidation-reduction properties of Pd also depend on the catalyst composition, for instance, the choice of support materials. As an example, PdO formation is facilitated by using support materials with a high capacity to store oxygen \[11\] and low acidity \[18,19\]. The Pd-to-PdO transition more readily occurs with small Pd particles. The reverse process of PdO decomposition is easier for larger particles because of the weaker interaction between oxygen and Pd in larger particles \[14\]. The strong Pd-O bonds in small crystallites (or those in closer contact with the support) has been suggested to lower the level of CH4 oxidation of small PdO crystallites than of slightly larger PdO particles \[4,20\]. The identification of the active phases as well the mechanism for this reaction is complex and has been the subject of various experimental and theoretical studies \[3,6,21,22\].

Palladium supported on metal oxides, most commonly Al2O3, is one of the most thoroughly investigated catalysts for CH4 oxidation. The Pd/Al2O3 can also be promoted with ceria and other rare-earth elements (La, Pr, Tb) \[23\]. However, Al2O3 supported catalysts severely deactivate in the presence of water vapour, which typically constitutes approx. 10–15 % of the exhaust gas in natural gas combustion \[2\]. Zeolite-supported Pd has gained attention as a CH4 oxidation catalyst, as it can be synthesized to provide high resistance against water deactivation \[17,24–27\]. For instance, the resistance to water deactivation can be significantly improved using low-alumina zeolites as the Pd \[24,26\] and Rh supports \[28\]. An increase in the SiO2/Al2O3 ratio (SAR) is commonly accompanied by a higher conversion of CH4 under dry conditions, which can be related to the formation of highly active PdO agglomerates, and the facilitated reduction and oxidation of Pd due to fewer ion-exchanged Pd2+ species \[24,26\]. It has been shown that the activity and stability of zeolite-supported Pd can be enhanced with the acid removal of extra-framework aluminium, the formation of mesopores upon desilication treatment, and with a decrease in acidity through full ion-exchange of the acidic sites with alkali metal cations \[17,25\]. Lim et al. \[27\] have compared various small-pore zeolites and the associated high activity of long-term CH4 oxidation was attributed to less Pd sintering of catalysts with strong interactions between PdO and zeolite frameworks with high acidity. The complexity of zeolite-supported catalysts is influenced by many factors, such as zeolite framework, SAR, the type of cation exchanged with the zeolite Bronsted acid sites, and the character of the acid sites \[17,24–27,29–31\]. By using zeolites, Pd can form particles either on the external surface of the zeolite and/or internally within its pores. In addition, Pd can form ion-exchanged Pd species on Bronsted acid sites in the zeolite framework. The current literature indicates that ion-exchanged Pd2+ species provide a lower level of CH4 oxidation than PdO particles \[24,26,32,33\], however, this should be confirmed with additional studies using detailed characterization methods. The formation of ion-exchanged Pd2+ species can be limited by using zeolites with high SAR \[24,34\]. The conditions for catalyst pre-treatment are another crucial factor for the formation of ion-exchanged Pd2+ species. The formation of ion-exchanged Pd2+ can be facilitated under lean conditions \[29–32,35\], at high temperatures, and for certain zeolite types, in the presence of water vapour \[36–36\]. Another complexity of zeolite-supported Pd catalysts is that Pd can rearrange to form large Pd+ particles on the external zeolite surface under rich conditions and then re-disperse into small PdO species and/or monodispersed Pd ions under lean conditions \[29–32,35\]. The latter feature is more pronounced for zeolites with a high content of Al; however, the type of zeolite framework can also influence Pd speciation \[29–31\]. Although the benefits of using zeolites over traditional metal oxide support materials for Pd-based CH4 oxidation catalysts have been demonstrated in the literature, there remain a number of open questions for these types of catalytic systems. The nature and distribution of the Pd species and their effect on catalytic activity remains a major and complex issue. Knowledge about this is essential to tailor the synthesis of highly efficient catalysts based on zeolite-supported Pd. The choice of zeolite support has a direct influence on the type of Pd species formed, which can strongly affect the level of CH4 oxidation. By combining catalytic activity measurements with a variety of characterization techniques (XANES, EXAFS, NH3-TPD, SAXS, STEM and NaCl-titrations), we can correlate the level of CH4 oxidation to the Si/Al ratio (SAR), the type of zeolite framework, and the reduction-oxidation behaviour and nature of Pd species.

2. Experimental

2.1. Catalyst preparation

Three catalyst samples were prepared with different support materials according to the methods described in our previous work \[24\]. One Pd/H-SSZ-13 (SAR = 43) sample and two Pd/H-beta samples (SAR = 40, 511) were prepared. The incipient wetness method was used to add 1 wt % of Pd to the support materials. The samples will be referred to hereinafter as PdB40 (Pd/H-beta, SAR = 40), PdB511 (Pd/H-beta, SAR = 511), and PdS43 (Pd/H-SSZ-13, SAR = 43). The highly siliceous beta zeolite PdB511 was chosen based on its high level of activity and stability in the presence of water as well as enhanced sulfur regeneration \[24\]. This catalyst was compared with low SAR Pd/zeolites (PdB40 and PdS43) to further understand its excellent properties.

It should be noted that the H-beta support used for the PdB511 sample was dealuminated with an oxalic acid solution prior to Pd deposition in order to increase its SAR. Details on the dealumination procedure are available in our previous work \[24\]. The contents of Pd, Al, and Si were measured with inductively coupled plasma sector field mass spectrometry (ICP-SFMS, performed by ALS Scandinavia AB) and are listed in Table 1. Parts of the prepared catalyst powders were washedcoated onto cordierite monoliths (diameter=15 mm, length=20 mm) \[24\]. The final monoliths contained 300 mg (±5 mg) of washcoat (95 wt% catalyst powder, 5 wt% boehmite binder, Sasol Disperal P2) and were used for temperature-programmed desorption of NH3 (NH3-TPD). The other methods were performed using the catalyst powder as prepared.

A Pd/Al2O3 reference sample was prepared and tested for comparison with the zeolite supported samples. The results for this sample are presented and discussed in Supplementary material (Figs. S11–15).

| Table 1 | Pd loading, SiO2/Al2O3 molar ratio (SAR), and Pd/Al molar ratio from ICP-SFMS measurements. |
|---------|--------------------------------------------------------------------------------------------------|
| Sample  | Pd loading (wt%) | SAR (SiO2/Al2O3, molar ratio) | Pd/Al (molar ratio) |
| PdB40   | Pd/H-beta        | 1.05                          | 511                   | 1.96                             |
| PdB511  | Pd/H-beta        | 0.92                          | 40                    | 0.14                             |
| PdS43   | Pd/H-SSZ-13      | 1.02                          | 43                    | 0.16                             |
2.2. Characterization and catalytic activity measurements

2.2.1. X-ray absorption spectroscopy (XAS) and catalytic activity measurements

Transmission QuickEXAFS measurements were performed at the SuperXAS beamline [39] at the Swiss Light Source (SLS) using 15 cm long ion chambers filled with 1 bar N\(_2\) and 200 mbar Ar. A Pd foil mounted between the second and third ion chambers was used to calibrate internal energy. The storage ring was operated at 2.4 GeV in top-up mode with a ring current of 400 mA. The polychromatic X-ray beam resulting from a 2.9 T superbend magnet was collimated with a Pt-coated mirror and subsequently monochromatised with a Si(111) channel-cut monochromator. The resulting monochromatic X-ray beam was focused using a Pt-coated toroidal mirror to a spot size of \(\sim 150 \times 150 \mu m\).

A sieved powder fraction (100–150 \(\mu m\)) was loaded between two plugs of quartz wool in a plug-flow reactor cell equipped with 250 \(\mu m\) thick graphite windows [40]. The weights of the loaded samples were 24.1 mg (PdB40), 24.6 mg (PdB511), and 26.6 mg (PdS43). The outlet gases were detected with a mass spectrometer (MS, Omnistar, Pfeiffer).

Prior to the rich-lean pulse experiments, the samples weredegreded and pretreated at 450 \(^\circ\)C according to the protocol in Table 2. The samples were then cooled to 360 \(^\circ\)C in 2 vol% O\(_2\)/Ar. This temperature was maintained while the catalyst powders were exposed to 2 vol% O\(_2\)/1000 ppm CH\(_4\)/Ar for 30 min. This was followed by a rich pulse (70 mL/min, 1430 ppm CH\(_4\)/Ar) for 1 min and then 9 min of lean conditions (100 mL/min, 2 vol% O\(_2\)/1000 ppm CH\(_4\)/Ar). The latter rich-lean pulse sequence was repeated 10 times, i.e., for a period of 100 min. This entire procedure, except for the degreeding steps i–iv in Table 2, was then repeated at 330 \(^\circ\)C, 300 \(^\circ\)C, and 270 \(^\circ\)C. After the pulse sequence at each temperature, the samples were cooled to the next temperature in 2 vol% O\(_2\)/Ar. The MS displayed sudden spikes in some of the gas concentration data. These spikes were caused by pressure fluctuations but did not influence the overall interpretation of the data.

X-ray absorption spectra were recorded at the end of the 30-min steady state CH\(_4\) oxidation, during the first rich and lean pulse, and the two last cycles of rich-lean pulses. The spectra were background corrected, normalized, and interpolated using ProXES data processing software [41]. The fraction of Pd\(^{2+}\) was determined with linear combination fitting (LCF) of the X-ray absorption near edge structure (XANES) spectra (24,250–24,450 keV) using the spectra of PdO and Pd\(^{+}\) metal foil as references. The extended X-ray absorption fine structure (EXAFS) analysis was performed on the Fourier transformed \(k^2\)-weighted EXAFS spectra (k range 3.0–10.25 \(\AA\)) using the DEMETER software package [42]. The amplitude reduction factor (S\(_{0}^2\)) was determined by fitting the first Pd-Pd coordination shell of a simultaneously collected Pd\(^{+}\) foil. The pseudo Debye-Waller factor (\(\sigma^2\)) for the first Pd-Pd shell of metallic Pd was estimated with the Debye model using a Debye temperature \(\Theta\) of 274 K. The spectra of the oxidized Pd samples were fitted to three scattering paths, i.e., Pd-O, Pd-(O)-PdI, and Pd-(O)-Pd2. The \(\sigma^2\) and \(\Delta R\) for these three shells were extracted from the fit of fully oxidized samples at each experimental temperature.

2.2.2. Temperature-programmed desorption of NH\(_3\) (NH\(_3\)-TPD)

The acidity of the samples was evaluated with NH\(_3\)-TPD. NH\(_3\) adsorption and desorption experiments were conducted using wash-coated monoliths and a flow reactor, described elsewhere [24], with a total flow of 800 mL/min (13 600 h\(^{-1}\)). The outlet gases were measured with a Fourier transform infrared (FTIR) spectrometer (MKS MultiGas 2030 HS FTIR). The monolith samples were first degreened and pretreated according to Table 2 and subsequently cooled to 100 \(^\circ\)C in 2 vol% O\(_2\)/Ar. This was followed by NH\(_3\) adsorption in 500 ppm NH\(_3\)/Ar at 100 \(^\circ\)C for 2 h. NH\(_3\) was then removed from the feed, and the samples were flushed with Ar for 1.5 h. The TPD was subsequently conducted in Ar by heating from 100 \(^\circ\)C to 500 \(^\circ\)C at 10 \(^\circ\)C/min.

2.2.3. Titration of ion-exchanged Pd\(^{2+}\)

The amount of ion-exchanged Pd\(^{2+}\) atoms was estimated with NaCl titration based on the method developed by Ogura et al. [43], demonstrating the high degree of ion-exchanged Pd\(^{2+}\) in Pd/ZSM-5 (98%) compared to Pd/SiO\(_2\) (0%), which was further verified with NO adsorption experiments. The sample powder used for this analysis was degreened, pre-treated (according to Table 2), and exposed to the experimental sequence at 360 \(^\circ\)C (i.e., 30-min steady state CH\(_4\) oxidation and 10 rich-lean pulse cycles). Sample powder (70 mg) was added to a 50 mL NaCl solution (0.1 M) at 80 \(^\circ\)C and heated and stirred for 1 h. This step was repeated one more time after removing the liquid phase with centrifugation, i.e., the powder was ion-exchanged in NaCl solution twice. Finally, the powder was washed with milliQ water and dried at room temperature. The remaining Pd content in the sample after the NaCl process was determined with ICP-SFMS.

2.2.4. Particle size estimation

Pd particle size was studied with scanning transmission electron microscopy (STEM) and small- and wide-angle X-ray scattering (SAXS/WAXS). The powder used for these techniques was degreened, pre-treated, and exposed to the experimental sequence at 360 \(^\circ\)C (i.e., 30-min steady state CH\(_4\) oxidation and 10 rich-lean pulse cycles). The STEM images were obtained with a FEI Titan 80–300 microscope using an acceleration voltage of 300 kV. Small- and wide-angle X-ray scattering (SAXS/WAXS) measurements were performed with the MOUSE, a customised Xeuss 2.0 (Xenocs, France). X-rays were generated from a microfocus X-ray tube with a copper target, and a multilayer optic was employed to parallelise and monochromatize the beam to the Cu K\(_\alpha\) wavelength of 0.1542 nm. The samples were examined as powders fixed between two pieces of Scotch Magic tape in the vacuum sample chamber. Data collection was performed using an in-vacuum Eiger I M detector (Dectris, Switzerland), which was placed at distances between 137–2507 mm from the sample. The resulting data was processed and scaled to absolute units using the DAWN software package according to

| Table 2 |
| --- |
| Conditions for degreening and pre-treatment at 450 \(^\circ\)C prior to the pulse operando XAS experiments. |
| Degreening | Pre-treatment |
| i) 2 vol% H\(_2\)/Ar, 20 min | v) 2 vol% O\(_2\)/Ar, 20 min |
| ii) 1000 ppm CH\(_4\)/2 vol% O\(_2\)/Ar, 10 min | |
| iii) 2 vol% H\(_2\)/Ar, 10 min | |
| iv) 1000 ppm CH\(_4\)/2 vol% O\(_2\)/Ar, 10 min | |

Fig. 1. Temperature-programmed desorption of NH\(_3\) (NH\(_3\)-TPD) for PdB511 (orange line), PdB40 (blue line), and PdS43 (green line) samples. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).
standardised procedures [44,45]. SAXS data analysis was performed using McSAS, a Monte Carlo fitting programme that utilises a form-free, minimum assumption methodology [46].

3. Results and discussion

3.1. Ex-situ characterization

The acidity of the samples was studied with NH₃ adsorption at 100 °C followed by NH₃-TPD (Fig. 1). The number of acid sites was proportional to the amount of desorbed NH₃ and the desorption temperature reflected the acid strength of the catalyst. The NH₃ desorption in the lower temperature region, around 200 °C, was ascribed to weakly adsorbed NH₃, for instance, on weak Lewis acid sites or physisorbed NH₃ [47–52]. The peaks at approx. 273–285 °C were assigned to Lewis acid sites of PdO [26,52,53]. Additional peaks at 339 °C for the PdB40 sample and at 420 °C for the PdB43 sample were assigned to the strong Bronsted acid sites of the zeolite framework, as reported in the literature for H-beta and H-SSZ-13 zeolites [35,47,54,55]. The use of different zeolite supports resulted in distinctly different NH₃-TPD profiles. The highest acidity was exhibited by the PdB40 and PdB43 samples, with a total NH₃ desorption amount of 0.141 and 0.134 mmol per monolith, respectively. However, the acid strength and amount of strong acid sites of the PdB40 and PdB43 were different. Despite similar SAR, the PdB43 sample exhibited a higher number of strong Bronsted acid sites than the PdB40 sample, which was seen in the deconvolution of the NH₃-TPD (see Fig. S1). 77 % of the desorbed NH₃ of the PdB43 was released in the high temperature peak associated with Bronsted acid sites. The value for the PdB40 sample was 63 %. Our results are consistent with the work reported by Zheng et al. [35]. By contrast, the dealumination treatment of the PdB511 sample caused a loss in the total acid sites on the sample (the total amount of NH₃ desorption was 13-fold lower than that of the PdB40 sample) and eliminated the strong Bronsted acid sites, which was reflected by the absence of NH₃ desorption at temperatures above 300 °C.

Pd particle size was estimated with SAXS (Table 3, Figs. 2 and S2 (Supplementary material)) using samples that had been degreened, pre-treated, and exposed to the experimental sequence used for operando XAS measurements at 360 °C (i.e. 30-min steady state CH₄ oxidation and 10 rich-lean pulse cycles). Note that the samples were subsequently cooled in 2 vol% O₂ to avoid any reduction of PdO. The results from SAXS were in good agreement with the STEM images in Fig. S3. However, the small PdO particles were difficult to distinguish in the STEM images, and consequently, the size of the small Pd particles was more accurately estimated with SAXS.

The PdB511 sample with the lowest Al content showed a broad range of Pd particle sizes with an average radius of 9 nm (Fig. 2a, Table 3). The formation of relatively large Pd particles in the PdB511 sample is not surprising as the interactions between the PdO and the B511 zeolite were expected to be weak due to the low acidity of this zeolite (Fig. 1). The strong acidity of zeolite frameworks with high Al content allows for the stabilization of well-dispersed PdO particles and/or monodispersed PdO clusters (Fig. 2c, Table 3). This is likely a result of strong PdO–zeolite interactions attributed to the high acidity of the B511 zeolite with the incipient wetness method used to deposit Pd onto the zeolite. In a recent publication, Khivantsev et al. [57] have associated the formation of large Pd particles in zeolites with high SAR to the incomplete impregnation of the hydrophobic pores of the zeolites with the aqueous Pd solution.

Table 3

| Pd particle radius measured with Small-Angle X-ray Scattering (SAXS) and portion of Pd exchangeable with Na⁺ obtained from NaCl titration. The full particle size distribution is shown in Figs. 2. The measurements were performed on the sample powder after rich-lean cycling at 360 °C and cooling to room temperature in the presence of O₂. |
|---|
| Total average radius (nm) | Average radius of particles < 10 nm (nm) | Average radius of particles > 10 nm (nm) | Portion of Pd exchangeable with Na⁺ (%) |
|---|---|---|---|
| PdB511 | 9.38 ± 0.54 | 5.39 ± 0.07 | 13.99 ± 0.09 | 4 |
| PdB40 | 18.39 ± 0.20 | 3.76 ± 0.01 | 22.65 ± 0.21 | 13 |
| PdB43 | 4.99 ± 0.37 | 4.24 ± 0.04 | 27.19 ± 14.12 | 2 |

Fig. 2. Pd particle size distribution of a) PdB511, b) PdB40, and c) PdB43 samples measured with Small-Angle X-ray Scattering (SAXS). The measurements were performed after rich-lean cycling at 360 °C and cooling to room temperature in the presence of O₂.

Pd ions [24,26,29,30,34,56]. Another factor that may also have contributed to the formation of large Pd particles in the PdB511 sample is the combination of the high hydrophobicity of the B511 zeolite with the incipient wetness method used to deposit Pd onto the zeolite. In a recent publication, Khivantsev et al. [57] have associated the formation of large Pd particles in zeolites with high SAR to the incomplete impregnation of the hydrophobic pores of the zeolites with the aqueous Pd solution.

The H-SSZ-13 zeolite in the PdB43 sample was able to stabilize the smallest PdO particles, on average with a radius of 5 nm, and a few large PdO clusters (Fig. 2c, Table 3). This is likely a result of strong PdO–zeolite interactions attributed to the high acidity of the S43 zeolite (Fig. 1) [24,26,29,30,34,56]. The PdB40 sample (Fig. 2b, Table 3) was dominated by large PdO particles (average radius 23 nm) in addition to a lower number of small particles (4 nm), although the SAR for the PdB40 sample was approximately the same as for the PdB43.

The amount of ion-exchanged Pd⁺⁺ in the zeolite-supported samples was estimated with NaCl titration, a method introduced by Ogura et al. [41] and utilized in several studies [35,37,58,59]. The results in Table 3
The catalyst samples were further evaluated under repeated rich (CH$_4$/Ar) and lean (CH$_4$/O$_2$/Ar) pulses. The rich-lean sequences were repeated ten times to ensure stable performance, which can be seen by

![Fourier transformed EXAFS spectra](image)

**Fig. 4.** Fourier transformed EXAFS spectra (non-phase shift corrected) recorded after 30 min of isothermal CH$_4$ oxidation at 360 °C (corresponding to the last minute in Fig. 3). Dashed lines represent EXAFS fits.

**3.2. Steady state CH$_4$ oxidation**

Steady state of CH$_4$ oxidation was tested at constant temperatures in a CH$_4$/O$_2$ feed for 30 min. The samples were degreased and pre-treated prior to this procedure (Table 2). The obtained CH$_4$ conversion at 360 °C is shown in Fig. 3. Both the PdB511 and PdS43 samples exhibited a similarly high level of CH$_4$ conversion. PdB40 was the least active sample with considerably less activity. This trend applied to all investigated temperatures (270–360 °C), however, the lower temperature was naturally accompanied by lower activity for all samples (Fig. S4).

![Methane conversion at 360 °C for 30 min in 1000 ppm CH$_4$/2 vol% O$_2$ using degreased and pre-treated samples.](image)

**Fig. 3.** Methane conversion at 360 °C for 30 min in 1000 ppm CH$_4$/2 vol% O$_2$ using degreased and pre-treated samples.

XANES spectra were collected after 30 min of isothermal CH$_4$ oxidation. The LCF of the PdB40 and PdS43 references for the XANES spectra showed that all the samples had been fully oxidized after 30 min of isothermal CH$_4$ oxidation (Fig. S5). In other words, the samples exhibited significantly different levels of activity despite that all samples were fully oxidized. This means that the level of CH$_4$ oxidation was not exclusively related to the oxidation state of the Pd but was also influenced by other factors.

To obtain more insight into this behaviour, the EXAFS spectra corresponding to the last minute of the time-on-stream shown in Fig. 3 were analysed. The Fourier transformed k$^2$-weighted EXAFS spectra shown in Fig. 4 correspond to the average of all spectra collected during the last minute of the 30-min sequence shown in Fig. 3. All fitting results are given in Figs. S6, S7, and Table S1. The best fit of the FT-EXAFS spectra showed that the coordination environment of the Pd-O scattering shell at 1.5 Å (all values non-phase shift corrected) was complete (CN$_{Pd-O}$=4). The LCF of the PdO and Pd$^+$ references for the XANES spectra showed that the coordination environment of the Pd-O scattering shell at 1.5 Å (all values non-phase shift corrected) was complete (CN$_{Pd-O}$=4).

The coordination number of this path (CN$_{Pd-(o)-Pd}$) strongly correlated particle to PdO particle size. However, EXAFS is a bulk technique, meaning that the recorded spectra represent the average coordination environment of the absorbing Pd atom. Thus, a relatively low CN$_{Pd-(o)-Pd}$ value can either indicate small PdO particles or a few large PdO particles combined with a high number of isolated Pd entities without neighbouring Pd atoms [36–38]. Based on the EXAFS analysis, CN$_{Pd-(o)-Pd}$ decreased in the order of PdB511 (CN$_{Pd-(o)-Pd}$ = 10.1, 360 °C) > PdS43 (CN$_{Pd-(o)-Pd}$ = 8.4, 360 °C) ≈ PdB40 (CN$_{Pd-(o)-Pd}$ = 8.1, 360 °C) as shown in Table 4. The trend was the same for all investigated temperatures (Table S1). This is in reasonable agreement with the aforementioned results of the characterization of Pd species from the SAXS and STEM techniques. The fact that the Pd-(o)-Pd shell of the PdB40 sample was not fully coordinated (CN$_{Pd-(o)-Pd}$ < 12) is likely because the PdB40 sample also contained monodispersed Pd ions without neighbouring Pd atoms, which lowered the average CN$_{Pd-(o)-Pd}$. The monodispersed Pd ions without neighbouring Pd atoms were also found in the NaCl titration (Table 3) results.

**3.3. CH$_4$ oxidation under alternating rich and lean pulses**

The catalyst samples were further evaluated under repeated rich (CH$_4$/Ar) and lean (CH$_4$/O$_2$/Ar) pulses. The rich-lean sequences were repeated ten times to ensure stable performance, which can be seen by
the nearly identical Pd$^{2+}$ profiles and CH$_4$ oxidation activity during cycle 9-10 in Figs. S8 and S9. The focus of the following analysis will therefore only concern the last two sequences (cycle 9-10), shown in Fig. 5.

### 3.3.1. Response to rich pulses

The rich pulses at 360 °C and 330 °C (Fig. 5a-b) resulted in nearly full reduction of all the samples (remaining oxidized fraction approx. 10 % Pd$^{2+}$). The extent of reduction was lower (Fig. 5c-d) at lower temperatures. The PdB40 sample could not be reduced at 300 °C (Fig. 5a) and remained nearly completely oxidized throughout the rich pulses. The PdB40 sample could not be reduced at 300 °C (Fig. 5a) and remained nearly completely oxidized throughout the rich pulses. The PdB511 (approx. 20 % Pd$^{2+}$) sample retained the oxide form (almost 100 % Pd$^{2+}$). The PdB511 sample was partially reduced (approx. 70 % Pd$^{2+}$). The responses to the rich pulses revealed that the samples had major differences in their inherent resistance to reduction. The order of reduction was PdB511 > PdS43 > PdB40, suggesting that both the SAR and the type of zeolite framework strongly influence the stability of Pd$^{2+}$. The above characterization with NaCl titration indicated that the PdB40 sample contained ion-exchanged Pd$^{2+}$ at the end of the lean pulse. However, it is not clear whether the monodispersed Pd ions remained in the same state during the rich pulse. Previous studies have shown that monoatomic Pd$^{2+}$ species and small PdO particles rearrange into PdO particles under rich conditions [29-32, 35]. It is possible that the monodispersed Pd ions in the PdB40 sample contributed to the stability of Pd against reduction (Fig. 5c-d).

The EXAFS data (Fig. 6) of the samples reduced under rich pulses at 360 °C and 330 °C were characterized by an intense Pd-Pd shell of metallic Pd$^0$ at 2.5 Å (non-phase shift corrected). Fitting this shell yielded CN$_{Pd-Pd}$ ≈ 10 for all samples. Large Pd$^0$ particles in about the same size range formed in all samples. Assuming spherical fcc Pd$^0$ particles with a density of 12.02 g/cm$^3$, the calculated particle diameter based on Hill’s function [60] was 2.3 nm for CN$_{Pd-Pd}$ = 10. CN$_{Pd-Pd}$ may be slightly underestimated but shows that the particles were located on the external zeolite surfaces. Fitting fully oxidized PdB40 (270–300 °C) and PdS43 (270 °C) samples returned CN$_{PdO-Pd}$ = 4 and CN$_{PdO-Pd}$ = 7, indicating that well-dispersed PdO particles and/or large PdO particles in combination with ion-exchanged Pd$^{2+}$ were present in the samples. Based on the ex-situ characterization of the oxidized samples, it is likely

![Fig. 5. Data acquired during the 9th rich-lean sequence and the 10th rich pulse at the indicated temperatures. (a-d) Fraction of Pd$^{2+}$ obtained from LCP of XANES spectra. (e-h) Corresponding outlet concentration of CH$_4$. The red dashed line illustrates the inlet CH$_4$ concentration. The oxidation rate (APd$^{2+}$ fraction/Δt, s$^{-1}$) at the beginning of the lean pulse is indicated in (a). Grey and white areas represent rich and lean pulses, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).](image-url)
that the PdB40 sample contained small PdO particles, monodispersed Pd ions, and large PdO particles, and the PdS43 sample probably contained mainly well-dispersed PdO particles.

### 3.3.2. Response to lean pulses

The introduction of oxygen was accompanied by the re-oxidation of Pd. The PdB511 and PdS43 samples displayed close to identical re-oxidation profiles at 360 °C, the PdB40 sample was the most difficult one to re-oxidize (approx. 59 % Pd retained at 300 °C), whereas the PdB40 and PdS43 samples remained completely oxidized at 300 °C and 270 °C. The PdB511 and PdS43 samples yielded about the same level of CH₄ oxidation increased when the Pd was being oxidized [5-7], which is in line with our results.

The PdB511 and PdS43 samples yielded about the same level of CH₄ oxidation and showed about the same state of oxidation throughout the pulse experiments. By contrast, the PdB40 sample, which was the most difficult sample to reduce and re-oxidize, had the lowest level of CH₄ oxidation. Approximately 10 % of the Pd in the PdB40 sample is estimated to have been ion-exchanged with the Brønsted acid sites in the zeolite at the end of the lean pulse at 360 °C, and large PdO particles were found using SAXS and STEM. This combination of large PdO particles and monodispersed Pd ions appears to have had a detrimental effect on the catalytic activity of CH₄ oxidation [7,12], which is in line with our results.

### 3.3.3. CH₄ oxidation activity under alternated rich and lean pulses

Fig. 5e–h shows the measured outlet concentrations of CH₄ during the rich-lean pulse experiments. A comparison of the CH₄ outlet concentration (Fig. 5e–h) and the Pd²⁺ fraction (Fig. 5a–d) recorded during exposure to rich and lean pulses shows that the CH₄ conversion can be directly related to the oxidation state of Pd. In other words, CH₄ oxidation increased when the Pd was being oxidized [5-7]. However, the CH₄ conversion was approximately the same for the completely oxidized catalysts prior to the first rich pulse (Fig. 3) and at the end of the lean pulse (Fig. 5e–h), where the catalysts had only been partially oxidized. This means that the same activity levels were obtained despite the different oxidation states. It has previously been reported that the CH₄ oxidation activity only increases up to a certain level of PdO formation, and that further oxidation of the Pd particles does not influence the catalytic activity of CH₄ oxidation [7,12], which is in line with our results.
increase, whereas a volcano-shaped correlation was found for weak interactions in Pd/MgO [63]. To gain further information regarding the effect Pd-particle sizes, the turnover frequencies (TOFs) were calculated at 270 °C where the CH₄ conversion over the three catalysts was low (4, 7, and 12.5 % for PdB40, PdB511, and PdS43, respectively), and differential reactor conditions can be applied. The TOF monotonously increased from 0.017 to 0.027 s⁻¹ as the particle size increased from 5.0–18.4 nm, as shown in Fig. S10. Thus, in term of the TOF, the larger the PdO particles, the larger CH₄ oxidation rate. This dependence of TOF on particle size suggests that there were strong metal-support interactions. It should be noted that because the Pd dispersion obtained from SAXS is used in the TOF calculations, the TOF is per active Pd site. However, many of the Pd sites are not accessible in the case of large particles, thus accounting for a low overall conversion.

Another possible explanation for the lower activity found for the PdB40 sample is related to the fact that complete CH₄ oxidation over PdO is considered to occur via the Mars van Krevelen mechanism [4,64,65], i.e. CH₄ molecules react with oxygen atoms of the PdO to form CO₂ and H₂O. Thus, the CH₂ oxidation reaction should benefit from a catalyst that easily forms catalytically active PdO. However, if the interactions between Pd and O are very strong, i.e. if the PdO is difficult to reduce, the CH₂ oxidation rate may decrease [4,20]. Consequently, CH₂ oxidation benefits from a catalyst with rapid transitions between oxidized and reduced states upon switching between lean and rich conditions. Since those rapid transitions between oxidized and reduced state were found for PdS43 and PdB511, but not for PdB40, our interpretation may explain why the CH₄ conversion level generally decreased in the order, PdS43 ≈ PdB511 > PdB40.

Lower sample temperature resulted in lower activity and less efficient reduction and re-oxidation. 270 °C and 300 °C were not sufficiently high temperatures to reduce PdB40, as shown in Fig. 5c-d, so Pd in PdB40 remained completely oxidized. This was reflected by the absence of CH₂ conversion for the PdB40 sample in Fig. 5g-h. 300 °C was not high enough to activate the C–H bond, which is required to initiate the PdO reduction by CH₄ and the catalysed CH₂ oxidation reaction [14]. The PdS43 sample exhibited the same behaviour at 270 °C. The PdB511 and PdS43 samples had only been partly reduced and re-oxidized at 300 °C, which was also reflected by the correspondingly low CH₄ conversion on these catalysts at 300 °C.

The formation of well-dispersed Pd particles in the PdS43 sample explains the facilitated reduction and re-oxidation of Pd, and the high level of activity for CH₂ oxidation shown in Fig. 5. It is, however, possible that the particles present in the PdB511 sample were small enough to provide the same effect, as the performance of the PdB511 sample was similar to that of the PdS43. By contrast, the coexistence of large Pd particles and monodispersed Pd ions in the PdB40 sample hampered Pd reduction and re-oxidation, thus providing a low level of CH₄ oxidation. This agrees with our previous findings [17,24,25], i.e. Pd/zeolites with dispersed and stabilized Pd particles have a higher CH₄ oxidation, impeded by Pd sintering and the formation of ion-exchanged Pd⁴⁺ species. It was demonstrated that Pd particles were stabilized by the formation of mesopores in the zeolite through mild desilication, the removal of extra framework alumina by selective dealumination [25], and the removal of acid sites in the zeolite either by full ion-exchange with alkali metals [17,25] or by using highly siliceous zeolites [24]. It should be noted that the zeolite pores of beta and SSZ-13 are much smaller (<1 nm [66]) than the particles observed with the characterization methods used in the present study. This suggests that the particles were located mainly on external zeolite surfaces or possibly within the mesopores of the zeolite. We suggest that the selection of the zeolite support is crucial in order to obtain highly active CH₂ oxidation catalysts. The selection should take into consideration the necessity to minimize the formation of monodispersed Pd atoms and to promote good dispersion of Pd particles simultaneously.

4. Conclusions

The effect of different zeolite supports for Pd-based CH₄ oxidation catalysts using activity tests and various characterization techniques was examined using three different zeolite supports, H-beta (SAR = 511 and 40) and H-SSZ-13 (SAR = 43). The results clearly show that both the type of zeolite framework and the SAR impact the type of Pd species formed, as well as the catalytic performance. Our findings suggest that the selection of a zeolite support is paramount to minimizing the formation of ion-exchanged Pd²⁺ species and the formation of well-dispersed Pd particles in order to obtain a highly active CH₄ oxidation catalyst. The formation of ion-exchanged Pd⁴⁺ and large Pd particles can be prevented by using highly siliceous zeolites with a low number of Bronsted acidic sites and small-pore zeolites, which decrease Pd mobility. By contrast, large pore size and low SAR facilitate the formation of ion-exchanged Pd²⁺ species and large Pd particles. This makes the catalyst more difficult to reduce and to re-oxidize and is accompanied by a low conversion of CH₄. The catalyst with the highest conversion of CH₄ was characterized by rapid transitions between metallic and oxidized Pd during switches between rich and lean pulses.

CRediT authorship contribution statement

Ida Friberg: Investigation, Formal analysis, Conceptualization, Writing - original draft. Adam H. Clark: Investigation, Formal analysis, Writing - review & editing. Phuoc Hoang Ho: Formal analysis, Writing - review & editing. Nadezda Sadokhina: Investigation, Supervision, Writing - review & editing. Glen J. Smale: Investigation, Formal analysis, Writing - review & editing. Jungwon Woo: Investigation, Writing - review & editing. Xavier Auvray: Investigation, Writing - review & editing. Davide Ferri: Supervision, Conceptualization, Methodology, Writing - review & editing. Maarten Nachttegaal: Supervision, Conceptualization, Methodology, Writing - review & editing. Olivier Krocher: Supervision, Conceptualization, Methodology, Writing - review & editing. Louise Olsson: Supervision, Conceptualization, Methodology, Writing - review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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