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Deciphering SO_2 poisoning mechanisms for passive NO_x adsorption: A kinetic modeling approach and development of a high-resistance catalyst

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Dawei Yao^a, Phuoc Hoang Ho^a, Johann C. Wurzenberger^b, Thomas Glatz^b, Wei Di^a, Rojin Feizie Ilmasani^a, Derek Creaser^a, Louise Olsson^{a,*}

^a Chemical Engineering division, Competence Center for Catalysis, Chalmers University of Technology, Göteborg SE-41296, Sweden ^b AVL List GmbH, Graz, Austria

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ABSTRACT

Passive NOx adsorption (PNA) is a promising technology aimed at reducing NO_x emissions from vehicles during the cold start phase of the engine. This work investigated the SO₂ poisoning mechanism of PNA through a combination of experimental research and kinetic modeling, leading to the development of a novel PNA sample with high resistance to SO₂ poisoning. Pd/SSZ-13 samples were synthesized using different drying conditions, revealing that samples dried at room temperature showed lower degradation (10 %) compared to those dried at 80 °C (26 %). Investigation into the degradation revealed that ion-exchanged Pd sites with a hydroxyl group were more resistant to SO₂ poisoning than other Pd sites. It is also found that SO₂ aids in NO_x storage on Pd sites, enhancing the PNA performance. A kinetic model was developed to describe the SO₂ poisoning behavior and its influence on NO_x storage. The model, which was verified under various conditions, effectively simulated the PNA behavior and SO₂ poisoning of Pd/SSZ-13.

1. Introduction

With the increasing global transportation sector, gasoline and diesel engines still have a great share in the production of new vehicles [1]. During the cold start of the engine, the nitrogen oxides (NO_x) are inevitably formed and become a main environmental issue. The cold start phase is characterized by the ignition of the engine and the subsequent thermal elevation of the engine block, tailpipe, and catalysts from their ambient temperature [2]. The efficiency of the exhaust after-treatment system in mitigating emissions experiences a substantial reduction during this phase owing to kinetic constraints. As emission regulations become increasingly stringent, coupled with the reduction in engine exhaust temperatures due to advancements in fuel efficiency, the need to address cold start emissions has been magnified [3].

Over recent years, Passive NO_x Adsorbers (PNAs) have emerged as a significant area of interest due to their potential to address the persistent challenge of NO_x emissions during the cold start phase in automotive exhaust systems [4]. PNAs exhibit a unique functionality due to their capacity to temporarily store NO_x at low temperatures. This capability enables the adsorption of NO_x during the cold start phase. Thereafter, when the temperature is further increased typically above ~ 200 °C the

urea can be dosed and the NH₃-Selective Catalytic Reduction (SCR) system can operate. Thus, at higher temperature when the NOx are desorbed from the PNA it can be reduced in the following SCR unit and thereby receiving a high NOx conversion both at low and high temperature. To thoroughly appraise the effectiveness and potential applicability of PNAs, several key parameters must be considered, including the NO_x storage at low temperatures, adsorption rate, NO_x desorption temperature, compatibility with real exhaust conditions, resistance to degradation, and cost-efficiency [5].

Like most after-treatment catalysts, PNAs often experience significant thermal aging and poisoning. During PNA performance evaluations, this degradation is typically reflected in a decrease in lowtemperature NO_x storage capacity and a shift in the desorption temperature into a non-optimal range. Therefore, understanding the material transformations that occur during aging and poisoning is essential for enhancing catalyst stability. There is growing interest in Pdimpregnated zeolites for PNA applications. Specifically, palladium loaded on small-pore zeolites, such as SSZ-13, have shown potential due to their comparatively appropriate NO_x desorption temperature window [6,7] and excellent tolerance to H_2O [5,8]. However, the stability of Pd/ SSZ-13 in the PNA process can be compromised by the presence of CO,

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^{*} Corresponding author. E-mail address: louise.olsson@chalmers.se (L. Olsson).

phosphorus, and sulfur. Previous reports indicate that high CO concentration can trigger Ostwald ripening of ion-exchanged Pd sites and particle migration of external PdO sites, consequently leading to the degradation of PNA performance [9,10]. Phosphorous is also recognized to cause both physical and chemical deactivation of Pd/SSZ-13, including the physical blockage of active Pd sites by P₂O₅ and dealumination of the zeolite framework caused by the formation of AlPO4 [11–13]. A recent patent from Johnson Matthey showed approximately 25 % degradation of PNA performance following sulfur poisoning [14].

However, among the current research, there is no conclusion on the mechanism of sulfur poisoning, or how to design a PNA sample that maintains stability in the presence of sulfur during the PNA process. In this work we, for the first time in literature, present a kinetic model that can describe the sulfur poisoning of Pd/SSZ-13, used as a PNA. In addition, we have developed the synthesis method for Pd/SSZ-13 and we have successfully enhanced the sulfur resistance of the PNA. In more detail, we synthesized a typical Pd/SSZ-13 using an impregnation method and then modified the drying process after impregnation to achieve different surface Pd site distributions. By varying the quantity of different surface Pd active sites, we were able to examine the stability of these Pd sites during sulfur poisoning in a multicycle PNA evaluation. The PNA experiments clearly showed that the modified synthesis method resulted in a PNA with higher sulfur resistance. The PNA samples were characterized before and after the multicycle test, and it was demonstrated that the different Pd active sites changed after sulfur poisoning. After collecting experimental data, we were able to develop a kinetic model to describe both the PNA performance and sulfur poisoning behavior, as well as the enhancement of NO_x storage by sulfur during the multicycle PNA test. This kinetic model can also simulate PNA performance under varying sulfur concentrations and adsorption temperatures.

2. Experiment

2.1. Synthesis of passive NO_x adsorber

2.1.1. Synthesis of SSZ-13

SSZ-13 zeolites with a SAR of 12 were synthesized using a hydrothermal method reported from our previous works [12,15-17]. In summary, a gel was formed by vigorously stirring a mixture of 1.6 g NaOH, 35.16 g TMAda-OH, 2.76 g Al(OH)₃, and 24 g SiO₂ in 132 mL of deionized water. The resulting mixture was then transferred into an autoclave and heated at 160 °C for 96 h with rotational stirring. After being cooled to ambient temperature, the mixture underwent a centrifugation washing process using 900 mL of water that had been deionized. The resulting product was then dried at 80 °C for 6 h and subsequently calcined at 600 °C for 8 h. NH₄-SSZ-13 was synthesized through an ion-exchange procedure. Specifically, a mixture was prepared by combining 15 g of Na-SSZ-13 with 300 mL of 1 M NH₄NO₃ solution, which was then heated up to a temperature of 80 °C. The resulting mixture was washed with 800 mL of deionized water. To ensure the complete removal of Na ions, the procedure of ion-exchange was performed two additional times. The resulting sample was subsequently dried at 80 °C for six hours.

The Pd/SSZ-13 samples were prepared using incipient wetness impregnation, where a Pd(NO₃)₂ solution was used to load 1 wt% Pd onto NH₄-SSZ-13. The resulting sample was subjected to different drying procedures (all drying was conducted in air). In one approach, the sample was dried at room temperature for 48 h. After drying, the sample underwent a calcination process at a temperature of 500 °C sustained for a duration of 5 h, and referred to as 'Roomdry'. In contrast, a different sample was subjected to an overnight drying process at 80 °C and a 5-hour calcination at 500 °C, a standard procedure for loading Pd on SSZ-13 [6,12,16]. This sample was subsequently termed as '80dry'.

2.1.2. Monolith preparation

The cordierite monoliths were structured in a honeycomb shape with a cell density of 400 cpsi. The monoliths were then sliced into small cylinders measuring 2 cm in length and 2.1 cm in diameter. To clean the channel surfaces, the monoliths were calcined at 550 °C for 2 h. After that, a mixture of 95 wt% Pd/SSZ-13 powder and 5 wt% Boehmite (Dispersal P2) was mixed in an ethanol/H₂O (1:1) solution. The mixture was then introduced into the channels of the monoliths and dried at 80 °C. This procedure was repeated several times until the desired weight of the washcoat was reached (700 \pm 7 mg). Finally, the loaded monoliths were calcined at 500 °C for 5 h.

2.2. Evaluation of the passive NO_x adsorber

2.2.1. Multicycling experiments

As we described in our previous studies [15,18–20], the PNA evaluation system consists of multiple mass flow controllers, a system dedicated to H_2O evaporation, a reactor tube equipped with a pair of thermocouples, along with a Fourier Transform Infrared (FTIR) spectrometer, specifically the MKS Multigas 2030 variant. The cordierite monolith was positioned within the reactor tube, and two thermocouples were positioned upstream of the monolith and inside the monolith channel, respectively.

2.2.2. Degreening process

To enhance the Pd sites generated during the ion-exchange phase and to ensure the stability of the PNA sample, a degreening process was carried out on all the samples prior to material characterization and PNA evaluation [21]. The monolith, once positioned in the reactor, was subjected to a treatment at 750 °C under a total flow rate of 900 mL/min with 400 ppm NO_x, 5 % H₂O, and 8 % O₂. After the treatment, the monolith was purged with Ar at 750 °C to eliminate any adsorbed species and then cooled down to either 80 °C for the subsequent evaluation or to room temperature for further characterizations. Notably, to confirm the absence of any pre-adsorbed species on the monolith, two test cycles under identical conditions were performed prior to SO₂ poisoning. Consistent NO_x storage behavior across these cycles indicates the successful removal of pre-adsorbed species following the degreening process.

2.2.3. Evaluation process

After undergoing the degreening process, the PNA samples were evaluated over 12 cycles to assess their performance with and without SO₂. Cycles 3–7 included the addition of 26–104 ppm SO₂ at 390 ppm NO_x, 4.6 % H₂O, and 6.9 % O₂, while cycles 1, 2, and 8 were conducted without SO₂. The evaluation process is shown in Fig. 1. Each evaluation cycle consisted of three stages that simulated the cold start period of an engine.

The first stage involved NO_x adsorption at a low temperature of 80 °C to 150 °C, with the gas mixture containing 200 ppm NO_x, 26–104 ppm SO₂ (added in cycles 3–7), 5 % H₂O, 8 % O₂, and the balance in Ar. The NO/NO_x ratio was 0.982. This stage was maintained for 45 min.

In the second stage, temperature-programmed desorption (TPD) was conducted. While keeping the gas conditions identical, the temperature was gradually increased to 550 $^\circ\rm C$ at a rate of 10 $^\circ\rm C/min$ and then held for 15 min.

In the third stage, the regeneration process was conducted. For cycles 1, 2, and 8–12, NO_x was first stopped. Following that, the system's temperature was kept at 550 °C for 5 min before being cooled down to the adsorption temperature for the next cycle. For cycles with SO₂, both NO_x and SO₂ were stopped, lowered to the adsorption temperature for the following cycle, without holding the temperature at 550 °C.

The total NO_x storage capacity is determined from the desorption branch of the NO_x profile, utilizing the equation below:



Fig. 1. Multicycle evaluation process was conducted on a PNA sample to assess its performance under different concentrations of SO_2 and at varying adsorption temperatures. The other reaction conditions included NO_2/NO_x at 1.8 %, 4.6 % H₂O, and 6.9 % O₂ in Ar, with a flow rate of 905 mL/min.

$$n = \int_{t=45\min}^{t=60\min} Q \times [c_{out}(t) - c_{in}(t)]dt$$
(1)

where *n* represents the total NO_x stored, *Q* is the total flow rate, *t* is the reaction time, c_{out} and c_{in} are the NO_x concentration at the outlet and inlet, respectively.

2.3. Characterization methods

An FEI Tecnai microscope was utilized for conducting Transmission Electron Microscopy (TEM), operated at a voltage of 200 kV. To set up the TEM sample, the powder was dispersed in ethanol and placed on a copper grid underpinned by a carbon film. To gather statistical data pertaining to the diameters and the average diameter of nearly 100 Pd particles in each specimen, the TEM images were analyzed.

Measurements using X-ray diffraction (XRD) were performed employing a Bruker D8 Discover apparatus, set at a voltage of 40 kV and a current of 40 mA. The radiation source was Cu K α radiation ($\lambda = 1.5418$ Å), which was filtered at ambient temperature. The resulting XRD pattern was achieved through a scanning range of 10° to 60°, at a rate of 1° per minute.

To analyze standard powder samples, a Scanning Electron Microscope (SEM, JEOL 7800F Prime) was utilized. Samples were goldsputter-coated for conductivity and examined under a high-resolution SEM at 20 kV. Imaging focused on surface morphology and particle size, with magnifications from 1000x to 25000x. Multiple areas were scrutinized, and representative micrographs were captured.

The nitrogen physisorption test for Pd/SSZ-13 was conducted using a Micromeritics ASAP 2020. The experiment aimed to determine the specific surface area of the palladium-loaded SSZ-13 zeolite. Prior to measurement, the sample was degassed under vacuum at 150 °C to remove moisture and other adsorbed gases. Nitrogen gas was then adsorbed onto the sample at liquid nitrogen temperature. The amount of gas adsorbed at various pressures was measured to construct an adsorption isotherm. The Brunauer-Emmett-Teller (BET) theory was applied to the isotherm to calculate the specific surface area of Pd/SSZ-13.

Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) was performed using a Bruker Vertex 70 spectrometer, with an in-situ diffuse chamber, an MCT detector and CaF windows. The gas mixture was controlled by mass flow controllers and an H_2O evaporator. The Pd/SSZ-13 samples, either degreened or tested, were scratched from the monolith wall and 20 mg of the powder was placed in the chamber.

For in-situ DRIFT of CO adsorption, the samples were then pretreated at 200 °C under pure Ar and cooled down to 30 °C. After cooling to 30 °C, the background was measured. CO adsorption was measured by scanning at 30 °C with a resolution of 4 cm⁻¹. During 20 min, 4000 ppm CO in Ar was added, followed by pure Ar for 5 min to flush out the gas phase CO. For the operando DRIFT of NO_x and SO₂ co-adsorption, NO_x adsorption was first conducted by exposing the samples to 390 ppm NO_x and 6.9 % O₂ in Ar for 30 min at 80 °C, with a subsequent addition of 104 ppm SO₂ in Ar for another 30 min. After exposure, the system was flushed with pure Ar for 5 min to remove residual gases.

The process of CO chemisorption was carried out utilizing an ASAP 2020 Chemi Plus instrument from the Micromeritics Instruments Corporation, which comes equipped with an additional chemisorption device. The powder was positioned in a tube and sealed within the device, which was followed by a pretreatment at 200 °C Celsius under a vacuum to eliminate any possible contaminants. Post-pretreatment, the tube's temperature was reduced to 35 °C, and CO was introduced ranging from 0.13 to 0.80 bar to capture the initial adsorption isotherm. The isotherm was then repeated after conducting an evacuation process to remove the CO from physisorption. The difference between the two isotherms provided the amount of chemisorbed CO.

2.4. Experimental results

2.4.1. Physicochemical properties of Pd/SSZ-13 samples

Scanning Electron Microscopy (SEM) images of SSZ-13 samples, prepared under room-dried and 80 °C-dried conditions, were captured at magnifications of x5000, x10000, and x20000, as shown in Fig. 2. The images consistently demonstrated similar morphologies for both drying conditions, highlighting the crystalline structure characteristic of the SSZ-13 zeolite. The SEM analysis provided a clear depiction of the zeolite's uniform and well-defined crystalline features.

Table 1 outlines the physicochemical properties of Pd/SSZ-13 samples dried at room temperature (Roomdry) and 80 °C (80dry). The surface area of the Roomdry sample is 764.2 m2/g, surpassing the 80dry's 681.4 m2/g, which suggests that higher drying temperatures may decrease surface area. However, pore volumes and sizes are closely matched, with the Roomdry at 0.068 cm³/g and 41.8 Å, and the 80dry at 0.060 cm³/g and 41.4 Å, indicating that drying conditions do not significantly alter the pore structure, maintaining the porous architecture essential for their performance in NO_x adsorption and release processes.



Fig. 2. SEM images of Pd/SSZ-13 sample. A-C: Images for Roomdry. D-F: Images for 80dry.

Table 1

Physicochemical properties of Pd/SSZ-13 samples.

Sample	Surface area (m2/ g) ^a	Pore volume (cm3/g) ^b	Pore size (Å) ^b
1 %Pd/SSZ-13 Roomdry	764.2	0.068	41.8
1Pd/SSZ-13 80dry	681.4	0.060	41.4

^a calculated by Brunauer-Emmett-Teller (BET) method.

^b calculated by Barrett-Joyner-Halenda method (BJH) method based on desorption branch.

2.4.2. No_x adsorption and release in multicycle PNA Test, including SO_2 poisoning

The NO_x storage and release performance, as well as SO₂ poisoning behavior, were examined over the Pd/SSZ-13 samples that were either dried after palladium impregnation at room temperature ("Roomdry") or at 80 °C ("80dry"). These two samples were evaluated over 12 PNA cycles, where cycles 3–7 included an addition of 104 ppm SO₂, while the other cycles were conducted without SO₂. The operating temperature on the adsorption branch of each cycle was set to 80 °C. To facilitate the comparison between different cycles, the time points of NO_x injection in

every cycle were considered as the beginning of each cycle (Time of stream = 0). We selected Cycle 2 (the last cycle before SO_2 addition), Cycle 3 (the first cycle with SO_2 addition), Cycle 7 (the last cycle with SO_2 addition), and Cycle 8 (the first cycle after SO_2 shut off) to represent the NO_x storage and release performance, and SO_2 poisoning behavior in this multicycle PNA test. Their respective NO and NO_2 profiles are displayed in Fig. 3. The full-cycle performance is shown in Fig. S1, Supplementary Information (SI). There was no degradation of PNA performance prior to the addition of SO_2 , as evidenced by the lack of change between cycles 1 and 2 (Fig. S1A, D). Similarly, no degradation of PNA performance was observed between cycles 9 and 12 (Fig. S1C, F), suggesting sample stability after poisoning. Therefore, we primarily focused on cycles 2–9 in subsequent experimental and modeling work.

In Fig. 3, we observe two NO release peaks at 290 °C and 460 °C, and NO oxidation to NO₂ peaks at 150–180 °C for both PNA samples. This suggests the presence of at least two Pd sites for NO storage and another Pd site for NO₂ formation. Notably, the peak around 2750 s marks the start of NO release at 290 °C. Based on previous kinetic modeling work on PNA [16,17,22,23], the NO oxidation to NO₂ peaks at 150–180 °C can be attributed to the reduction of ion-exchanged Pd sites with a hydroxyl group (Z⁻[Pd(II)OH]⁺) to lower valence ion-exchanged Pd sites (Z⁻Pd(I)), as this reaction has relatively low activation energy compared

.



Fig. 3. PNA and SO₂ poisoning performance of Roomdry (A) and 80dry (B) in selected cycles. Reaction conditions: 290 ppm NO_x, NO₂/NO_x = 1.8 %, 104 ppm SO₂ (only in Cycle 3–7), 4.6 % H₂O, 6.9 % O₂ in Ar with the flow rate of 905 mL/min. Adsorption temperature: 80 °C. 'Roomdry' and '80dry' denote the Pd/SSZ-13 samples which were dried at room temperature or 80 °C after Pd impregnation.

to the reduction of other Pd site reactions [24]. The formation and decomposition of Pd nitrates on PdO also contribute to these NO oxidation to NO_2 peaks [25].

With the presence of SO₂ from cycles 3 to 7, it is clear that the NO₂ generation at 150–180 °C was markedly suppressed, while the NO consumption peak at the same temperature persisted. These results suggest that SO₂ can inhibit the reduction of Z⁻[Pd(II)OH]⁺ or directly cover the surface of Z⁻[Pd(II)OH]⁺. Simultaneously, SO₂ may also hinder the formation of Pd nitrates but provide another site for NO adsorption at the same temperature.

The NO desorption peaks at 290 $^\circ C$ and 460 $^\circ C$ can be attributed to

the release of NO species from Z⁻Pd(II)Z⁻ and Z⁻Pd(I), respectively, in accordance with the adsorption energies of NO on these different Pd sites [16,17,24]. The Z⁻Pd(II)Z⁻–NO peak in the 80dry sample is much larger than that in the Roomdry sample, indicating that drying at 80 °C generates more ion-exchanged Pd sites without a hydroxyl group, while room-drying tends to produce more hydroxyl groups on ion-exchanged Pd sites.

Upon the addition of SO₂, both Z⁻Pd(II)Z⁻–NO peaks on Roomdry and 80dry samples decreased significantly, with a more pronounced decrease in the 80dry sample due to its larger initial quantity. This suggests that Z⁻Pd(II)Z⁻ sites are susceptible to SO₂ poisoning, either by being covered by SO₂ species or converted to another inactive Pd species. In contrast, the Z⁻[Pd(II)OH]⁺–NO (released at 460 °C) is considerably more stable than Z⁻Pd(II)Z⁻–NO, as evidenced by its peak intensity remaining unchanged before and after SO₂ poisoning (in cycle 2 and cycle 8). During cycles 3 to 7, the existing SO₂ even appeared to aid NO adsorption on Z⁻[Pd(II)OH]⁺–NO as the related peak increased. This finding also suggests that the remaining NO adsorption peak at 150–180 °C after adding SO₂ may be attributed to the increase of the NO desorption peak at 460 °C, since the mass of NO_x adsorption and desorption should be balanced. Notably, the slight peak increase at approximately 3300 s in cycle 8 results from minor disturbance due to unstable H₂O injection.

Fig. 4A illustrates the calculated NO_x storage amount and the NO_x/Pd ratio for each cycle of the Roomdry sample under varying SO₂ concentrations. Interestingly, upon addition of 104 ppm SO₂, the NO_x adsorption amount rose from 3.0×10^{-4} mol per monolith to 4.2×10^{-4} mol per monolith, then gradually declined in the subsequent cycles. With the decreasing concentration of SO₂, the adsorption amount of NO_x also proportionally decreased. Conversely, changes in SO₂ concentration did not have a pronounced effect on the degradation in the multicycle test, especially in cycle 7 when the SO₂ inlet was shut off.

These results suggest that SO_2 can both enhance NO_x adsorption and deactivate the PNA. For the enhancement aspect, SO_2 seems to offer additional adsorption sites for NO_x or allow co-adsorption of NO_x on the same sites, as NO_x demonstrated predominantly singular adsorption on Pd sites in the presence of H₂O prior to SO_2 addition [6,16]. The results indicate that the SO₂-induced poisoning, after SO₂ is removed, is likely due to the blockage of Pd active sites by SO_2 , since an increase in SO_2 concentration only slightly influences the poisoning behavior.

Fig. 4B presents the calculated NO_x desorption amounts and the NO_x/Pd ratios for the Roomdry and 80dry samples. In the first two cycles without SO₂, the NO_x storage amount is similar for both Roomdry and 80dry, suggesting comparable amounts of exposed Pd sites in both samples. However, as seen from the NO & NO2 profiles in cycle 2 in Fig. 5A, 80dry contains more Z⁻Pd(II)Z⁻ sites capable of releasing NO at 290 °C, while Roomdry has more $Z^{[Pd(II)OH]^+}/Z^{Pd(I)}$ sites that can release NO at 460 °C. Upon the addition of SO₂ from cycle 3, the NO_x storage in 80dry does not increase as significantly as in Roomdry. We suggest that this is due to the lower quantity of Z⁻[Pd(II)OH]⁺ sites in 80dry, which are the adsorption sites that can be strongly enhanced by SO₂. The two samples, Roomdry and 80dry, are compared in Fig. 5, for different cycles. As depicted in Fig. 4B and 4C, the NO release peak from $Z^{[Pd(II)OH]^+}/Z^{Pd(I)}$ (460 °C) in Roomdry is much higher than that in 80dry during sulfur exposure. Following the shutting of SO₂ inlet from cycle 8, degradation in 80dry (26 %) is substantially more severe than in Roomdry (10 %) (Fig. 4B), because 80dry contains more Z⁻Pd(II)Z⁻ sites that are vulnerable to SO2. Moreover, the NO desorption peak area in Roomdry is higher than that in 80dry, as Roomdry contains more

$Z^{-}[Pd(II)OH]^{+}/Z^{-}Pd(I)$ sites (Fig. 5D).

2.4.3. Characterization of Pd sites during SO₂ poisoning

To further validate the SO₂ poisoning mechanism inferred from the PNA evaluation results, in situ DRIFT of CO adsorption was conducted to determine the distribution of exposed Pd sites in Roomdry and 80dry samples before and after SO₂ poisoning, as depicted in Fig. 6A. CO is well-established as a probe molecule stems from its specific binding characteristics, which allow for the precise identification of various Pd active sites on the zeolites. These sites are pivotal in the adsorption process, influencing the catalyst's performance regarding NO. By examining how CO interacts with Pd sites, we can infer the dispersion, oxidation state, and coordination environment of Pd, which are critical factors affecting the adsorption and catalytic behavior towards NO [26,27]. The Pd/SSZ-13 sample was first exposed to 4000 ppm CO/Ar for 20 min at 30 °C, then purged with pure Ar for 5 min to collect the spectra. Peaks at 2211 cm^{-1} and 2190 cm^{-1} was assigned to CO adsorbed on Z⁻Pd(II)Z⁻ sites [28,29], while the peak at 2111 cm⁻¹ was attributed to linear adsorption on Pd/PdO_x clusters [13,30]. The peak at 1895 cm⁻¹ is attributed to bridged carbonyls of Pd⁰ and multi-bridged carbonyls on small Pd clusters [31,32]. Peaks at 2127 cm⁻¹ and 2146 cm^{-1} are assigned to CO adsorption on ion $Z^{-}[Pd(II)OH]^{+}$ complexes [28,32,33], and the peak at 2164 cm⁻¹ can be assigned to CO adsorption on the surface of Z⁻[Pd(II)OH]⁺ [13,34,35]. The relative intensity proportion of the CO adsorption peaks, which correspond to different Pd sites, are calculated and listed in Table 2. Compared to the Roomdry sample, more Z⁻Pd(II)Z⁻ sites exist in the 80dry sample; however, they are vulnerable to SO_2 poisoning, as evidenced by the decrease in CO-Z⁻Pd(II)Z⁻ peaks at 2211 cm⁻¹ and 2190 cm⁻¹ after poisoning. Following SO₂ poisoning, the relative intensity of linear CO-Pd⁰/PdO_x (2211 cm⁻¹) decreased in both Roomdry and 80dry, suggesting a reduction in the surface area of Pd/PdOx after poisoning. This could indicate that Pd/PdOx sites may be covered by SO2 or that SO2 induced the agglomeration of Pd/PdOx into larger nanoparticles. The CO-Z⁻[Pd (II)OH]⁺ peaks (2127 and 2146 cm⁻¹) are dominant both before and after SO₂ poisoning for both Roomdry and 80dry samples, suggesting that the $Z^{-}[Pd(II)OH]^{+}$ sites are more stable than other Pd sites. The peak proportion analysis reveals that Z⁻[Pd(II)OH]⁺ is the predominant site across all samples. However, the Roomdry sample contains more stable Z⁻[Pd(II)OH]⁺ sites compared to the 80dry sample. Conversely, the 80dry sample has a higher quantity of Z⁻Pd(II)Z⁻ sites than those found on Roomdry. These observations align with the findings from the PNA evaluation.

To explore the reaction mechanism spectrographically, operando DRIFT spectroscopy of NO and SO₂ co-adsorption was performed, as depicted in Fig. 6B. The adsorption process was carried out at 80 $^{\circ}$ C, using gas concentrations of 390 ppm NO_x, 6.9 $^{\circ}$ O₂, and 104 ppm SO₂, which align with the conditions used in the PNA tests. It is important to



Fig. 4. Stored amount of NO_x per monolith (700 mg washcoat) and NO_x/Pd in each cycle of PNA test with SO₂ poisoning. A: Roomdry sample tested under different SO₂ concentrations. B: Roomdry and 80dry sample tested under 104 ppm SO₂. 'Roomdry' and '80dry' denote the Pd/SSZ-13 samples which were dried at room temperature or 80 °C after Pd impregnation.



Fig. 5. PNA performance of Roomdry and 80dry in Cycle 2 (last cycle without SO₂), Cycle 3 (first cycle with SO₂), Cycle 7 (last cycle with SO₂) and Cycle 8 (first cycle shut SO₂). Reaction conditions: 290 ppm NO_x, NO₂/NO_x = 1.8 %, 104 ppm SO₂ (only in Cycle 3–7), 4.6 % H₂O, 6.9 % O₂ in Ar with the flow rate of 905 mL/min. Adsorption temperature: 80 °C. 'Roomdry' and '80dry' denote the Pd/SSZ-13 samples which were dried at room temperature or 80 °C after Pd impregnation.

note that H₂O was omitted from the system to prevent the formation of sulfuric acid, which could potentially damage the DRIFT chamber. Analysis of Fig. 6B reveals the presence of peaks at 1360 cm⁻¹, 1376 cm⁻¹, and 1202 cm⁻¹, indicative of bulk sulfate species on aluminum sites [36,37], while the peak at 1256 cm⁻¹ is associated with PdSO₄ species [36,38,39]. Additionally, the peaks at 1277 cm⁻¹ and possibly 1292 cm⁻¹ suggest the chemisorption of SO₂ on Pd sites [36,37], implying that SO₂ chemisorption occurs on ion-exchanged Pd sites. The peak at 1303 cm⁻¹ is attributed to physisorbed SO₂ [36,37,40]. In conclusion, the DRIFT study indicates two primary forms of SO₂ adsorption on the Pd-SSZ-13 samples: chemisorbed SO₂ on ion-exchanged Pd sites and PdSO₄, both of which are crucial in understanding the SO₂ poisoning behavior towards the PNA sample.

XRD was performed on the Pd/SSZ-13 samples to assess the condition of both the SSZ-13 support and Pd sites, as illustrated in Fig. 7. Both the Roomdry and 80dry samples clearly contain well-crystallized SSZ-13. In addition, the diffraction peaks of SSZ-13 can be discerned from the XRD patterns post-PNA test, implying that the SO₂ contamination did not notably affect the SSZ-13 structure. The diffraction peak seen at 2θ $= 33.8^{\circ}$ is a characteristic peak of the PdO(101) phase. To amplify the differentiation between the peaks in various samples, the XRD patterns within this area are magnified, as demonstrated in the right panel of Fig. 7. The PdO(101) diffraction peaks are observed at $2\theta = 33.8^{\circ}$ for fresh Roomdry and 80dry completely overlap each other and in addition, the peaks are broad suggesting that the Pd sites are well dispersed. After the PNA experiments, both the PdO(101) peaks for Roomdry and 80dry increased, indicating that SO₂ may induce agglomeration of Pd/ PdOx sites during the PNA process. These results are consistent with the in-situ DRIFT of CO adsorption (Fig. 6A). In addition, the results are supported by TEM (Fig. S2), where the particle size increased from 3.0 nm for degreened Roomdry sample, to 7.7 nm after poisoning with 26 ppm SO₂ to 6.7 nm after exposure to 54 ppm SO₂.

3. Kinetic modelling

The kinetic model used in this work was built upon our previous PNA model that described the PNA performance [17]. We utilized AVL BOOSTTM, embedded with an aftertreatment module, along with the AVL User Coding Interface for constructing the simulations. The parameters relating to mass transfer and heat transfer were established according to the property database in AVL CRUISETM M. Further information on the equations and assumptions can be found in our previous works on PNA modeling [15–17].

3.1. Reactor model

The design of our reactor model incorporates three critical components: the inlet boundary, the monolith, and the outlet boundary. A single channel model is applied, which is very common for kinetic models of monoliths [16,41-45]. Across the system, we have set 20 grid points: 18 of these are arranged along the axial path within the monolith channels, and the remaining two are designated for the inlet and outlet boundaries. The model consists of both the internal mass transfer within the washcoat as well as the external mass transfer in the gas phase. The film model [46] serves as the basis for our description of external mass transfer. In terms of internal mass transfer, we have employed a pore diffusion model with a constant effective diffusivity, in line with the research advocated by Chatterjee et al [47] and consistently applied in our earlier studies [16,41-45]. Given the low concentration of NO_x, the reaction heat associated with adsorption/desorption could be neglected. This simplifying assumption aligns with practices commonly seen in a multitude of kinetic modeling studies [41,42,44,48,49].

3.2. Kinetic model

The reaction rate constants are defined by the Arrhenius equation as



Fig. 6. A: in situ DRIFT of CO adsorption on PNA samples prior to and post SO_2 contamination. Test condition: 80 °C, 4000 ppm CO, balanced in Ar. B. operando DRIFT of NO and SO_2 adsorption on PNA samples. Test conditions: 80 °C, 390 ppm NO_{x_2} 6.9 % O_2 and 104 ppm SO_2 , balanced in Ar.

Table 2

Peak intensity proportion of all of the ion-exchanged Pd sites $(Z^Pd(II)Z^-/Z^-[Pd(II)OH]^+)$ and Pd/PdO_x calculated from in situ DRIFT of CO adsorption as shown in Fig. 6. (Roomdry' and '80dry' denote the Pd/SSZ-13 samples which were dried at room temperature or 80 °C after Pd impregnation.

	Z ⁻ Pd(II)Z ^{-a}	Z ⁻ [Pd(II)OH] ⁺	Pd/PdO _x
Roomdry	0.34	0.52	0.14
Roomdry 26 ppm SO ₂	0.18	0.65	0.17
Roomday 52 ppm SO ₂	0.19	0.68	0.12
Roomdry 106 ppm SO ₂	0.20	0.65	0.14
80dry	0.43	0.44	0.13
80dry 106 ppm SO ₂	0.20	0.62	0.17

^a The proportion of peak intensity was determined in each spectrum by dividing the specified peak intensity by the total peak intensity.



Fig. 7. XRD patterns of PNA samples before and after SO₂ poisoning. 'Roomdry' and '80dry' denote the Pd/SSZ-13 samples which were dried at room temperature or 80 °C after Pd impregnation.

follows:

$$k_j = A_j e^{\frac{-E_{A_j}}{(RT_s)}} \tag{2}$$

Here, for reaction j, A_j represents the pre-exponential factor (s^{-1}), E_{A,j} stands for the activation energy (J/mol), and k_j refers to the rate constant j (s^{-1}). The Temkin isotherm was employed to linearly associate the activation energy with adsorption coverage [50]:

$$E_{des} = E_{des}^0 (1 - \alpha \theta) \tag{3}$$

In this equation, E_{des}^0 symbolizes the desorption energy at zero coverage, α is the coverage dependence constant, and θ signifies the fractional coverage.

This reaction model comprises two elements: one describes the basic adsorption and release behaviors of NO_x , and the second part outlines the effects of SO_2 poisoning, which is the focus of the current work.

3.2.1. Kinetic model for describing PNA performance before SO₂ poisoning

Based on our previous kinetic model on PNA [17], we used a series of basic reactions to describe the NO_x adsorption and release within the first two PNA test cycles. These were conducted in the presence of NO_x, H₂O, and O₂. Since no SO₂ was present in these initial cycles, no degradation of PNA was observed, as illustrated in Fig. S1 A, D. Consequently, no reactions involving SO₂ poisoning were included in the basic reaction model. The NOx adsorption and release profiles (Fig. 3), along with several prior studies [6,13,15,22,23], indicate that there are three initial Pd active sites: two ion-exchanged Pd sites (with and without a hydroxyl group, Z⁻Pd(II)Z⁻, Z⁻[Pd(II)OH]⁺), and one Pd/ PdO_x cluster on the external surface of zeolites. We categorized the reactions into three groups based on the initial Pd sites involved (Table 3). The first group, 'Reaction series Z', comprises reactions occurring at Z⁻Pd(II)Z⁻ sites, including the adsorption and release of NO and NO₂ species (Reactions Z1 and Z2). 'Reaction series ZH' includes reactions at Z⁻[Pd(II)OH]⁺ sites. This involves the adsorption and desorption of NO (Reaction ZH1), reduction of Z⁻[Pd(II)OH]⁺ to Z⁻Pd(I) (Reaction ZH2), adsorption and desorption of NO on Z⁻Pd(I) (Reaction ZH3), and re-

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Basic reaction model of	f passive NO _x	adsorption	without SO ₂	poisoning
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Reactions	Reaction Step	Rate Expressions
Z1	$Z^{Pd}(II)Z^{+}NO \rightleftharpoons Z^{Pd}(II)$	$r_{Z1} = k_{Z1f} \Psi_Z \theta_Z y_{NO} \textbf{ - } k_{Z1b} \Psi_Z \theta_{Z\text{-}NO}$
Z2	Z = NO $Z^{-}Pd(II)Z^{-} + NO_2 \rightleftharpoons Z^{-}Pd(II)$ $Z^{-}NO_2$	$r_{Z2} = k_{Z2f} \Psi_Z \theta_Z y_{NO2} \text{ - } k_{Z2b} \Psi_Z \theta_{Z\text{-}NO2}$
ZH1	$Z^{-}[Pd(II)OH]^{+}+NO \rightleftharpoons Z^{-}[Pd$	$r_{ZH1} = k_{ZH1f} \Psi_{ZH} \theta_{ZH1} y_{NO} \mbox{ - }$
	(II)OH] ⁺ –NO	$k_{ZH1b}\Psi_{ZH}\theta_{ZH1-NO}$
ZH2	$2Z^{-}[Pd(II)OH]^{+}-NO \rightleftharpoons$	$r_{ZH2} = k_{ZH2f} (\Psi_{ZH} \theta_{ZH1-NO})^2$
	$Z^Pd(I) + Z^Pd(I)-NO + NO_2 + H_2O$	$-k_{ZH2b}\Psi_{ZH}\theta_{ZH2}\Psi_{ZH}\theta_{ZH2-NO}y_{NO2}y_{H2O}$
ZH3	$Z^{-}Pd(I) + NO \rightleftharpoons Z^{-}Pd(I) - NO$	$\mathbf{r}_{\text{ZH3}} = \mathbf{k}_{\text{ZH3f}} \Psi_{\text{ZH}} \theta_{\text{ZH2}} \mathbf{y}_{\text{NO}} - \mathbf{k}_{\text{NO}} \mathbf{y}_{\text{NO}} \mathbf$
7114	27=D-1(1) + U O + 0 EO	$K_{ZH3b} = ZH^{0}ZH^{2}-NO$
ZH4	$2Z Pd(1) + H_2O + 0.5O_2 \rightleftharpoons$	$\mathbf{r}_{\text{ZH4}} = \mathbf{\kappa}_{\text{ZH4f}} (\Psi_{\text{ZH}} \Theta_{\text{ZH2}}) \mathbf{y}_{\text{H2O}} \mathbf{y}_{\text{O2}}$
	2Z ⁻ [Pd(II)OH] ⁺	$-k_{ZH4b}(\Psi_{ZH}\theta_{ZH1})^2$
01	$Pd(II)O + 2NO + 1.5O_2 \rightleftharpoons Pd$	$r_{O1} = k_{O1f} \Psi_O \theta_{O1} y_{NO}^2 y_{O2}^{1.5} - k_{O1b} \Psi_O \theta_{O2}$
	(NO ₃) ₂	
02	$Pd(NO_3)_2 + NO \Rightarrow Pd(II)O +$	$r_{O2} = k_{O2f} \Psi_O \theta_{O2} y_{NO}$ -
	3NO ₂	$k_{O2b}\Psi_{O}\theta_{O1}y_{NO2}^{3}$

oxidation of Z⁻Pd(I) to Z⁻[Pd(II)OH]⁺ (Reaction ZH4). These reactions on Z⁻[Pd(II)OH]⁺ form a complete NO oxidation process, with the kinetic parameters aligning with thermodynamic reactions (Table 4). It is worth noting that the activation barrier for the reduction of Z⁻[Pd(II) OH]⁺ is significantly lower than for Z⁻Pd(II)Z⁻ reduction [24]. As a result, with increasing temperature, NO is unlikely to reduce Z⁻Pd(II)Z⁻ prior to desorption. The final reaction group, 'Reaction series O', contains the formation (Reaction O1) and decomposition (Reaction O2) of Pd(NO₃)₂. The kinetic parameters of these reactions also conform to thermodynamic constraints (Table 4).

Given the similar synthesis processes of Roomdry and 80dry samples (excluding the drying process), reactions at the Pd active sites in both samples should have identical kinetic parameters. However, the quantities of each Pd site are expected to vary between the two samples. The kinetic parameters of the basic reaction model are adjusted according to the NO and NO₂ profiles from the second cycle of both Roomdry and 80dry samples, as detailed in Table S1 and S2. Fig. 8 presents the NO and NO₂ profiles from the second cycle of both Roomdry and 80dry samples, as detailed in Table S1 and S2. Fig. 8 presents the NO and NO₂ profiles from the experimental evaluation. There is a strong correlation between the simulated and experimental results. Notably, the ratio between Z⁻Pd(II)Z⁻ and Z⁻[Pd(II)OH]⁺ derived from the kinetic model matches the ratio calculated from in-situ DRIFT of CO adsorption, as outlined in Table 5. These findings suggest that the basic kinetic model and parameters effectively describe PNA performance prior to the introduction of SO₂ into the system.

3.2.2. Kinetic model for SO_2 poisoning of $Z^-Pd(II)Z$

Based on the evaluation and characterization results, the number of exposed Z⁻Pd(II)Z⁻ sites decreases after SO₂ poisoning, primarily due to SO₂ covering on the Z⁻Pd(II)Z⁻ site and deactivating its ability to store NO_x species. Consequently, the adsorption of SO₂ on the Z⁻Pd(II)Z⁻ site is incorporated into the reaction model, as depicted in Reaction Z3 (Table 6). SO₂ anchoring on Z_Pd(II)Z⁻ forms an inactive Z⁻Pd(II)Z⁻-SO₂ site, which cannot contribute to NO_x adsorption. There is no reverse reaction for Z3 because the NO desorption peak on Z⁻Pd(II)Z⁻ (at 290 °C) remains stable after terminating SO $_2$ exposure in cycles 8–12, as shown in Fig. 1C, F. This indicates that Z⁻Pd(II)Z⁻-SO2 is stable throughout the PNA multicycle test. The kinetic model results, incorporating Reaction Z3, are presented in Fig. 9. We selected the first and last cycles with SO₂ (cycles 3 and 7) and the first cycle after shutting off SO₂ (cycle 8) to represent the overall kinetic model. It should be noted that the kinetic modeling results for cycles 1 and 2 are unaffected by the addition of Reaction Z3, as no SO₂ is present in these cycles.

It can be seen in Fig. 9 that the NO release peak from $Z^-Pd(II)Z^-$ (at 290 °C) drastically decreased in the first cycle when adding SO₂ (cycle 3) in the kinetic modeling results, which also consists with our evaluation results. After shutting off SO₂ in cycle 8, all the NO desorption and conversion peaks as well as the NO₂ generation peak match well with the experimental results. From this result it also can be deduced that the Z-Pd(II)Z- sites were completely poisoned by SO₂, while Z-[Pd(II)OH]⁺ can still maintain a similar NOx uptake as before SO₂ poisoning, because the peak at 460 °C did not change. On the contrary, in Cycle 3 and Cycle 7 we can see that the kinetic modeling result deviate from experimental results, mainly for the NO desorption at 460 °C and NO₂ formation at ~

Table 4

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Reactions	Overall reaction	Restriction on parameters
2ZH1 + ZH2- ZH3 + ZH4	$\begin{array}{l} NO + 0.5O_2 \\ \rightleftharpoons NO_2 \end{array}$	$\begin{split} \Delta H_{NOox} &= 2[E_{ZH1f}E_{ZH1b}(1-\alpha_{ZH1b}\theta_{ZH1-NO})] + \\ (E_{ZH2f}E_{ZH2b}) - [E_{ZH3f}E_{ZH5f}(1-\alpha_{ZH5f}\theta_{ZH2-NO})] + \\ (E_{ZH4f}E_{ZH4b}) \\ \Delta S_{NOox} &= 2Rln(A_{ZH1f}/A_{ZH1b}) + Rln(A_{ZH2f}/A_{ZH2b}) - Rln(A_{ZH3f}/A_{ZH5f}) + Rln(A_{ZH4f}/A_{ZH4b}) \end{split}$
02 + 03	$\begin{array}{l} 3\mathrm{NO} + 1.5\mathrm{O}_2 \\ \rightleftharpoons 3\mathrm{NO}_2 \end{array}$	$\begin{split} \Delta H_{\text{NOox}} &= (E_{\text{O2f}} \cdot E_{\text{O2b}}) + (E_{\text{O3f}} \cdot E_{\text{O3b}}) \\ \Delta S_{\text{NOox}} &= \text{Rln}(A_{\text{O2f}} / A_{\text{O2b}}) + \text{Rln}(A_{\text{O3f}} / A_{\text{O3b}}) \end{split}$



Fig. 8. Experimental and basic modeling results in Cycle 2 of Roomdry sample (A) and 80dry sample (B) in Cycle 2 (last cycle without SO₂). Reaction conditions: 290 ppm NO_x, NO₂/NO_x = 1.8 %, 4.6 % H₂O, 6.9 % O₂ in Ar with the flow rate of 905 mL/min. 'Roomdry' and '80dry' denote the Pd/SSZ-13 samples which were dried at room temperature or 80 °C after Pd impregnation.

Table 5

Ratio between Z⁻Pd(II)Z⁻ and Z⁻[Pd(II)OH]⁺ in DRIFT and kinetic model.

Samples	DRIFT	Model
Roomdry	0.66	0.68
80dry	0.98	0.95

Table 6

Reactions in kinetic model for describing SO₂ poisoning of Z⁻Pd(II)Z⁻.

Reactions	Reaction Step	Rate Expressions
Z3	$\rm Z^-Pd(II)Z^- + SO2 \rightarrow Z^-Pd(II)Z^-SO_2$	$r_{Z3}=k_{Z3f}\Psi_Z\theta_Z y_{SO2}$

165 °C. The NO₂ generation peak at ~ 165 °C drastically decreased after adding SO₂, indicating that the reduction of Z-[Pd(II)OH]⁺ to Z-Pd(I) is greatly suppressed. Given the derivation of these profiles, more reaction on Z-[Pd(II)OH]⁺ should be added into the kinetic model.

3.2.3. Kinetic model for describing SO₂ enhanced NO_x adsorption on Z^{-} [Pd (II)OH]⁺

Evaluations depicted in Figs. 3 and 3, along with the kinetic model in Fig. 9, reveal that the NO desorption peak at 460 °C increases after adding SO₂. This clearly illustrates that SO₂ can enhance NO adsorption on Z⁻[Pd(II)OH]⁺/Z⁻Pd(I) sites. Moreover, Fig. 3 shows a gradual decrease in the NO release peak at 460 °C during the multicycle test with SO₂ (from cycle 3 to cycle 7), suggesting that SO₂ also slightly poisons the Z⁻[Pd(II)OH]⁺/Z⁻Pd(I) sites. However, after the termination of SO₂ (cycle 8), this peak maintains the same intensity as before SO₂ poisoning, indicating that the poisoned Z⁻[Pd(II)OH]⁺/Z⁻Pd(I) sites can be regenerated in the absence of SO₂. Furthermore, SO₂ is more likely to initially cover Z⁻[Pd(II)OH]⁺, thereby inhibiting the reduction of this site to Z⁻Pd(I). Based on these findings, we have introduced additional reactions (Table 7) showing that Z⁻[Pd(II)OH]⁺ can also be covered by SO₂ in a reversible reaction (reaction ZH5). The adsorbed SO₂ can further facilitate NO adsorption, as demonstrated in reactions



Fig. 9. Selected cycles of kinetic modeling results of 80dry Pd/SSSZ-13 sample with SO₂ poisoning reactions on Z⁻Pd(II)Z⁻ sites. Cycles: Cycle 3 (first cycle with SO₂), Cycle 7 (last cycle with SO₂) and Cycle 8 (first cycle shut SO₂). Reaction conditions: 290 ppm NOx, $NO_2/NO_x = 1.8$ %, 104 ppm SO₂ (only in Cycle 3–7), 4.6 % H₂O, 6.9 % O₂ in Ar with the flow rate of 905 mL/min. '80dry' denote the Pd/SSZ-13 sample which was dried at 80 °C after Pd impregnation.

Table 7

Reactions in kinetic model for describing SO_2 enhanced NO_x adsorption reactions on $Z^-[Pd(II)OH]^{+,}$

Reactions	Reaction Step	Rate Expressions
ZH5	$Z^{-}[Pd(II)OH]^{+}+SO2 \rightleftharpoons Z^{-}[Pd(II)OH]^{-}SO_{2}$	$r_{ZH5} = k_{ZH5f} \Psi_{ZH} \theta_{ZH1} y_{SO2}$ - $k_{ZH5b} \Psi_{ZH} \theta_{ZH1,SO2}$
ZH6	$Z^{-}[Pd(II)OH]^{-}SO_{2} + NO \rightleftharpoons$ $Z^{-}[Pd(II)OH]^{-}SO_{2} - NO$	$r_{ZH6} = k_{ZH6f} \Psi_{ZH} \theta_{ZH1-SO2} y_{NO} - k_{ZH6b} \Psi_{ZH} \theta_{ZH1-SO2-NO}$
ZH7	$Z^{-}[Pd(II)OH]^{-} NO + SO_{2} \rightleftharpoons$ $Z^{-}[Pd(II)OH]^{-} NO - SO2$	$\begin{split} r_{ZH7} &= k_{ZH7} / \Psi_{ZH} \theta_{ZH1-NO} y_{SO2} - \\ k_{ZH7b} \Psi_{ZH} \theta_{ZH1-SO2-NO} \end{split}$

ZH6 and ZH7.

The updated kinetic modeling results, accounting for the SO₂enhanced NO_x adsorption reactions on Z⁻[Pd(II)OH]⁺, are displayed in Fig. 10. With the reactions ZH5 to ZH7, the simulated NO and NO₂ profiles exhibit a better alignment with experimental results compared to the previous kinetic model. This is particularly evident at ~ 165 °C, where NO oxidation is significantly suppressed, leaving only small NO consumption and NO₂ generation peaks. However, there is still an issue remaining: at ~ 165 °C, the NO consumption peak area is actually larger than the NO₂ generation peaks. This suggests that another Pd site may be adsorbing NO at this temperature. It is important to note that NO_x



Fig. 10. Selected cycles of kinetic modeling results of 80dry with SO₂ enhanced NO_x adsorption reactions on Z⁻[Pd(II)OH]⁺. Cycles: Cycle 3 (first cycle with SO₂), Cycle 7 (last cycle with SO₂). Reaction conditions: 290 ppm NOx, NO₂/NO_x = 1.8 %, 104 ppm SO₂ (only in Cycle 3–7), 4.6 % H₂O, 6.9 % O₂ in Ar with the flow rate of 905 mL/min. '80dry' denote the Pd/SSZ-13 samples which was dried at 80 °C after Pd impregnation.

adsorption is a non-activated reaction [45,48,51], and therefore should occur at the beginning of the PNA process when NO_x is introduced. Consequently, we can deduce that a new Pd site is generated at \sim 165 °C, which adsorbs NO_x at a relatively high reaction rate. The stored NO on this newly generated Pd site is also released at 460 °C, subsequently increasing the peak intensity.

3.2.4. Kinetic model for describing the formation of PdSO₄

According to our kinetic model, which takes into account SO₂ poisoning of Z⁻Pd(II)Z⁻ and enhancement of NO_x storage on Z⁻[Pd(II) OH]⁺, it becomes evident that new Pd sites are generated during the temperature rise in the TPD branch. These sites subsequently adsorb more NO and release it at higher temperatures. Numerous reports indicate that PdO_x clusters can be transformed into PdSO₄ in the presence of gaseous SO₂, providing an additional adsorption site for NO_x storage [52-54]. Consequently, we have incorporated two more reactions on PdO_x sites to describe the formation of PdSO₄ (Reaction O3) and its adsorption of NO (Reaction O4), as detailed in Table 8. Note that Reaction O3 is reversible, given that PdSO₄ can decompose at higher temperatures [53]. The adjusted kinetic modeling results, alongside with the experimental results for the 80dry sample, are shown in Fig. 11 and Fig. S3. We have also validated the kinetic model based on the PNA performance for the Roomdry sample, as depicted in Fig. 12 and Fig. S6. Notably, while the initial Pd site amounts differ in the two PNA samples, all other kinetic parameters remain constant when simulating PNA performance across different samples. From the comparison results in Roomdry and 80dry, it is clear that the kinetic model can accurately describe PNA performance and SO₂ poisoning performance, regardless

Table 8

Reactions in	kinetic mod	el for o	describing	the f	formation	of P	dSO₄
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Reactions	Reaction Step	Rate Expressions
03 04	$\begin{array}{l} PdO + SO_2 + 0.5O_2 \rightleftharpoons PdSO4 \\ PdSO4 + NO \rightleftharpoons PdSO4 \text{-NO} \end{array}$	$\begin{split} r_{O3} &= k_{O3j} \Psi_O \theta_{O1} y_{SO2} y_{O2}^{0.5} \text{-} k_{O3b} \Psi_O \theta_{O3} \\ r_{O4} &= k_{O4j} \Psi_O \theta_{O3} y_{NO} \text{-} k_{O4b} \Psi_O \theta_{O3} \text{-} \text{NO} \end{split}$



Fig. 11. Multi-cycle experimental and modeling results of 80dry sample in Cycle 2 (last cycle without SO₂), Cycle 3 (first cycle with SO₂), Cycle 7 (last cycle with SO₂) and Cycle 8 (first cycle shut SO₂). Reaction conditions: 290 ppm NOx, $NO_2/NO_x = 1.8$ %, 104 ppm SO₂ (only in Cycle 3–7), 4.6 % H₂O, 6.9 % O₂ in Ar with the flow rate of 905 mL/min. Adsorption temperature: 80 °C. Full cycle results are shown in Fig. S3. '80dry' denote the Pd/SSZ-13 samples which was dried at 80 °C after Pd impregnation.



Fig. 12. Multi-cycle experimental and modeling results of Roomdry sample in Cycle 2 (last cycle without SO₂), Cycle 3 (first cycle with SO₂), Cycle 7 (last cycle with SO₂) and Cycle 8 (first cycle shut SO₂). Reaction conditions: 290 ppm NOx, $NO_2/NO_x = 1.8$ %, 104 ppm SO₂ (only in Cycle 3–7), 4.6 % H₂O, 6.9 % O₂ in Ar with the flow rate of 905 mL/min. Adsorption temperature: 80 °C. Full cycle results are shown in Fig. S6. 'Roomdry' denote the Pd/SSZ-13 samples which was dried at 80 °C after Pd impregnation.

of how the initial Pd sites change with different synthesis methods.

3.2.5. Verifying the kinetic model under different PNA operation conditions

Our developed model was further tested under a range of PNA operational conditions, applied to both Roomdry and 80dry samples, as depicted in Fig. S3 to S8. These conditions largely involved adjustments to the operational temperature in adsorption branch, ranging from 80 °C to 110 °C (Fig. S4) and up to 150 °C (Fig. S5), and reductions in the SO₂ concentration from 104 ppm to 52 ppm (Fig. S7) and down to 26 ppm (Fig. S8). The primary influence of the adsorption temperature variations can be seen in the NO oxidation peaks on PdO sites and the NO release peaks from Z⁻Pd(II)Z⁻. An increase in adsorption temperature results in that these peaks occuring earlier in time, while the SO₂ poisoning behavior stays relatively stable. Notably, the NO_x desorption peak from Z⁻[Pd(II)OH]/Z⁻Pd(I) at 460 °C does not change when the adsorption temperature is set to 150 °C. This finding supports our kinetic model, which describes that SO₂ mainly enhances NO_x adsorption by generating new PdSO₄ sites, rather than aiding NO_x adsorption on Z⁻[Pd (II)OH]/Z⁻Pd(I). The concentration of SO₂ predominantly influences the desorption of NO_x from PdSO₄ at 460 °C. With an increase in SO₂ concentration, more PdSO₄ forms, leading to an increase in NO_y adsorption and its subsequent release at 460 °C. The NO adsorption on Z⁻Pd(II)Z⁻ remains unaffected by changes in SO₂ concentration due to the relatively high poisoning rate of Z⁻Pd(II)Z⁻.

In order to delve deeper into the contribution of various reactions and kinetic parameters to the kinetic modeling of sulfur poisoning in PNA, we calculated the normalized sensitivity coefficients of the kinetic parameters, based on model validation. It is important to note that normalized sensitivity is a dimensionless constant. A higher value of normalized sensitivity suggests that the kinetic model is more susceptible to the corresponding kinetic parameter, which in turn is more accurate in predicting experimental behavior. To calculate the normalized sensitivity, we incremented only one kinetic parameter by 1 % at each time in the simulation to obtain the deviated simulation result. This process incorporated 666,000 data points from the NO profile and another 666,000 data points from the NO₂ profile under various operation conditions, as demonstrated from Fig. S3 to Fig. S8. The normalized sensitivity coefficients were then determined using the following equation:

$$S = \beta_0 \frac{\sum \left(\frac{\Delta y^{im}}{y_0^{im}}\right)^2}{\Delta \beta} \tag{4}$$

where β_0 represents the optimized parameter, $\Delta\beta$ is the deviated value of the changed parameter from the optimized parameter. y_0^{sim} is the NO/NO₂ concentration at one time point from the optimized value. Δy^{sim} is the deviation in NO/NO₂ concentration between the optimized result and the result with the changed parameter.

We arranged the kinetic parameters in descending order, from the highest to the lowest normalized sensitivity, as depicted in Fig. 13. The parameters with the greatest influence on the NO profiles are Af-O4, Af/ Ab-ZH3, and Af/Ab-ZH5. Reaction O4, which describes the NO adsorption and desorption on PdSO₄ (Table 8), influences both the NO consumption peak at approximately 165 °C and the NO desorption peak at 460 °C following the addition of SO₂. Due to its effect on these two peaks in the NO profiles, Af-O4 emerges as the most sensitive parameter for the kinetic model. Reactions ZH3 (Table 3) and ZH5 (Table 7) predominantly describe the NO adsorption on Z-Pd(I), which is reduced from Z⁻[Pd(II)OH]⁺, and SO₂ adsorption on Z⁻[Pd(II)OH]⁺. These reactions are sensitive to the kinetic model as they contribute to the dominant NO desorption peak at 460 °C, with or without SO₂. Although Af-Z3, Af-Z2, Af-O1, and Af-O2 display the least sensitivity towards the kinetic modeling results, they remain crucial to the overall reaction model. Af-Z3 describes the SO₂ poisoning on Z⁻Pd(II)Z⁻ sites (Table 6). The rate of this reaction needs to be high enough to ensure that the Z⁻Pd $(II)Z^{-}$ are poisoned in the first cycle with SO₂, thereby reducing the sensitivity of Af-Z3. Af-Z2 contributes to the NO2 profile, so its sensitivity on the NO profile can be neglected. The relatively lesser sensitivity of O₁ and O2 is due to the slow formation of Pd(NO₃)₂ compared to other reactions because the Pd(NO₃)₂ only partially formed in the first two cycles without SO₂ adding. However, these reactions must be retained because we have observed the formation of Pd(NO₃)₂ in our previous work [17].

The sensitivity of parameters on NO₂ profiles is lower than on NO profiles due to the significantly lower NO₂ concentration in the PNA process. The parameters that most significantly influence the NO₂ profiles are Af-O3 and Af-Z1. Reaction O3 strongly suppresses the formation of Pd(NO₃)₂ at ~ 165 °C, leading to its subsequent decomposition and NO₂ release. Given the competitive adsorption of NO and NO₂ on Z-Pd (II)Z-, the adsorption rate of NO significantly influences the adsorption of NO₂, which explains why reaction Z1 also greatly influences the NO₂ profile. The less sensitive parameters based on the NO₂ profiles are more sensitive towards simulated NO profiles, indicating that those corresponding reactions are also critical to this kinetic model.



Fig. 13. Normalized sensitivity co-efficient of kinetic parameters based on NO profiles (A) and NO₂ profiles (B).

4. Conclusions

In this study, we explored the SO₂ poisoning mechanism through a combined experimental and kinetic modeling approach, and also developed an innovative PNA sample exhibiting strong resistance to SO₂ poisoning. Pd/SSZ-13 samples were synthesized using the impregnation method but under varying drying conditions—room temperature drying and 80 °C drying—to create different surface distributions of Pd active sites. Our multicycle PNA tests with SO₂ poisoning revealed that the Roomdry sample experienced a 10 % degradation, while the 80dry sample experienced a considerably greater degradation of 26 %.

Upon conducting in-situ DRIFT of CO adsorption, XRD, and TEM for the degreened and poisoned samples, we discovered that the ionexchanged Pd sites with a hydroxyl group $(Z^{-}[Pd(II)OH]^{+})$ demonstrated strong resistance to SO₂ poisoning, whereas $Z^{-}Pd(II)Z^{-}$ was notably vulnerable to SO₂ poisoning. By reducing the drying temperature after the Pd/SSZ-13 loading process, we were able to generate more ion-exchanged Pd sites with a hydroxyl group, which explains the high SO₂ tolerance of the Roomdry sample. Moreover, we found that SO₂ also aids in NO_x storage on both $Z^{-}[Pd(II)OH]^{+}$ and PdO_x sites, resulting in improved PNA performance.

Based on these experimental findings, we developed a kinetic model that can describe both the SO₂ poisoning behavior of PNA and the role of SO₂ in assisting NO_x storage. The kinetic model incorporates three initial Pd sites, Z⁻Pd(II)Z⁻, Z⁻[Pd(II)OH]⁺, and PdO, with 14 reactions. The model accounts for the SO₂ poisoning of Z⁻Pd(II)Z⁻ and Z⁻[Pd(II)OH]⁺, as well as the enhancement of NO_x storage on Z⁻[Pd(II)OH]⁺ and PdO due to SO₂. After being verified under different reaction conditions, this model has proven its robustness in simulating the PNA behavior and the influence of SO₂ in multicycle PNA tests, including varying SO₂ concentrations (26 ~ 104 ppm), different operation temperatures in adsorption branch (80 ~ 150 °C), and even different reaction parameters by varying the kinetic parameters and comparing them with the optimized results. This analysis demonstrated that all the reactions in this kinetic model are crucial for accurately describing PNA behavior.

CRediT authorship contribution statement

Dawei Yao: Writing – original draft, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. Phuoc Hoang Ho: Writing – review & editing, Formal analysis, Data curation. Johann C. Wurzenberger: . Thomas Glatz: Writing – review & editing, Supervision, Software. Wei Di: Data curation, Formal analysis. Rojin Feizie Ilmasani: . Derek Creaser: Writing – review & editing, Supervision, Methodology, Conceptualization. Louise Olsson: Writing – review & editing, Supervision, Methodology, Funding acquisition, Conceptualization.

Declaration of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Johann C. Wurzenberger and Thomas Glatz reports a relationship with AVL List GmbH that includes: employment. Louise Olsson is one of the Editors for Chemical Engineering Journal.

Data availability

Data will be made available on request.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.cej.2024.150406.

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