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Operando sulfur speciation during sulfur poisoning-regeneration of Ru/SiO₂ and Ru/Al₂O₃ using non-resonant sulfur $K\alpha_{1,2}$ emission†

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A periodic oxidative regeneration of a sulfur-poisoned methanation catalyst is an alternative to the expensive state-of-the-art process of syngas cleaning using wet scrubbers. Here we have employed operando X-ray emission spectroscopy (XES) to study sulfur speciation on Ru/SiO₂ and Ru/Al₂O₃ during methanation in the presence of H_2S and subsequent regeneration in dilute O_2 at 360 °C. XES allowed us to obtain semi-quantitative sulfur speciation and to monitor changes in the absolute sulfur concentration. It was established that Al_2O_3 , in contrast to SiO₂, forms sulfite/sulfate species by reacting with SO_2 , which is released from the poisoned Ru surface upon oxidative treatment. The concentration of sulfite/sulfate species is reduced upon switching the feed to H_2/CO while no simultaneous increase in sulfide concentration is observed. For both catalysts, the regenerative treatment removes adsorbed sulfur as SO_2 only partially, which we propose is the main reason for the incomplete activity recovery of the poisoned catalyst after regeneration.

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Introduction

The production of methane from dry biomass consists of four main process steps: biomass gasification, syngas cleaning to remove catalyst poisons such as sulfur-containing compounds (H_2S , COS, and thiophene), methanation ($CO + 3H_2 \rightleftharpoons CH_4 + H_2O$) and removal of H_2O and CO_2 . The second step, syngas cleaning, is currently energy inefficient since it requires cooling of syngas to ambient temperature between the gasification and methanation steps, which are performed at around 850 °C and 400 °C respectively. We have been exploring an alternative process, where the low temperature syngas cleaning step is skipped and the methanation catalyst is periodically regenerated after sulfur poisoning which would make the whole dry biomass to methane process more energy efficient.

We have reported recently that the ability to regenerate sulfur poisoned ruthenium nanoparticles, as measured by its methanation activity, is improved when ruthenium nanoparticles are supported on a sulfur-inert support, SiO_2 compared to Al_2O_3 .⁴ This was attributed to a higher Ru dispersion and lower sulfur storage

of SiO₂ upon multiple poisoning-regeneration cycles. With the

In that context, X-ray spectroscopy would be a possible solution since it allows for detecting all sulfur species. Our group has recently reported a study on the mechanism of sulfur poisoning on Ru/Al₂O₃ by using *in situ* sulfur K-edge X-ray absorption (XAS),⁵ where we proposed that sulfate species, formed during oxidative regeneration, under subsequent reducing conditions convert to H₂S, which re-poisons Ru nanoparticles. However, S K-edge XAS only provided qualitative information. On the other hand, semi-quantitatively sulfur speciation under *operando* conditions should be accessible by sulfur X-ray emission spectroscopy (XES). Indeed, while XAS probes the unoccupied states and has many overlapping features, reflecting not only the sulfur oxidation state but also

help of *in situ* diffuse reflectance infrared spectroscopy, we were able to monitor the speciation of some of the sulfur species, where the formation of sulfate and sulfite species was observed when the poisoned sulfided catalyst was exposed to dilute oxygen. During subsequent exposure to methanation conditions (H₂/CO) the intensity of the sulfate and sulfite peak decreased, but we could not determine if this was related to a reduction of sulfate and sulfite species to gas phase SO₂ or to the sulfate reduction and subsequent adsorption of sulfides since diffuse reflectance spectroscopy did not allow detecting sulfides. Another open question is how efficient the regeneration process is, *i.e.* what percentage of initially adsorbed sulfur is removed from the catalyst surface *via* oxidative treatment. To answer these questions, it would be valuable to find ways to quantitatively trace sulfur speciation under *operando* conditions.

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the local coordination of the sulfur atom,6 XES gives information on the occupied states and primarily reflects the nuclear charge. The sulfur $K\alpha_{1,2}$ emission signal appears in the XES spectrum as a doublet, which can be readily fitted with a convolution of two pseudo-Voigt function peaks. For different sulfur oxidation states, only a systematic shift in peak energy position and a slight change in the ratio of $K\alpha_1$ to $K\alpha_2$ intensity have been observed.7 This allows obtaining quantitative sulfur speciation from XES using, for example, a linear combination fitting approach, quick and straightforward. In addition, since the X-ray emission signal intensity is directly proportional to the number of sulfur atoms in the beam, the changes in signal intensity can be used to estimate the changes in sulfur concentration. We have recently reported the development of a dedicated non-resonant XES setup including operando cell that allows to study sulfur speciation at low concentrations and in a time-resolved manner.8 Here, by employing non-resonant sulfur X-ray emission spectroscopy, we investigate the poisoning and regeneration mechanisms of sulfur-poisoned Ru nanoparticles supported on SiO₂ and Al₂O₃. The key difference between the two catalysts arises from the ability of Al₂O₃ to form sulfates by reacting with SO₂, which is released from the Ru surface upon oxidative treatment. By using operando nonresonant S XES, we show that for both catalysts, oxidative treatment at 360 °C only partially removes adsorbed sulfur as SO_{2.} Sulfur remains adsorbed on the metal surface as sulfite/ sulfate species. When the oxygen supply is removed, these oxidized species are reduced back to sulfide, which is the reason for the incomplete activity recovery.

2. Experimental

Non-resonant sulfur XES measurements were carried out using a dedicated S XES von Hamos spectrometer including an operando spectroscopic cell⁸ installed at the SuperXAS beamline of the Swiss Light Source (SLS), Villigen, Switzerland. The SLS operates at 400 mA and 2.4 GeV. The polychromatic beam from a 2.9 Tesla superbend magnet was collimated by a Si-coated mirror at 2.5 mrad and subsequently monochromatized by a Si (111) channel-cut crystal. Downstream of the monochromator, a Rh-coated toroidal mirror was used to focus the incident X-rays to a spot size of 100 \times 100 μm at the sample position. The incoming X-ray energy was calibrated by using an Fe foil in transmission mode at the start of the experiment. The dedicated XES spectrometer9 was enclosed in a stainless steel chamber evacuated by a turbo pump to 10^{-3} to 10^{-4} mbar. In this vacuum chamber a 15 cm bending radius Si (111) von Hamos crystal was placed on a motorized stage allowing to fine tune the crystal position in focusing direction. A charge-coupled device (CCD) detector (Andor DO440, 2048 × 512 array with 13.5 \times 13.5 μ m pixels), which was thermoelectrically cooled to -30 °C, was used as a position sensitive detector. The chip length, consisting of 2048 pixels, allows measuring the X-ray emission energy range from 2.255 to 2.348 keV in a single acquisition. An aluminized 6 µm thick Mylar window was placed in front of the CCD chip to prevent visible light from reaching the chip. According to Bragg's law, the center of the Si

(111) von Hamos crystal diffracts not only the $K\alpha$ emission but also triple the energy of the sulfur $K\alpha$ emission, *i.e.* 6.921 keV. Hence, the incident energy for non-resonant XES was set to be around 6.9 keV since this allowed us to observe both sulfur $K\alpha$ emission and elastic scattering of the incident 6.9 keV beam in the same spectrum. The elastic scattering peak position was subsequently used for CCD pixel to energy calibration.

For performing *operando* experiments, the *operando* cell⁹ was filled with *ca.* 15 mg of 3% Ru/SiO₂ and Ru/Al₂O₃, prepared according to ref. 4. The *operando* cell is operated at atmospheric pressure and is equipped with a thermocouple close to the catalyst bed. A 7 μ m thin Kapton film was used as window material. The cell was heated to 360 °C (10 °C min⁻¹ ramp) while He was flowing through the cell. Subsequently, the catalyst was treated with the following gas sequence: methanation in the presence of inorganic sulfur (0.5 vol% CO, 2.5 vol% H₂, 100 ppm H₂S), He flush, oxidative regeneration (1 vol% O₂), He flush and methanation without sulfur poisons (0.5 vol% CO, 2.5 vol% H₂). The gases were dosed with mass flow controllers (Bronkhorst) and the gas phase after the reactor cell was analyzed with a quadrupole mass spectrometer (Max 300-LG, Extrel).

Reference compounds (Ag₂S, FeS₂, S, Na₂SO₃ and Na₂SO₄) were purchased from Sigma Aldrich and used as received. The powders were crushed using a pestle and mortar and subsequently pressed in a 0.5 cm diameter pellet without addition of any binder. Emission spectra were recorded for 60 seconds, apart for the elemental S sample, where an acquisition time of 5 seconds was sufficient to obtain a high signal-to-noise ratio.

The raw spectra were baseline corrected and normalised by the total area under the peak using an in-house written Python script. The normalised spectra of the reference compounds were used for fitting *operando* spectra. The following procedure for fitting the reference non-resonant XES spectra was followed: pseudo-Voigt peak functions were fitted to $K\alpha_1$ and $K\alpha_2$ emission peaks of the reference compounds and the extracted values are peak center, full width at half maximum (FWHM). The peak profile of the reference compounds was used for fitting the unknown *operando* spectra. The data treatment was performed using Origin software.

Results and discussion

3.1. Reference sulfur compounds

The spectra of various sulfur reference compounds Ag_2S (S^{2-}), FeS_2 (S^{1-}), S (S^0), Na_2SO_3 (S^{4+}) and Na_2SO_4 (S^{6+}) were previously measured by our group⁸ and the published spectra are shown in Fig. 1 to help the readers with *in situ* spectra interpretation. In these spectra, the $K\alpha_1$ and $K\alpha_2$ lines can be clearly resolved, and an energy shift for various sulfur oxidation states is also observed, similar to previously reported values.^{7,11,12} Table S1 in ESI† gives the parameters (peak center, FWHM *etc.*) extracted from fitting the reference and used for fitting of the *operando* spectra described below.

3.2. Ru/SiO₂

To investigate the relationship between the catalyst methanation activity and its structure during sulfur poisoning and Paper

2.5 - Na₂SO₄ (S⁶⁺)

Na₂SO₃ (S⁴⁺)

1.5 - S⁰

FeS₂ (S¹)

0.5 - Ag₂S (S²⁻)

2304 2306 2308 2310 2312

Energy, eV

Fig. 1 Sulfur $K\alpha_{1,2}$ emission spectra (dotted black line) and the corresponding pseudo-Voigt peak function fit (colored full line) of reference compounds: Ag₂S, FeS₂, S, Na₂SO₃, and Na₂SO₄.⁹

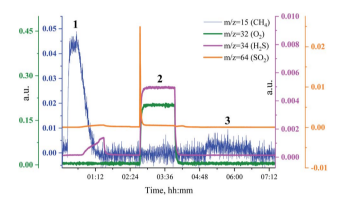


Fig. 2 MS traces for CH₄ (m/z=15, blue), O₂ (m/z=32, green and m/z=34, purple), H₂S (m/z=34, purple) and SO₂ (m/z=64, orange) during poisoning-regeneration of Ru/SiO₂ at 360 °C. H₂S breakthrough is observed.

regeneration, we have carried out H2S poisoning, oxidative regeneration and subsequent sulfur-free methanation of Ru/ SiO₂ while simultaneously measuring non-resonant X-ray emission spectra. Fig. 2 shows the mass spectrometer (MS) traces for CH₄ (m/z = 15), O₂ (m/z = 32 and m/z = 34), H₂S (m/z = 32)34) and SO_2 (m/z = 64) recorded during poisoning ($H_2/CO/H_2S$ labelled 1), regeneration (O₂ labelled 2) and subsequent sulfurfree methanation (H2/CO labelled 3) of Ru/SiO2. Note that the increase in m/z = 34, that accompanies the increase in m/z = 32during oxidation, is solely due to oxygen, because the O atom has two important isotopes: ¹⁶O and ¹⁸O and hence the O₂ molecule produces m/z = 32 and m/z = 34 fragments. The initial level of methane production, when Ru/SiO₂ is treated with H₂/ CO/H₂S, decreases gradually because of H₂S adsorption on the active sites of Ru nanoparticles. The breakthrough of H2S is observed as the catalyst loses its methanation activity (m/z = 34in Fig. 2). After methanation activity dropped to zero, the catalyst is treated with dilute oxygen to remove adsorbed sulfur atoms as SO_2 , which was detected as a spike in the MS data (m/z= 64 in Fig. 2). In the subsequent sulfur-free H_2/CO treatment, we see a small recovery of methanation activity (m/z = 15 in

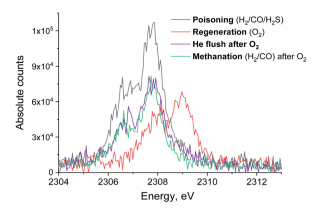


Fig. 3 S $K\alpha$ XES spectra of Ru/SiO₂ in H₂/CO/H₂S (black), O₂ (red) and subsequent pure He (purple) and H₂/CO atmospheres (green). Integration time for each spectrum is 13 minutes.

Fig. 2). The recovered methanation activity is smaller than that reported in our previous studies probably due to the lower regeneration temperature (360 $^{\circ}$ C vs. 430 $^{\circ}$ C) used in this operando experiment.

Fig. 3 shows the non-resonant XES spectra for all stages of the sulfur poisoning-regeneration cycle for methanation over a Ru/SiO2 catalyst. First, we observe a peak corresponding to reduced sulfur species (Fig. 3, black), which likely results from dissociative adsorption of H₂S on the ruthenium surface since SiO₂ is not known to form sulfides. In situ DRIFTS is not able to detect this reduced species4 likely because of the low IR intensity of this species and/or their vibration frequency being beyond the detectable range.13 Upon exposure to dilute oxygen, we see that the emission peak shifts to higher energy and its intensity decreases by ca. a factor of two (Fig. 3, red). The intensity decrease suggests that around half of the sulfidic species leave the metal surface as gas phase SO2, consistent with the observation of a spike of SO₂ in the MS trace signal (Fig. 2, orange). The remaining oxidized species could be located on the support, SiO2, or on the Ru nanoparticles. We have previously shown4 using in situ DRIFTS that sulfur species were only observed upon treatment of sulfur poisoned Ru/SiO2 as a small peak at 1425 cm⁻¹ (which was assigned to sulfate species on SiO₂ (ref. 14–17)) when the O₂ concentration is increased from 1 to 5% (Fig. S1†). Importantly, due to absorption of IR radiation below 1250 cm⁻¹ by bulk Si-O bonds, we were only able to examine the region above 1250 cm⁻¹ for in situ DRIFTS experiment. This implies that all species appearing below 1250 cm⁻¹ will be invisible. This could have potentially masked sulfate species, as they could appear below 1250 cm⁻¹, 18 formed when sulfur poisoned Ru/SiO₂ was treated with 1% O₂ in Ar.

When the flow of oxygen is switched to pure He, the nonresonant XES spectra show that S^{4+}/S^{6+} species are fully converted to S^{2-} (Fig. 3, purple), indicating that S^{4+}/S^{6+} species are not stable in the absence of oxygen. No significant changes to the XES spectrum are observed when CO and H_2 are introduced in the feed (Fig. 3, green). We propose that the sulfidic species present after the oxidative regeneration (and flushing with inert gas) are blocking the active sites for methanation and cause the

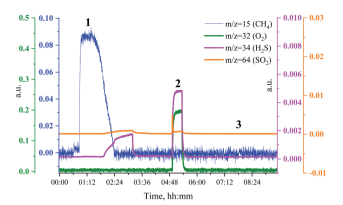


Fig. 4 MS trace for CH₄ (m/z=15, blue), O₂ (m/z=32, green and m/z=34, purple), H₂S (m/z=34, purple) and SO₂ (m/z=64, orange) during poisoning-regeneration of Ru/Al₂O₃ at 360 °C. H₂S breakthrough is observed at 1 h 15 min.

loss of Ru/SiO_2 methanation activity. We also observe that when the gas feed is switched to O_2 , the intensity of the emission peak decreases by half, indicating that only half of the sulfur species at the position where the X-ray beam impinges the sample desorb as SO_2 . This means that at the temperature of this experiment, 360 °C, only half of the poisoning species are removed from the surface. Increasing the temperature of the oxidative treatment could improve the regeneration efficiency because more sulfur atoms could desorb as SO_2 .

3.3. Ru/Al_2O_3

To see if the methanation activity of the Ru/Al₂O₃ catalyst recovers after sulfur poisoning, we have carried out H2S poisoning, oxidative regeneration and subsequent sulfur-free methanation of Ru/Al₂O₃. In addition, non-resonant X-ray emission spectra were acquired simultaneously at each stage. Fig. 4 shows the MS traces for CH_4 (m/z = 15), O_2 (m/z = 32, m/z= 34), H_2S (m/z = 34) and SO_2 (m/z = 64) recorded during poisoning (H₂/CO/H₂S labelled 1), regeneration (O₂ labelled 2) and subsequent sulfur-free methanation (H2/CO labelled 3) over a Ru/Al₂O₃ catalyst. When treated with H₂/CO/H₂S, Ru/Al₂O₃ also experienced methanation activity loss. However, deactivation of Ru/Al₂O₃ (and for H₂S to break through) is ca. twice slower than observed for Ru/SiO₂ (cf. Fig. 2 and 4). As we have previously proposed,4 this could result from two causes. First, since Ru/Al₂O₃ has a smaller initial particle size than Ru/SiO₂, there are more active sites and hence the deactivation is slower. Secondly, H₂S adsorbs on to the Al₂O₃ surface, which also slows down deactivation of Ru nanoparticles. After regeneration however, no methanation activity is observed probably because of the long H₂S treatment resulting in a high sulfur coverage that made oxygen adsorption on to the poisoned surface difficult or due to a low regeneration temperature. In addition, the SO₂ signal during oxidative regeneration for Ru/Al₂O₃ is much smaller than for Ru/SiO₂ (cf. Fig. 2 and 4, the scale for the SO₂ signal is kept the same in Fig. 2 and 4 to simplify comparison), suggesting that the Al₂O₃ support reacts with SO₂ released from Ru nanoparticles upon oxidative regeneration.

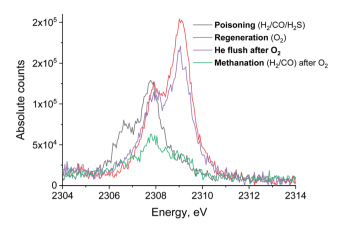


Fig. 5 S XES spectrum of Ru/Al_2O_3 in $H_2/CO/H_2S$ (black), O_2 (red) and subsequent pure He (purple) and H_2/CO atmosphere (green). Integration time for each spectrum is 13 minutes.

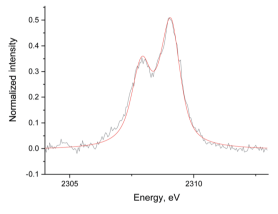


Fig. 6 The data (black) and fit (red) of the Ru/Al $_2$ O $_3$ S XES spectrum in He without S 2 reference added. The data was background corrected and normalized by integral intensity.

Fig. 5 shows the non-resonant XES spectra for all stages of the sulfur poisoning-regeneration cycle for Ru/Al₂O₃. As for Ru/ SiO₂, after H₂/CO/H₂S treatment we observe a spectrum, which corresponds to sulfidic species (Fig. 5, black). When the poisoned catalyst is contacted with oxygen, SO2 is removed from the surface of the metal and reacts with Al₂O₃, forming S⁴⁺ and S⁶⁺ species. When looking at raw counts (Fig. 5, cf. black and red), we observe that the signal intensity of these oxidized species is higher than the intensity of the sulfide. The observed signal intensity change can be explained by the fact that the Xrays are probing the middle of the reactor: SO₂ desorbed from the metal surface at the inlet of the reactor reacts with Al₂O₃ in the middle (also probably the outlet of the reactor) increasing the overall signal intensity. When the flow is switched from 1% O_2 to pure He (Fig. 5, *cf.* red and purple), we observe the appearance of a small shoulder at lower energies. Fig. 6 shows the fit of this spectrum with a linear combination of S⁴⁺ and S⁶⁺ references. When adding 10% of the S2- reference to the linear combination of S4+ and S6+ references, the fit also accounts for the low energy shoulder-like feature as is shown in Fig. 7. This

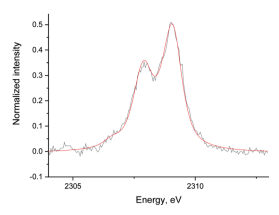


Fig. 7 The data (black) and fit (red) of the Ru/Al_2O_3 S XES spectrum in He with S^{2-} reference added. The data was background corrected and normalized by integral intensity.

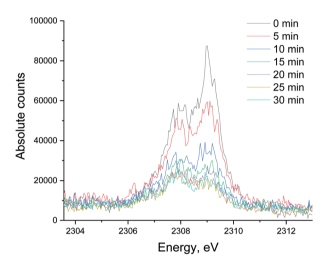


Fig. 8 The evolution of the sulfur XES signal during H_2/CO treatment after oxidative regeneration of Ru/Al_2O_3 . Integration time for each spectrum is 5 minutes.

can be explained by the presence of two different oxidized species - one on the surface of the metal and another one on the support. The former as in the case of Ru/SiO2 (Fig. 3) are converted to S²⁻ when the oxygen supply is switched off, while the latter, presumably, in the form of $Al_2(SO_4)_3$, are stable even without oxygen in the feed. After 30 minutes of subsequent H₂/ CO treatment (Fig. 5, green) a decrease of the sulfate signal intensity is observed. Fig. 8 shows the sulphur XES signal temporal evolution during H₂/CO treatment and Fig. 9 shows the peak area of each species every 5 minutes derived from fitting (Fig. S2-S7 in ESI†) of the spectra using S²⁻, S⁴⁺, S⁶⁺ references. The fitting parameters of the references and the percentage of each reference component in the fit of the in situ spectra are given in Tables S1 and S2† respectively. From Fig. 9, we can conclude that the absolute concentration of S⁴⁺ and S⁶⁺ is decreasing while the concentration of S²⁻ stays constant during H₂/CO treatment.

In a separate experiment, Ru/Al_2O_3 was exposed to $H_2/CO/H_2S$ at the same temperature and gas flow for a shorter period,

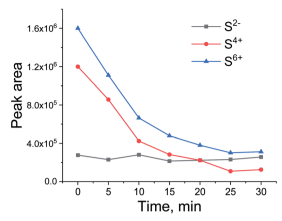


Fig. 9 Peak area of S²⁻, S⁴⁺, S⁶⁺ during H_2/CO treatment after oxidative regeneration of Ru/Al_2O_3 .

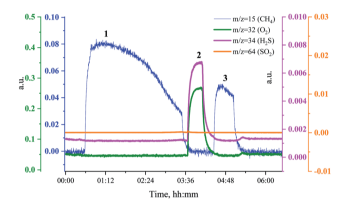


Fig. 10 MS traces for CH₄ (m/z=15, blue), O₂ (m/z=32, green and m/z=34, purple), H₂S (m/z=34, purple) and SO₂ (m/z=64, orange) during poisoning-regeneration of Ru/Al₂O₃ at 360 °C. No H₂S breakthrough is observed.

which resulted in an incomplete deactivation and methanation activity being observed in the MS traces after oxidative treatment. Fig. 10 shows the MS traces for CH₄ (m/z=15), O₂ (m/z=32 and m/z=34), H₂S (m/z=34) and SO₂ (m/z=64) recorded during poisoning (H₂/CO/H₂S labelled 1), regeneration (O₂ labelled 2) and sulfur-free methanation (H₂/CO labelled 3). The methanation activity is not fully lost due to poisoning and, importantly, no H₂S breakthrough is seen (Fig. 10, pink). Further, no or only a very small SO₂ peak is visible upon O₂ addition.

The evolution of XES spectra during H_2/CO treatment is shown in Fig. 11 and the peak area for each species is shown in Fig. 12 (fitting in S8–S10 in ESI†). Overall, we observe a decrease in the total area of the peak as H_2/CO treatment proceeds, the intensity of the oxidized species (S^{4+} only could be fitted) peak decreases and there is a small increase in the peak area of S^{2-} . This suggests that there is possibly a small chance of a partial sulfite reduction to sulfide for a partially poisoned catalyst. Most of the oxidized species however are reduced to gas phase SO_2 rather than to adsorbed sulfidic species.

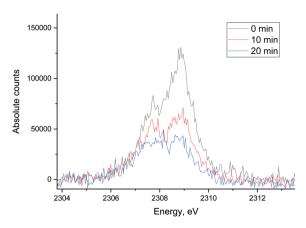


Fig. 11 The evolution of sulfur XES signal during H_2/CO treatment after oxidative regeneration of Ru/Al_2O_3 . Integration time for each spectrum is 10 minutes.

3.4. Proposed mechanism

By combining all results presented here, we propose the following mechanisms of sulfur transport on Ru/SiO2 and Ru/ Al_2O_3 (Fig. 13). On Ru/SiO₂, around half of the S^{2-}/S^{1-} species, formed on the surface of the Ru metal during poisoning via dissociative adsorption of H2S, are oxidized upon oxygen introduction at 360 °C and released as gas phase SO₂ while the other half remains bound to the metal surface as a mixture of S⁴⁺/S⁶⁺ species. As oxygen is removed from the feed, S⁴⁺/S⁶⁺ species on the Ru nanoparticle surface are reduced back to S²⁻/ S¹⁻ and subsequent H₂/CO treatment does not change the speciation significantly. The main difference for Ru/Al₂O₃ is that oxidation also produces S4+/S6+ species bound to the support, Al₂O₃, which are stable when oxygen is removed from the feed and are partially removed (likely reduced to gas phase species) when H₂/CO is introduced. In addition, for Ru/Al₂O₃, the mechanism also depends on the sulfur surface coverage established during H2S poisoning. For high coverage, no increase in S²⁻ species concentration accompanies S⁴⁺/S⁶⁺ partial removal, while for low coverage we see a small increase in S²⁻ species as the number of S⁴⁺/S⁶⁺ species decreases. This can be explained by the difference in the availability of

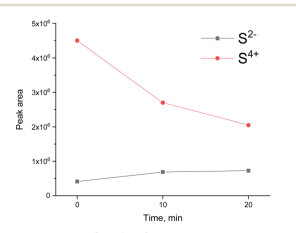


Fig. 12 Peak area of S $^{2-}$, S $^{4+}$, S $^{6+}$ during H $_2$ /CO treatment after oxidative regeneration of Ru/Al $_2$ O $_3$.

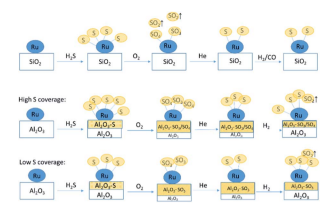


Fig. 13 Proposed mechanism of sulfur poisoning and regeneration of Ru/SiO₂ (top) and Ru/Al₂O₃ (middle and bottom).

adsorption sites on Ru: a heavily poisoned catalyst has less sites which are available for sulfide re-adsorption.

4. Conclusions

In order to understand the influence of periodic oxidative regeneration of sulfur-poisoned methanation catalysts as alternative to cost intensive low temperature scrubbers, mechanistic studies are indispensable. Based on the previous study of sulfur poisoningregeneration of a Ru/Al₂O₃ catalyst using operando sulfur K-edge XAS, the reduction of sulfate species, formed during oxidative regeneration, under subsequent reducing conditions to sulfides was proposed. However, since no exact reference compound spectra were available, only qualitative information was obtained. We have shown here that operando non-resonant X-ray emission spectroscopy is an alternative powerful technique for studying sulfur speciation. In contrast to XAS, XES allowed us to monitor the changes in the absolute concentration of various sulfur species during poisoning-regeneration and determine the proportion of each oxidation state species by a straightforward linear combination fitting of the references.

We have established that oxygen treatment of both poisoned catalysts removes adsorbed sulfur as SO_2 , but only partially. The remaining part stays adsorbed on to Ru nanoparticles as S^{4+}/S^{6+} species. These species are only stable under oxygen flow and reduce back to sulfide when no oxygen is in the feed. The loss of methanation activity is due to an incomplete removal of sulfur species from the Ru metal surface during oxidative regeneration and an increase of the oxidative regeneration temperature could help in removing more sulfur species from the surface to the gas phase.

Conflicts of interest

There are no conflicts to declare.

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