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Structural dynamics in Ni-Fe catalysts during CO₂ methanation - role of iron oxide clusters†

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Bimetallic Ni-Fe catalysts show great potential for CO₂ methanation concerning activity, selectivity and long-term stability even under transient reaction conditions as required for Power-to-X applications. Various contrary suggestions on the role of iron in this system on CO₂ activation have been proposed, hence, its actual task remained still unclear. In this study, we used X-ray absorption spectroscopy (XAS) combined with X-ray diffraction (XRD), XAS in combination with modulation excitation spectroscopy (MES) and density functional theory (DFT) to shed detailed light on the role of iron in a bimetallic Ni-Fe based CO₂ methanation catalyst. During catalyst activation we observed a synergistic effect between nickel and iron that led to higher fractions of reduced nickel compared to a monometallic Ni-based catalyst. By XAS-XRD combined with DFT, we found formation of FeO_x clusters on top of the metal particles. Modulation excitation coupled XAS data complemented with DFT calculations provided evidence of a $Fe^0 \Rightarrow Fe^{2+} \Rightarrow$ Fe³⁺ redox mechanism at the interface of these FeO_x clusters. This may promote CO₂ dissociation. This is the first time that this highly dynamic role of iron has been experimentally confirmed in bimetallic Ni-Fe based catalysts with respect to CO2 activation during the methanation reaction and may also be at the origin of better performance of other CO₂-hydrogenation catalysts. The insight into the structural surface changes reported in this study show the dynamics of the Fe-Ni system and allow the development of realistic surface models as basis for CO₂ activation and possible intermediates in these bimetallic systems.

Introduction

In future, the amount of renewable energy generated from wind and solar power will grow significantly. In order to maintain grid stability on the one hand and not to waste excess electricity on the other, there is the demand for efficient energy storage technologies.² The "Power-to-X" concept describes an ideal case for short and long-term storage by sector coupling of electricity, heat, gas, chemistry and transport.^{3,4} Ideally, the (chemical energy) carriers suggested in this approach can be produced using CO2 as feedstock in order to close the carbon cycle. One part of this concept is "power-to-gas" in which methane is catalytically synthesized from CO2 and renewable hydrogen via the Sabatier reaction (eqn (1)):^{5,6}

$$CO_2 + 4H_2 \Rightarrow CH_4 + 2H_2O \quad \Delta H_{298K} = -165 \text{ kJ mol}^{-1}$$
 (1)

Methane can be stored and transported cost efficiently to the end consumer via the already existing gas grid without the need of a new infrastructure. Furthermore, CH₄ provides a three times higher volumetric energy density than H2 and can, in contrast to H2, be directly used in conventional gas power plants for reconversion into electrical power.⁷ However, the weather related generation of power and the thereto related generation of renewable H₂ might lead to challenging operation conditions of the respective applied catalyst systems. This is critical, as catalysts are highly dynamic systems.8 For the industrial methanation of CO2 cheap and yet efficient Ni-based catalysts are widely used. 9-15 Mutz et al. 15-17 reported e.g. that γ-Al₂O₃ supported nickelbased catalysts tend to strongly deactivate during methanation of CO2 in the event of fluctuations.

Assuming that CO dissociation is the rate-determining step in CO2 methanation on metallic surfaces, Nørskov, Christensen and co-workers 18,19 predicted and verified various alloy surfaces which might provide an enhanced CO and thus also CO2 methanation. Among them, Ni-Fe alloy catalysts appeared very attractive. In fact, recent studies demonstrate

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that Ni-Fe catalysts show both a high CO2 methanation activity and an improved long-term stability, even under dynamic conditions. 20-25 Surprisingly, recent studies reported that nickel remains mainly reduced while iron seems to be oxidized during CO2 methanation, which contradicts other studies and requires modifications in the assumptions made in computational screenings. 26-29 Various additional hypotheses are thus reported concerning the role of iron on enhanced CO2 activation, e.g. synergistic effects due to the alloy or the Ni-Fe(Ox) interface such as faster CO dissociation, 18-20 enhanced reduction properties of Ni or improved CO₂ dissociation. ^{26,27} Further, cascade mechanisms have been considered, i.e. Fe_xC as additional active sites,²⁸ enhanced but limited CO2 dissociation on Fe2+ under irreversible formation of Fe^{3+} , 28 or protection of the active Ni^0 centers by iron. 21,30 Recently, Burger et al. suggested via ex situ studies that segregation of Fe to the particle surface under formation of Fe2+ might provide redox active sites for enhanced CO₂ activation.²⁹

In conclusion, the detailed role of iron in bimetallic Ni-Fe catalysts during CO2 methanation remains unclear and requires monitoring of the structure of iron under reaction conditions. This is challenging, as the structure of both iron and nickel are very dynamic. We shed here more light on structural changes of these catalysts in detail by applying advanced synchrotron-based operando tools. Combined X-ray absorption spectroscopy (XAS) and X-ray diffraction (XRD) was used to monitor changes in the amorphous and crystalline bulk phases as they can be applied under realistic reaction conditions. 21,31-33 Further, surface sensitivity of these methods was enhanced by coupling modulation excitation spectroscopy with XAS, a technique that has received strong attention in catalysis recently.³⁴⁻³⁶ With this method, "spectator species", i.e. species not taking part in the reaction, are filtered out after periodically stimulating active species by alternatingly applied external conditions (e.g., concentration, reactants, temperature, pressure, pH). 34-40 To further complement and substantiate the experimental findings, we used density functional theory DFT calculation and developed structural models. With this we were able to get new insight into the role and dynamic surface changes of iron within the bimetallic Ni-Fe CO₂ methanation catalyst under realistic reaction conditions.

Experimental

Catalyst preparation

A 17 wt% Ni/ γ -Al $_2$ O $_3$ and 17 wt% Ni $_3$ Fe/ γ -Al $_2$ O $_3$ catalyst were synthesized via homogeneous precipitation with urea similar as in previous publications. To ensure an appropriate absorption step during the synchrotron radiation experiments, the catalysts have been diluted with γ -Al $_2$ O $_3$ (1/8" pellets, Alfa Aesar, crushed to <100 μ m, calcined at 600 °C (5 K min $^{-1}$), 4 h) using a ratio of 1:1.3.

Operando setup for the combined X-ray based characterization at synchrotron radiation facilities

A schematic drawing of the setup used in the experiments presented in this study is given in Fig. S1.† A micro quartz capillary reactor (1.5 mm diameter, 20 μm wall thickness, 1 cm catalyst bed length, 100–200 μm catalyst sieve fraction) heated by an Oxford GSB-1300 hot air blower was used for all experiments. The gases were dosed with mass flow controllers by Bronkhorst that were calibrated prior to the experiments using a gas flow calibrator by Mesa Labs. For the MES experiments, a pneumatic 4-way valve by Swagelok was used to ensure fast switches (down to 1 s) between the two gas mixtures that were adjusted separately on two different gas lines.

Combined operando XAS-XRD (BM31, ESRF)

The combined XAS & XRD experiments were performed at the BM31 beamline⁴¹ at the European Synchrotron Radiation Facility (ESRF). XAS spectra were recorded in transmission mode at the Ni K-edge (8333 eV) and Fe K-edge (7112 eV) using a double-crystal Si(111) monochromator. The beam spot size at the sample position was 0.5 mm \times 4 mm ($\nu \times h$). XRD data was collected with a 2D DEXELA detector using a Si(111) channel-cut monochromator, set at a wavelength of 0.4943 Å (beam size 1.0 mm \times 1.0 mm). The XAS-XRD measurements were performed quasi-simultaneously in loop: first XRD data were recorded for 3 minutes (5 diffraction patterns), then X-ray absorption spectra. Switching between XAS and XRD measurements took 2 minutes before and after the XRD. Due to the low signal-to-noise quality at the Fe K-edge two X-ray absorption near edge spectra (XANES) were recorded for 5 minutes, followed by four Ni K-edge extended X-ray absorption fine structure (EXAFS) spectra for 4 minutes, respectively. For data analysis, the spectra were averaged resulting in a time resolution of \approx 29 minutes.

The catalyst was activated by temperature-programmed reduction in 50% $\rm H_2/N_2$ ($\rm H_2\text{-}TPR$), 2 h at 500 °C, heating rate 10 K min⁻¹ at 6 bar. The total gas flow in the capillary was adjusted to 20 mL min⁻¹. The product feed was additionally diluted after the reactor with a constant flux of 40 mL min⁻¹ $\rm N_2$ in order to ensure an adequate gas flow for the $\rm \mu\text{-}GC$ sampling. Before and after the $\rm H_2\text{-}TPR$, XAS and XRD were recorded at room temperature (RT). The catalyst was heated under TPR conditions to 250 °C at 1 bar, and afterwards the gas feed was changed to 25 vol% $\rm H_2$: $\rm CO_2$ = 4:1 in $\rm N_2$ to start the methanation of $\rm CO_2$. To exclude any temperature effects, the catalyst was cooled down to RT after 60 minutes of reaction and structural changes were monitored at RT by XAS–XRD. This procedure was repeated for the methanation of $\rm CO_2$ at 350 °C and 450 °C, respectively.

For calibration and fingerprinting, XAS spectra of reference compounds NiO, NiCO₃, FeO, γ -Fe₂O₃, as well as Fe and Ni foils were measured. Using the Athena software of the Demeter package (version 0.9.25)⁴² the XAS spectra at Ni and Fe K-edges were energy calibrated to the respective metal foils,

background subtracted and normalized. The normalized XANES spectra were analyzed by the linear combination fitting (LCF) module of Athena in the energy interval from -20 to +50 eV using the spectrum of the fully oxidized sample after calcination at 500 °C in air and the spectrum of a Ni foil as references. For EXAFS analysis, Artemis software was used to fit the models to the experimental data obtained during different stages of catalytic treatment. Model structures of Ni/Fe oxides, Ni/Fe metals and Ni-Fe alloy have been used to fit the EXAFS data in R-space for determining the structural parameters. These parameters include energy shift of the path (ΔE_0) , change in the half path length (ΔR) , amplitude reduction factor (S_0^2) , number of identical paths (N) and relative mean-square displacement of the atoms included in path (Debye-Waller factor, σ^2). Further details about the EXAFS data analysis are given in the ESI,† section 3.2.1.

The 2D-XRD diffraction patterns were averaged and azimuthally integrated using the software pyFAI (v 0.15.0).43

Powder XRD (PXRD) patterns were refined by the Rietveld method using the FullProf software package.44 An instrumental resolution file was first obtained by profile fitting of a LaB₆ NIST 640b standard to correct instrumental line broadening. The wavelength used for refinements is 0.049434 nm. The refinement was conducted using a Thompson-Cox-Hastings pseudo-Voigt description of the profile. The background was defined by linear interpolation between background points with refinable heights. A polynomial convolution was initially tested but could not describe the background well. A stack of the full range of all five refined diffraction patterns is shown in ESI† in section 3.2.2. Due to the broad feature at about 7° caused by the capillary the refinement was performed for a range from 10 to 30°. Initial structural models for γ -Al₂O₃ by Zhou et al. 45 and Ni by Rouquette et al.46 were used from literature data. Further information on XRD evaluation and calculations based on Vegard's law is provided in the ESI.†

Modulation excitation spectroscopy during QEXAFS experiments (SuperXAS, SLS)

MES was performed in quick scanning EXAFS (QEXAFS)^{47,48} mode at the Super-XAS beamline^{49,50} at the Swiss Light Source Synchrotron (SLS), Paul Scherrer Institute (PSI). The 17 wt% Ni/Al₂O₃ and 17 wt% Ni-Fe/γ-Al₂O₃ catalysts were activated by H2-TPR in 20 mL min⁻¹ 50% H2/N2, 2 h at 500 °C, 10 K min⁻¹ heating rate at 1 bar. The MES experiments were performed at 250 °C, 350 °C and 450 °C for each catalyst. After the activation, the catalysts were heated to the desired reaction temperature under reducing conditions. Subsequently, the reaction was started by switching to methanation conditions (50 mL min⁻¹ 25 vol% H_2 : $CO_2 = 4:1$ in N2). The catalytic performance was monitored for 1 h under steady state conditions. Afterwards, modulations were applied by periodically switching between 50 mL min⁻¹ 25 vol% $H_2:CO_2 = 4:1$ (60 s) and 20 vol% H_2 in N_2 (60 s) using a pneumatic 4-way valve while simultaneously recording the corresponding XAS data and writing the valve position in the files (details see ESI,† section 2.1). For collimation of the X-ray beam, prior to monochromatisation, a Si coated mirror at 2.9 mrad was used. The QEXAFS monochromator was equipped with the Si(111) channel-cut crystal scanning at 4 Hz. The subsequent monochromatic beam was focused onto the sample position using a toroidal Rh coated mirror. XAS spectra were collected in transmission geometry using a simultaneously measured Ni foil for energy calibration. One period including both conditions took 120 s in which 480 EXAFS spectra were recorded. 30 periods were applied and recorded at each temperature step. The energy range of the recorded transmission mode XAS spectra was adjusted to cover both the Fe and Ni K-edge in one scan.

After the MES experiment, methanation conditions were applied for 1 h to follow changes in the catalytic activity under steady state conditions after the modulations. Subsequently, the MES experiment was repeated at 50 °C as blind test.

The recorded spectra were energy calibrated, normalized and exported as normalized $\mu(E)$ files using the "ProXAS-GUI" software (version 2.9).51 As in case of the bimetallic catalyst the Ni and Fe K-edges were recorded in one scan, the corresponding data files were cut in two after calibration and normalization. One period was divided into 24 sections (resolution 5 s). Each section contained 20 spectra representing a step of 0.25 s, respectively. These 20 spectra per section were averaged to obtain one "time-resolved spectrum" per section. Subsequently, to enhance the signalto-noise ratio, the time-resolved spectra were averaged over all 30 periods. Phase-resolved spectra were obtained by correlation to periodic modulations using eqn (2). 34,35

$$\mu(E,\Delta\varphi) = \frac{2}{T} \int_{0}^{T} \mu(E,t) \sin\left(\frac{360^{\circ}}{T}t + \Delta\varphi\right) dt$$
 (2)

The time-resolved spectra $\mu(E, t)$ were demodulated into phase-resolved spectra $\mu(E, \Delta \varphi)$, by using a sine function of period T = 120 s and a phase shift of $\Delta \varphi$, $0^{\circ} \leq \Delta \varphi <$ 360°. The obtained phase-resolved spectra were interpreted by comparison with difference spectra of reference compounds obtained experimentally as well as simulated ones by FEFF9⁵² that were expected to be formed as intermediates during the modulation (details cf. ESI,† section 3.2.1).

Product analysis

All experiments presented in this study were performed operando. For this purpose, the gaseous products were analyzed on-line using a ThermoStar GSD 320 mass spectrometer by Pfeiffer Vacuum and an Agilent 490 μ-GC (channel 1: 10 m PoraPLOT Q, 0.25 mm diameter, carrier gas helium; channel 2: 10 m mole sieve column with 5 Å, 0.25 mm diameter, carrier gas argon). N2 was used as internal standard during all experiments. The micro-GC measured with a time-resolution of 4 minutes per scan

(1 minute sampling, 3 minutes analyzing). The conversion $X(CO_2)$ and selectivity $S(CH_4)$ were calculated using eqn (3) and (4):

Conversion:
$$X(CO_2) = \left(1 - \frac{CO_{2,\text{out}} \cdot N_{2,\text{in}}}{N_{2,\text{out}} \cdot CO_{2,\text{in}}}\right) \cdot 100\%$$
 (3)

$$\mbox{Selectivity: } \mbox{S(CH$_4$)} = \frac{\mbox{CH$_4$}}{\mbox{CH$_4$} + \mbox{CO} + \mbox{C}_2\mbox{H}_6} \cdot \mbox{100\%} \eqno(4)$$

DFT calculations on the stability of NiFe and FeO_x on Ni and NiFe

Periodic DFT calculations were performed with the projector-augmented-wave (PAW) method using the Vienna *ab initio* simulation package (VASP) in version 5.4.1 as well as the standard VASP-PAWs^{53,54} in connection with the Atomic Simulation Environment (ASE).⁵⁵ The Bayesian error estimation functional with van-der-Waals correlations (BEEF-vdW)⁵⁶ together with *a* plane-wave cutoff energy of 450 eV was used. The GGA+U method^{57,58} was used in order to improve the description of delocalized Fe d states. Using the reduction energy of FeO we obtained a U of 2.7 eV (See ESI†). Lattice constants for Ni, Ni₃Fe and FeO were calculated with a 9 × 9 × 9 Monkhorst–Pack

k-point grid.⁵⁹ The optimized lattice constants are 3.538, 3.599 and 4.363 Å for Ni, Ni₃Fe and FeO, respectively. Infinite slab models with a thickness of four layers and separated by more than 15 Å of vacuum were used to represent Ni_3 Fe(111). The slabs were of the size 2 × 2 in x and y-direction and sampled with a Monkhorst-Pack k-point grid of $6 \times 6 \times 1$. A slab with 5×2 atoms in x and y-direction with a k-point sampling of $2 \times 6 \times 1$ was used to model the Ni₄Fe(111) surface containing 2 Fe and 8 Ni atoms in each layer of the slab. (FeO)₄/Ni₄Fe(111) and (FeO)₆/Ni₄Fe(111) contain FeO periodically repeated in x direction with four and six FeO units per unit cell, respectively. In all calculations, the bottom two layers were kept fixed at the bulk positions whereas all other atoms were allowed to relax during geometry optimization. Convergence criteria of 10⁻⁷ eV and 0.01 eV Å⁻¹ were cycles and geometry optimization, applied to SCF respectively. Spin polarization was considered in all calculations. Vibrational analyses were carried out in the harmonic approximation using a finite difference with a magnitude of displacements of 0.01 Å. Ab initio XANES calculations were performed using the DFT optimized model structures employing the FEFF9 code.⁵² More details can be found in the ESI.†

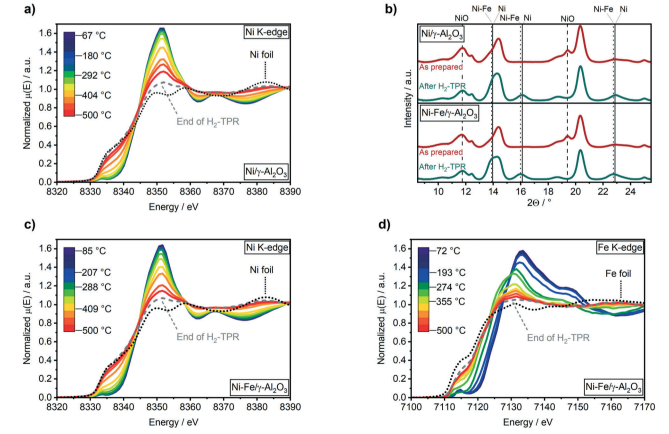


Fig. 1 H_2 -TPR recorded in transmission mode at the Ni K-edge of the 17 wt% Ni/ γ -Al₂O₃ catalyst (a) and at the Ni (c) and Fe (d) K-edge of the 17 wt% Ni-Fe/ γ -Al₂O₃ catalyst using 50% H_2 in N₂ with a total flow of 20 mL min⁻¹ from RT to 500 °C (10 K min⁻¹). (b) XRD patterns (λ = 0.4943 Å) of the 17 wt% Ni/ γ -Al₂O₃ catalyst (top) and 17 wt% Ni-Fe/ γ -Al₂O₃ catalyst (bottom) as prepared (red) and after (green).

Results and discussion

Catalyst activation and Ni-Fe alloy formation

To follow a possible alloy formation, the activation of the Ni-Fe/γ-Al₂O₃ catalyst was monitored by combined XAS-XRD during temperature-programmed reduction in H₂. Further, the results of the Ni-Fe/γ-Al₂O₃ catalyst were compared to a monometallic Ni/y-Al₂O₃ catalyst to elucidate an influence of iron on the reduction properties of nickel (Fig. 1).

The reduction of the Ni/γ-Al₂O₃ catalyst was initiated between 290 °C and 330 °C, as depicted by the decrease in the white line intensity at 8352 eV and the shift to lower energies of the pre-edge feature in the Ni K-edge XANES spectra (Fig. 1a). Contribution of about 86% of Ni^o after the reduction step was estimated by linear combination analysis (LCA) after the H_2 -TPR. Ni reflections at $2\Theta = 13.9^{\circ}$ and 16.1° were observed in the XRD patterns (Fig. 1b) at temperatures above 390 °C (Fig. S2a, ESI†). Simultaneously, NiO reflections at $2\Theta = 11.7^{\circ}$ and 19.5° declined. The formation of Ni⁰ in the Ni– Fe/γ-Al₂O₃ catalyst was also observed around 290 °C to 330 °C (Fig. 1c), while bulk formation was found above 380 °C in the XRD data at $2\Theta = 13.8^{\circ}$ and 16.0° (Fig. S2b† and 1b). About 96% of reduced nickel were obtained after 2 h reduction at 500 °C on the bimetallic Ni–Fe catalyst (86% Ni⁰ for Ni/ γ -Al₂O₃).

EXAFS fitting results at Ni K-edge for Ni-Fe catalyst (ESI,† Table S1) also confirms presence of stable Ni metal phase in bimetallic catalyst after H2-TPR and methanation cycles at different temperatures. Hence, the presence of Fe close to Ni(O) atoms improved reduction of nickel during the activation procedure due to a lower activation energy on mixed Ni-Fe-oxides, as proposed by Unmuth et al. 60 As there was no shift observed in the reduction temperature of nickel, iron might especially enhance the reduction of the smallest nickel particles that are harder to reduce.

Starting from Fe³⁺ state in γ-Fe₂O₃ (Fig. S2c†), the reduction to Fe2+ was visible in XANES from around 85 °C (Fig. 1d). Further reduction of Fe²⁺ to Fe⁰ was observed above 315 °C, as depicted by the shift of the pre-edge feature from 7115 eV to lower energies (Fig. 1d). A maximum reduction of up to 75% compared to the metallic state of iron was found after catalyst activation (H2-TPR). No reflections for any iron or other phase were found in the XRD data. However, a shift of the Ni reflections to lower angles was found in presence of Fe (Fig. 1b). Applying Vegard's law⁶¹ on lattice parameters obtained by Rietveld refinement of XRD data, this shift evidences the formation of a Ni_{4.5}Fe alloy.

This is in contrast to previously reported ex situ HR-TEM and EDX mappings after catalyst activation under similar conditions by which the formation of a Ni₃Fe alloy was reported.²⁰ A Ni-Fe alloy formation could be also observed in fitting of EXAFS data in R space (Fig. 2 and ESI,† S2d) where presence of a Fe-Ni/Fe shell at around 4 Å has been observed.⁶² Corresponding EXAFS fitting results at Fe K-edge are given in ESI,† Table S2. It is worth mentioning here that the detection of FeO backscattering is difficult in EXAFS region due to weak contribution from Fe-O species as well as lower data quality at

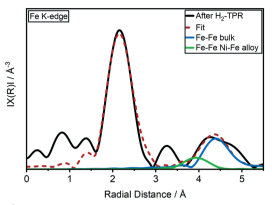


Fig. 2 k^2 -Weighted Fourier transformed EXAFS (black) and the fitted spectrum (red) at the Fe K-edge of the 17 wt% Ni-Fe/γ-Al₂O₃ catalyst after H_2 -TPR in 20 ml min⁻¹ 50 vol% H_2/N_2 at atm (EXAFS fitting results in Table S1†).

elevated temperature. Also, the FeO_x clusters are of amorphous nature which, in combination with the smaller particle size, adds to the loss of the backscattering contribution. The crystallite sizes of both catalysts were estimated to 2.7 \pm 0.1 nm using the Rietveld refined XRD data after the H2-TPR (see ESI†). These were markedly smaller than the estimated particle size of around 3.9 \pm 0.9 nm from a previous study based on scanning transmission electron microscopy (STEM).²⁰

Active state of the Ni-Fe catalyst under steady state conditions

A comparison of the catalytic activity of Ni and Ni-Fe catalysts including detailed structural information about monometallic Ni catalysts during CO2 methanation is given in other studies. 23,29,30,63,64 Hence, the focus in this study is on gaining information about structural composition of the Ni-Fe catalyst with operando techniques. The Ni-Fe/γ-Al₂O₃ catalyst was already active at 250 °C, providing a CO2 conversion of 5% (Table 1) and a selectivity to methane of 75%. The only observed byproduct was CO. Based on these results, the presence of Fe led to a higher formation of CO at 250 °C.20 This indicates an enhanced CO2 dissociation on the Ni-Fe catalyst at low temperatures which would be in agreement to the results obtained by Yan et al.27 on a Ni-FeO_x catalyst. Compared to the laboratory results, 20 the overall activity was lower in this combined operando XAS & XRD synchrotron study. This was due to differences in space

Table 1 CO₂ conversion (X_{CO_2}) and CH₄-selectivity (S_{CH_4}) of the 17 wt% $Ni-Fe/\gamma-Al_2O_3$ catalyst at various temperatures in 20 mL min⁻¹ 50 vol% $H_2:CO_2$ = 4:1 in N_2 at atm; m_{cat} = 3.4 mg of 1:1.3 in γ -Al $_2O_3$ diluted catalyst

| | 250 °C | 350 °C | 450 °C | 350 °C II |
|------------------------|--------|--------|--------|-----------|
| X_{CO_2} /% | 5 | 61 | 66 | 59 |
| $S_{\mathrm{CH_4}}/\%$ | 75 | 96 | 92 | 96 |
| $S_{\rm CO}$ /% | 25 | 4 | 8 | 4 |

velocity caused by catalyst dilution and limitations in catalyst mass that can be loaded in a micro capillary reactor.

No changes were found during CO2 methanation at 250 °C compared to the oxidation state after H2-TPR in the Ni K-edge XANES spectra (Fig. 3a). However, in the Fe K-edge, the pre-edge feature shifted to higher energy and the intensity of the white line at 7130 eV immediately increased (Fig. 3b). These changes represent an oxidation of Fe⁰ to Fe²⁺, e.g. by formation of Fe oxide or carbide. In a recent study it was suggested that iron carbide might be an active species during CO2 methanation, similar as in Fischer-Tropsch synthesis.²⁸ Kureti et al. investigated such an iron carbide formation during CO2 methanation on bulk Fe catalysts and found it only for iron crystallites bigger than 23 nm.65 As the crystallite sizes of the catalysts used in our study were around 2.7 nm, a bulk formation of carbides is unlikely. Further, iron carbide formation was reported mainly for CO methanation, while Fe₃O₄ (mixture of Fe²⁺ and Fe³⁺) was reported to form during CO₂ methanation on pre-reduced iron catalysts.⁶⁶ Finally, a comparison of the Fe K-edge XANES spectra obtained during CO2 methanation with the partially reduced FeOx spectra during H2-TPR provided evidence on FeO_x formation (cf. Fig. S3†). As oxygen free gases were used in this study, this clearly demonstrates that the oxygen must originate from CO2 activation.

By increasing the reaction temperature to 350 °C a CO₂ conversion of 61% with a selectivity to CH₄ of 96% was achieved (Table 1). While the elevated temperature had no effect on the Ni K-edge XANES spectrum (Fig. 3), iron was further oxidized up to 46% FeO (estimated from LCA) as depicted by a further shift in the feature at 7117 eV and an increase in the intensity at 7130 eV in the Fe K-edge spectrum (Fig. 3b). In contrast to the temperature step at 250 °C, a slight shift of the Ni(200) reflection to a slightly higher diffraction angle at $2\theta = 16.10^{\circ}$ was observed in the XRD patterns (Fig. 3c). According to analysis of lattice parameters from Rietveld refinement (Table 2) using Vegard's law, this shift represents an enrichment of Ni in the alloy from 4.8 to about 6.0. Hence, the increased temperature and CO₂ conversion led to a segregation of iron to the surface including formation of a higher fraction of FeO. Such a structural change was also reported by Yan et al.27 based on situ ADF-STEM and EELS elemental maps after methanation of CO₂. However, in contrast to their results, the higher fraction of FeO did not lead to a higher CO selectivity in the present study (Table 1). A further increase in temperature to 450 °C resulted in a CO2 conversion of 66% (Table 1). The selectivity to methane declined from 96% to 92% due to the thermodynamically favored water-gas-shift

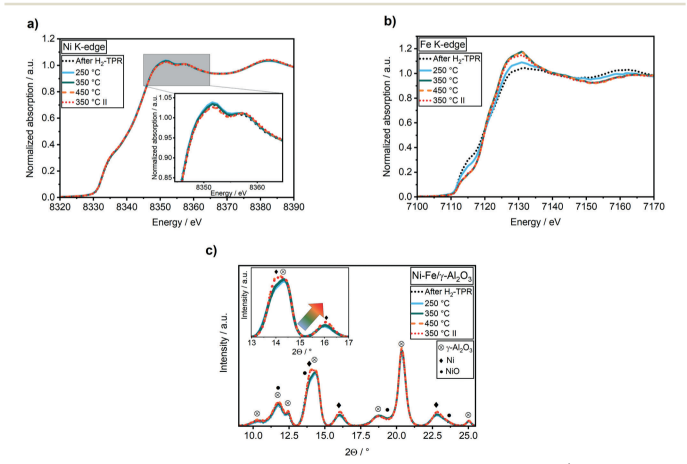


Fig. 3 XANES spectra recorded in transmission mode at the Ni (a) and Fe (b) K-edge and (c) XRD patterns (λ = 0.4943 Å) of the 17 wt% Ni-Fe/γ- Al_2O_3 catalyst at 50 °C after applying methanation conditions of 20 mL min⁻¹ 50 vol% H_2 : $CO_2 = 4:1$ in N_2 at atm and various temperatures for 1 h.

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Table 2 Lattice parameters, crystallite sizes and via Vegard's law calculated amount of Ni in the Ni-Fe alloy of the Ni phase obtained from Rietveld refinement of the XRD patterns (λ = 0.4943 Å) of the 17 wt% Ni-Fe/γ-Al₂O₃ catalyst recorded at 50 °C after applying methanation conditions of 20 mL min⁻¹ 50 vol% $H_2:CO_2 = 4:1$ in N_2 at atm and various temperatures for 1 h

| | 250 °C | 350 °C | 450 °C | 350 °C II |
|--|--------------------------------|--------------------------------|--------------------------------|----------------------------|
| Lattice parameter/pm | 354.3 ± 0.3 | 353.9 ± 0.2 | 353.6 ± 0.2 | 353.6 ± 0.2 |
| Crystallite size/nm Bulk: ratio Ni:Fe/- | 2.5 ± 0.1 4.8 ± 0.8 | 2.7 ± 0.1 6.0 ± 0.7 | 3.2 ± 0.1 7.3 ± 0.9 | 3.3 ± 0.1 7.5 + 1.0 |
| Bulk: oxidized Fe/% | 19.5 ± 0.8 | 45.8 ± 1.2 | 51.0 ± 1.4 | 49.5 ± 1.4 |

reaction. Interestingly, no further significant increase in FeO was observed (Fig. 3b). This provides evidence, that the effect of FeO formation correlates rather to catalytic activity than temperature. Concurrently, an enrichment of Ni in the alloy from 6.0 to 7.3 was found, while the crystallite size increased from 2.7 nm to 3.2 nm (Table 2). Hence, alloy stability and sintering depend on reaction temperature.

To investigate the influence of dealloying and sintering on the catalyst performance, a second methanation step was performed for comparison at 350 °C. According to the Ni and Fe K-edge XANES spectra (Fig. 3a and b) and the results obtained by XRD data (Table 2) the changes were not reversible. A slightly lower CO2 conversion of 59% compared to 61% was observed (Table 1). This might be caused either by the higher Ni: Fe ratio of 7.5 compared to 6.0 or by the increase in crystallite sizes from 2.7 nm to 3.3 nm.

Surface changes and FeO_x cluster formation

The segregation of iron to the particle surface might lead to formation of, e.g. a core-shell like structure or to formation of clusters on the surface. To obtain further information about the structural composition at the catalyst surface, we calculated the number and composition of the surface atoms. For this purpose, we assumed that nickel and iron were both in fcc structure (premise for alloy formation). A particle size of 3.9 \pm 0.9 nm was determined for this catalyst via STEM in a previous study.²⁰

For calculation of volume v_p (eqn (5)) and surface area a_s (eqn (6)) of one particle, we assumed hemispherical particles on the support and a linear correlation of changes in crystallite (operando XRD data) with particle size.

$$\nu_{\rm p} = \frac{2}{3} \pi \cdot r_{\rm p}^3 \tag{5}$$

$$a_{\rm s} = 2\pi \cdot r_{\rm P}^2 \tag{6}$$

The average number of atoms n_S per m² and the area occupied by a surface atom a_{m} were calculated as described by Bergeret and Gallezot.⁶⁷ For this, it was assumed that the molar volume $\nu_{\rm m}({
m Ni-Fe})$ for a Ni-Fe "model" atom can be estimated by the fraction X_{Ni} of nickel in the alloy multiplied with the respective standard molar volume for nickel and iron⁶⁸ (eqn (7) and (8)).

$$X(Fe) = 1 - X(Ni) \tag{7}$$

$$\nu_{\rm m}(\text{Ni-Fe}) = X(\text{Ni}) \cdot \nu_{\rm m}(\text{Ni}) + X(\text{Fe}) \cdot \nu_{\rm m}(\text{Fe})$$
(8)

The total number of atoms in one particle N_p and at the surface N_s were subsequently obtained by eqn (9) and (10).

$$N_{\rm p} = \frac{\nu_{\rm s}}{\nu_{\rm m}} \tag{9}$$

$$N_{\rm s} = \frac{a_{\rm s}}{a_{\rm m}} \tag{10}$$

The number of iron or nickel atoms in the particle or at the surface were determined by multiplying N_p and N_s with the fractions X of nickel and iron in the alloy, respectively.

Assuming the number of atoms is constant at each temperature step "T", iron that formally leaves the Ni-Fe alloy must agglomerate at the surface. The amount of segregated iron $X_{\text{seg}}(\text{Fe})$ out of total number of iron atoms after H₂-TPR ("TPR") was calculated by eqn (11):

$$X_{\text{seg}}(\text{Fe}) = 1 - \frac{X_{\text{T}}(\text{Fe})}{X_{\text{TPR}}(\text{Fe})}$$
(11)

Subsequently, the surface composition R was obtained according to eqn (12) and (13):

$$N_{\rm T}({\rm Fe}) = X_{\rm TPR}({\rm Fe}) \cdot N_{\rm s} + X_{\rm seg}({\rm Fe}) \cdot N_{\rm p}({\rm Fe})$$
 (12)

$$R_{\rm T}({\rm Ni:Fe}) = \frac{N_{\rm s} - N_{\rm T}({\rm Fe})}{N_{\rm T}({\rm Fe})} \tag{13}$$

Assuming the amount of oxidized bulk iron atoms OBA(Fe) that was estimated by LCA originated mainly from the surface atoms, the fraction of oxidized surface atoms OSA(Fe) was given by eqn (14):

$$\mathrm{OSA_T(Fe)} = \frac{\mathrm{OBA_T(Fe)} \cdot \left[X_{\mathrm{T}}(\mathrm{Fe}) \cdot N_{\mathrm{p}} \right]}{N_{\mathrm{T}}(\mathrm{Fe})} \tag{14}$$

The results are displayed in Table 3 and S8.† The big error margin results from the error of ±0.9 nm in particle size that gives a significant variation in the total number of atoms which is reflected in all values due to the propagation of uncertainty.

Nevertheless, assuming that the model particle is 3.9 nm without any variation, the values given in Table 3 can be considered as relevant to observe and discuss trends. In future, this may be substantiated by in situ XPS under selected model conditions.

We observed that up to 35% of the total iron moved from bulk (cf. Table 2) to the particle surface. This resulted in changes in the Ni:Fe surface ratio from 4.9 up to 1.9 at 450 °C. An amount of \approx 60% oxidized surface iron atoms were determined at 250 °C, ≈87% at 350 °C and ≈78% at 450 °C. Hence, we can conclude that the iron surface species were in a mainly oxidized state. The segregation and

Table 3 Calculations on the surface composition and changes with respect to iron. For further details see text and ESI[†]

| | ATPR | 250 °C | 350 °C | 450 °C | 350 °C II |
|---|---------------|--------------------------------|---------------------------------|---------------------------------|------------------------------|
| Iron atoms segregated/% Surface: ratio Ni:Fe/- | 4.5 ± 0.8 | 4.4 ± 1.4 3.7 ± 0.8 | 20.8 ± 1.8 2.3 ± 0.9 | 32.8 ± 1.8 1.7 ± 0.8 | 35.0 ± 1.0 1.6 ± 0.8 |
| Surface: oxidized Fe/% | _ | 59.5 ± 14.5 | 86.6 ± 15.2 | 78.4 ± 11.2 | 74.2 ± 9.7 |

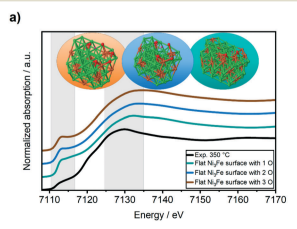
enrichment of iron on the surface under formation of oxide species might either lead to a "core-shell"-like structure of an FeO_x layer or to formation of FeO_x clusters on top of the Ni-Fe alloy particles. We simulated several structures for these cases with DFT and calculated the respective XANES spectra using the FEFF9-code⁵² to compare them to our experimental data (Fig. 4). The spectra were simulated by averaging the core-holes in Ni/Fe atoms at all possible positions on the surface to get the final spectrum. Details about the model generation and simulation procedure with parameters used in FEFF9 are given in ESI,† section 3.2.1.

The simulated XANES spectra of the oxidized "core-shell"like flat Ni₃Fe surface (Fig. 4a) were not in a good agreement to the experimentally obtained Fe K-edge XANES spectrum during methanation of CO₂ at 350 °C (Fig. 4a, black curve): the white line feature was shifted to 7133 eV instead of 7130 eV and the pre-edge feature at \approx 7112 eV was too sharp. In contrast, a good match was found for FeOx clusters on both, a Ni₃Fe and a Ni surface (Fig. 4b): the white line feature appeared at the same energy of 7130 eV as the experimental data and the pre-edge feature at ≈7112 eV provided a similar shape. Concerning the latter, the best match was found for the FeO_x cluster I on a Ni_xFe_y layer.

Dynamic changes of the active centres at the surface (MES)

To investigate the influence of these FeO_x clusters and to understand their role on CO2 activation at the particle surface, we coupled the bulk technique XAS with modulation excitation spectroscopy (MES)35 to enhance its surface sensitivity. For this purpose, periodic switches (modulation) between H_2/N_2 (0-60 s) and $H_2/CO_2/N_2$ (60-120 s) in the gas feed were conducted at various temperatures while monitoring the dynamic responses at the Ni and Fe K-edges during these CO2 modulations with QEXAFS. Subsequently, "spectator" species (those which do not respond to the atmosphere) were filtered out.³⁴ In our case, we decided to use a sinusoidal function as approximation for the stimuli we applied, similar to other catalytic studies. 36,69 Note, that external stimuli in a real experiment do not perfectly match a mathematical sinusoidal wave. Hence, in some cases, the signal processing can be enhanced by using square wave modulation and/or corrections terms for amplitude.34,70 These were not applied in our study, as the focus was laid on relative phase changes rather than kinetics or quantitative analysis. Prior to the modulation experiments, the catalyst was activated by H2-TPR and treated under methanation conditions until a steady state was reached. As these structural changes were discussed in detail in the previous section and are the same as during the XAS-XRD experiments, the data are only shown for completeness in the ESI.† In addition, we focus here on the role of iron, hence only a short summary on the MES results at the Ni K-edge will be given.

At 250 °C, the shape of the demodulated Ni K-edge spectrum was similar for both catalysts (cf. Fig. S18 and S19†). This demonstrates that the origin in the difference in catalytic activity at 250 °C (Table 1) can be traced back to reactions on iron sites. None of the difference spectra obtained from measured and simulated references matched precisely to experimental demodulated MES data obtained at the Ni K-edge. Hence, surface changes at Ni atoms are very complex. Nevertheless, we can conclude with certainty that we have no indication of Ni-O species formed during CO2



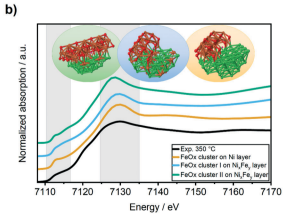
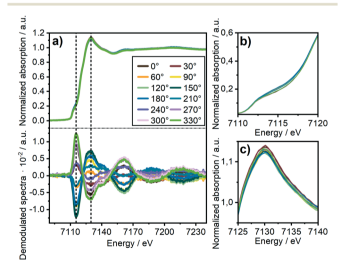


Fig. 4 Fe K-edge XANES spectra of the experimental data obtained from the bimetallic Ni-Fe/Al₂O₃ catalysts after methanation of CO₂ at 350 °C compared to FEFF calculated XANES spectra of iron-rich surface with increasing amount of oxygen (a) and to FeOx clusters-like structures (b).

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modulation, and that the same surface species were formed on Ni on both catalysts. For further details and discussion see ESI.† Fig. 5a (top) shows time-resolved XANES spectra of the Ni-Fe catalyst at the Fe K-edge during CO₂/H₂ vs. H₂ cycling obtained from averaging the complete 30 periods. There are some slight changes observed for the pre-edge feature at ≈7116 eV and for the white line feature at 7129 eV as the zoomed spectra in Fig. 5b and c show, respectively. However, in conventional XAS spectra these changes are too small to predict some phase transitions. In contrast, using demodulation uncovers striking changes. The corresponding demodulated XANES spectra during cycling at various phase angles are shown in Fig. 5a (bottom) where noticeable changes were both observed at the energy positions mentioned for pre-edge and white line region in the Fe K-edge spectra. An asymmetry can be observed in these demodulated spectra, especially at 7140 eV. This demonstrates that complex changes occurred at the Fe K-edge corresponding to changes of more than two species.³⁷ To get further information about these changing species, the demodulated spectrum with the highest amplitude is in a next step compared to experimentally obtained difference spectra from reference spectra of Fe metal, FeO and γ-Fe₂O₃ (Fig. 6a). A very good correlation to oxidized iron species, such as Fe₂O₃-FeO and Fe₂O₃-Fe, was found. This demonstrates that during CO2 cycling Fe provided highly dynamic changes concerning its oxidation state even up to Fe³⁺. Note, this does not mean that a major Fe₂O₃ phase is formed during modulation. Fe³⁺ is known to fit well into the FeO-lattice. In addition, Fe²⁺ atoms at the interface between FeO_x and the Ni-Fe surface may be partially oxidized to Fe3+ due to CO2 activation. As this is the only species that can be periodically correlated to the modulations in the gas phase seen in MES,



these changes can be clearly correlated to CO2 activation.

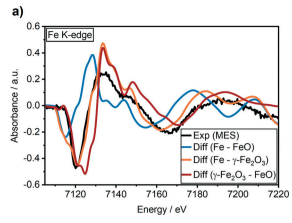
Fig. 5 (a) Normalized time-resolved Fe K-edge XANES consisting of a total of 24 spectra covering 5 seconds each for the 17 wt% Ni₃Fe/γ- Al_2O_3 catalyst during $CO_2/H_2/N_2$ (0-60 s) vs. H_2/N_2 (60-120 s) cycling and 30 periods average (top) and corresponding demodulated spectra (bottom) at selected values of phase angle $(\Delta \phi)$. (b) Zoomed edge region, (c) zoomed white line region.

The temperature dependency is given in Fig. 6b. At 250 °C a very low intensity in the features of the demodulated spectrum was observed. As we observed a higher selectivity to CO for the bimetallic Ni-Fe sample at 250 °C (Table S9†), we can assume that CO₂ is only slowly activated at the Fe⁰ centers under formation of CO and FeO. In this temperature range, Ni provides a comparatively high affinity to CO, as observed in the MES at the Ni K-edge (cf. ESI† section 3.4.7). DFT calculations demonstrated that the presence of Fe does not influence the CO adsorption energy on Ni (cf. section 3.5.3). Hence, the Ni surface is covered by CO probably hampering the hydrogen activation that led to the high fraction of CO formation. An increase in temperature from 250 °C to 350 °C and 450 °C showed large increase in intensity of the demodulated spectrum at Fe K-edge as shown in fig. 6b. This provides evidence of strong dynamic responses of Fe to the CO₂ modulations at elevated temperature which can be correlated well to the determined catalytic activity (see Table 1).

However, there are various iron positions where these dynamic changes might occur, e.g. on top of the cluster or at the Ni-FeO_x interface (cf. Fig. 4b). We observed that the amplitude of the response signal obtained from stimulated Ni (see ESI,† Fig. S20) and Fe atoms was in the same range for each temperature step and that it increased with temperature. As this correlates well with the degree of CO2 conversion (Table 1), this indicates that stimulated atoms represent the most active centers and that the "redox"-like behavior they provide is important for CO₂ activation and the overall catalyst performance.

Stability of Ni-Fe alloy and formation of FeOx clusters

In order to shed light on the driving forces of the iron oxide cluster formation and on the stability of the Ni-Fe alloy during reaction, DFT calculations of this process have been performed. The calculated heat of formation of the Ni₃Fe alloy is -0.60 eV. We tested the surface segregation of Fe using a Ni(111) slab. Under vacuum, this energy is 0.16 eV, such that Fe is more stable in the subsurface. When CO2 and H₂O are present in the reaction atmosphere, there is a change of the oxygen chemical potential. We calculated the oxidation of a Ni-Fe alloy with Ni₄Fe stoichiometry. Here, we considered the segregation of Fe on top of the Ni₄Fe(111) surface where it forms FeO (this is calculated by a corresponding decrease of the iron content in the alloy, see ESI† for further details). The result is depicted as a phase diagram in Fig. 7 for two different FeO structures the stability of whose is calculated as a function of the H₂O/H₂ pressure and temperature (see Fig. S24 in the ESI† for the corresponding diagram with CO₂/CO pressure). As can be seen from Fig. 7, Fe is according to the calculation predicted to segregate on top of the Ni₄Fe(111) surface where it is oxidized to small FeO clusters, as also observed experimentally. Note that these structures were used to calculate the XANES spectra in Fig. 4, which showed a



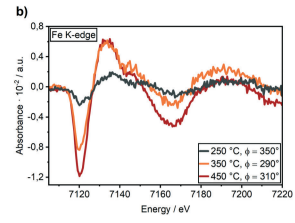


Fig. 6 (a) Demodulated Fe K-edge spectra of the 17 wt% Ni-Fe/γ-Al₂O₃ catalyst at 450 °C (black) compared to difference spectra obtained from experimental Fe reference spectra and (b) demodulated Fe K-edge spectra at 250 °C (black), 350 °C (orange) and 450 °C (red) during CO₂/H₂/N₂ $(0-60 \text{ s}) \text{ vs. } H_2/N_2 \text{ (60-120 s)}$ cycling with 30 periods average.

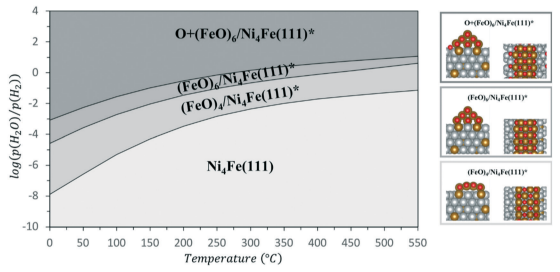


Fig. 7 Phase diagram of $Ni_4Fe(111)$ and $(FeO)_x/Ni_4Fe(111)$ system as a function of the pH_2O/pH_2 ratio and for temperature range between 0 and 550 °C. see ESI† for a phase diagram using pCO₂/CO and for further details.

reasonable agreement. We considered oxidation of (FeO)₆/ Ni₄Fe(111) such that there is now an extra oxygen atom at the interface between FeO and the Ni₄Fe(111) surface (see Fig. 7). This oxygen, however, is slightly unstable under the applied conditions and we therefore speculate that CO2 methanation could take place via a redox-mechanism at the FeO-alloy interface, that is, CO2 splits at the interface into CO (adsorbed on the Ni₄Fe(111) facet) and O* located at the interface. We calculated the energy to remove this O* via hydrogenation with H₂ to be downhill by -0.61 eV in energy. Removal of stoichiometric oxygen of the FeO with H2 is about thermoneutral (-0.02 eV), again pointing towards a possible redox mechanism at the FeO-alloy interface.

Mechanism of CO₂ activation on Ni-Fe alloy catalysts

In our operando studies on the bimetallic Ni-Fe alloy catalyst, we observed partial segregation of iron under formation of FeO_x at the particle surface during CO_2 methanation. The

increasing amount of FeOx at the surface led to formation of FeO_x cluster on top of the Ni-Fe particles, which is schematically shown in Fig. 7. These clusters offer new actives sites for CO2 activation and provide, as we demonstrated by MES, a redox cycle between $Fe^0 \Rightarrow Fe^{2+} \Rightarrow Fe^{3+}$. Such a redox cycle was suggested e.g. on bulk Fe₃O₄ catalysts during CO₂ activation 71-73 under distinct formation of carbonaceous surface species which led to a superior CH₄ yield compared to iron carbide based catalysts.71,74 However, we were not able to detect any surface carbon species on the bimetallic Ni-Fe catalyst during our operando studies, probably due to the high CO2-pressure. Previous operando Raman studies on a Ni-Fe based catalyst from our group¹⁶ did not provide any hints for formation of carbonaceous surface species. Besides CH4, CO remains the main product on FeOx catalysts in the gas-phase during CO₂ methanation.⁷⁵ The formation of CO can be attributed to the RWGS activity of FeO_x and is, according to our experimental observations and calculations, another key step in enhanced CO_2 activation of Ni-FeO_x catalysts (cf. Fig. 8). On

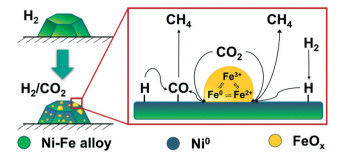


Fig. 8 Scheme of the CO₂ activation mechanism on Ni-Fe alloy-based catalysts during methanation reaction under realistic conditions.

monometallic Ni catalysts, CO2 dissociation proceeds fast, but CO is strongly bound on reduced Ni⁰ centers and therefore adsorption sites for CO₂ are blocked.⁷⁶⁻⁷⁹ These steps can proceed separately on the Ni-FeO_x system: CO₂ can be activated and hydrogenated either directly to CH4 or dissociated under formation of CO at the interface between the FeO_r clusters and the Ni-Fe surface (Fig. 8). The strong adsorption of CO on Ni⁰ might subsequently attract it to the Ni centers where it can be further hydrogenated via the CO methanation route. This RWGS activity of the FeOx clusters on the Ni-FeOx catalyst further explains the high CO selectivity at low temperatures: while the FeO_x clusters are already active in RWGS activity, the subsequent hydrogenation step at the Ni^o centers does not proceed at low temperature. These two suggested mechanisms are in line with the MES data obtained at the Ni K-edges of both catalysts which demonstrated the formation of identic intermediates on the Ni and the Ni-Fe catalyst (ESI,† Fig. S22).

To summarize, the $Fe^0 \Rightarrow Fe^{2+} \Rightarrow Fe^{3+}$ redox cycle provided by the $Fe(O_r)$ entities at the interface between the clusters and the metal surface improved both, dissociative and associative CO2 activation. Beneficial in the case of the bimetallic Ni-Fe catalyst is that different active sites are available on the reduced Ni surface, the interface and the iron oxide clusters for the adsorption of the respective reactants and, thus, always free sites are available for both CO₂ activation and hydrogenation in contrast monometallic Ni catalysts.

Conclusions

In this study, we investigated in detail the role of iron in a highly active Ni-Fe/γ-Al₂O₃ catalysts during CO₂ methanation. For this purpose, advanced synchrotron-based tools, such as XAS-XRD and XAS coupled MES have been applied and the findings further substantiated by DFT calculations.

We noticed a synergistic effect of iron on nickel which led already during catalyst activation by H2-TPR to a higher fraction of active Ni⁰ species. First, surface atoms or amorphous species were reduced followed by the bulk phase up to formation of nanocrystalline nickel species at elevated temperature. The reduction of Fe2+ to Fe0 occurred simultaneously to the reduction of Ni2+ to Ni0 which led to the formation of the desired Ni-Fe alloy. Notably, no reflections were found in the XRD data for any iron phases during the experiments, which, supported by the changes in the lattice parameters, evidences an incorporation into an alloy.

During stoichiometric CO₂ methanation in the temperature range from 250 °C to 450 °C, a good comparability of the operando synchrotron studies to laboratory studies could be achieved concerning CO2 conversion and selectivity to CH4. Under all applied CO2 methanation conditions nickel remained in its reduced metal state as obtained after H2-TPR. In contrast, an oxidation of Fe⁰ to Fe²⁺ was immediately observed upon addition of CO₂. The amount of Fe²⁺ thereby correlated to CO2 conversion, either due to CO2 dissociation or from other intermediates/products, such as H₂O. Increasing the reaction temperature from 250 °C up to 450 °C resulted in sintering, and agglomeration of Fe at the particle surface under formation of FeOx clusters, as demonstrated by XAS-XRD and DFT calculations.

By enhancing the sensitivity of XAS with MES, it was shown that iron exhibits a highly dynamic behavior during CO_2 activation. A $Fe^0 = Fe^{2+} = Fe^{3+}$ redox cycle, highly likely located at the interface between the FeOx clusters and the surface of the metal particles, promotes CO2 dissociation during the methanation reaction. Combination of these findings with DFT calculation conclude that FeOx clusters on reduced Ni particles are at the origin of the high catalytic activity of Fe-Ni catalysts and, by this, CO2 activation and the following hydrogenation are improved.

Conflicts of interest

There are no conflicts to declare.

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