Surface modification processes during methane decomposition on Cu-promoted Ni–ZrO₂ catalysts

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The surface chemistry of methane on Ni–ZrO₂ and bimetallic CuNi–ZrO₂ catalysts and the stability of the CuNi alloy under reaction conditions of methane decomposition were investigated by combining reactivity measurements and in situ synchrotron-based near-ambient pressure XPS. Cu was selected as an exemplary promoter for modifying the reactivity of Ni and enhancing the resistance against coke formation. We observed an activation process occurring in methane between 650 and 735 K with the exact temperature depending on the composition which resulted in an irreversible modification of the catalytic performance of the bimetallic catalysts towards a Ni-like behaviour. The sudden increase in catalytic activity could be explained by an increase in the concentration of reduced Ni atoms at the catalyst surface in the active state, likely as a consequence of the interaction with methane. Cu addition to Ni improved the desired resistance against carbon deposition by lowering the amount of coke formed. As a key conclusion, the CuNi alloy shows limited stability under relevant reaction conditions. This system is stable only in a limited range of temperature up to ~700 K in methane. Beyond this temperature, segregation of Ni species causes a fast increase in methane decomposition rate. In view of the applicability of this system, a detailed understanding of the stability and surface composition of the bimetallic phases present and the influence of the Cu promoter on the surface chemistry under relevant reaction conditions are essential.

Introduction

Hydrogen is known as a promising source of energy and fuel. Up to now hydrogen is mostly produced out of hydrocarbons such as methane. Via steam reforming¹–⁴ or partial oxidation¹⁵ of methane, CO or CO₂ is formed as a by-product. For many applications such as the use of hydrogen in PEM fuel cells CO needs to be removed because it is a catalyst poison. One possibility to overcome the problem of CO or CO₂ formation is the direct production of hydrogen out of methane by catalytic decomposition in the absence of air or oxygen. The by-product of this process is carbon in the form of layers, filaments, fibres or nanotubes. Especially, the formation of useful nanocarbons⁷–¹⁴ is of high interest. Li et al.¹⁵ published a comprehensive review about this topic discussing the effect of the nature of the catalyst and feed composition on the catalytic activity, stability and the influence on the morphology of the produced carbon.

During methane decomposition, methane is adsorbed on the catalyst followed by C–H bond breaking, deposition of carbon and generation of hydrogen. Transition metals such as Ni,¹⁶–¹⁸ Fe¹⁷ and Co¹⁹ are known to be effective for methane decomposition to hydrogen and carbon nanofibres. Ni catalysts are known for their high activity but are easily deactivated due to carbon deposition. However, nickel-based catalysts have been proven to be sensitive to doping and structure modification by other metals, such as Cu, which affects the structure and morphology of the produced carbon.⁸ Improved stability against deactivation has been achieved by adding a second metal component; for example, Au²⁰–²¹ and Cu¹³,²⁴–³² have been studied for this aim. The Ni–Au surface alloy was active for steam reforming and more resistant toward carbon formation than the pure Ni, which was explained by blocking of highly reactive Ni edge and kink sites by Au atoms.²¹ The addition of Cu to nickel-based catalysts has several effects such as an easier reducibility of Ni. Cu is rich in d-electrons which may exert an electronic effect on Ni and furthermore affect the affinity of the metal surface to graphite.¹⁵ Ashok et al. observed that addition of Cu to Ni
enhanced the methane decomposition activity of hydrotalcite-like Ni–Cu–Al catalysts when added in an appropriate amount.\textsuperscript{26} Cunha \textit{et al.} showed that doping of Cu has a strong influence on the dispersion of nickel and enhances the catalyst activity.\textsuperscript{30} Similarly, Monzón found that promoting Ni with a small amount of Cu improves the activity and stability of the catalyst.\textsuperscript{31} Cunha \textit{et al.} reported that doping with Cu improves the catalyst stability.\textsuperscript{32} Li \textit{et al.} observed that in Cu$_{0.25}$Ni$_{0.75}$Al$_2$O$_4$ Cu is acting as a dispersing agent to prevent Ni from growing\textsuperscript{33} and enhances the catalyst stability when employed in a suitable amount.\textsuperscript{13} Lazaro \textit{et al.} found that the presence of copper in NiCu–SiO$_2$ catalysts led to higher methane conversion,\textsuperscript{28} and Park \textit{et al.} observed that on Cu–Ni–YSZ anode materials for the direct use of methane fuel in solid oxide fuel cells (SOFC) carbon deposition decreased compared to the amount of carbon on Ni–YSZ.\textsuperscript{29} DFT-based computational studies by Liu \textit{et al.} compared methane dissociation on NiCu(111) bimetallic alloy surfaces. Their calculations indicated that C is easily formed on a uniform 1:1 NiCu surface but unfavourable on a Cu-rich NiCu surface.\textsuperscript{33} Therefore, the surface composition on a NiCu alloy catalyst under reaction conditions is expected to play the key role in determining the catalytic activity and stability.

One prerequisite for a successful promoter is in situ stability under relevant reaction conditions, \textit{i.e.} high temperatures and reactive surrounding gas atmosphere. Therefore, understanding the stability of the bimetallic phases and the influence of the promoter on the surface chemistry are essential. We have studied CH$_4$- and methanol reforming activities and selectivities on Ni-, Cu- and Pd-based catalyst systems by combining in situ spectroscopic techniques such as XPS, FTIR and XAS.\textsuperscript{25,34-39} In the present work, we have selected Cu as an exemplary compound for modifying the reactivity of Ni in methane decomposition and enhancing the resistance against coke formation.

In this work we focus on the surface chemistry of methane on Ni–ZrO$_2$ catalysts during methane decomposition at temperatures up to 773 K and study the stability of the CuNi alloy under reaction conditions. By \textit{in situ} synchrotron-based near-ambient pressure (NAP-)XPS we determined the surface composition of the catalysts and the nature of the carbonaceous species present and correlated these data with the catalytic performance. Cu addition to Ni improved the desired reduction of carbon deposition,\textsuperscript{40} but only in a limited range of temperature due to the limited stability of this system.

\section*{Experimental}

\subsection*{Sample preparation}

Ni–ZrO$_2$ and bimetallic CuNi–ZrO$_2$ catalysts were synthesized by impregnation of ZrO$_2$ using nitrate precursor salts. Prior to impregnation, commercial Zr(OH)$_4$ (MEL Chemicals XZO 880/01) was calcined at a heating rate of 2 K min$^{-1}$ from room temperature to 973 K and kept at this temperature for 2 h. After calcination the zirconia support exhibits a specific surface area of 36.6 m$^2$ g$^{-1}$.

Bimetallic CuNi samples were prepared by coimpregnation. Cu(NO$_3$)$_2$·3H$_2$O (Fluka, p.a.) and Ni(NO$_3$)$_2$·6H$_2$O (Merck, p.a.) were mixed to obtain the following Cu:Ni molar ratios: 1:0, 3:1, 1:1, 1:3 and 0:1. The nitrates were dissolved in water, and in each case 4.75 g of ZrO$_2$ powder was suspended in these solutions. Every solution contained as much metal nitrate to obtain 5 wt% metal loading in the final catalyst powder. Additionally, a 5 wt% Cu–ZrO$_2$ reference material was synthesized by the same procedure.

For \textit{in situ} X-ray photoelectron spectroscopy (XPS) investigations an additional sample was prepared via the same synthesis route with a higher overall metal loading of 50 wt% in order to prevent sample charging.

All catalysts and their respective compositions are summarized in Table 1.

\begin{table}[h]
\centering
\caption{Composition of CuNi–ZrO$_2$ catalysts}
\begin{tabular}{llll}
\hline
Sample name & ZrO$_2$ (wt\%) & Cu (wt\%) & Ni (wt\%) \\
\hline
Ni–ZrO$_2$ & 95 & — & 5 \\
13CuNi–ZrO$_2$ & 95 & 1.25 & 3.75 \\
11CuNi–ZrO$_2$ & 95 & 2.5 & 2.5 \\
31CuNi–ZrO$_2$ & 95 & 3.75 & 1.25 \\
Cu–ZrO$_2$ & 95 & 5 & — \\
13CuNi–ZrO$_2$50 & 50 & 12.5 & 37.5 \\
\hline
\end{tabular}
\end{table}

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For the quantitative determination of the reaction rates the reaction products were analyzed under steady-state conditions by a gas chromatograph equipped with a thermal conductivity detector (TCD) and flame ionization detector (FID).

**In situ XPS measurements**

In situ synchrotron-based near-ambient pressure (NAP-)XPS measurements were carried out at the ISSIS-PGM beamline at the Helmholtz-Zentrum Berlin using a high-pressure XPS station constructed at the FHI Berlin. Details of the setup are described elsewhere. Briefly, the setup uses a differentially pumped electrostatic lens system and a SPECS hemispherical electron analyser.

The sample was placed inside a reaction cell with adjustable gas flow and a total pressure up to the millibar range. The composition of the gas phase was analysed by MS. Heating was done by an IR laser from the back side. The sample was in situ oxidized in 0.1 mbar O$_2$ at 573 K and reduced in 0.25 mbar H$_2$ at 673 K prior to exposure to 0.25 mbar CH$_4$ at temperatures ranging from 523 up to 723 K.

The catalyst 13CuNi–ZrO$_2$50 with 50 wt% total metal loading was used for these experiments which prevented sample charging. The calcined powder was pressed onto a copper plate and reduced in 0.25 mbar H$_2$ at 773 K prior to placing it into the in situ reaction cell. Measurements were performed at photon energies of 1100 eV for the Cu 2p, 1010 eV for the Ni 2p and 425 eV for the C 1s XP spectra, yielding a photoelectron kinetic energy of 150 eV. For depth profiling the photon energies were varied to obtain additional photoelectron kinetic energies of 350 eV and 550 eV. All spectra were corrected for synchrotron beam current, incident photon flux and energy-dependent photo-ionization cross sections.

Binding energies (BEs) were referenced to the Fermi edge recorded after each core-level measurement. The measured signal envelopes were fitted to Gauss–Lorentzian peaks by using Casa XPS software after subtraction of a Shirley background.

**Results**

**Transmission electron microscopy**

Fig. 1 shows transmission electron micrographs of Ni–ZrO$_2$, Cu–ZrO$_2$ and 11CuNi–ZrO$_2$ after ex situ reduction.

The size of the ZrO$_2$ particles was between 50 and 100 nm. In general the metal particles were found to be evenly distributed on the oxide support. The Ni particles in Ni–ZrO$_2$ exhibit a homogeneous size distribution of approximately 20 nm mean size and are of polyhedral shape. The Cu particles in Cu–ZrO$_2$ are spherical and about 5 nm in size. In contrast, the metal particles on the bimetallic 11CuNi–ZrO$_2$ with average sizes of around 20 nm are polycrystalline and rather irregular in shape. EDX spectra showed that those particles contained both Cu and Ni, although not in the expected 1:1 ratio, but they were enriched in nickel. To get a better understanding of the Cu and Ni distribution on the bimetallic catalyst, elemental maps were recorded. The results are displayed in Fig. 2.

While Ni was always accompanied by the simultaneous presence of Cu as demonstrated with full arrows in Fig. 2, Cu was additionally found finely distributed in regions where no or hardly any nickel was detected (dashed arrows in Fig. 2). Thus, the elemental maps revealed the presence of Ni-rich bimetallic particles as well as monometallic Cu particles. XANES spectra of the same catalyst during reduction confirmed the presence of bimetallic CuNi particles.

The observation that the presence of copper improves the reducibility of nickel and lowers its reduction temperature is in agreement with TPR experiments reported in the literature.

**Temperature-programmed methane decomposition**

The reactivity of the catalysts for decomposition of methane was investigated by TPMD measurements. The results obtained over Ni–ZrO$_2$ are shown in Fig. 3a).

The production of hydrogen started at around 625 K, increased further with increasing temperature and decreased again reversibly when the sample was cooled down to 573 K. Upon heating again to 773 K, the hydrogen production rate remained approximately the same as in the first cycle. Thus, no deactivation of H$_2$ production activity was observed. Besides H$_2$ formation, carbon dioxide evolution occurred during the first heating cycle in methane up to ~700 K. In the second and third heating cycles a small CO$_2$ desorption peak was observed with a maximum at 630 K. The detected CO$_2$...
likely originates from the reaction of methane with surface oxygen and carbonates of zirconia or with NiO present due to incomplete reduction.

Fig. 3b) shows the rate of hydrogen production over 13CuNi–ZrO₂ under the same conditions. Over this sample no catalytic activity was found up to 735 K. At this temperature, the amount of produced hydrogen increased rapidly up to a level similar to that over Ni–ZrO₂. The onset of H₂ production was accompanied by a small amount of CO₂ evolving which occurred only in the first heating cycle. During the second and third reaction cycles about the same amount of hydrogen was produced above 690 K but decreased again during the isothermal period at 773 K. A small amount of CO₂ was also detected during the first heating cycle. Afterwards, Cu–ZrO₂ was completely inactive. The low initial activity can likely be attributed to the reaction of methane with surface oxygen and hydroxide species from zirconia or incompletely reduced Cu species and are not available anymore in the second and third reaction cycles. Often CO is reported as a by-product of the reaction between methane and oxygen species on the support or the incompletely reduced metal oxides.⁶,⁷ However, formation of various surface carbonate species was observed from CO on zirconia and all zirconia-supported catalysts by IR spectroscopy, which likely decompose to CO₂.

In order to find out whether hydrogen evolution occurs because of further reduction of the catalyst in methane, the measurement was repeated with the same sample but after reduction at 773 K instead of 673 K. The sudden onset in hydrogen formation accompanied by carbon dioxide evolution was also observed, but the onset temperature was lowered to 650 K.

For quantitative comparison of the catalytic activity the amount of produced hydrogen was determined by GC-TCD.

Table 2 shows the hydrogen yields and reaction rates for hydrogen production during steady-state methane decomposition at 773 K.

The initial reaction rate \( r \) calculated per gram catalyst decreased with decreasing overall Ni content. However, the rate \( R \) normalized per gram Ni is of about the same order of magnitude for Ni and all bimetallic samples containing 5 wt% total loading. In agreement with TPD measurements, similar rates of hydrogen formation were detected on Ni-rich 13CuNi–ZrO₂ compared to those on Ni–ZrO₂. Cu is almost inactive and the reaction rate decreases fast to zero with time on-stream while it stays constant for the other catalysts for at least 1 hour. The rate \( r \) of the catalyst containing 50 wt% metals, 13CuNi–ZrO₂50, is about the same as that of 13CuNi–ZrO₂ with 5 wt% metal loading, which results in a reaction rate \( R \) per gram nickel of one order of magnitude lower. This can probably be explained by a lower Ni dispersion at such high loading. In general, we did not observe an enhancement of the catalytic activity by addition of copper to the Ni–ZrO₂ catalyst.

**Temperature-programmed oxidation**

To obtain insights into coke formation, temperature-programmed oxidation (TPO) was performed after reaction in methane. The amount of carbon dioxide formed during TPO provides information about the amount of coke formed during the reaction, and the temperature, which is needed to burn off the carbon species, may be characteristic of the carbon bond strength to the catalyst’s surface. Fig. 4 shows the production of carbon dioxide observed during TPO in 20% O₂ in Ar performed directly after methane decomposition (described in the previous section). Over Ni–ZrO₂, the highest amount of coke was formed and temperatures of 770 K were
needed to oxidize most of it. On the nickel-rich bimetallic catalyst 13CuNi–ZrO₂ the amount of formed CO₂ was lower by about 50% compared to that on Ni–ZrO₂. Furthermore, much lower temperatures of around 700 K were required to remove carbon by oxidation. At about 750 K a second smaller CO₂ evolution peak was observed in the TPO. Over 11CuNi–ZrO₂ the amount of coke was reduced to 30% compared to that over Ni–ZrO₂, and the maximum of CO₂ formation was detected at 650 K. Over the Cu-rich catalyst 31CuNi–ZrO₂ the amount of carbon was reduced to only 20% compared to that over Ni–ZrO₂. Here, one needs to keep in mind the lower activity of the Cu-rich compositions compared to that of Ni–ZrO₂. The maximum of the CO₂ formation rate was observed at 660 K. When considering the differences in H₂ production rate (Table 2) and the CO₂ peak areas observed in the TPO measurements, the addition of Cu lowered the amount of coke formed. A major effect of Cu was, however, the down-shift of the carbon oxidation temperature. Since hardly any methane decomposition reaction took place over the Cu–ZrO₂ reference material, no coke was burnt off during TPO over this catalyst.

The TPO curves show large CO₂ desorption peaks in a narrow temperature range as well as broader peaks of CO₂ formation over a broad temperature range. The narrow peaks were always accompanied by a small peak of water (m/z 18) which indicates oxidation of CH₄ species in this temperature range. The much broader peak underneath might originate from other carbonaceous species such as filamentous carbon.

The presence of CH₄ species indicates that part of the adsorbed methane dissociated incompletely.

In summary, the highest C and CH₄ oxidation temperature was observed on monometallic nickel at 750–770 K. Since the oxidation temperature decreased on the bimetallic catalysts we conclude that C species are less strongly bonded to the bimetallic surface. A similar trend with the composition was noticed as for the activity onset temperature described in the section about TPd. The highest temperature required to burn off coke was observed for 13CuNi–ZrO₂ with a CO₂ evolution maximum at 700 K, while 11CuNi–ZrO₂ exhibited the lowest carbon oxidation temperature with a CO₂ evolution maximum at 650 K.

**In situ X-ray photoelectron spectroscopy**

In order to learn more about the surface composition, oxidation states and carbonaceous species present under the reaction atmosphere, *in situ* XPS at millibar pressures was employed. The bimetallic Ni-rich 1 : 3 composition which showed the most pronounced increase in methane decomposition activity was investigated further by synchrotron-based *in situ* XPS aiming at understanding the nature of the irreversible surface modification process. To avoid further complication of the (already complex) spectra by charging effects, a catalyst with a higher overall metal loading of 50 wt% was utilized after carefully checking the comparability of this material to the respective 5 wt% sample.

**XP spectra during reduction in H₂**

Fig. 5 shows the Cu 2p₃/₂ binding energy region of 13CuNi–ZrO₂₅₀ during a) oxidation and b) reduction. The peak positions and full width at half-maximum (FWHM) are listed in Table 3.

The broad peak at 933.3 eV is attributed to CuO⁴⁸–⁵³ with two characteristic shake-up satellite peaks at 940.7 and 943.3 eV. The structure seen in the satellite line is due to the multiplet splitting in the 2p 3d final state.⁵⁰ The small peak located at 932.3 eV is characteristic of Cu₂O and/or metallic Cu.⁴⁸,⁵⁰,⁵²,⁵¹ During reduction at 673 K the CuO peak decreased in intensity while the signal of reduced Cu species strongly increased. Since the shift of alloyed Cu in CuNi has been reported to be only about 0.2 eV compared to that in monometallic Cu⁴⁴–⁵⁶ we could not resolve this species.

Fig. 6 shows the Ni 2p₃/₂ core levels of 13CuNi–ZrO₂₅₀ measured during a) oxidation and b) reduction. The peak at

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**Table 2** Hydrogen yields and initial reaction rates during methane decomposition at 773 K in 5% CH₄ and 95% Ar with a total flow of 50 ml min⁻¹

<table>
<thead>
<tr>
<th>Sample</th>
<th>H₂ yield</th>
<th>Reaction rate r per g catalyst (mol H₂/(g catalyst s⁻¹))</th>
<th>Reaction rate R per g nickel (mol H₂/(g nickel s⁻¹))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni–ZrO₂</td>
<td>63.3%</td>
<td>2.4 × 10⁻³</td>
<td>4.8 × 10⁻⁴</td>
</tr>
<tr>
<td>13CuNi–ZrO₂</td>
<td>49.8%</td>
<td>1.9 × 10⁻⁵</td>
<td>5.0 × 10⁻⁴</td>
</tr>
<tr>
<td>11CuNi–ZrO₂</td>
<td>16.9%</td>
<td>6.4 × 10⁻⁶</td>
<td>2.6 × 10⁻⁴</td>
</tr>
<tr>
<td>31CuNi–ZrO₂</td>
<td>13.6%</td>
<td>5.1 × 10⁻⁶</td>
<td>4.1 × 10⁻⁴</td>
</tr>
<tr>
<td>Cu–ZrO₂</td>
<td>1.5%</td>
<td>5.6 × 10⁻⁷</td>
<td>—</td>
</tr>
<tr>
<td>13CuNi–ZrO₂₅₀</td>
<td>40.0%</td>
<td>1.5 × 10⁻⁵</td>
<td>3.9 × 10⁻⁵</td>
</tr>
</tbody>
</table>
854.3 eV is attributed to the main line of NiO.23,53,57 In a reducing atmosphere an additional peak appeared at 852.6 eV, which is ascribed to metallic nickel.53,60,61 Again, it is not possible to distinguish between nickel interacting with copper and non-interacting with copper due to the small chemical shift of about 0.2 eV54–56 which would be expected for Ni alloyed with Cu. In agreement with previously reported findings by IR spectroscopy of CO adsorption on reduced
CuNi both oxidized and reduced Ni species are present at the surface after reduction at 673 K, although the bulk consists predominantly of reduced Ni species according to XANES measurements.

Due to the high complexity of the NiO spectrum consisting of shake-up peaks and multiplet states, we fitted Ni 2p\textsubscript{3/2} spectra by the linear combination of reference spectra of NiO and completely reduced Ni. The resulting Ni-to-NiO ratios are summarized in Table 4.

Overall, the XP spectra reveal the coexistence of reduced Cu and a mixture of oxidized and metallic Ni species in the near-surface region after reduction.

### XP spectra during methane decomposition

Fig. 7 shows a) the Cu 2p\textsubscript{3/2} region and b) the Ni 2p\textsubscript{3/2} region of 13CuNi–ZrO\textsubscript{2} during exposure to 0.25 mbar methane at temperatures from 523 to 723 K directly after reduction in H\textsubscript{2}.

At 523 K mainly metallic copper and small amounts of CuO were present in the Cu 2p\textsubscript{3/2} region. By increasing the temperature the metallic Cu peak became broader and shifted to a slightly higher binding energy indicating further alloy formation between copper and nickel. The amount of nickel oxide decreased with temperature, while the signal of metallic nickel increased. At 723 K a new Cu species at 933.6 eV appeared at the highest temperature. A potential assignment of this species is Cu interacting with surface carbonates; however, a larger shift to higher binding energies would be expected compared to CuO. CuO core levels have been observed over a broad binding energy range, but in the XP spectrum at 723 K no satellite peaks characteristic of CuO are found, which rather excludes the assignment of the peak at 933.6 eV to CuO. Currently, the nature of this peak remains an open question.

Between 698 and 723 K the biggest changes were observed in the Ni 2p\textsubscript{3/2} region as well. The amount of NiO decreased significantly, while the concentration of metallic nickel increased in the near-surface region. Note that in the temperature range between 698 and 723 K where the strongest spectral changes occurred the catalytic activity of this catalyst increases dramatically, as shown by TPMD and in agreement with the simultaneously recorded mass spectrometry data.

Fig. 8 shows the C 1s core level region during exposure of 13CuNi–ZrO\textsubscript{2} to methane at temperatures increasing from 523 to 723 K. A number of different carbon-containing surface species were present in methane. Polycarbonates, carbonates and carbon oxygenates above 287 eV hardly changed with temperature. Various carbonate species were also observed by infrared spectroscopy on ZrO\textsubscript{2}. C–O\textsuperscript{63–66} or C–OH\textsuperscript{64,65} at 286.3 eV increased with temperature, indicating an interaction of carbon with the oxide support or NiO and CuO which had not been reduced up to that temperature.

### Table 4 Proportion of NiO and Ni in Ni 2p\textsubscript{3/2} spectra fitted by linear combination of reference spectra

<table>
<thead>
<tr>
<th>Atmosphere and temperature (K)</th>
<th>NiO (%)</th>
<th>Ni (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>O\textsubscript{2} – 573</td>
<td>100%</td>
<td></td>
</tr>
<tr>
<td>H\textsubscript{2} – 573</td>
<td>77 ± 3%</td>
<td>23 ± 3%</td>
</tr>
<tr>
<td>CH\textsubscript{4} – 523</td>
<td>79 ± 3%</td>
<td>21 ± 3%</td>
</tr>
<tr>
<td>CH\textsubscript{4} – 573</td>
<td>80 ± 2%</td>
<td>20 ± 2%</td>
</tr>
<tr>
<td>CH\textsubscript{4} – 623</td>
<td>79 ± 3%</td>
<td>21 ± 3%</td>
</tr>
<tr>
<td>CH\textsubscript{4} – 673</td>
<td>77 ± 3%</td>
<td>23 ± 3%</td>
</tr>
<tr>
<td>CH\textsubscript{4} – 697</td>
<td>75 ± 3%</td>
<td>25 ± 3%</td>
</tr>
<tr>
<td>CH\textsubscript{4} – 723</td>
<td>66 ± 3%</td>
<td>34 ± 3%</td>
</tr>
</tbody>
</table>

**Fig. 7** a) Cu 2p\textsubscript{3/2} and b) Ni 2p\textsubscript{3/2} core levels regions of 13CuNi–ZrO\textsubscript{2} in 0.25 mbar CH\textsubscript{4} at rising temperature from 523 to 723 K. The incident photon energies were 1100 eV (Cu) and 1010 eV (Ni).
The amount of graphitic carbon at 284.3 eV binding energy increased until 673 K but then decreased again at 723 K. The peak at 284.8 eV attributed to C having sp² hybridization indicates the formation of carbon nanotubes (CNTs) and increased with increasing temperature. The signal at 285.4 eV corresponds to CₓHᵧ species. These species increased upon heating as well. Alternatively, the signal could indicate the presence of structural defects on carbon nanotubes. NiₓC₆⁴,₆₅ appeared at 283.4 eV and increased at first with rising temperature but then completely vanished at 723 K.

In summary, the amount of several carbon species increased during methane decomposition. In particular, the growth of CNTs is significantly enhanced at the highest temperature at which catalytic activity sets in at the same time, while graphitic carbon decreases, either by transformation to CNTs or dissolution in the bulk of the particles.

Depth profiling

To get more insights into the distribution of Ni, Cu and C species in the surface and subsurface region XP spectra were recorded using different photon energies. The photon energies were varied to obtain photoelectron kinetic energies of 150 eV, 350 eV and 550 eV for each set of spectra of the relevant core level regions (Ni 2p, Cu 2p, C 1s). These depth profiles were performed at 523, 673 and 723 K in methane.

Fig. 9 visualizes the relative core level peak areas of the various Cu, Ni and C species based on the respective total areas of Cu, Ni and C signals. Upon heating from 523 to 673 K,
the relative amount of metallic Ni increases mainly in the deeper (near-surface) layers. After further heating from 673 to 723 K an increased amount of reduced Ni is present on the surface as well. The amount of metallic Cu increases with temperature. At 723 K the new Cu species appeared which cannot be identified yet (as discussed in the results section). Interestingly, this species is related to the surface of the catalyst and appears only when using photon energies yielding photoelectrons of 150 eV kinetic energies, i.e. when measuring with the highest surface sensitivity.

In general, the total amount of carbon species decreases with increasing kinetic energy, indicating that the carbon species are present mostly at the surface of the catalyst. As shown in Fig. 9c), the relative proportion of CNTs increases significantly at 723 K, in particular in the surface-sensitive measurement, as shown by the strongly decreasing amount of sp\(^2\) hybridized carbon with increasing kinetic energy. Nickel carbide disappeared at 723 K in the most surface-sensitive spectra recorded at 150 eV kinetic energy, while a small amount of carbide is still present in the somewhat deeper near-surface layers. The relative amount of graphitic carbon is highest at 673 K reaction temperature and then decreases considerably at 723 K. This is observed for all photoelectron kinetic energies that were analyzed. The proportion of carbon present as carbonates, carbon-oxygenates and hydrocarbon species does not depend much on reaction temperature and photon energy.

**Discussion**

CH\(_4\) decomposition was studied on bimetallic CuNi–ZrO\(_2\) catalysts by temperature-programmed reaction, TPO and NAP-XPS in order to get insights into the in situ stability of CuNi compounds and the effect of Cu addition on the carbon chemistry. In contrast to monometallic Ni–ZrO\(_2\), an irreversible activation step was observed in methane at around 700 K, with the exact onset temperature depending on the composition, reduction temperature and metal loading. In the following we will discuss the potential origin and the factors affecting this surface modification process responsible for the onset of catalytic activity and take a closer look at the bimetallic Cu–Ni system.

The question arises as to what is the origin of the observed activation process occurring in methane between 650 and 735 K and leading to an irreversible modification of the catalytic performance of all bimetallic catalysts towards a Ni-like behaviour. Previous work on these materials allowed us to conclude that after reduction the surface of the bimetallic catalysts appears enriched in Cu\(^{25}\) as indicated by selective H\(_2\) chemisorption on Ni and by FTIR spectroscopy of adsorbed CO. Since Cu is practically inactive in methane decomposition a change in surface composition, e.g. by Ni segregating to the surface, is a potential reason for the drastic change in CH\(_4\) decomposition activity. The interaction of Ni with C and/or CH\(_4\) would be a likely driving force for the proposed surface segregation of Ni.

In contrast to H\(_2\) chemisorption experiments at room temperature\(^{25}\) showing that the surface is enriched in copper after reduction, the surface Cu : Ni ratio observed by XPS at 673 K was nearly similar to the overall Cu : Ni ratio of the catalyst. According to the literature, a copper-enriched top surface layer is in agreement with the lower surface free energy of copper compared to that of nickel.\(^{72}\) According to Sachtler et al., the Gibbs free energy of a copper–nickel mixture at 475 K has a maximum at about 30% copper and a minimum at 80% and at 2% copper\(^{73}\) in the bulk. Therefore, we expect coexisting phases to be present at lower temperatures. Since the surface energy is lower for Cu compared to that for Ni the Cu-rich alloy phase is likely present at the surface and the Ni-rich alloy phase underneath (core–shell-like structure). For Cu surfaces with Ni monolayers deposited on top it was reported in ref. 74 and 75 that Ni segregates into the first subsurface layer of Cu(100) as well as Cu(111).\(^{76}\) By theoretical calculations, Pourovskii et al.\(^{77}\) found a rather sharp minimum of the surface energy corresponding to the arrangement of a Ni layer in the second layer of Cu(100).

Anyway, this surface composition is only present at low temperature. At elevated temperature the two phases become miscible. According to the Cu–Ni phase diagram, CuNi forms a single-phase alloy above 600 K.

As revealed by XPS measurements under reaction conditions, the increase in activity is clearly accompanied by an increase in the concentration of metallic Ni at the surface. Catalytic activity for methane decomposition therefore depends on the alloy surface composition, specifically on the presence of a sufficiently high concentration of metallic Ni at the surface.

Studt et al.\(^{78}\) reported Ni segregation to the surface of CuNi alloys due to the higher bond strength towards CO. Li et al.\(^{79}\) reported that the dissociative adsorption of methane and the formation of carbon flakes would be a plausible cause for their observation of reconstruction of Ni–Cu metal particles during methane decomposition reaction.\(^{79}\) In a remarkable study, Li et al. observed a recrystallization of the Ni–Cu alloy on copper–nickel–alumina catalysts during induction of carbon formation in methane decomposition at 773 K.\(^7\) In a similar way, Ni could be segregated to the very surface layer at elevated temperatures induced by the presence of methane and/or carbon, which consequently leads to an onset of catalytic activity for methane decomposition. Thus, the surface initially enriched in Cu at low temperatures exhibits low methane decomposition activity until the activation step changes the surface concentration of reduced Ni.

Ni segregation to the top surface is a potential explanation for the observed activation step. However, this could not be confirmed by quantitative analysis of the core level spectra, which is complicated by the (very likely) preferential decoration of Ni by carbon species formed over Ni during the reaction of methane. Nevertheless, there is clear evidence by the XPS data that the sudden increase of reactivity can be linked to the higher concentration of reduced Ni at the surface.
When TPMd was carried out after applying a higher reduction temperature of 773 K an activation step was observed as well, although the activation step temperature was lower compared to that for the catalyst reduced at 673 K. Since reduction at 673 K leads to only partially reduced Ni, the Ni : NiO ratio is increased after reduction at higher temperature. The higher amount of metallic Ni (at or near the surface) after applying a higher reduction temperature could be a likely reason for the down-shift of the activation temperature.

Cu addition to Ni helps in the desired lowering of carbon deposition. TPO measurements showed a reduction of the amount of coke burnt off as CO2 by about 50% when comparing Ni and the Ni-rich 13CuNi-ZrO2 catalyst exhibiting similar catalytic activity (Table 2). Another effect of the Cu modification, which influences the carbon chemistry and coking behavior, is the decrease in the carbon oxidation temperature, probably due to a weaker bonding of carbon on the CuNi alloy surface.

However, the CuNi system is stable only in a limited range of conditions (temperatures up to around 700 K depending on the composition and pretreatment). Therefore, a key result of this work is the detection of the instability of CuNi under hydrocarbon-rich conditions at higher temperatures (> ~700 K).

Conclusions

In this work we have investigated the surface chemistry of methane on Ni–ZrO2 and bimetallic CuNi–ZrO2 catalysts with a focus on the \emph{in situ} stability under relevant reaction conditions, \emph{i.e.} high temperatures and reactive surrounding gas atmosphere methane.

The activity of bimetallic CuNi–ZrO2 catalysts for methane decomposition strongly increased after an irreversible activation step occurring at different temperatures depending on the Cu : Ni ratio. By \emph{in situ} synchrotron-based near-ambient pressure (NAP)-XPS we determined the surface composition of the catalysts and the nature of the carbonaceous species present and correlated these data with the catalytic performance. The sudden increase in catalytic activity could be explained by an increase in the concentration of reduced Ni atoms at the catalyst surface in the active state, likely as a consequence of the interaction with methane. XPS showed the appearance of an additional Cu species on the catalyst surface, when catalytic activity sets in. This species could not be conclusively identified up to now, but is related to the catalyst surface according to the depth profiles recorded.

C 1s core level spectra revealed the presence of various carbonaceous species. The signal attributed to carbon nanotubes increased when activity sets in, while the amount of graphite and carbide decreased again. Cu addition to Ni improved the desired resistance against carbon deposition by lowering the amount of coke formed. A lower temperature was required to burn off carbon from the bimetallic catalysts compared to Ni–ZrO2.

As a key result on the Cu-modification of Ni catalysts, we conclude that the CuNi alloy shows limited stability under relevant reaction conditions. This system is stable only in a limited range of temperature up to ~700 K in a hydrocarbon atmosphere. Beyond this temperature, the alloy phase becomes unstable, and carbon-induced segregation of Ni species induces a fast increase in methane decomposition rate. In view of the applicability of this system, a detailed understanding of the stability and surface composition of the bimetallic phases present and the influence of the Cu promoter on the surface chemistry under actual reaction conditions are essential.

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Notes and references