Effects of dealumination on the performance of Ni-containing BEA catalysts in bioethanol steam reforming

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The effects of dealumination of BEA zeolite on the formation of nickel active sites and the performance of Ni-containing BEA zeolite catalysts in the steam reforming of ethanol have been studied. Ni-containing BEA zeolite catalysts were prepared by the impregnation of unmodified and dealuminated BEA zeolites with Ni(NO3)2 precursor. The properties of Ni10HmBEA and Ni10SiBEA zeolite catalysts were studied by means of X-ray diffraction, 1H, 27Al and 29Si magic-angle spinning nuclear magnetic resonance, Fourier-transform infrared and Raman spectroscopy, transmission electron microscopy, temperature-programmed reduction, temperature-programmed ammonia and hydrogen desorption methods. High initial activity and selectivity of Ni10HmBEA to hydrogen and carbon dioxide with unmodified BEA zeolite support in the steam reforming of ethanol reaction performed at 500 °C was observed. However, fast deactivation of Ni10HmBEA catalyst, manifested in the decrease of water conversion, drop of selectivity to H2 and CO2, and increase in the selectivity to ethylene with the time-on-stream, was observed. In contrast, Ni10SiBEA zeolite catalyst showed lower initial activity but higher durability and resistance for carbon deposition. It was stated that dealumination of BEA zeolite led to the slight structural changes and simultaneously pronounced decrease of acidity. Formation of the large nickel crystallites was hindered on Ni10SiBEA zeolite catalyst. TEM and Raman spectroscopy studies indicated that deactivation of Ni10HmBEA was related to formation of nickel mediated filamentous, graphitic and amorphous carbon deposits. Much smaller amounts of filamentous carbons were observed on the Ni10SiBEA zeolite catalyst prepared by the use of dealuminated zeolite support.

1. Introduction

Bioethanol has been commonly used as solvent, valuable chemical and energy carrier. It is currently produced mainly from sugar and starch crops, however, it seems that bioethanol can be also commercially synthesized in the near future from wood residues, wastes or energy crops [1–4]. This may increase the potential applications of bioethanol for production of biofuels, biopolymers, and energy carriers, such as hydrogen which can be subsequently utilised for generation of electricity in the fuel cell systems. Ethanol, as a liquid fuel can be easily transported and stored, therefore directly used as hydrogen carrier for stationary and mobile units. One of the most attractive ways of the production of hydrogen from renewable sources is steam reforming of ethanol (ESR): \( \text{C}_2\text{H}_5\text{OH} + 3\text{H}_2\text{O} \rightarrow 6\text{H}_2 + 2\text{CO}_2 \), \( \Delta H_{\text{REA}}^0 = 173.2 \text{ kJ mol}^{-1} \). Reaction can be carried out in the presence of supported noble and transition metal catalysts, including rhodium, nickel and cobalt in the range of 300–500 °C [5–11]. Bio-CO2 - the product of the steam reforming of bioethanol can be easily removed from the product stream or even supplied together with hydrogen to the fuel cells. Depending on the properties of catalysts and reaction conditions, additional side products of ESR are often observed in the product stream, including carbon monoxide, acetaldehyde, acetone, acetic acid, methane, ethane, ethylene and higher hydrocarbons [5–10]. The catalysts may deactivate with the time-on-stream due to the sintering and formation of carbon deposits. Several routes have been proposed to improve the activity and durability of catalysts. Large-surface area supports, such as γ-Al2O3, silica mesoporous materials, mixed oxides, e.g. CeZr1-xO2, usually allow for the formation of strongly dispersed surface precursors and retard sintering processes. Similar effect can be achieved by the introduction of certain modifiers or promoters, which change metal-support interactions. However, large number of acid or redox sites presented on the surface of supports may intensify dehydration reaction.
with formation of ethylene or condensation/ketonization reactions, leading to the production of aceton and acetic acid. Ethylene is commonly regarded as coke precursor. Acetone can be also transformed to coke through mesityl oxide formation and further aldol condensation [12-15]. On the other hand, an introduction of redox components into the catalysts or modification of the acidity by some means, may improve coking resistance. It is commonly accepted opinion that ethanol molecules in the steam reforming reaction are dissociatively adsorbed on the surface of catalysts with formation of ethoxy groups. Such species can be subsequently transformed into various intermediates by successive dehydrogenation and dehydration reactions, C-C bond scission, oxidative surface reactions with contribution of hydroxyl groups or oxygen species in the catalysts. The intensification of surface reactions between carbonaceous surface intermediates - coke precursors with surface oxygen species or hydroxyl groups, bounded to the acid sites may retard fast development of filamentous or encapsulating deposits.

In spite of vast literature devoted to the ESR reaction in the last years, zeolite supported catalysts have not been widely studied. The micro-mesoporous structure of zeolites may promote formation of the small metal nanoparticles [16-21]. On the other hand, high acidity of zeolites may drive to the extensive formation of carbon deposits in the ESR reaction. Several methods of changing acid-base properties of zeolite supports have been proposed, mainly by the introduction of alkali metals or alkali treatment [17,22,23]. Dealumination of zeolites has been described in the literature as the useful method of modification of structural, as well as acid-base properties [23-29]. The advantage of the use of dealuminated ITQ-2 zeolites as supports of nickel and cobalt catalysts for the ESR reaction has been recently pointed out by Chica et al. [28,29].

The removal of aluminum ions from the zeolite framework may not only drive to the changes of acidity, but also facilitate formation of peculiar adsorptive sites for anchoring metal precursors and stabilization of metallic nanoparticles. The aim of our studies was determination of the effects caused by the dealumination of BEA zeolite supports on the nickel dispersion, structural and surface properties of catalysts, catalytic performance in the steam reforming of ethanol and the nature of carbon deposits.

2. Experimental

2.1. Catalysts preparation

\[ \text{Ni}_{10}\text{HAlBEA} \] and \[ \text{Ni}_{10}\text{SiBEA} \] zeolite catalysts were prepared by conventional wet impregnation and a two-steps postsynthesis procedure, as reported earlier [30]. A tetraethylammonium BEA (TEABEA) zeolite provided by RIPP (China) was divided for two parts. The first fraction was calcined in air at 550 °C for 15 h in order to obtain an organic - free HAlBEA zeolite (Si/Al = 17). The Ni_{10}HAlBEA zeolite catalyst was prepared by impregnation of HAlBEA with an aqueous solution of Ni(NO_3)_2·6H_2O (pH = 2.6) under aerobic conditions, as described earlier [30]. Then the suspension was stirred for 2 h at 80 °C until water evaporation. The resulting solid was dried in air at 80 °C for 24 h and then calcined in air at 400 °C for 2 h.

The second fraction of TEABEA was treated in a 13 mol dm⁻³ HNO_3 aqueous solution at 80 °C for 4 h to obtain a dealuminated and organic - free SiBEA support (Si/Al = 1000) with vacant T-atom sites (T = Al). SiBEA was then recovered by centrifugation, washed with distilled water and dried at 80 °C. To incorporate nickel ions in vacant T-atom sites, 2 g of SiBEA was stirred under aerobic conditions for 24 h at 25 °C in 200 mL of Ni(NO_3)_2·6H_2O aqueous solution (pH = 2.6). The resulting solid was dried in air at 80 °C for 24 h. Then solid was calcined in air at 400 °C for 2 h and labelled as Ni_{10}SiBEA.

2.2. Materials characterization

X-ray diffraction studies (XRD) of materials after calcination and after reduction (prior to the ESR reaction) were performed with Empyrean (PAAnalytical) diffractometer using Cu-Kα radiation (0 = 1.5418 Å). Rietveld method was used to estimate mean crystallite size of nickel oxide (d_{NiO}) and metallic nickel (d_{Ni}^{0}). The samples prior to the determination of d_{Ni}^{0} were reduced in the microreactor in the flow of hydrogen with the ramp rate of 10 °C min⁻¹ up to 600 °C followed by the isothermal reduction for 2 h. After cooling down to the room temperature, the samples were passivated in the flow of the mixture of 5% O_2 in He.

Nickel content in the catalysts was determined by the application of X-ray fluorescence method using ED-XRF Canberra Packard 1510 spectrometer.

The porous structure and specific surface area of zeolites and modified with nickel after calcination at 400 °C were determined from the measurements of nitrogen adsorption/desorption isotherms. The isotherms were obtained volumetrically at −196 °C using ASAP 2405 N analyzer (Micromeritics Instrument Corp.). The samples were outgassed under ~10⁻² Pa at 200 °C. The standard Brunauer-Emmett-Teller (BET) method was used to calculate specific surface area (S_{BET}). The total pore volume (V_p) was estimated from single point adsorption at p/p_0 = 0.98. Mean pore diameter (D_{p,4,3}) and pore size distribution were determined by applying the Barret-Joyner-Halenda (BJH) method from the desorption branch of isotherms [31,32].

29Si MAS NMR spectra were recorded at 99.3 MHz with Bruker AVANCE500 spectrometer at 11.7 T and 7 mm (external diameter) zirconia rotors. Chemical shifts of silicon were measured by reference to tetramethysilane (TMS). 29Si MAS NMR spectra were obtained with 5 kHz rotors spinning speed, 3 μs excitation pulse duration and 10 s recycle delay. 27Al and 1H MAS NMR experiments were performed on a Bruker AVANCE500 spectrometer at 11.7 T in 4 mm zirconia rotors spinning at 12 kHz. The resonance frequency of 1H and 27Al were 500.16 MHz and 130.33 MHz, respectively. Chemical shifts, δ, were reported relative to TMS and aqueous Al(NO_3)_3·11 H_2O solution. 1H MAS NMR spectra are performed with a 90° pulse duration of 2.6 μs and a recycle delay of 5 s and 16 accumulations. 27Al MAS NMR spectra were recorded by small-flip-angle technique with a pulse of 0.9 μs (π/8), 0.5 s for the recycle delay and 2048 accumulations.

The FTIR/PAS spectra were recorded by means of a Nicolet 8700 (Thermo Scientific) spectrometer equipped with PA301 Gasera photo-acoustic detector in the range of 400–4000 cm⁻¹ with the resolution of 1 cm⁻¹ and maximum source aperture. A standard carbon black was used as background. A sample was placed to the holder and inserted into the analyzer purged with helium. Interferograms of 1024 scans were averaged for each spectrum.

Acidity of the supports and catalysts was determined by the temperature-programmed desorption of ammonia using Altamira AMI-1 system coupled with mass spectrometer HAL 201RC (Hiden Analytical). The intensities of selected ions were monitored. Samples (0.1 g) were introduced into the plug flow microreactor and were reduced at 600 °C for 2 h. Samples after cooling down were flushed with helium at 100 °C for 0.5 h, next flushed with the mixture of 0.5% NH_3 in He for 0.5 h, and again with He for 0.5 h. The flow rate of gases was equal to 30 mL min⁻¹. Next the temperature was increased with the ramp rate 10 °C min⁻¹ to 600 °C.

Additionally the surface properties were investigated by the application of Diffuse Reflectance Infrared Fourier-Transform Spectroscopy (DRIFTS). DRIFTS spectra were recorded using FTIR spectrometer (Nicolet 6700, Thermo Scientific) equipped with LN_2 cooled MCT/A detector. Samples were introduced into the Praying Mantis High-Temperature Reaction Chamber (Harrick), which was covered by a dome with ZnSe windows. The samples were heated up in the flow of H_2 (30 mL min⁻¹) from 20 to 600 °C with the ramp rate 10 °C min⁻¹, and after 2 h of isothermal heating were cooled down to 100 °C. DRIFTS...
spectra were collected at selected temperatures with the resolution of 0.48 cm$^{-1}$ and maximum source aperture. Interferograms of 256 scans were averaged for each spectrum. The spectra were corrected by applying the background signal recorded for KBr powder.

Analysis of the acidic properties of samples was also performed by adsorption of pyridine (Py) followed by infrared spectroscopy. For analysis, the samples were pressed at ~1 ton cm$^{-2}$ into thin wafers of ca. 10 mg cm$^{-2}$ and placed inside the IR cell. Before Py adsorption/desorption experiments, the wafers were activated by calcination in static conditions at 450 °C for 3 h in O$_2$ (1.6 × 10$^3$ Pa) and then outgassing under secondary vacuum at 300 °C (10$^{-3}$ Pa) for 1 h. These wafers were contacted at room temperature with gaseous Py (133 Pa) via a separate cell containing liquid Py. The spectra were then recorded following desorption at 150 °C for 1 h with a Bruker Vector 22 spectrometer (resolution 2 cm$^{-1}$, 128 scans). The reported spectra were obtained after subtraction of the spectrum recorded before Py adsorption. The amount of Brønsted and Lewis acidic centers titrated by pyridine was obtained using a molar extinction coefficient value of $\varepsilon = 1.67$ cm mol$^{-1}$ cm$^{-1}$ for the $v_{19b}$ vibrations of protonated pyridine (Py-H$^+$) at ~1540 cm$^{-1}$, and of $\varepsilon = 2.22$ cm mol$^{-1}$ cm$^{-1}$ for the $v_{19b}$ vibrations of coordinated pyridine (Py-L) at ~1455 cm$^{-1}$ [33].

The nature of carbon deposits in the samples after ESR reaction tests performed at 500 °C/Ch 22 h was investigated by Raman spectroscopy. Raman spectra were recorded with the resolution of 2 cm$^{-1}$ using a Raman microscope (inVia Reflex, Renishaw) with the Raman dispersive system, using a semiconducting laser 785 nm with 0.5 mW of power to avoid sample overheating. Spectra were deconvoluted with the Lorentzian function available at the PeakFit SeaSolve software.

The catalysts for microscopic studies were grinded in an agate mortar to a fine powder. The resulting powders were poured with 99.8% ethanol (POCH) to form a slurry which subsequently were inserted into the ultrasonic homogenizer for 20 s. Then, the slurry containing the catalyst was pipetted and supported on a 200 mesh copper grid covered with lacy formvar and stabilized with carbon (Ted Pella Inc.) and left on the filter paper until the ethanol has evaporated. Subsequently, the samples deposited on the grid were inserted to holder and moved to electron microscope. The electron microscope Tecnai G2 20 X-TWIN FEI Company, equipped with an LaB$_6$ source, HAADF detector and EDS spectrometer (Energy Dispersive X-Ray Spectroscopy) was used to display the prepared catalysts. Microscopic studies of the catalysts were carried out at an accelerating voltage of the electron beam equal to 200 kV. The elements mapping was carried out in the STEM mode by collecting point by point EDS spectrum of each of the corresponding pixels in the map. The collected maps were presented in the form of a matrix of pixels with the colour mapped significant element and the intensity corresponding to the percentage of the element. The average size of nickel particles ($d_{Ni}^{STEM}$) was calculated from the equation:

$$d_{Ni}^{STEM} = \frac{\sum n_i d_i}{\sum n_i}$$

where: $n_i$ – the numbers of metal crystallites in a specific size range, $d_i$ – the average diameter in each diameter range.

Reducibility of nickel catalysts was investigated by the temperature-programmed reduction method (TPR) using Autochem II 2920 (Micromeritics Instrument Corp.), equipped with thermal conductivity detector (TCD). The samples (m = 0.05 g) were reduced in the mixture of 5% H$_2$ in Ar with the total flow rate of 30 mL min$^{-1}$ using 10 °C min$^{-1}$ ramp rate. The evolved water was removed from the stream in a cold trap with the mixture of liquid nitrogen and isopropyl alcohol at ~89 °C.

The nature of active surface sites was determined by hydrogen temperature-programmed desorption (TPD) method. Studies were carried out in the Autochem II 2920 (Micromeritics Instrument Corp.). The samples (0.1 g) were initially reduced at 600 °C for 2 h and subsequently were cooled down in the flow of hydrogen to ~60 °C. After flushing with Ar (30 mL min$^{-1}$) and stabilization of TCD signal, the temperature was increased with the ramp rate 10 °C min$^{-1}$. The evolved water was removed in a cold trap in a similar manner as in the TPR procedure. The active surface area of nickel catalysts ($S_{Ni}$) was determined by the integration of TPD curves, assuming chemisorption stoichiometry H:Ni = 1:1 and that the surface area occupied by one atom of hydrogen was equal to 0.0649 nm$^2$.

The activity and selectivity of zeolite supports and nickel-containing zeolite catalysts were determined in the steam reforming of ethanol reaction (ESR) under atmospheric pressure using Microactivity Reference unit (PID Eng & Tech.), equipped with fixed-bed continuous-flow quartz microreactor. The catalyst sample of the weight 0.1 g was diluted with quartz grains (0.3–0.6 mm) at the weight ratio of 1/25. The catalysts prior to the reaction were reduced in hydrogen at 600 °C for 2 h. The aqueous solution of ethanol with molar ratio of water/ethanol = 12/1 was supplied by a mass controller (Bronkhorst) to an evaporator maintained at 150 °C, and the reaction mixture, without diluting with any inert gas, was fed to the reactor at a flow rate of 100 mL min$^{-1}$. Weight hourly space velocity (WHSV$_{EthOH}$) was equal to 9.5 EthOH $h^{-1}$ EthOH$^{-1}$. The temperature was measured in the centre part of the catalytic bed. The tests were performed for 22 h at 500 °C. The catalysts after ESR reaction tests were cooled down to the room temperature under the flow of an inert gas. The analysis of the reaction mixture and the reaction products (all in the gas phase) was carried out on-line by means of two gas chromatographs, i.e. Bruker 450-GC and Bruker 450-GC. The first one was equipped with a TCD detector and two columns, filled with a porous polymer Porapak Q (for all organics, CO$_2$ and H$_2$O vapor) and capillary column CP-Molsieve 5 Å (for CH$_4$ and CO analysis) with helium as a carrier gas. The second chromatograph equipped with a Molsieve 5 Å column and TCD detector, argon as a carrier gas was used for determination of hydrogen concentration. The conversion of ethanol (X$_{EthOH}$), water (X$_{H_2O}$), and conversion of ethanol into particular carbon-containing products (X$_{EthOH}$) were calculated on the basis of their concentrations before and after the reaction, with a correction introduced for the change in gas volume during the reaction, from the following equations:

$$X_{EthOH} = \frac{C_{in}^{EthOH} - C_{out}^{EthOH} \times K}{C_{in}^{EthOH}} \times 100\%,$$

(2)

$$X_{H_2O} = \frac{C_{in}^{H_2O} - C_{out}^{H_2O} \times K}{C_{in}^{EthOH}} \times 100\%,$$

(3)

$$X_{EthOH} = \frac{C_{in}^{EthOH} - C_{out}^{EthOH} \times K}{(n/2)C_{EthOH}} \times 100\%,$$

(4)

where: $C_{in}^{EthOH}$ - is the molar concentration of ethanol in the reaction mixture (mol %), $C_{out}^{EthOH}$ - the molar concentration of ethanol in the post-reaction mixture (mol %), $C_{in}^{H_2O}$ - the molar concentration of water in the reaction mixture (mol %), $C_{out}^{H_2O}$ - the molar concentration of water in the post-reaction mixture (mol %), $C_{in}^{C}$ - the molar concentration of the carbon-containing products in the post-reaction mixture (mol %), n – the number of carbon atoms in the carbon-containing molecule of the reaction product, K – the volume contraction factor ($K = C_{CIP} / C_{in}^{EthOH}$), where $C_{CIP}$ and $C_{in}^{EthOH}$ are the molar concentrations of carbon in the ethanol feed to the reaction and in all carbon-containing compounds, which were present in post reaction gases, respectively). The selectivity of ethanol into individual carbon-containing products was expressed as:

$$S_{EthOH} = \frac{X_{EthOH}}{X_{EthOH} \times 100\%}.$$

(5)

The selectivity of hydrogen formation was determined from the equation:

$$S_{H_2} = \frac{C_{out}^{H_2}}{C_{in}^{EthOH} + 2C_{out}^{CO_2} + 2C_{out}^{CH_2CHO} + 2C_{out}^{CH_4} + 3C_{out}^{CO} + 4C_{out}^{C_2H_6}} \times 100\%.$$

(6)
3. Results and discussion

3.1. Phase composition

X-ray diffraction patterns of zeolite samples are presented in the Fig. 1a. Typical BEA zeolite structure can be inferred from the diffraction lines of HAIBE and SiBEA [34]. The shape of the most intensive peak (302) of BEA zeolites is similar in both samples. Thus it is difficult to find any differences of the crystallinity of samples. However, more detailed inspection of the XRD curves reveals, that most significant diffraction line, i.e. (302) located at $2\theta = 22.53^\circ$ for HAIBE zeolite is observed at $2\theta = 22.71^\circ$ for SiBEA sample (Fig. 1b). Similar shifts are visible for other diffraction lines, e.g. (304) line located at $2\theta = 25.38^\circ$ is shifted to $2\theta = 25.57^\circ$, (306) line positioned at $2\theta = 28.75^\circ$ moves to $2\theta = 29.10^\circ$. This points out the slight lattice contraction caused by the dealumination. It is interesting that positions of some diffraction lines are the same in both samples, e.g. (006) located at $2\theta = 27.12^\circ$.

Similar patterns are also observed for zeolites modified with nickel, both recorded for the samples after calcination and reduction. However, a minute shift of (302) diffraction line to $2\theta = 22.45^\circ$ may indicate slight expansion of HAIBE zeolite upon introduction of nickel and subsequent calcination of the sample. This effect may result from the incorporation of nickel ions into the lattice framework or location of nickel oxide nanoparticles in the zeolite channels. The covalent radius of $r(Ni^{2+}) = 128$ pm is larger than the covalent radius $r (Al^{3+}) = 121$ pm. However, well evidenced diffraction lines of NiO (PDF-4 No. 04-001-9373), visible for Ni10HAIBE zeolite simultaneously indicate the presence of relatively large nickel oxide crystallites. Mean size of NiO crystallites is equal to 22.3 nm (Table 1). Hence, one can assume that Ni10HAIBE zeolite may contain both Ni2+ species embedded within zeolite framework and large NiO nanoparticles surrounded by the zeolite material. Although the positions of zeolite lattice diffraction lines of SiBEA and Ni10SiBEA are almost the same, wider shape of the most significant peak at $2\theta = 22.71^\circ$ indicates a slight loss of zeolite crystallinity. Note that the size of NiO in Ni10SiBEA is equal to 4.1 nm, in spite of similar content of nickel in zeolites (9.86 and 9.57 wt. % in Ni10HAIBE and Ni10SiBEA, respectively) and it is much smaller than in the Ni10HAIBE catalyst (Table 1).

The crystallinity of Ni10HAIBE zeolite is not distorted by the reduction process. The position and shape of XRD peaks observed for reduced Ni10HAIBE are almost the same as for calcined one. The reduction of Ni10HAIBE zeolite at 600 °C leads to the formation of Ni° crystallites of the mean size of 30.2 nm (determined by the XRD method, Table 2). This process include transformation of large NiO nanoparticles and Ni(II) species located in the zeolite framework into the metallic nickel nuclei, followed by their agglomeration. The size of Ni° nanoparticles formed under the same reduction conditions in Ni10SiBEA is equal to 4.3 nm (Table 2). Simultaneously stronger broadening of BEA(302) peak indicates more evident changes of Ni10SiBEA zeolite crystallinity. The obtained results point out that removal of Al during dealumination process led to the formation of the specific sites which may strongly interact with nickel precursor and

![Fig. 1. X-ray diffraction curves of zeolites and nickel catalysts after calcination and reduction at 600 °C for 2 h; wide (a) and short (b) range of 2 theta.](image)

### Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>Ni content (wt. %)</th>
<th>$S_{BET}$ (m²/g)</th>
<th>$V_p$ (cm³/g)</th>
<th>$D_p$ (nm)</th>
<th>$d_{NiO}$ (nm)</th>
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<tr>
<td>HAIBE</td>
<td>–</td>
<td>540.5</td>
<td>0.53</td>
<td>15.1</td>
<td>–</td>
</tr>
<tr>
<td>SiBEA</td>
<td>–</td>
<td>421.4</td>
<td>0.51</td>
<td>18.0</td>
<td>–</td>
</tr>
<tr>
<td>Ni10HAIBE</td>
<td>9.86 ± 0.30</td>
<td>481.7</td>
<td>0.48</td>
<td>15.5</td>
<td>22.3</td>
</tr>
<tr>
<td>Ni10SiBEA</td>
<td>9.57 ± 0.30</td>
<td>406.8</td>
<td>0.42</td>
<td>16.2</td>
<td>4.1</td>
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</tbody>
</table>

### Table 2

<table>
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<tr>
<th>Sample</th>
<th>$d_{XRD}$ (nm)</th>
<th>$d_{TEM}$ (nm)</th>
<th>$S_a$ (m²/g)</th>
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<tr>
<td>Ni10HAIBE</td>
<td>30.2</td>
<td>22.5</td>
<td>1.43</td>
</tr>
<tr>
<td>Ni10SiBEA</td>
<td>4.3</td>
<td>7.3</td>
<td>6.02</td>
</tr>
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Fig. 2. a) TEM images of Ni10HAIBEA catalyst after reduction, b) STEM images of Ni10HAIBEA catalyst after reduction, c) TEM images of Ni10SiBEA catalyst after reduction, d) STEM-EDS images of Ni10SiBEA catalyst after reduction, e) nickel crystallite size distribution in catalysts.
hinder agglomeration of nickel species into the large nanoparticles during calcination and reduction.

3.2. Structural and surface properties

TEM images of Ni10HAlBEA and Ni10SiBEA zeolite catalysts, presented in the Fig. 2, show that nickel crystallites are dispersed on the regular individual zeolite support grains of the size from about 30 to 60 nm. Studies confirm general trends outlined in the previous section.

In the case of Ni10HAlBEA zeolite, both large (the size above 50 nm) and small (below 5 nm) nickel crystallites can be distinguished. The function of crystallite size distribution is relatively wide (Fig. 2e). The predominant contribution is observed for nickel crystallites in the range of 20–30 nm ($d_{20-30}^{NN} = 22.5$ nm). STEM images (Fig. 2b) indicate that large crystallites are located on the external part of HAlBEA zeolite grains. Detailed inspection of the TEM images of higher resolution (Fig. 2a) points out that some very small nickel crystallites can be located in the inner part of zeolite particles.

In contrast, Ni10SiBEA catalyst contains very small nickel crystallites of the predominant size 2–8 nm, $d_{2-8}^{NN} = 7.3$ nm (Fig. 2e). Nickel crystallites are uniformly distributed in the Ni10SiBEA zeolite (Fig. 2c and d). STEM images (Fig. 2b and d) evidence also uniform distribution of silicon, aluminum and oxygen atoms within HAlBEA zeolite particles, and only silicon and oxygen atoms within SiBEA particles.

Nitrogen adsorption-desorption isotherms and pore size distribution of zeolite supports and Ni-containing zeolite catalysts prior to the reduction are presented in Fig. 3. Large increase of the adsorption values at high partial pressures and the occurrence of slight changes of the shape of hysteresis loops point out slight differences of meso- and micro-porous structure of materials. The decrease of adsorption value at low relative pressures (below $p/p^* = 0.05$) upon dealumination may indicate the loss of microporosity. Such effect can be related to the partial distortion of initial BEA pore structure. An introduction of nickel to HAlBEA zeolite with subsequent calcination leads to the slight decrease of microporosity. This effect can be ascribed both to the partial incorporation of Ni$^{2+}$ to the zeolite framework, and thus shrinkage of the pores, as well as partial blocking of zeolite channels by NiO crystallites. Such changes are less evident in the catalysts with SiBEA support.

Fig. 4 shows FTIR/PAS spectra of the samples. Several characteristic O–H stretching vibrations bands are observed for HAlBEA and SiBEA zeolites. Note that the spectra were recorded for the samples at room temperature, hence some individual vibration bands could be overlapped with intensive vibration bands of H-bonded hydroxyl groups and physically adsorbed water, located at around ~3520, ~3420 cm$^{-1}$ and ~3200 cm$^{-1}$. The spectra for both zeolite supports show slight vibration bands located at 3747 cm$^{-1}$ which can be attributed to the internal isolated Si–OH groups. The intensity of the strong vibration bands located between 3700 and 3600 cm$^{-1}$ become weaker upon dealumination. This can be attributed to the removal of Al–OH groups (with vibration bands at ~3660 cm$^{-1}$), and bridging hydroxyls Al-O (H)-Si (~3610 cm$^{-1}$). The relative increase of the vibration bands intensity located at ~3520 cm$^{-1}$ can be attributed to the H-bonded Si–OH groups [26]. Ni10HAlBEA and Ni10SiBEA after drying show weaker vibration bands of isolated Si–OH and Al–O(H)–Si groups in the range 3760–3650 cm$^{-1}$. The presence of nickel precursor in the Ni10SiBEA catalyst drives to the increase in the vibration intensity of H-bonded hydroxyl bands with maximum at ~3560 cm$^{-1}$, whereas in the case of Ni10HAlBEA, the vibration bands in this region are much weaker, and moreover new maxima can be found at ~3520 and 3445 cm$^{-1}$. The bending hydroxyl group vibrations ($\delta_{OH}$) with maximum located at 1628 cm$^{-1}$ for HAlBEA are more intensive than similar vibrations in SiBEA sample (with maximum located at 1631 cm$^{-1}$). The peaks representing $\delta_{OH}$ vibration bands in the nickel containing samples after drying, especially for Ni10SiBEA dried sample, are much wider. Bentaleb et al. argued that an impregnation of silica support with an aqueous solution of nickel nitrate can lead to the development of crystalline nitrate phases [Ni(H$_2$O)$_6$](NO$_3$)$_2$ and [Ni(H$_2$O)$_5$](NO$_3$)$_2$ [35].

The development and transformation of such phases into the zeolite lattice bonded Ni(II) species or bulk NiO crystallites can be related to the surface properties of zeolite support. The vibration bands of nickel nitrate in the Ni10HAlBEA dried sample is visible as the doublet with maximum at 1409 and 1339 cm$^{-1}$. This picture is similar to the bulk nickel nitrate [36]. In the case of Ni10SiBEA the wide peaks with maxima located at 1478, 1440, 1393, and 1325 cm$^{-1}$ can be found. In the samples after calcination such maxima are not observed, which confirm complete transformation of nitrates into the suitable Ni–O or ~Ni–O–Si~ species.

Wide absorption band between 1250 and 1000 cm$^{-1}$ can be ascribed to the presence of asymmetric stretching vibrations of Si–O–Si bridges, regarded often as “external” (1250–1200 cm$^{-1}$) and “internal” (1100–1000 cm$^{-1}$) [37]. The presence of distinct peaks in the range of 1250 and 1000 cm$^{-1}$ in the HAlBEA and Ni10HAlBEA may indicate the occurrence of Q$^0$ silica oligomeric species. Upon dealumination one can observe a slight widening of such peaks, which can be attributed to the appearance of Q$^0$ silica oligomeric species [37]. The absorption bands at 780–830 cm$^{-1}$, and around 550–50 cm$^{-1}$ are the symmetric stretching vibrations of Si–O–Si bridges, while the band at 955–80 cm$^{-1}$ can be attributed to the stretching vibrations of the Si–OH groups. The bands located at 440–480 cm$^{-1}$ can be assigned to the Si–O bending vibration [38]. In the pure silica material (SiBEA sample) the intensity of the stretching vibrations of the Si–OH groups is much stronger than in the HAlBEA and nickel modified zeolites. It is interesting that such vibrations bands are observed (although with weaker intensities) in the samples after nickel incorporation prior to the calcination.

The general conclusion outlined in the XRD and FTIR/PAS studies on the state of nickel catalysts with BEA zeolites as supports before and after dealumination well correspond to the detailed studies performed by the application of $^1$H, $^{29}$Si and $^{27}$Al MAS NMR studies.

The $^{27}$Al NMR spectra of HAlBEA zeolite have 4 resonances (Fig. 5). The resonances at 54.8 and 52.5 ppm correspond to two types of aluminum atoms in the tetrahedral environment of the framework. The resonances at ~1.7 and ~10 ppm correspond to two types of aluminum atoms in an octahedral environment. The first is a thin peak, then it can be attributed to aluminum in a tetrahedral environment surrounded by water molecules. The second is a broad peak that may be attributable to extraframework species. Due to dealumination, the $^{27}$Al NMR spectrum of SiBEA zeolite (Fig. 5) has only one small contribution of framework aluminum atoms in the tetrahedral environment [27,39].

![Fig. 3. Nitrogen adsorption-desorption isotherms on nickel catalysts, solid lines – adsorption branch, dashed lines – desorption branch. Inset - pore size distribution calculated from the desorption branch of isotherms.](image-url)
The $^{27}$Al MAS NMR spectrum of Ni$_{10}$AlBEA is a little different from that of HAIBEA, loss of resolution and increase the width at mid-height of the peaks. These modifications come from the paramagnetic properties of NiO, which indicates an antiferromagnetic oxide at low temperature but becomes paramagnetic at higher temperatures. This temperature depends on the size of the particles [40,41]. It is known that the presence of paramagnetic species strongly disrupts the NMR spectra [42,43]. There are also changes in the $^{29}$Si (spinning sidebands, Fig. S1) and $^1$H (increased the width, Fig. S2) spectra when nickel is introduced into the HAIBEA zeolite. The $^{27}$Al MAS NMR spectrum (Fig. 5) of Ni$_{10}$SiBEA is very similar to spectrum of SiBEA.

The $^{29}$Si MAS NMR spectrum of HAIBEA (Fig. S1) shows two peaks at −114.5 and −111.0 ppm due to framework Si atom in a Si(OSi)$_4$ (Q$^4$) environment, located at different crystallographic sites. The Q$^3$ species show two contributions, one signal at −102.2 ppm for (OSi)$_3$Si(OH) groups and one signal at −106.1 ppm for (OSi)$_3$Si(Al) groups. After dealumination, $^{29}$Si MAS NMR spectrum of SiBEA (Fig. S1) is composed of three resonances, −114.5 and −111.0 ppm for Q$^4$ species and −102.2 ppm for Q$^3$ species. The contribution of (OSi)$_3$Si(Al) group (−106.1 ppm) has disappeared [43–45]. The $^{29}$Si MAS NMR spectra of Ni$_{10}$AlBEA and Ni$_{10}$SiBEA (Fig. S1) show that incorporation of nickel causes a decrease in the signal corresponding to the (OSi)$_3$Si(OH) group (−102.2 ppm), which suggests an interaction between the nickel and the silanols of the both zeolites.

The $^1$H spectra of the HAIBEA zeolite (Fig. S2) shows a main resonance at 4.7 ppm due to bridging hydroxyl protons (Si(OH)Al). The low contribution at 1.37 ppm corresponds to Si(OH). In the $^1$H MAS NMR spectrum of SiBEA (Fig. S2) two main peaks appear at 4.5 and 1.37 ppm, due to the protons of H−bonded Si−OH groups present at vacant T-atom sites and isolated Si−OH groups respectively [46,47]. Two peaks at 7 and 3.22 ppm are also present and correspond to protons of water molecules and of H−bonded Si−OH groups located in a second type of crystallographic site [48]. In the $^1$H MAS NMR spectra of Ni$_{10}$HAIBEA and Ni$_{10}$SiBEA (Fig. S2), signals at 4.63 and 4.50 ppm are not observable. The signal is probably too broadened by a paramagnetic effect, which means an interaction between the nickel and the bridging hydroxyl and silanols of the zeolite.

Fig. 6 shows temperature-programmed desorption of ammonia from the surface of reduced samples. Two irregular ammonia desorption peaks (m/e = 16) are observed for HAIBEA and Ni$_{10}$HAIBEA samples. The first peak, located between 100 and 300 °C evidences the presence of the large number of weak acid sites. Slight shift of the main maximum to lower temperatures is observed for Ni$_{10}$HAIBEA zeolite catalyst. The second peak spreads from 300 to 500 °C. The removal of Al from zeolite framework leads to the strong decrease of the number of acid sites both in the SiBEA zeolite and Ni-containing zeolite catalyst (Ni$_{10}$SiBEA). The samples prior to the adsorption of ammonia were reduced. Hence ammonia during desorption from the strong acid sites, i.e. above 300 °C can be decomposed to nitrogen and hydrogen on
nickel crystallites. Such effect is manifested in the appearance of desorption peaks of hydrogen (m/e = 2) and nitrogen (m/e = 28), partially overlapped with desorption peaks of ammonia from Ni10HAIBEA zeolite catalyst [49]. The lack of high-temperature desorption peaks of hydrogen in the curves of Ni10SIBEA catalyst can be connected with low acidity of the support. Small amount of adsorbed ammonia molecules can be converted to nitrogen and hydrogen, which recreates Brønsted hydroxyl on surface of zeolite. It is also interesting that very low desorption peaks of hydrogen at low temperatures (100 – 300 °C) may in- suggest presence of strong Brønsted acidic centers related to proton of Si–O(H)–Al groups. After high temperature treatment and cooling down to 100 °C, the band appears in the similar position as before treatment, indicating high mobility of such groups in zeolite materials under different reaction conditions. Similar shift is observed for nickel catalyst with dealuminated zeolite support Ni10SIBEA. However the bands are located at slightly lower wavenumbers, and hence the shift is noticed from 3735 cm$^{-1}$ to 3725 cm$^{-1}$. Two well developed vibration bands are observed at elevated temperatures for Ni10HAIBEA catalyst, located at around 3660 and 3600 cm$^{-1}$, and ascribed to extra-framework AlO–H and bridging acidic hydroxyls Si–O (H)–Al groups. In the spectra of the samples cooled down after high reduction temperature to 100 °C, such vibration bands are observed at similar positions. Moreover weak vibration band located at 3705 cm$^{-1}$ can be detected. The increase of treatment temperature leads to the decrease of the intensity of hydrogen bonded SiO–H groups in the Ni10SIBEA catalyst, located at around 3520 cm$^{-1}$. Strong vibration band on the spectra of this sample recorded at room temperature is observed at around 3630 cm$^{-1}$. It can be attributed to the interaction of SiO–H groups with Ni(II) species [39,50]. The intensity is gradually decreased with an increase in reduction temperature. After cooling down the sample to 100 °C, only strong vibration band of isolated silanol groups is observed, indicating very low acidity of Ni10SIBEA catalyst.

The FTIR spectrum of activated HAIBEA zeolite before pyridine adsorption (Fig. 8) exhibits five characteristic bands in OH stretching region: at 3780 and 3661 cm$^{-1}$ attributed to AlO–H groups of extra-framework aluminum species [51,52], at 3742 cm$^{-1}$ associated to isolated external SiO–H groups [53], at 3608 cm$^{-1}$ related to bridging hydroxyls Si-O(H)-Al and at around 3550 cm$^{-1}$ broad band related to H–bonded SiO–H groups. The introduction of nickel ions into HAIBEA leads to disappearance of the band at 3780 cm$^{-1}$ (Fig. 8) suggesting that nickel ions interacts with corresponding hydroxyl groups of HAIBEA.

The treatment of TEABEA with high concentrated nitric acid solution leads to removal of aluminum atoms and appearance in the spectrum of resulted SIBEA a very intense narrow band at 3735 cm$^{-1}$ related to isolated internal SiO–H groups (Fig. 8) and an intense broad band at 3521 cm$^{-1}$ due to hydrogen bonded silanol groups suggesting formation of vacant T-atom sites, as reported earlier [54]. The impregnation of SIBEA with aqueous solution of nickel precursor results in decreasing of intensity of OH bands of isolated internal SiO–H groups at 3735 cm$^{-1}$ and hydrogen bonded silanol groups at 3521 cm$^{-1}$ (Fig. 8) suggesting that nickel ions react with these silanol groups leading to the incorporation of nickel ions into framework of SIBEA, in line with earlier report [55].

To determine the nature, number and strength of acidic centers of supports and Ni-containing catalysts the FTIR spectra of adsorbed pyridine as probe molecule were recorded. The FTIR spectra after adsorption of pyridine at room temperature and then outgassed at 150 °C are given in Fig. 9. For HAIBEA sample very intense, characteristic bands of pyridinium cations are seen at 1636 and 1546 cm$^{-1}$ suggesting presence of strong Brønsted acidic centers related to proton of Si–O(H)–Al groups. Furthermore, the band at 1455 cm$^{-1}$ is related to
interaction between pyridine molecules and strong Lewis acidic centers (Al\(^{3+}\)) while the band at 1622 cm\(^{-1}\) may be ascribed to pyridine coordinately bonded to Lewis acidic centers [56,57]. Finally, the band at 1490 cm\(^{-1}\) corresponds to C–C oscillation of pyridine aromatic ring chemisorbed on both Brønsted and Lewis acid centers. Additionally, presence of small band at 1575 cm\(^{-1}\) may be associated with pyridine coordinately bonded to weak Lewis acidic centers [58].

Introduction of nickel ions into HAlBEA leads to formation of additional amount of Lewis acidic centers related to Ni(II) species (Fig. 9).
Table 3
The amounts of the Brønsted and Lewis acidic centers in Ni10HAIBEA, HAIBEA, Ni10SIBEA and SIBEA.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Brønsted acidic centers (μmol g⁻¹)</th>
<th>Lewis acidic centers (μmol g⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HAIBEA</td>
<td>268</td>
<td>130</td>
</tr>
<tr>
<td>Ni10HAIBEA</td>
<td>154</td>
<td>227</td>
</tr>
<tr>
<td>SIBEA</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Ni10SIBEA</td>
<td>3</td>
<td>120</td>
</tr>
</tbody>
</table>

* Quantification of number of acidic centers in zeolite was done as reported earlier by Emeis [33].

and Table 3) confirmed by appearing of intense bands at 1610 and 1450 cm⁻¹ and to decrease of quantity of Brønsted acidic centers evidenced by lower intensity of the bands at 1637 and 1546 cm⁻¹ for Ni2.0HHAIBEA than HAIBEA (Fig. 9). These phenomena may be explained by some degree of ion exchange of proton of bridging hydroxyls Si–O(H)–Al with nickel ions. As we have recently reported [39], SIBEA contains very small amount of the both Brønsted and Lewis acidic centers due to presence of traces of aluminum atoms remaining after nitric acid treatment. The impregnation of SIBEA with the aqueous Ni(NO₃)₂ solution leads to increase of amount of Lewis acidic centers evidenced by appearance of FTIR bands at 1610 and 1450 cm⁻¹ probably due to pyridine bonded to isolated nickel(II) species. The results presented in Fig. 9 and Table 3 show that HAIBEA and Ni10HAIBEA contain much higher number of Brønsted and Lewis acidic centers than SIBEA and Ni10SIBEA.

3.3. Reducibility and active surface area of nickel catalysts

The temperature-programmed-reduction curves are presented in the Fig. 10. Reduction of Ni10HAIBEA catalyst goes through two well separated maxima located in the range 300–430 °C and 430–600 °C. In the case of Ni10SIBEA catalyst, the first maximum can be distinguished between 250–350 °C, the second maximum is located between 350 and 600 °C. Small amounts oxide phases are reduced above 600 °C. Complex shape of the TPR curves has been usually regarded as indication of the reduction of nickel oxide phases of different metal-oxygen interactions in accordance to the reaction equation: NiO + H₂ → Ni° + H₂O. Low temperature peaks may indicate the reduction of large NiO crystallites (regarded as “free” or “bulk-like” NiO). While high-temperature reduction peaks can be ascribed to the reduction of nickel aluminate or nickel silicate species. The interfacial border between NiO particles and the support increases with the decrease of the size of NiO crystallites, which may facilitate diffusion of nickel species to the support during thermal treatment [59]. A complex shape of TPR curves of nickel zeolites has been also explained by the successive reduction of Ni(II) and Ni (I) species in accordance to the reaction equations Ni²⁺ + ½ H₂ → Ni⁺ + H⁺; followed by Ni⁺ + ½ H₂ → Ni° + H⁺), in which the evolved protons take part in the reaction with lattice oxygen, giving hydroxyl group n(Si)AlOH + nH⁺ → n(Si)AlO⁻H [60].

The temperature programmed desorption curves of hydrogen from the surface of catalysts are presented in the Fig. 11. Wide desorption peak of low intensity observed for Ni10HHAIBEA catalyst is located between −60 and 500 °C. Desorption of hydrogen from the surface of Ni10SIBEA catalyst is much higher. Irregular and large desorption peak is located in the similar temperature region as in the Ni10HHAIBEA catalyst. The complex shape of TPD curves is usually ascribed to the presence of sites with different Ni-H bond strength [61]. One can observe pronounced contribution of weak adsorption sites on the surface of Ni10SIBEA catalyst. The active surface area of Ni10HHAIBEA catalyst (Sₐ), calculated by the integration of the low-temperature peak (−60 to 500 °C) is equal to 1.4 m²/g, while in the case of Ni10SIBEA, it is equal to 6.1 m²/g. High temperature peaks (in the range 500–800 °C) are observed on the TPD curves of both catalysts. Their presence can be attributed to the formation of hydrogen in the reaction of the oxidation of nickel with hydroxyl groups located on the support [62]. Taking into account similar content of nickel, much smaller active surface area of Ni10HHAIBEA catalyst, and the same the presence of larger nickel crystallites, as well as shorter Ni-support border line in the Ni10HHAIBEA catalysts, one can suppose that HAIBEA zeolite, i.e. the support of high acidity and activity of hydroxyl groups, can influence the oxidation degree of nickel surface atoms to a much greater extent than SIBEA. High activity of the HAIBEA support in the surface reoxidation processes of nickel crystallites may to some extent facilitate the formation of coke precursors.

3.4. Catalytic performance in the steam reforming of ethanol

Preliminary studies of HAIBEA and SIBEA zeolites showed their high activity in ethanol conversion under applied reaction conditions, i.e. at 500 °C in the reaction mixture of high molar ratio H₂O/C₂H₅OH = 12/1 (Fig. 12). Almost complete ethanol conversion (X_eth = 100%) is observed for long time-on-stream for both zeolites (Fig. 12a). However practically no water conversion is seen (Fig. 12b). The main reaction product in the presence of HAIBEA and SIBEA zeolites, in spite of their
different acidity, is ethylene ($S_{\text{EtOH}} = \sim98\%$) (Fig. 12f), which indicates the predominant role of ethanol dehydration reaction: $\text{C}_2\text{H}_5\text{OH} \rightarrow \text{C}_2\text{H}_4 + \text{H}_2 \text{O}$ [63]. Small amount of propane ($S_{\text{C}_3\text{H}_8} = \sim6\%$) in the presence of HAIBEA zeolite is initially formed, which probably results from the consecutive reactions of ethylene. Direct production of C2-C4 olefins and higher hydrocarbons (gasoline-range) by the conversion of bioethanol (ethanol to gasoline, ETG) has recently gained great interests [64–68]. Formation of propane can be directly related to the higher acidity of HAIBEA zeolite. Such processes were less evident in the presence of dealuminated zeolite, and only small amounts of acetaldehyde were initially formed ($S_{\text{CH}_3\text{CHO}} = \sim2\%$). One can suppose that the gradual drop of selectivity to propane with the time-on-stream for HAIBEA zeolite was related the deposition of coke on the acid sites. Thus partial deactivation of zeolite led to the blocking of the sites which were active in the subsequent transformation of ethylene to hydrocarbons. Detailed discussion of the nature of carbonaceous deposits formed in such process is presented in the next section. One can state that dealumination of BEA zeolite can be regarded as useful method of the development of catalysts for ethylene production by the use of cheap dilute aqueous solutions of ethanol in biorefineries [69,70]. We have confirmed that introduction of large amounts of water to the bioethanol feed can suppress formation of higher hydrocarbons and large amounts of carbon deposit at high temperatures, especially on the SiBEA zeolite.

Nickel catalysts Ni$_{10}$HAIBEA and Ni$_{10}$SiBEA show high activity in the ESR reaction performed at 500 °C. Complete ethanol conversion is observed over both catalysts for long time-on-stream (22 h tests). Initial water conversion ($X_{\text{H}_2\text{O}}$), regardless of the type of catalysts is equal 40%. Lower conversion of water than ethanol ($X_{\text{EtOH}}$) results from the excess of water in relation to the reaction stoichiometry and occurrence of parallel reactions, such as ethanol dehydration and dehydrogenation ($\text{C}_2\text{H}_5\text{OH} \rightarrow \text{CH}_3\text{CHO} + \text{H}_2$). The drop of water conversion to about $X_{\text{H}_2\text{O}} = 10\%$ in the presence of Ni$_{10}$HAIBEA catalyst occurs during the initial 5 h time-on-stream. Whereas dealuminated nickel catalyst (Ni$_{10}$SiBEA) shows higher durability, and $X_{\text{H}_2\text{O}}$ remains on the initial level for more than 20 h. Ni$_{10}$HAIBEA catalyst shows higher initial selectivity to $\text{H}_2$, $\text{CO}_2$, and lower selectivity to $\text{CH}_4$ than Ni$_{10}$SiBEA. However, strong decrease of selectivity of Ni$_{10}$HAIBEA catalyst to $\text{H}_2$ (from 90% to about 60%), $\text{CO}_2$ (from 65% to 30%), slight decrease of selectivity to $\text{CH}_4$ (from 25% to 2–3%), CO (from 10% to ~3%), and pronounced increase in selectivity to ethylene (from zero to ~65%) is observed with the time-on-stream. Moreover, additional compounds appear in the reaction product mixture, such as acetaldehyde, ethane and propane. Although initial selectivity of Ni$_{10}$SiBEA catalyst to $\text{H}_2$ ($S_{\text{H}_2} = 75\%$) is smaller than Ni$_{10}$HAIBEA catalyst, it is not so strongly decreased with the time-on-stream. Similar trend is observed for $\text{CO}_2$ selectivity. Ni$_{10}$SiBEA catalyst shows slightly smaller initial selectivity to $\text{CO}_2$ than Ni$_{10}$HAIBEA ($55$ and 65%, respectively). $\text{CO}_2$ selectivity remains on the same level for long-time-on-stream, and only small drop is observed after 20 h of the reaction test. Ni$_{10}$SiBEA catalyst, in spite of lower initial selectivity to $\text{H}_2$ ($S_{\text{H}_2} = \sim65\%$) and $\text{CO}_2$ ($S_{\text{CO}_2} = \sim55\%$) shows better durability than Ni$_{10}$HAIBEA. No ethylene and acetaldehyde production is observed for relatively long time. Ni$_{10}$SiBEA catalyst, in contrast to the Ni$_{10}$HAIBEA, displays high selectivity to methane ($S_{\text{CH}_4} = \sim35\%$) and carbon monoxide ($S_{\text{CO}} = \sim10\%$). Partial deactivation of is observed after 20 h of the time-on-stream.

The catalysts were reduced before ESR reaction, hence the presence of nickel metallic nanoparticles influenced the route of catalytic reactions. The product selectivity was different from that observed for pure or unreduced nickel-modified zeolites in the conversion of diluted bioethanol solutions, where reaction was conducted towards ethylene or higher hydrocarbons [70,71]. The main initial reaction products in the steam reforming of ethanol in the presence of reduced Ni$_{10}$HAIBEA and Ni$_{10}$SiBEA catalysts were hydrogen and carbon dioxide. Formation of side reaction products, such as methane and carbon monoxide can be ascribed to the direct decomposition of ethanol (C$_2$H$_5$OH $\rightarrow$ H$_2$ + CH$_4$)

Fig. 12. The results of ethanol steam reforming performed at 500 °C for 20 h tests over bare zeolites and nickel catalysts; conversion of ethanol (Fig. 12a) and water (Fig. 12b), selectivity to hydrogen (Fig. 12c), carbon dioxide (Fig. 12d), carbon monoxide (Fig. 12e), ethylene (Fig. 12f), acetaldehyde (Fig. 12g), methane (Fig. 12h), ethane (Fig. 12i), and propane (Fig. 12j).
demonstrated, that both Ni\textsubscript{10}HAlBEA and Ni\textsubscript{10}SiBEA catalysts showed is almost not visible on the Ni\textsubscript{10}SiBEA images after ESR reaction test (Fig. 2a). Most of the catalysts with the small crystallites in the ESR reaction [72,73]. It was demonstrated, that both Ni\textsubscript{10}HAIBE A and Ni\textsubscript{10}SiBEA catalysts showed high initial activity in the ethanol steam reforming reaction. Both catal-ysts contained very small crystallites, below 10 nm (Fig. 2). However crystallite size distribution was much wider for Ni\textsubscript{10}HAIBE A, and con-sequently ESR reaction occurred both in the presence of small and large crystallites. It is interesting that in spite of lower mean nickel crystallite size, narrower crystallite size distribution, and much higher active surface area, Ni\textsubscript{10}SiBEA catalyst showed a little lower initial selectivity to \( \text{H}_2 \) (75% for Ni\textsubscript{10}SiBEA, in contrast to 90% for Ni\textsubscript{10}HAIBE A) and to \( \text{CO}_2 \) (55% for Ni\textsubscript{10}SiBEA and 66% for Ni\textsubscript{10}HAIBE A). This effect can be explained by the influence of support. The presence of Bronsted acid centres increased ability of Ni\textsubscript{10}HAIBE A catalyst to ethanol and water activation and their interaction with the surface intermediates, formed on the nickel crystallites. Simultaneously, the enhanced activation of ethanol on the sites of the Ni\textsubscript{10}HAIBE A catalyst led to the dehydrogenation reaction, formation of ethylene and then development of coke with the time-on-stream on the acid sites. The presence of carbonaceous deposits in zeolite pores can also induce some diffusion limitation. Such effects may explain gradual drop of selectivity to \( \text{H}_2 \) and \( \text{CO}_2 \) during a few hours of the reaction test in the case of Ni\textsubscript{10}HAIBE A. In contrast this phenomenon is not observed for Ni\textsubscript{10}SiBEA resisted to formation such coke because of the absence of strong acidity and characterised by high stability in formation of \( \text{H}_2 \) and \( \text{CO}_2 \). Similar phenomena were observed by Sánchez-Sánchez et al. who achieved lower acidity and better nickel dispersion of Ni/Al\textsubscript{2}O\textsubscript{3} by the introduction of Mg [76]. The inhibition of ethanol dehydrogenation reaction to ethylene, increase in \( \text{CO}_2 \) selectivity and improved coking resistance of catalysts in ESR reaction was in si-milar way achieved by Denis et al. through the introduction of sodium [77].

3.5. The nature of carbon deposits

TEM and STEM images of nickel catalysts after ESR reaction tests performed at 500 °C are presented in the Fig. 13. Several forms carbon deposits can be found on the images of Ni\textsubscript{10}HAIBE A catalyst (Fig. 13a). Filaments of different diameter, length and internal structure are well visible. Some nickel crystallites are located on the top of filaments or surrounded by carbonaceous material. The inset in the Fig. 13a shows the image of the filaments with regular, fishbone-like structure of ex-ternal parts and less ordered inner parts. It is difficult to estimate nickel crystallite size from TEM images, part of them seems to be disintegrated into the smaller particles. The microstructure of zeolite particles is less regular than in the sample before the ESR reaction test (Fig. 2a). Most of them seem to be surrounded by amorphous carbon. Filamentous carbon is almost not visible on the Ni\textsubscript{10}SiBEA images after ESR reaction test (Fig. 13c,d). One can find only on the bottom part of the image (Fig. 13c) small amounts of thin rather uniform filaments, with the size corresponding to the size of the small nickel nanoparticles.

FTIR/PAS spectra of the supports and nickel catalysts after reaction tests are presented in the Fig. 14. It is difficult to find strong C–H vibra-tions, typical for organic compounds. Hence one can assume that samples contain carbon deposits with high C/H ratio. Small absorption bands located between 2000 and 1400 cm\(^{-1}\) can be ascribed to the stretching vibrations of carbonyl groups in surface intermediate species and bending vibrations of hydroxyl groups (\( \delta_{\text{OH}} \)), as well as conjugated olefins, polycondensed aromatics, and deformation vibrations of \( \text{CH}_x \) groups [78] Low intensities of such vibrations in the Ni\textsubscript{10}HAIBE A cata-lyst can indicate partial removal of hydroxyl groups, resulted from the presence of carbon deposits. Slight differences of the shape of absorption bands located between 1250 and 1000 cm\(^{-1}\), related with stretching vibrations of Si–O–Si bridges and the bands between 955 and 980 cm\(^{-1}\), ascribed to the vibrations of the Si–OH groups after activation (Fig. 4) and ESR reaction (Fig. 14) can indicate, that the microstructure of most zeolites has not been strongly changed during reaction. However more pronounced changes, visible for Ni\textsubscript{10}SiBEA catalyst may confirm partial breakdown of original microstructure. It is worth to note that the catalytic tests were performed under severe conditions to intensify deactivation phenomena, performing tests at high temperatures, using high ethanol weight hourly space velocity (\( \text{WHSV}_{\text{EOH}} = 9.5 \, \text{gEOH h}^{-1} \, \text{gcat}^{-1} \)) and undiluted reaction mixture with inert gas.

The Raman spectra of the supports and nickel catalysts after ethanol steam reforming reaction performed at 500 °C are shown in Fig. 15. Several peaks are distinguished on the Raman spectra of unmodified HAIBE A zeolite. Weak vibrations near 2700–3000 cm\(^{-1}\) were detect-able. Such vibrations can be ascribed to the C–H bonds in carbon de-posits and to overtone of D–band [79]. The irregular shapes of peaks with maxima at ca. 1604, 1365, and 1200 cm\(^{-1}\) indicate superposition of vibrations of different carbon-containing groups. The asymmetric peak located between 1670 and 1500 cm\(^{-1}\) can be ascribed to the C=C stretching vibrations of sp\(^2\) C=C bonds, in the conjugated olefin and aromatic species - near 1600 cm\(^{-1}\), and to E\(_2g\) carbon-carbon in plane-in-plane stretching vibration in graphite-like deposits - close to 1570 cm\(^{-1}\), designated as G-band. Wide peaks in the range 1500–1100 cm\(^{-1}\) has been often ascribed in the literature to the vibrations of CH\(_x\) group, vibration of olefin and aromatic species, and also to some defects in the structural units of graphite-like deposits, referred as the D–band [80]. D–band has been has been often correlated with breathing modes of sp\(^2\) atoms in carbon rings [81]. The lack of such strong vibration bands for SiBEA zeolite points out that carbon was not formed during ESR reaction test. The colour of the SiBEA sample was not changed during the ESR reaction test, while HAIBE A sample became grey. Studies confirm high durability of dealuminated SiBEA zeolites in the conversion of diluted solution of ethanol to ethylene at 500 °C, and this reaction can be proceed without formation of carbonaceous deposits with the long-time-on-stream. In turn, the presence of water does not eliminate formation of discussed above olefin and aromatic species, as well as graphite-like deposits resulted from the ethylene oligomerization, aromatization and then cracking of hydrocarbons on the HAIBE A zeolites.

Strong Raman vibration bands are observed for nickel catalysts. The shape and position of the peaks is slightly different from that presented for pure HAIBE A zeolite. In part, the nature of carbon deposits in the Ni\textsubscript{10}HAIBE A results from similar reactions, initiated by ethanol dehy-dration on the acidic sites of the support to ethylene, and then oligo-merization and transformation to coke. The peaks in the range 1600–1500 cm\(^{-1}\) can indicate the presence of the conjugated olefin and aromatic species, and more graphic-like structures, deposited on the support (G–band). However taking into account wide discussion in the literature [82–85] and presented above TEM results, one can assume that vibrations in this region may also indicate sp\(^3\) vibrations in gra-phitic carbon structures of different disorder, i.e. highly ordered structures in carbon filaments and surface films (at ca. 1580, 1570 cm\(^{-1}\), nanocrystalline graphite (at ca. 1600 cm\(^{-1}\)), amorphous carbon and CH\(_x\) groups (at ca. 1490 cm\(^{-1}\)) [81]. Similar vibrations bands are observed for Ni\textsubscript{10}SiBEA catalyst. However, several features of the spectra indicate more diverse nature of carbon structures in the Ni\textsubscript{10}HAIBE A catalyst. G-peak in the Ni\textsubscript{10}HAIBE A is wider than in the
Ni10SiBEA, which is often regarded as indication of disorder [81]. Similar conclusions can be derived comparing the ratio of intensities of D to G bands, \( I(D)/I(G) = 1 \) for Ni10HAlBEA and \( I(D)/I(G) = 0.7 \) for Ni10SiBEA catalysts, respectively.

We have evidenced that the use of dealuminated BEA zeolite as support for nickel catalysts led to the pronounced increase of durability in the ESR reaction. However, the key question, what is the most important factor responsible for improved performance of obtained catalysts in the ESR reaction seems to be still open. Carbon deposit formation is regarded as one of the most crucial reasons of catalysts deactivation in the ESR reaction. In general it can be formed as result of Boudouard reaction (\( 2\text{CO} \rightarrow \text{CO}_2 + \text{C} \)), methane decomposition (\( \text{CH}_4 \rightarrow \text{C} + 2\text{H}_2 \)), decomposition or polymerization or of ethylene (\( \text{C}_2\text{H}_4 \rightarrow 2\text{C} + \text{H}_2 \)), aldol condensation of acetaldehyde (\( \text{CH}_3\text{CHO} \rightarrow \text{c}oke \)) or condensation of acetone to mesitil oxide (\( 2\text{CH}_3\text{CCH}2\text{OH} \rightarrow (\text{CH}_3)\text{C(OH)CH}_2\text{COCH}_3 \rightarrow (\text{CH}_3)\text{CCHCOCH}_3 + \text{H}_2\text{O} \)) [13,14,86]. The

**Fig. 13.** Microscopic images of the catalysts after ESR reaction performed at 500 °C for 20 h; a) Ni10HAlBEA - TEM (Fig. 13a) and STEM (Fig. 13b); Ni10SiBEA - TEM (Fig. 13c) and STEM (Fig. 13d).

**Fig. 14.** FTIR/PAS studies of the zeolites and nickel catalysts after ESR reaction.

**Fig. 15.** Raman spectra of zeolites and nickel catalysts after ESR reaction.
growth of filamentous carbon deposits starts from the migration of carbon atoms on the surface of nickel crystallite and formation of graphitic layers, which can expel crystallite from the support. The retardation of such initial processes can be achieved by intensification of surface gasification processes, e.g. with participation of redox sites in the support, interaction with promoters (K, Ce), changing of electronic properties of metal particles and their shape or enhancing surface oxygen mobility. Similar effects can be obtained by the decrease of metal crystallite size, which result in the increase of metal-support border-line, and increase of the rate of surface reactions with participation of hydroxyl groups or redox sites on the support. The decrease of the rate of filamentous carbon deposition formation by the decrease of nickel crystallite size in ESR reaction in the Ni/CeO2 catalysts has been recently evidenced by Slowik et al. [73]. Fig. 2e reveals dominant contribution of very small nickel crystallites in the Ni10SiBEA catalyst. Whereas Ni10HAIBEA catalyst contains both small and large nickel crystallites, the crystallite size distribution is very broad. Hence the observed differences of the nature of carbon deposits in the catalysts after ESR reaction (Fig. 13 and 15) can be explained by the participation of both small and large crystallites in the formation of carbon deposits. Moreover, due to the different size of nickel crystallites and different rate of carbon deposition, such process may lead to the development of carbon filaments of different diameter and graphitization degree.

Another reason of improved resistance of Ni10SiBEA catalyst towards coking is lower acidity of the SiBEA support. TPD NH3, DRIFTS and pyridine FTIR studies indicated the presence of the large number of Brønsted acid sites on the HAIBEA support (Figs. 6–9, Table 3). It seem very probable that high initial activity and selectivity of Ni10HAIBEA catalyst can be ascribed to the enhanced adsorption of ethanol and water on the acid sites on the support, fast migration of surface species to the metal-support border line and C–C bond cleavages on nickel crystallites. Due the presence of Brønsted acid sites on the HAIBEA support, ethanol can be easily decomposed to ethylene, then partially converted to hydrocarbons and carbon deposit. Hence the role of hydroxyl groups in activation of water and ethanol molecules in ESR reaction can be partially suppressed with the time–on–stream. In the subsequent stages of Ni10HAIBEA catalyst operation, the reaction may exclusively occur on the surface of nickel crystallites, located on the top of carbon filament or partially covered by carbon deposit, as well as independently on the zeolite acid sites, leading to the gradual transformation of both forms of carbon deposits. In the case of Ni10SiBEA catalyst, which showed very low acidity, the processes of the formation of heavier hydrocarbons and their transformation to coke on the support are less pronounced. Surface species formed after adsorption of ethanol or water, due to the presence of small nickel crystallites, can easily diffuse and participate in surface reactions occurred on nickel crystallites or close to the interfacial region metal-support, preventing fast transformation of filaments or surface carbonaceous films towards encapsulating carbons.

4. Conclusions

The effects of dealumination of BEA zeolite on the properties of nickel catalyst and its performance in the steam reforming of bioethanol were studied. It was indicated that dealumination of BEA zeolite led to the formation of specific sites in the zeolite framework, which facilitated particular adsorption of nickel precursors, hindering agglomeration of nickel ionic species during calcination into the large NiO crystalline species and subsequent formation of metallic nickel crystallites during reduction. Nickel-SiBEA catalyst obtained by the application of dealuminated BEA zeolite showed higher active surface area in comparison to that prepared using unmodified BEA support. It was observed that dealumination led to the slight decrease of crystallinity and simultaneously pronounced decrease of acidity of zeolite. Pure zeolites and obtained nickel catalysts showed complete conversion of ethanol during long operation time of more than 20 h. However, the main reaction product of the reaction performed in the presence of zeolite supports was ethylene. Small amounts of propane were initially formed in the presence of HAIBEA zeolite. The decrease of selectivity to propane was ascribed to the partial deactivation of zeolites by carbon deposit formation. Nickel modified HAIBEA zeolite (Ni10HAIBEA) showed higher initial selectivity to hydrogen and carbon dioxide than the nickel catalyst with dealuminated support (Ni10SiBEA), however the selectivity decreased during the first 5 h with the time-on-stream, and the reaction led to the formation of large amounts of ethylene. In contrast, nickel catalyst with less acidic support showed stable selectivity to CO2 on the level of 60% for long time-on-stream. Microscopic and Raman spectroscopy studies of the samples after SRE reaction evidenced different forms of deposits on the surface of catalysts. Large amounts filamentous carbon and less ordered carbon deposit forms were identified in the catalysts with unmodified BEA zeolite support. The presence of highly dispersed nickel nanoparticles, and simultaneously the decreased acidity of the support as shown for SiBEA support, facilitated high conversion of ethanol and water, as well as good selectivity to H2 and CO2 for long operation time.

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Appendix A. Supplementary data

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References

[16] I. Rossetti, M. Compagnoni, E. Finocchio, G. Ramis, A. Di Michele, A. Zucchini,


