Synthesis of Na-LSX type zeolite from Polish fly ash

Introduction

The coal combustion process is the basic method of electricity production in Poland. The wastes from this process are slag and exhaust fumes which contain fly ash and gases (N₂, H₂O, SO₄, NOₓ, CO, CO₂, carbohydrates, and aldehydes). The main disadvantage of the currently used combustion technologies is the emission of pollutants, mainly CO₂, into the atmosphere, as well as the low efficiency of power units (Kordylewski 2005; Ściążko 2009). In Poland, the wastes resulting from the energetic combustion of coal are mainly neutralized by landfilling and they are therefore seen as a threat to the environment (Łączny 2002). On the other hand, fly ash can be used in environmental protection, including air protection (for desulphurization and removal of CO₂ from exhaust gas), surface waters (for removing the causes of eutrophication of lakes), groundwater (for creating protective barriers) and for the recultivation of degraded areas (Łączny 2002; Lee et al. 2005; Prasad et al. 2012; Łączny et al. 2015; Mishra et al. 2019).
The composition of fly ash depends mainly on the type of coal, temperature and the type of atmosphere in the furnace, as well as the flue gas purification technology (SCR or SNCR reactor for NO\textsubscript{x} reduction, desulphurization installations, mercury removal, etc.). The final chemical and mineral composition of the ash is formed in the exhaust stream. In the case of fly ash from coal combustion in pulverized-fuel boilers, the content of the crystalline phase is from 15% to 50%. The vitreous phase, which is a residue, is composed mainly of SiO\textsubscript{2} and Al\textsubscript{2}O\textsubscript{3} (Łączny 2002).

Zeolites are crystalline, hydrated metal aluminosilicates (mainly groups I and II). These minerals belong to the group of spatial aluminosilicates (skeletal). Their structure consists of tetrahedrons [AlO\textsubscript{4}] and [SiO\textsubscript{4}], which form rings, and open channels with strictly defined diameters are found between them. The characteristic feature of zeolites associated with their construction is the ability to give away water without changing the shape of the crystal or grain during heating to 400°C and taking it back during cooling. The channels sizes in zeolites range from about 3Å to 10Å. A different degree of silicon and aluminum oxidation gives zeolites an electronegative nature, the size of which is determined by the Al/Si ratio. Therefore, these minerals are characterized by high adsorption capacity and a high ion exchange capacity. In addition, strictly defined channel diameters give the possibility of the selective adsorption of specific ions and molecules (Bolewski and Manecki 1993). Zeolites are used, among others, to remove heavy metals, ammonium ions, radioactive elements, hydrocarbons, organochlorine compounds and the separation of selected gases, among others H\textsubscript{2}S, SO\textsubscript{2}, CH\textsubscript{4}, CO\textsubscript{2}, Hg BTEX, as well as an addition to fertilizers, soil cleaning, medicine (Erđem et al. 2004; Franus and Wdowin 2010; Wdowin et al. 2014; De Smedt et al. 2015; Sadeghi et al. 2016; Bandura et al. 2017; Channabasavaraj and Ramalinga 2017; Chen et al. 2017; Collins et al. 2020; Costa et al. 2020; Czuma et al. 2020a, 2020b; Eisenwagen and Pavelić 2020; Ferretti et. al. 2020; Kunecki et al. 2020; Pambudi et al. 2020; Vaezihir et al. 2020). Due to their chemical composition (high content of aluminum and silicon), fly ash from coal combustion in pulverized-fuel boilers is a substrate for the synthesis of zeolites and thus finds application in environmental protection. Considering that the main disadvantage of currently used combustion technologies is the emission of large quantities of CO\textsubscript{2} into the atmosphere (for example, in Poland in 2017, the emission was 308.6 mln Mg (BP 2018), the use of zeolites synthesized on the basis of ash for the sorption of this gas is of particular application importance. Among others, zeolites of the faujasite group have appropriate channel sizes to use them for CO\textsubscript{2} sorption (Prats et al. 2017).

The synthetic X-type zeolites belong to the faujasite group (FAU code), the structure of which is as follows: β-cages and hexagonal prisms are connected in such a way that large internal super cages (α -cages) are created. Molecules, including CO\textsubscript{2}, can enter the α-cages through 12 Membered Ring (12MR). The windows have a diameter of 7.4Å (Deams et al. 2006). Table 1 presents the chemical formulas of selected synthetic X-type zeolites. These zeolites can basically be divided into two groups: low silicon content (LSX type zeolites) and medium silicon content zeolites (X-type zeolites). The established chemical formulas as well as the chemical composition of X-type zeolites reveal their considerable diversity.
The main cations in them can be Na⁺, K⁺ and Ca²⁺, which is important for the content of SiO₂ and Al₂O₃.

The basic method of synthesis of zeolites based on the ash from coal combustion is a hydrothermal conversion (Querol et al. 2001, 2002; Bukhari et al. 2015; Lee et al. 2017; Belviso 2018; Czuma et al. 2020b; He et al. 2020; Lankapati et al. 2020; Miricioiu and Niculescu 2020; Todorova et al. 2020; Verrecchia et al. 2020; Wiśniewska et al. 2020). Kotova et al. (Kotova et al. 2016) studied the effect of synthesis duration (2–12 hours), the concentration of activation solution (1.5–4.5M NaOH) and temperature (80–180°C) on the type of zeolites synthesized on the base of fly ash. The zeolite X was obtained in the temperature range of 90–100°C, and the increase in reaction temperature caused the formation of zeolites with medium (zeolite P – 140°C) and small (analcime and cancrinite – 180°C) channel size. It has also been shown that the type of zeolite and its content in the reaction products are significantly affected by reaction time and alkali concentration. A series of syntheses were carried out at 140°C. Under these conditions, the X zeolite was obtained applying a short reaction time (4 hours) and a high concentration of activating solution (4.5M NaOH). Longer reaction time led to the disappearance of the X zeolite phases and crystallization of the more thermodynamically stable phases (zeolite P followed by analcime).

So far, few syntheses of X-type zeolites from fly ash has been done in Poland (Derkowski et al. 2006, 2007; Franus and Wdowin 2011; Franus 2012; Bukalak et al. 2013; Franus et al. 2014). Zeolitization was carried out on fly ashes from the Kraków CHP plant as well as the Koziene, Rybnik and Stalowa Wola power plants, in which hard coal is the basic fuel.

Table 1. Characterization of selected synthetic zeolites, author’s own study based on (Yang et al. 2010)

Tabela 1. Charakterystyka wybranych syntetycznych zeolitów

<table>
<thead>
<tr>
<th>Mineral name</th>
<th>Chemical formula</th>
<th>SiO₂</th>
<th>Al₂O₃</th>
<th>SiO₂ + Al₂O₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zeolite X</td>
<td>Na₈₈(Al₈₈Si₁₀₄O₃₈⁴)(H₂O)₁₇₂,₁</td>
<td>37.72</td>
<td>27.09</td>
<td>64.81</td>
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<tr>
<td>Zeolite X</td>
<td>Ca₄₀Al₈₀Si₁₂₅O₃₈⁴(H₂O)₃₁₆</td>
<td>44.44</td>
<td>26.94</td>
<td>71.38</td>
</tr>
<tr>
<td>Zeolite X</td>
<td>K₉₀,(Al₂Si₁₀₀O₃₈₄)</td>
<td>40.21</td>
<td>31.39</td>
<td>71.60</td>
</tr>
<tr>
<td>Zeolite Na-LSX</td>
<td>Na₉₆(Al₉₆Si₉₆O₃₈⁴)(H₂O)₃₈₄,₃</td>
<td>28.05</td>
<td>23.80</td>
<td>51.85</td>
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<tr>
<td>Zeolite Na-K-LSX20</td>
<td>Na₇₇K₁₀Al₉₆Si₉₆O₃₈⁴(H₂O)₉₆₉,₉</td>
<td>27.34</td>
<td>23.20</td>
<td>50.54</td>
</tr>
<tr>
<td>Zeolite Na-K-LSX42</td>
<td>Na₉₆K₄₀(Al₉₆Si₉₆O₃₈₄)(H₂O)₄₀₈,₇</td>
<td>26.65</td>
<td>22.61</td>
<td>49.26</td>
</tr>
<tr>
<td>Zeolite Na-K-LSX80</td>
<td>Na₂₄.₄K₇₁.₆(Al₉₆Si₉₆O₃₈₄)(H₂O)₂₉₄,₂</td>
<td>24.34</td>
<td>20.65</td>
<td>44.99</td>
</tr>
<tr>
<td>Zeolite Na-K-LSX80</td>
<td>Na₁₉K₇₇(Al₉₆Si₉₆O₃₈₄)(H₂O)₃₈₅,₇</td>
<td>26.42</td>
<td>22.42</td>
<td>48.85</td>
</tr>
<tr>
<td>Zeolite K-LSX</td>
<td>K₉₀Al₉₆Si₉₆O₃₈⁴(H₂O)₃₆₂,₃</td>
<td>26.56</td>
<td>22.54</td>
<td>49.11</td>
</tr>
</tbody>
</table>
The synthesis processes were carried out under various laboratory conditions and a semi-technical scale:

- amount of fly ash – 10 g, concentration of sodium hydroxide solution – 3M, temperature – 75°C, synthesis duration – 24 h (Derkowski et al. 2006),
- amount of fly ash – 10 g, ratio of sodium hydroxide with sodium chloride in the ratio of solid to liquid 25–100 g/dm³, temperature 21–22°C, synthesis duration – 2–20 months (Derkowski et al. 2007),
- amount of fly ash – 10–15 kg, amount of sodium hydroxide – 11 kg, amount of water – 0.9 m³, temperature – 80°C, synthesis duration – 24 h (Franus and Wdowin 2011),
- amount of fly ash – 20 g, concentration of sodium hydroxide solution – 2–4M or 2–7M, temperature – 50–100°C or 21–22°C, synthesis duration 6–36 h or 1–6 months (Franus 2012),
- sodium hydroxide was fused with fly ash in a 1.2:1 ratio at 550°C, after grounding the sinter was mixed with water in a ratio of 1:4, synthesis temperature – 90°C, synthesis duration – 24 h,
- amount of fly ash – 20 g, concentration of sodium hydroxide solution – 3M or 5M, temperature – 75°C or 95°C, synthesis duration – 24 h (Franus et al. 2014).

In the synthesis products, admixtures of sodalite were sometimes found and there were phases that are relics of fly ash (quartz, mullite, amorphous phase, calcite).

The aim of the study was to carry out zeolitization of fly ash using a conventional hydrothermal process in such conditions as to obtain an X-type zeolite in the synthesis products. The process was carried out using various NaOH / fly ash mass ratio – 3.0 (sample A), 4.0 (sample B) and 6.0 (sample C). The process was carried out under the following permanent conditions: temperature: 90°C, time – 16 h, water solution of NaOH (L)/fly ash (g) ratio – 0.025.

The products obtained as a result of zeolitization were the subject of qualitative testing to determine new phases resulting from the process. The tests included observations in a mineralogical microscope, identification of phases by X-ray diffraction, observations and measurements in a scanning electron microscope and thermal analysis.
The microscopic observations of fly ash and products of synthesis were carried out on the Zeiss Axioscope polarizing microscope in the transmitted light. Powder preparations were prepared for this testing.

The identification of phases by X-ray diffraction (XRD) for fly ash and products of synthesis was performed on the Malvern Panalytical Aeris 1 diffractometer with the CuKα lamp. The conditions of measurement were as follows: voltage – 40 kV, current – 8 mA, time – 4.84 s, increment of the 2-theta angle – 0.003°, range of 2-theta angle – 4–74°. The HighScore Plus software with the database was used to interpret the XRD spectra.

The measurements of mass loss of the tested fly ash and the products of synthesis were made by thermal analysis using the analyzer SDT Q600 by TA Instruments. The conditions of measurement were as follows: mass sample approx. 20 mg, temperature range – 22–1000°C, temperature increase – 10°C/min, atmosphere – air.

Grain morphology and measurements of the chemical composition in the micro-area for the fly ash and the products of synthesis were made using the electron microscope (SEM) SU3500 by Hitachi, equipped with the UltraDry energy dispersive spectrometer (EDS) detector by ThermoFisher Scientific.

2. Results

The chemical composition of the tested fly ash sample, as indicated by multi-year studies (1999–2019), did not show any significant differentiation. The main ash components were SiO₂ and Al₂O₃, the contents of which were respectively 51.62% by mass and 28.14% by mass (in this study), while so far these contents varied respectively in the range of 49.20–50.70% by mass for SiO₂ and 25.24–26.35% by mass for Al₂O₃ (Table 2). One of the more important chemical components was Fe₂O₃, the content of which was 6.34% by mass (in this study), and in earlier studies, it ranged from 5.18 to 8.10% by mass. The contents of TiO₂, MgO, CaO, Na₂O, and K₂O ranged from approx. 1% by mass to approx. 3% by mass, while the content of other chemical components (MnO, SO₂, P₂O₅) did not exceed 1% by mass (in this study), and these components were determined at a similar level in the earlier studies (Ratajczak et al. 1999; Wala and Rosiek 2008; Klupa et al. 2017a). The loss on ignition (LOI) in the studied ash was 2.13% by mass and was mainly associated with the unburnt organic matter (carbon), although in the earlier studies it was even 6.80% by mass. The low content of rare earth elements was also found (Klupa et al. 2017b).

The results of chemical composition studies indicate that the SiO₂/Al₂O₃ ratio is 1.83 (mol ratio 3.12), which predisposes this fly ash to zeolite conversion (Querol et al. 2002; Bukhari et al. 2015; Belviso 2018; Ren et al. 2020).

Studies on the mineral composition of fly ash also do not indicate a large variation. The microscopic observations in transmitted light revealed that the ash contained mainly mullite and quartz, as well as a significant share of amorphous substance (glass and unburnt organic substance). Mullite and spherical enamel often formed joint aggregates (Fig. 1). In fly ash,
Table 2. Variability of the chemical composition of fly ash in one of the Polish power plants in the years 1999–2018 (% mass)

Tabela 2. Zmienność składu chemicznego popiołu lotnego w jednej z polskich elektrowni w latach 1999–2018 (% masy)

<table>
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<tr>
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<tbody>
<tr>
<td>SiO₂</td>
<td>49.60</td>
<td>49.92</td>
<td>50.70</td>
<td>51.62</td>
</tr>
<tr>
<td>TiO₂</td>
<td>1.00</td>
<td>–</td>
<td>0.99</td>
<td>1.21</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>26.35</td>
<td>25.82</td>
<td>25.24</td>
<td>28.14</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>8.10</td>
<td>5.18</td>
<td>5.68</td>
<td>6.34</td>
</tr>
<tr>
<td>MnO</td>
<td>0.10</td>
<td>–</td>
<td>0.06</td>
<td>0.07</td>
</tr>
<tr>
<td>MgO</td>
<td>2.68</td>
<td>1.82</td>
<td>2.58</td>
<td>2.37</td>
</tr>
<tr>
<td>CaO</td>
<td>4.10</td>
<td>3.92</td>
<td>3.41</td>
<td>3.02</td>
</tr>
<tr>
<td>Na₂O</td>
<td>2.94</td>
<td>1.10</td>
<td>0.97</td>
<td>1.32</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.86</td>
<td>3.00</td>
<td>3.71</td>
<td>3.35</td>
</tr>
<tr>
<td>SO₃</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.28</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.48</td>
<td>–</td>
<td>0.57</td>
<td>0.63</td>
</tr>
<tr>
<td>LOI</td>
<td>3.01</td>
<td>6.80</td>
<td>4.79</td>
<td>2.13</td>
</tr>
<tr>
<td>Total</td>
<td>99.21</td>
<td>97.56</td>
<td>98.71</td>
<td>100.48</td>
</tr>
<tr>
<td>Mol ratio Si/Al</td>
<td>3.20</td>
<td>3.29</td>
<td>3.41</td>
<td>3.12</td>
</tr>
</tbody>
</table>

Fig. 1. Microscopic image of the tested fly ash, light passing through one Nicol prism, 200× magnification, (A) in the center a spherical enamel aggregate with mullite, (B) elongated grains of opaque organic matter

Rys. 1. Obraz mikroskopowy badanego popiołu lotnego, światło przechodzące jeden nikol, pow. 200× (A) w centrum kuliste agregaty szkliwa z mullitem, (B) wydłużone ziarna nieprzezroczystej substancji organicznej
in addition to quartz and mullite, hematite was also found by the X-ray method. A high background on the diffractogram in the 2-theta angle range of 16–31° proves the presence of an amorphous substance (Fig. 2). Studies by other authors have shown that spinels (magnetite, magnesioferrite, hercynite, pleonaste), wustite, rutile, and florencite may occur in smaller amounts (Ratajczak 1999; Adamczyk and Makosz 2014; Klupa et al. 2017a; Cempa et al. 2018).

The identification of phases by X-ray diffraction revealed the phase composition of the products of synthesis. However, it should be emphasized, that all the synthesized materials contain the mineral components of fly ash (mullite with quartz, as well as magnetite) unreacted in the synthesis. The newly formed mineral components are:

- in sample A – Na-LSX type zeolite and hielscherite,
- in sample B – Na-LSX type zeolite, hielscherite and hydrosodalite,
- in sample C – hydrosodalite (Fig. 2).

This indicates the presence of basic reflections originating from these phases appearing on the diffractograms:

- Na-LSX type zeolite (Å) – 14.46; 8.86; 7.55; 5.74; 4.43; 3.82; 3.34; 3.06,
- hielscherite (Å) – 9.62; 5.55; 4.91; 4.60; 3.81; 3.43; 2.74; 2.62; 2.52; 2.17; 2.12,
- hydrosodalite (Å) – 6.30; 3.63; 2.82; 2.57; 2.10; 1.81; 1.74; 1.57.

![Fig. 2. X-ray diffraction pattern of fly ash (sample IS) and synthesis products (samples: A, B and C)](image-url)
The chemical formulas of the new phases in the products of synthesis are as follows:

- Na-LSX type zeolite (X) – Na₉₆(Al₉₆Si₉₆O₃₈₄) · 348.3 H₂O,
- hielscherite (Hi) – Ca₃Si(OH)₆(SO₄)(SO₃) · 11 H₂O,
- hydrosodalite (S) – Na₆(AlSiO₄)₆ · 8 H₂O.

They indicate that the theoretical loss of mass associated with the dehydration of these phases is respectively: X – 33.64% by mass, Hi – 39.38% by mass, and S – 14.46% by mass.

The results of the TGA analysis revealed that the weight loss of fly ash is 1.90%, which corresponds well with the chemical analysis (2.13%). This weight loss was probably mainly associated with the oxidation of unburned coal in ash. The mass stabilizes at a temperature of approx. 700°C.

In the products after synthesis, a much larger mass loss was visible compared to the initial fly ash sample and amounted to a maximum of 17.65% for sample B and a minimum for sample C – 14.62%. The mass losses in these three samples were therefore similar to each other, as was the course of TG curves.

Thermogravimetric studies indicate (Esposito et al. 2004; Bukalak et al. 2013; Fan et al. 2015) that the Na-LSX zeolite has a mass loss of several degrees. The first effect up to about 220°C corresponds mainly to the desorption of physisorbed water, and another one occurred from 220 to 480°C, which was attributed to the dehydration of chemisorbed water. The TG curves of the synthesis products tested have an almost identical course. However, the last effect of mass loss on these curves is noteworthy ending at 700°C, after which the mass stabilizes. It is identical to the ash sample and should be associated with the oxidation of unburned coal in the sample.

The reason for the lower weight loss in sample C was mainly the presence of hydrosodalite synthesis in this product. As indicated by thermogravimetric studies (Majchrzak-Kucęba and Nowak 2004), the weight loss of sodalite is 13.9%, which was similar to the weight loss of this sample.

The share of new phases was calculated in the products after synthesis, taking the mass loss recorded on the TG curves of the tested samples (Fig. 3) and the theoretical dehydration of the new phases into account. These calculations indicate that the share of Na-LSX zeolite in A and B samples is comparable and amounts to 42.9% by mass and 41.0% by mass, respectively (Table 3).

The share of hielscherite in the analyzed samples is also similar and amounts to 6.1% by mass and 5.0% by mass, respectively. However, the share of hydrosodalite varies greatly in the tested samples and a clear increase is seen with the increasing concentration of activating NaOH solution.

Grain morphology of the products after synthesis, revealed in the electron microscope observations, confirms the new phases identified by X-ray diffraction (Khemthong et al. 2007; Musyoka et al. 2011; Pekov et al. 2012; Franus et al. 2014; Kwakye-Awuah et al. 2014; Zgureva and Boycheva 2015; Donkor and Buamah 2016; Salih et al. 2019) in the form of:
cubes – na-lSX type zeolite (Fig. 4A–C),
rods – hielscherite (Fig. 4A and 5C),
spherical agglomerates – hydrosodalite (Fig. 4D).

The Na-LSX type zeolite forms crystals with a size of 0.7–4 μm. The smallest ones often form agglomerates, consisting of several or a dozen crystals. The hielscherite rods show a length of 0.4–6 μm, with the sample A being much smaller compared to sample B. The spherical agglomerates of hydrosodalite rarely exceed 3 μm, it is noteworthy that the largest agglomerates occur in sample C. The microscopic observations indicate that the newly formed phases crystallize on the unreacted mineral components of fly ash.
In particular, this is evident in samples A and B, in which common aggregates of mullite, quartz, and enamel are formed with the new phases of tens of micrometers size, and even exceeding 100 μm.

The new phases in products most often create mutual outgrowths. Observations of grain morphology in the electron microscopy reveal a certain order of crystallization of these phases, which is best seen in the sample B in which all three new phases are present. The first to crystallize was hielscherite onto which successively formed the Na-LSX type zeolite, and the whole was covered with hydrosodalite crystallizing at the end.

Mutual outgrowths of the new phases in products are also revealed in the results of tests on the chemical composition in the micro-area (Table 3). These results clearly indicate that the stoichiometric composition of Na-LSX type zeolite, hielscherite or hydrosodalite was not found in any of the tested grains (measuring points). The tested grains are always poly-mineral, and usually one of the new phases dominates in them (Fig. 5). The results of the tests

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**Fig. 4** SEM images of aggregates of the products of synthesis:
A) in sample A; B) in sample B; C) in sample B; D) in sample C

Hi – hielscherite, S – hydrosodalite, X – Na-LSX type zeolite

**Rys. 4.** Obrazy sem skupień produktów syntetyz:
A) w próbce A; B) w próbce B; C) – w próbce B; D) w próbce C

Hi – hielscherite, S – hydrosodalit, X – zeolite typu Na-LSX
on the chemical composition in micro-areas reveal that apart from hydrosodalite, sample C also contains another newly formed phase – hielscherite.

In each of the tested samples of the synthesis products, two grain populations can be distinguished in terms of chemical composition:
- containing SO3,
- not containing SO3 (Table 4).

However, the main chemical components of the tested grains, irrespective of the samples, are SiO2, Al2O3 and Na2O. Their average contents range accordingly 37.51–43.57% by mass, 31.46–37.94% by mass and 9.23–18.10% by mass. The highest SiO2 content occurs in grains of sample B, Al2O3 – in grains of sample A, and Na2O in – in grains of sample C.

The average contents of other chemical components of the grains do not exceed 7% by mass, and in the case of K2O and SO3 rarely exceed 1% by mass. The tendency to decrease the average contents of Fe2O3 and K2O with an increase in the concentration of NaOH of the solution used in the synthesis is noteworthy. Such a variation in the chemical composition of the tested grains may indicate an increase in the share of phases containing sodium with an increase in the concentration of NaOH in the solution, which contributes to an increase in the dissolution of fly ash components containing SiO2 and Al2O3 (mainly enamel and mullite) and overgrowing of the unreacted ash phases with the new phases (Na-LSX type zeolite, hielscherite and hydrosodalite).

The differentiation of SiO2, Al2O3 and Na2O contents according to samples A, B and C are presented on the diagrams (Fig. 6A and 6B). The content ranges of these chemical compo-
Table 4. Basic statistical parameters of the chemical composition of the studied micro-areas

<table>
<thead>
<tr>
<th>Chemical composition</th>
<th>Sample</th>
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<tbody>
<tr>
<td></td>
<td>A (without SO₃)</td>
<td>A (with SO₃)</td>
<td>B (without SO₃)</td>
<td>B (with SO₃)</td>
<td>C (without SO₃)</td>
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<tr>
<td>TiO₂</td>
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The distribution of potassium contents in the studied grains is interesting in the diagram in the Na₂O–K₂O system according to the samples (Fig. 6C). There is a clear tendency to decrease the K₂O contents in the tested grains while increasing the NaOH concentration in the solutions used for synthesis. The geochemical affinity of these two elements is manifested by mutual substitution in many minerals. Therefore, it can be concluded that in the case of a lower concentration of NaOH in the solution (sample A), potassium probably enters the structure of new phases containing sodium (Na-LSX type zeolite) in larger amounts compared to the new sodium phases formed at a high NaOH concentration in the solution (hydrosodalite).

The substitution of potassium in place of sodium affects the morphology of Na-LSX type zeolite (Basaldella and Tara 1995; Hui et al. 2014). The substitution of Na by K favors, as Basaldella and Tara (Basaldella and Tara 1995) prove, that with the increase in the amount
of K in the LSX type zeolite, the sizes of crystals increase, while their morphology changes from octagons to round shapes. Meanwhile, Hui et al. (Hui et al. 2014) claim that with an increase in the amount of K in place of Na in the LSX type zeolite the octagonal crystals are formed in place of round ones, although at the same time the size of the crystals increases and their walls are smoother. Therefore, it seems that the more important factor determining the morphology of the LSX type zeolite are the conditions of the synthesis. In the first case, the hydrothermal synthesis was carried out at 56°C in various time periods (maximum 196 hours) but it was preceded by calcination of the reagents at 800°C for two hours. In the second case, the reagents were aged at 50°C for 24 hours, followed by the hydrothermal synthesis at 100°C, at various time intervals (maximum 10 hours).

A significant correlation occurs between Fe$_2$O$_3$ and TiO$_2$ ($R^2 > 0.82$), regardless of the concentration of the NaOH solution used for the synthesis (Fig. 6D). As it is known, both of these components are mainly found in Fe oxides (magnetite, hematite) in fly ash. A higher contents of Fe$_2$O$_3$ in the A and B samples may indicate that the Fe oxides at such NaOH concentrations of the solution used for synthesis are more stable than the other ash phases. However, the contents of Fe$_2$O$_3$ are lower in sample C, which, however, does not necessarily mean the loss of their stability. The crystallization of more new phases (mainly hydroxosodalite) is so intense that the share of Fe oxides decreases.

Tests on the chemical composition of the grains in the micro-area revealed that each of them is poly-mineral. The unreacted ash phases (quartz, mullite and Fe oxides) are always found in the grain composition. The average number of these phases in grains decreases with increasing NaOH concentration in the solution. The highest average amounts of quartz, mullite and Fe oxides occurred in the grains of the A sample, their total average amount was between 41% by mass and 53% by mass, with a higher share observed in the grains containing SO$_3$ compared to the grains without this component (Fig. 7A).

![Graph](image1.png)  
![Graph](image2.png)

**Fig. 7.** Average shares of the unreacted fly ash phases (A) and new phases (B) in the products of synthesis by population of sample (samples A, B and C)

**Rys. 7.** Średnie udziały nieprzereagowanych faz popiołu lotnego (A) i nowych faz (B) w produktach syntezy według populacji próbek A, B i C
Fig. 8. Differentiation of the contents of the selected chemical components and phases in the studied micro-areas for samples: A, B, C: (A) phases containing sodium and K$_2$O; (B) phases containing sodium and MgO; (C) sum of the phases containing sodium and mullite; (D) Fe spinels and the sum of the phases containing sodium.

Rys. 8. Zróżnicowanie zawartości wybranych składników chemicznych i faz w badanych mikroobszarach dla próbek A, B i C: (A) fazy zawierające sód i K$_2$O; (B) fazy zawierające sód i MgO; (C) suma faz zawierających sód i mullit; (D) spineli Fe i suma faz zawierających sód.
Lower average shares were observed in the grains of sample B (28–30% by mass) and, similarly to sample A, a higher average share was found in the grains containing SO$_3$ compared to the grains without this component. The lowest average amounts of the unreacted components occurred in the grains of sample C, their total amount was about 18% by mass.

The unreacted fly ash phases are covered with the new ash phases. In the tested grains of samples A and B, the dominant new phase is the Na-LSX type zeolite (Fig. 7B). The average share of this component in the grains of sample B is clearly higher than the average share in sample A. It is noteworthy that in the grains containing SO$_3$ the share of Na-LSX type zeolite is slightly lower (44.38% by mass for sample A, 64.86% by mass for sample B) than in the grains not containing this component (58.10% by mass for sample A, 66.33% by mass for sample B).

In the grains of sample C, the dominant new phase is hydrosodalite, the average share of which was above 81% by mass. In the case of this sample, as in the case of samples A and B, a slightly less dominant phase are in the grains containing SO$_3$.

The composition of the new phases in all samples is supplemented by hielscherite, present in small amounts, containing SO$_3$. The average amount of hielscherite tends to decrease phases in synthesis products as the concentration of NaOH in solution increases (Table 4, Fig. 7B).

The new phases containing sodium in the studied grains of the products of synthesis (Na-LSX type zeolite and hydrosodalite) micro-areas show a tendency to decrease the content of potassium and magnesium together with the increase of the share of these phases. The exception is hydrosodalite found in grains of sample B. In this case, with an increase in the share of this component in the grains, both potassium and magnesium tend to increase their quantity (Fig. 8A and 8B).

The decrease in the mullite contents with the increase in the share of the sum of the Na-LSX type zeolite and hydrosodalite is also noteworthy (Fig. 8C). Therefore, it can be assumed that the source of SiO$_2$ and Al$_2$O$_3$ is not only the enamel found in the studied ash but also mullite.

**Conclusions**

In the studied fly ash, originating from the hard coal combustion, the dominant chemical components were SiO$_2$ and Al$_2$O$_3$, while the main phase components were mullite, quartz and hematite, and the significant share of an amorphous substance (glass and unburnt organic substance).

In the products after hydrothermal synthesis, the presence of unreacted fly ash phases was found, as well as new phases which quality and quantity depends on the concentration of NaOH in the solution used for synthesis:

- sample A – Na-LSX type zeolite and hielscherite,
- sample B – Na-LSX type zeolite, hielscherite and hydrosodalite,
- sample C – hydrosodalite and hielscherite.
A new aspect at work is the crystallization of hielscherite, which accompanies both Na-LSX zeolite and sodalite. In hydrothermal conversion of fly ash to zeolites, hielscherite has not yet been reported in products.

The grains in all synthesis products are poly-mineral. However, it was found that the new phases, overgrowing the unreacted phase components of fly ash, crystallize in a certain order. Hielscherite is the first crystallizing phase on which the Na-LSX type zeolite crystallizes and the whole is covered with hydrosodalite.

The share of sodium-containing phases (Na-LSX type zeolite and hydrosodalite) in the products of synthesis increases with increasing NaOH concentration in the solution used for the process. A tendency to decrease potassium and magnesium content was observed along with the increase of the share of these phases in grains. The exception is hydrosodalite present in the products formed in sample B because with the increase in the share of this component in grains, both potassium and magnesium contents increase.

It was also found that at a lower concentration of NaOH solution (sample A), sodium can be replaced with potassium in the Na-LSX type zeolite.

The decrease in the mullite content with an increase in the share of the total amount of Na-LSX type zeolite and hydrosodalite may indicate that the source of SiO2 and Al2O3 is not only the glass found in the studied ash but also the mullite.

The results obtained in this study encourage authors to increase the scale of synthesis in order to have more zeolite material available. This material will be used in tests as an addition to concrete to optimize its care and as an admixture to degraded soils for water retention and as a carrier of plant nutrients.

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**REFERENCES**


The paper presents the results of hydrothermal zeolitization of fly ash from hard coal combustion in one of the Polish power plants. The synthesis was carried out using various NaOH fly ash mass ratio (3.0, 4.0 and 6.0) and the effect of NaOH concentration in the activating solution on composition of synthesized sample was tested. The process was carried out under the following permanent conditions: temperature: 90°C, time – 16 hours, water solution of NaOH (L)/fly ash (g) ratio – 0.025. In the studied fly ash the dominant chemical components were SiO₂ and Al₂O₃, while the main phase components were mullite, quartz and hematite, and a significant share of amorphous substance (glass and unburnt organic substance). After hydrothermal synthesis, the presence of unreacted fly ash phases was found in the products, as well as new phases, the quality and quantity of which depend on the NaOH to fly ash mass ratio used for synthesis:

- for ratio 3.0 – Na-LSX type zeolite and hielscherite,
- for ratio 4.0 – Na-LSX type zeolite, hielscherite and hydrosodalite,
- for ratio 6.0 – hydrosodalite and hielscherite.

The grains in all products of synthesis are poly-mineral. However, it was found that the new phases, overgrowing the unreacted phase components of fly ash, crystallize in a certain order. Hielscherite is the first crystallizing phase, on which the Na-LSX type zeolite crystallizes then, and the whole is covered by hydrosodalite. In the products of synthesis, the share of sodium-containing phases (the Na-LSX type zeolite and hydrosodalite) increases with the increasing concentration of NaOH in the solution used for the process.
SYnteza zeolitu Na-LSX na bazie polskiego popiołu lotnego

Słowa kluczowe
popiół lotny, Na-LSX zeolit, hydrosodalit, hielscherit

Streszczenie
W pracy przedstawiono wyniki badań hydrotermalnej zeolityzacji popiołu lotnego pochodzącego ze spalania węgla kamiennego w jednej z polskich elektrowni. Syntezę przeprowadzono przy różnych stosunkach wagowych NaOH/popiół lotny (3,0, 4,0 i 6,0) i badano wpływ stężenia NaOH w roztworze aktywującym na skład zsyntetyzowanej próbki. Proces był prowadzony w następujących warunkach: temperatura syntez – 90°C, czas syntez – 16 godzin, stosunek roztworu NaOH (L)/popiół lotny (g) – 0,025. W badanym popiele lotnym dominującymi składnikami chemicznymi były SiO₂ i Al₂O₃, natomiast głównymi składnikami fazowymi były mullit, kwarc, hematyt oraz stwierdzono znaczny udział substancji amorficznej (szkliwo i nieprzepalona substancja organiczna). W produktach po hydrotermalnej syntezie stwierdzono obecność nieprzerząganych faz popiołu lotnego, a także nowe fazy, których jakość i ilość uzależnione są od stosunku masowego NaOH do popiołu lotnego:
- dla stosunku 3,0 – zeolit typu Na-LSX i hielscherite,
- dla stosunku 4,0 – zeolit typu Na-LSX, hielscherite i hydrosodalit,
- dla stosunku 6,0 – hydrosodalit i hielscherite.
Ziarna we wszystkich produktach syntetyz są poliminalne. Stwierdzono jednak, że nowe fazy, obrastające nieprzerzągane składniki fazowe popiołu lotnego, krystalizują w określonej kolejności. Hielscherite jest pierwszą krystalizującą fazą, na którym krystalizuje zeolit typu Na-LSX i całość obejmuje hydrosodalit. W produktach syntetyz udział faz zawierających sól (zeolit typu Na-LSX i hydrosodalit) wzrasta wraz ze wzrostem stężenia NaOH w roztworze użytym w procesie.