CO₂ sorption and regeneration properties of fly ash zeolites synthesized with the use of differentiated methods

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Production of fly ash zeolites may be an attractive method for the utilization of solid wastes from the energy sector. Different methods of synthesis often yield a variety of zeolite types, thereby affecting the properties of the resulting materials. The attention paid to carbon dioxide emission reduction technologies fully justifies the study of the sorption behaviours of fly ash zeolites synthesized by different methods. This work investigates the sorption properties of fly ash zeolites synthesized with different methods using CO₂. Sorption capacity and adsorption isotherms were determined following the volumetric method and textural parameters were resolved according to the Dubinin-Astakhov (DA) method. The CO₂ sorption capacity was in the range 0.24–4.16 mmol/g. The relationships between structure and sorption behaviour were studied for each synthesis method. Some strong similarities between commercial zeolites and fly ash zeolites were found. The mechanism for sorption was proved to be physisorption which is fully reversible under selected conditions. The observed trends were used to identify the best sorbent.

It is believed that increasing the amount of CO₂ caused by anthropogenic emission (recently observed to exceed the 400 ppm) is leading to global climate change. Currently, many political and ecological actions are aimed at limiting CO₂ emissions into the atmosphere. Difficulty avoiding emissions in many sectors fully justifies the efforts made to capture carbon dioxide. The most common method is based on the utilization of liquid amines for the removal of CO₂ from flue gases. However, this process has some drawbacks due to amine degradation leading to low CO₂ capture efficiency, high regeneration energy requirements and corrosive effects on separation equipment units¹. Therefore, exploration of differentiated materials for CO₂ capture is needed. One possibility is the use of technologies based on adsorption processes. Adsorbents need to be cost-effective (low cost raw material), have low heat capacity, fast kinetics, high CO₂ adsorption capacity and selectivity, and thermal, chemical and mechanical stability under extensive cycling². The most interesting adsorbents include activated carbons, zeolites and MOFs. MOFs are currently of great importance for CO₂ capture as they have high microporosity, unlimited chemical adjustability and surface functionality compared to conventional porous zeolites and activated carbon. Although these adsorbents work well in CCS technologies, they may not be sufficient in post-combustion processes due to very high temperatures. Additionally, MOFs are extremely expensive relative to zeolites and activated carbon; the price is estimated to be approximately $130–200/kg³. Activated carbon is one of the most widely used solid adsorbents for the capture of CO₂. This is due to its microporous character, adequate pore size distribution, low cost and easy regeneration. However, activated carbon CO₂/N₂ selectivity is relatively low. For the purpose of adjusting the properties of activated carbons to the carbon dioxide sorption, modifications to the surface may be needed, increasing the cost of those materials. Zeolites have been proposed as potential physical adsorbents of carbon dioxide and their adsorption efficiencies depend on size, charge density, and chemical composition of cations in their porous structures⁴. Carbon dioxide capture using zeolites has certain advantages over capture processes that use carbon-based sorbents. One advantage is the higher selectivity of carbon dioxide over nitrogen, which leads to the production of highly purified carbon dioxide streams. The price of commercial zeolite starts at $100/kg. Natural zeolites are the cheapest of all the above mentioned materials, however, it should

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Zeolites were synthesized from fly ash raw materials following three different methods: hydrothermal (Zarębska et al., 2015), fusion (Czuma et al., 2017) and a modified two-step procedure (Czuma et al., 2019). Major and minor element contents were determined by X-ray fluorescence (XRF) analysis with a Philips PW 1404 spectrometer. Mineral identification was carried out by powder X-ray diffraction (XRD) using a Philips X’pert APD diffractometer (with a PW 3020 goniometer), with Cu cathode and a graphite monochromator (in a 20 degree range of 5–50°, with a step-size of 0.02° and measuring time of 1 s per step). A Thermo Scientific Nicolet 6700 FT-IR spectrometer equipped with a DTGS-CsI detector and operating across a spectra range from 400 cm$^{-1}$ to 4000 cm$^{-1}$, 64 scans and 4 data point spacing was used for infrared spectroscopy characterization. Scanning electron microscopy was performed with a Neon 40 Crossbeam workstation, operating at 15 keV.

Specific surface area ($S_{BET}$) measurements of raw fly ash and synthesized zeolites were performed with a Micrometrics ASAP 2020 analyser and calculated using the linear form of the Brunauer-Emmett-Teller (BET) equation, assuming an area of 0.162 nm$^2$ for the N$_2$ adsorbed molecule. Nitrogen adsorption-desorption isotherms were determined at liquid nitrogen temperature (77 K). Sorption experiments using the volumetric method were carried out using equipment for precise physical sorption and chemisorption measurements, with a vapoport adapter and an Autosorb-1-C mass spectrometer (Quantachrome Instruments, USA). Samples were pre-heated to 473 K for 12 hours using a degasser equipped with a vacuum system with a turbomolecular pump and an Autosorb-1-C mass spectrometer (Quantachrome Instruments, USA). Samples were method were carried out using equipment for precise physical sorption and chemisorption measurements, with a vapoport adapter and an Autosorb-1-C mass spectrometer (Quantachrome Instruments, USA). Samples were pre-heated to 473 K for 12 hours using a degasser equipped with a vacuum system with a turbomolecular pump. Measurements were carried out at room temperature. Thermogravimetric analyses (TGA) were performed with a TA Instruments Q50 analyser. Samples of about 10 mg were placed in the furnace, heated to 458 K (10 K/min) in an inert nitrogen atmosphere, kept at constant temperature for 20 minutes and then cooled down to 323 K. When the temperature reached 323 K the gas flow was switched to carbon dioxide and kept for 40 minutes under isothermal conditions. This procedure was repeated for three sorption-desorption cycles.

Table 1. Oxide composition of raw fly ash from the Polish power plant used for zeolite synthesis.

| Component      | Concentration, % |
|----------------|------------------|
| SiO$_2$        | 38.9             |
| Al$_2$O$_3$    | 20.3             |
| Fe$_2$O$_3$    | 9.0              |
| CaO            | 7.7              |
| MgO            | 3.3              |
| Na$_2$O        | 3.0              |
| K$_2$O         | 2.9              |
| SO$_3$         | 1.8              |
| TiO$_2$        | 1.2              |

Results

Materials characterization. Fly ash resulting from the combustion of a pulverized hard coal were collected at the cumulative container of the Polish power plant. These materials can be classified as a standard F class fly ash, according to ASTM C618 (2019). The elemental composition determined by XRF is shown in Table 1.

The mineralogical (XRD) study of the raw materials shows that the fly ash is mainly composed of mullite and quartz, with some calcite (Fig. 1a). The increase of the intensity of the XRD halo pattern in the 20 range between 20° and 35° suggests the presence of a significant amorphous component in the sample$^10$. Treatment with different synthesis methods yielded the formation of at least three different types of zeolites: A and P1 in the hydrothermal method, A and X with the fusion method, P1 and sodalite when fly ash samples were subjected to the two-step zeolite synthesis method (Fig. 1b–d, respectively).

Characterization of the materials by FTIR (Fig. 2) shows the differences between the starting fly ash and the materials synthesized, where the following zeolite characteristic bands can be observed: bands at around 3500 cm$^{-1}$ are associated with the stretching vibration of the hydroxyl groups –OH; bands at 1600 cm$^{-1}$ are characteristic of the bending mode in the water molecule; bands at 1000 cm$^{-1}$ are related to the asymmetric internal
T–O (T = Si, Al) stretching vibrations of the TO4 primary building units; bands at 561 cm$^{-1}$ can be attributed to the D6R T–O–T symmetric stretching; a sharp band at 460 cm$^{-1}$ corresponds to the Si–Al–O bending mode$^{11-15}$. SEM imaging of the fly ash (Fig. 3a) reveals the predominance of aluminosilicate glass spherical particles. All post-synthesis samples display significant changes in their microstructures: cubic crystals characteristic of zeolite A (Fig. 3b), needle-shaped zeolite P1 (Fig. 3b,d) and complex zeolite X octahedra (Fig. 3c), as identified by Fotovat et al., (2009), Koshy and Singh (2016) and Shigemoto and Hayashi (1993), respectively. It is evident that not many unreacted fly ash particles remain after any of the synthesis processes and that the surface of post-synthesis materials is much rougher than the raw material.

Low temperature nitrogen adsorption isotherms are presented in Fig. 4. The following textural parameters were calculated from low-temperature (77 K) nitrogen adsorption isotherms: specific surface area (BET model), micropore volumes (Dubinin Radushkevich model-DR) and total pore volume at p/p$_0$ = 0.95. The results are shown in Table 2.

Figure 1. XRD analyses of raw fly ash (a) and samples synthesized by the H (b), F (c) and TS (d) methods (labels: m = mullite; q = quartz; Ca = calcite; X = zeolite; A = zeolite A; P = zeolite P1; s = sodalite).

Figure 2. IR analysis of starting fly ash and synthesized zeolites.
The specific surface area parameter indicates that all synthesis methods yield zeolites with higher values of \( \text{SBET} \) compared to the raw fly ash (by about a factor of 20 in the H and TS methods and 200 times more than the original surface area after the F synthesis). Micropore pore volumes (MPV) follow a similar trend, with the value for sample F being about 10 times higher than for H and TS. The MPV/SBET ratios are fairly constant, \((3.2–4.1 \times 10^{-4})\), suggesting that dissolution (melting) and precipitation of single crystals (as opposed to, for example, diffusional growth) is most likely to account for the conversion of fly ash into zeolite in all cases.

The adsorbents should have an optimum pore size distribution for the best possible access of the gas molecules through the nano-pores. The most common adsorbents are composed of a combined network of micro, meso and macropores. The only exception is zeolites; their pores are precisely defined.

The affinity between adsorbate and adsorbent is estimated based on the slope of isotherms at low pressures. The best material should have a steep isothermal slope which indicates high CO\(_2\) sorption capacity at low concentrations. A milder slope angle of the carbon dioxide adsorption isotherms is associated with lower capture.

**CO\(_2\) sorption experiments.** Carbon dioxide sorption experiments were carried using two different methods providing complementary information. The aim was to carry out a comparative study of sorption capacity, textural parameters and preliminary regeneration possibilities between synthesized materials and two commercial zeolites. Such a comparison enabled the evaluation of their potential utilization.

The pore size distribution (PSD) of samples was determined through measurement of a CO\(_2\) adsorption isotherm. PSD measurements are commonly performed using nitrogen or argon at 77 K and 87 K, respectively. At these low temperatures, kinetic limitations can have an effect on the process reaching equilibrium during isotherm measurements.
The study of the adsorption of a wide range of microporous materials is commonly addressed following the equation proposed by Dubinin and Astakhov (1971) (Eq. 1):

\[
W = W_0 \exp \left[ - \left( \frac{-RT \ln P/P_0}{E} \right)^\varphi \right]
\]

where:
- \( W \) – weight adsorbed at \( P/P_0 \) and \( T \)
- \( W_0 \) – total weight adsorbed
$E$ – characteristic energy of adsorption ($E = E_0 \beta$)

$E_0$ – characteristic energy of adsorption for a standard vapour

$\beta$ – coefficient of affinity

$n$ – non-integer value (typically between 1 and 3)

This approach requires the calculation of $n$ and $E$ parameters by non-linear curve fitting.

The values can then be used to determine the pore size distribution of the material through the application of the expression proposed by Medek:

$$\frac{d(\frac{W}{W_0})}{dr} = 3n \left( \frac{k}{E} \right)^n r^{-(3n+1)} \exp \left[ - \left( \frac{k}{E} \right)^n r^{-3n} \right]$$

(2)

where $k$ is the interaction constant for CO$_2$ and N$_2$, the values are typically 1.5 and 2.96 kJ·nm$^{-3}$·mol$^{-1}$, respectively, with a pore radius $r$.

The results of volumetric tests are presented on the adsorption isotherm in Fig. 5, where $a$ is the adsorption capacity value and $p/p_0$ is relative pressure.

Carbon dioxide isotherms were used to calculate sorption capacities (Table 3) and the differential pore distribution $dV/dD$ was calculated using the Dubinin-Astakhov method (Fig. 6).

Characterization of sorption properties was carried out by TGA, the results are presented in Fig. 7. A sorption temperature of 323 K was chosen based on previous work on zeolite 4 A. Desorption temperature (458 K) was chosen on the assumption that such a temperature can easily be attained in industrial applications using waste heat.

**Discussion**

Zeolites synthesized from fly ash by different methods showed different conversion efficiencies and carbon dioxide sorption capacities. Volumetric measurements indicated that the fusion method (F) yielded the best sorption capacity, approximately three times higher than the hydrothermal (H) method and almost 20 times higher than the two-step synthesis (TS). The sorption capacity of both F and H samples increases rapidly at low $p/p_0$ values, indicating the microporous character of the sample surface. This is confirmed by the high surface area calculated from both N$_2$ and CO$_2$ measurements, accounting for the observed high conversion ratio of fly ash into crystalline zeolite. The isotherms of samples H and F are similar to those of commercial zeolites A and X. Unprocessed fly ash does not show any evidence of carbon dioxide sorption properties.

| Sample symbol     | CO$_2$ sorption capacity, mmol/g | DA Micropore volume, cm$^3$/g | S$_{BET}$ Surface area, m$^2$/g | DA Pore diameter, nm |
|-------------------|---------------------------------|-------------------------------|---------------------------------|----------------------|
| fly ash           | 0.03                            | 0.006                         | 11                              | 1.2                  |
| H                 | 1.20                            | 0.070                         | 162                             | 0.86                 |
| F                 | 4.16                            | 0.227                         | 549                             | 0.84                 |
| TS                | 0.24                            | 0.043                         | 50                              | 1.06                 |
| Commercial zeolite A | 3.88                          | 0.202                         | 506                             | 0.84                 |
| Commercial zeolite X | 5.03                          | 0.261                         | 652                             | 0.84                 |

Table 3. Comparison of results from volumetric sorption of carbon dioxide test with textural parameters calculated from carbon dioxide isotherms.
The values of specific surface area calculated from low temperature nitrogen sorption $S_{BET}$ and the values calculated from the carbon dioxide sorption $S_{BETCO2}$ cannot be compared. However, common trends can be observed. In the case of $S_{BET}$, the values obtained for samples H and TS are of the same order. Once the analogical comparison is made for $S_{BETCO2}$, the value obtained for H is four times higher. Such observation proves the micro-porous character of sample H. It can be stated that in the case of microporous materials, the $S_{BET}$ analysis with the use of low temperature nitrogen sorption may not give sufficient data for comparison.

Differential pore size distribution (PSD) calculated from the DA method presented in Fig. 6 represents the trend of decreasing sorption properties along with the decreasing number of micropores. This proves that the presence of the micropores determines the effectiveness of the sorbent. DA pore mean diameter values presented in Table 3 show that in the case of commercial zeolites and samples H and F, the mean diameter is relatively low. In the case of unprocessed fly ash, the value of 1.2 nm should be interpreted as intergrain spaces, since fly ash (as will be proven in the following section), does not exhibit porous characteristics. The same approach could be applied to sample TS.

Mass change evolution in sorption experiments monitored by TGA shows an initial drop for all zeolites analysed, mainly due to desorption of water molecules present in intercrystalline voids. It may be assumed that a

Figure 7. Temperature (dotted) and mass change (solid) evolution of sorption experiments performed on unprocessed fly ash (a), H zeolite (b), F zeolite (c), TS zeolite (d), commercial zeolite A (e), and commercial zeolite X (f).
higher mass decrease in the initial stages of the experiment indicates a higher content in the zeolite phase. Sample 
F, together with commercial zeolites, shows the larger desorption (more than 15% of the original mass), followed 
by samples H and TS (both at about 10% mass loss), indicating the higher conversion rate of fly ash into zeolite 
by the fusion method.

TGA of samples H and F showed a relatively high increase in mass after the gas inlet was switched to CO₂. No 
such an increase was observed in sample TS. This may be due to the low conversion rate of fly ash into zeolite 
of the two-step method, and/or the presence of different types of zeolites in the samples. As evidenced by XRD analysis, 
sample TS contains type P1 zeolites and sodalite. Due to the presence of very small pores, neither of these 
substances is believed to have a good sorption capacity18,19.

On the other hand, when CO₂ is introduced in the system, commercial zeolite X experiences a lower mass 
gain compared to commercial zeolite A despite the fact that its micropores are larger than those in zeolite A. This 
bias has also been reported in previous work (e.g., Bukalak et al., 2012).

It should be noted that both the increasing amount of micropore share in the pore volume and the presence of 
different metal phases of fly ash zeolites, provides stronger CO₂/N₂ selectivity (Zgureva, 2016). It is confirmed in 
the literature that commercial zeolites A and X present CO₂ selectivity over N₂ (Myers, 1973).

There are some indications that the CO₂ affinity of zeolites may also be connected with their Si/Al ratio20; the 
lower value of this parameter the better carbon dioxide sorption properties21.

In TGA experiments with samples F and H, very small decreases in mass sorbed (~2%) were observed at each 
cycle, suggesting good regeneration possibilities. Although the mass seemed to remain essentially unchanged 
after regeneration, the slight decrease may be associated either with remains of physisorbed CO₂ (which may be 
responsible for blocking entrances to the pore network), or to slight structural changes taking place during the 
heating process. It must be taken into account that chemisorption may also occur, resulting in the formation of 
carbonates and carboxylates21. This has been attributed to the presence of unreacted calcium oxide, however, no 
such phases have been identified in this work.

Analogous experiments were performed using a heating temperature of 573 K (see Supplementary 
Appendix 1). The outcome of these experiments proved that the use of a higher temperature avoids decreases in 
sorption capacity, as the mass of samples in the following cycles did not change. This is complementary informa-
tion indicating that the use of higher temperatures allows for full regeneration and maximum cycling capacity, 
also confirmed that the process of binding of CO₂ is physisorption.

The results from the experimental procedure presented in this work indicate that unprocessed fly ash does not 
have any significant sorption capacity. Samples H, F, commercial zeolite A and commercial zeolite X, seem to have 
relatively short CO₂ saturation times22.

The SEM study revealed clear textural differences among the zeolites synthesized by different methods. 
Although this is difficult to correlate quantitatively with the physical parameters determined by the other experi-
mental and characterization procedures, it is evidence that the microstructural basis is the explanation for the 
observation that the fusion method yields the highest conversion efficiency of all the methods studied in this 
work.

Conclusions

Compared to unprocessed fly ash, fly ash zeolites exhibit sorption capacity related to carbon dioxide. The sorp-
tion capacity was the lowest for the TS sample (0.24 mmol/g), the H sample reached a value of 1.2 mmol/g, and 
the highest for fly ash zeolite was observed in the F sample with a value of 4.16 mmol/g. The changes in the 
value occur according to the synthesis method and can be associated with textural parameters like the S BET value 
(sample TS 34 m²/g, sample H 39 m²/g, sample F 414 m²/g). Similarities of fusion synthesized materials and com-
mercial zeolites can be observed, proving that this synthesis method is highly efficient. Specific surface area cal-
culations with the use of two gases provide information on the microporous character of the sample, indicating 
that nitrogen low temperature sorption may be insufficient to compare samples. Pore size distribution proves that 
the presence of the micropores determines the efficiency of the sorbent. Both volumetric and thermogravimetric 
tests indicate that a fusion synthesized sample, followed by hydrothermally synthesized sample present very good 
and good sorption capacity characteristics, two step synthesized sample sorption properties are much lower. The 
reason for this is the presence of different types of zeolites in the samples, the Si/Al ratio and the yield of zeolite 
present in the sample. Thermogravimetric tests provide information about regeneration properties of analysed 
samples, regarding the energy penalty for the purpose of regeneration. This information also confirms that a 
physical adsorption process is taking place. Comparing structure with the sorption results provides evidence of a 
correlation between the amount of crystals and sorption capacity.

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Author contributions
N.C. and P.B. conducted the experiments. J.S., N.C. and I.C. analysed the results and interpreted the data. K.Z. and I.C. conceived the experiments. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

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