Investigation of the Sorption Properties of Ore Materials for the Removal of Sulfur Dioxide from Exhaust Flue Gases of Power Plants

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Abstract

The prospects of using a natural material – ferromanganese nodules (FMN) from the Gulf of Finland – as the SO₂ adsorbent are discussed. The starting material was studied as pellets and powder using X-ray fluorescence spectroscopy, XRD, BET, and mechanical strength analysis; dependences of physicochemical parameters of the material on heat treatment at 100-1000 ºC were found. FMN samples were tested in the process of SO₂ sorption. The sorptive capacity of FMN samples for SO₂ was found to increase with humidity of adsorbents; FMN samples with a humidity above 40% were shown to be promising for the removal of sulfur dioxide from gases.

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1. Introduction

Generation of electrical and heat energy is accompanied by a large-scale interchange of matter and energy with the environment, thus producing a detrimental effect that should be remediated.

Catalytic combustion is the way to radically improve the efficiency of power production and minimize its detrimental effect on the environment [1-4]; however, electrical and heat energy is now produced mostly at conventional power plants, with coal playing the dominant role in the fuel balance.

Modern power plants with an output of 2.4 million kW consume up to 20 thousand tons of coal per day and emit to the atmosphere up to 680 tons of SO₂ and SO₃ (at a 1.7% sulfur content in the inlet fuel).

The content of sulfur dioxide can be decreased by chemisorption purification, which is able to completely remove sulfur dioxide from the exhaust flue gases.

The materials containing oxides of alkali-earth elements, CaO and MgO, are conventionally used as sorbents [5-7]; however, such materials are quite expensive, which stimulates a search for alternative sorption systems.

Of particular interest for the removal of sulfur dioxide from flue gases is the use of inexpensive and harmless natural sorbents containing metal oxide and hydroxide compounds. In a series of works [8-11], T.J. Bandoz and co-authors studied the adsorption of sulfur dioxide on inexpensive sorbents obtained by the pyrolysis of sewer sludge at 400-900 ºC. These sorbents contain 20-40% carbon, up to 3-5% Fe and 4-5% Ca as oxides as well as oxides of other metals: Mg, Cu, Zn, Cr, Al. High specific surface area (up to 880 m²/g) and pore volume (up to 0.46 cm³/g) along with the presence of metal oxides in the sorbent composition provide quite a high dynamic capacity for sulfur dioxide (up to 50 mg/g). Investigation of sulfur dioxide adsorption (0.3% in air) at room temperature revealed that humidification of the gas stream (up to a relative humidity of 80%) results in a more efficient adsorption of SO₂; under such conditions, dynamic capacity of the sorbent is approximately two times higher as compared to the dry gas [12]. The beneficial effect of water on SO₂ adsorption is attributed to the formation of surface hydroxyl groups in metal oxides. Manganese dioxide in the composition of manganese ores and specially prepared adsorbents serves as an efficient adsorber of sulfur dioxide and hydrogen sulfide [13-17]. A comprehensive review of the
catalytic and sorption properties of ferromanganese ores was made by M. Nitta [18], who demonstrated that ferromanganese ore is a promising material for purification of gases from toxic impurities, in particular sulfur dioxide, hydrogen sulfide, ammonia, and heavy metal ions (Hg, Pb, Cd and others).

For the first time a method for the application of manganese ore as a sorbent of sulfur dioxide was reported in 1967 as a US patent [19]. It was shown that the ore is an efficient adsorber of SO$_2$ as a component of waste gases from boilers and furnaces at a concentration of 0.04-1.2% and WHSV of 1000 h$^{-1}$ and can be used for purification of ambient air from SO$_2$ at the initial concentration of 25 ppm or H$_2$S (10 ppm) at WHSV of ca. 10000 h$^{-1}$. The removal of SO$_2$ from gases is accompanied by accumulation of sulfates in the sorbent, which decreases its adsorption activity. When more than 15 wt.% SO$_2$ with respect to the starting ore weight is adsorbed, the adsorbent is regenerated to remove the accumulated sulfates by washing with water; after that the sorbent can be reused. Therewith, efficiency of the sorbent remains high for a long time. As shown in [20], the adsorption of sulfur dioxide by a water pulp containing 3-20% of the ore provides a 95-99% purification and virtually a complete sulfation of nickel, cobalt and manganese.

Thus, the information available from scientific and patent literature makes it possible to estimate the prospects of using ferromanganese ore as a SO$_2$ sorbent. In this connection, of particular interest are the shelf ferromanganese nodules (FMN), whose rich deposits have been found in the water areas of Russian seas. 50% of the ore component of such nodules is represented by complex composites of Mn and Fe hydroxides and oxides. Up to 90% of the ore substance is in the X-ray amorphous state. Manganese minerals in the nodules include birnessite, buserite, cryptomelane and the X-ray amorphous mass of a wad-psimelane composition. The ferric component of the nodules is represented by ferrihydrite, hydrogoethite and lepidocrocite [21].

Our work was aimed to study physicochemical characteristics of ferromanganese nodules from the bottom of the Gulf of Finland and to assess the possibility of their use as adsorbents of sulfur dioxide.

2. Experimental

2.1. Materials

Sorbent of SO$_2$ were prepared using the pelletized and powdered material “Ore of ferromanganese nodules from the Gulf of Finland” Specs 0731-001-50855050-2005”.

2.2. Methods

2.2.1. Determination of the chemical composition

The elemental composition of the samples was determined by X-ray fluorescence spectroscopy on a VRA-30 analyzer with the Cr anode of X-ray tube. Prior to measurements, pellets of the test sample with the addition of boric acid or cellulose were prepared. A result of the analysis was obtained as the arithmetic mean of two parallel measurements diverging from one another by no more than 2%. The allowed relative total measurement error was ±0.05% at a confidence level of 0.95.

2.2.2. Determination of the phase composition

Phase composition of the samples was determined by analyzing the diffraction patterns obtained on an HZG-4 diffractometer using monochromatic Cu K$_\alpha$ source. The observed phases were identified with the use of JCPDS radiographic database.

2.2.3. Specific surface area and pore volume measurements

Pore structure of the samples was examined by the low-temperature nitrogen adsorption on an ASAP-2400 (Micromeritics, USA) instrument at the liquid nitrogen temperature, 77 K, after pretreatment of the samples at 150 ºC and a residual pressure below 0.001 mm Hg; the samples were subjected to a standard pretreatment by the BET and Barrett-Joyner-Halenda (BJH) methods.

2.2.4. Bulk density measurements

Bulk density was measured according to GOST 16190. A dried sample was placed in a 100 cm$^3$ volumetric flask and packed by shaking down. After that, weight of the sample in this volume was determined. Bulk density ($\rho$) expressed in g/cm$^3$ was calculated by the formula: $\rho = m/V$, where $m$ is the sample weight, g; and $V$ is the sample volume, cm$^3$. A result of the measurement was obtained as the arithmetic mean of two parallel measurements diverging from one another by no more than 0.01 g/cm$^3$. The allowed total measurement error was ±0.005 g/cm$^3$ at a confidence level of 0.95.

2.2.5. Mechanical strength measurements

Mechanical strength was measured using an MP-9C instrument intended for measuring the strength of porous dispersed solids under static conditions.
The essence of the method consists in measuring the breaking force applied to the generating line of each pellet of the catalyst sample between two parallel planes. For testing, no less than 30 pellets were selected from each sample. The pellets should not have visible damages and shape defects. The length and diameter of the pellets were measured once using a caliper; after that, the pellets were dried at a temperature of 110 ± 10 ºC at least for 2 h. The dried sample was transferred into an excicator and cooled to room temperature. For measurements, a metal template represented by a plate with a cylindrical pit in the middle was placed on a stage. The prepared pellet was mounted in the pit; the lower stage was moved to the upper plane of the instrument using a hoisting gear. The pellet was uniformly loaded until its failure occurred. The moment of the pellet failure was fixed visually from the reverse motion of the pointer. A load that destructs the sample was indicated by the detector at the moment of failure. Mechanical strength of the pellets (P_i) expressed in MPa (or in kg/cm^2 * 10) was calculated as a ratio of the breaking force to the sectional area using the formula

\[ P_i = \frac{N \cdot A}{S} \cdot 10, \]

where \( N \) is the indicated value, scale division; \( A \) is the calibration factor equal to 0.6025; \( S \) is the sectional area of the test pellet, cm^2, which is calculated as

\[ S = L \cdot D, \]

where \( L \) is the length of a cylindrical pellet, cm; and \( D \) is the pellet diameter, cm.

A result was represented by the arithmetic mean of 30 parallel measurements.

2.2.6. Moisture capacity measurements

Moisture capacity characterizes the pore volume of a sample accessible to water. To measure moisture capacity with respect to water, the weighed sample \( (m_0) \) was placed in a capped glass weighing bottle and poured with distilled water so that the water level was 1 cm above the pellets. After 30 min, water was decanted from the pellets. The pellets were placed on a plastic sieve and subjected to air blasing to remove excess moisture. The sample impregnated with water was weighed \( (m_1) \); after that, moisture capacity was calculated by the formula

\[ \text{Moisture capacity} = \frac{(m_1 - m_0)}{m_0}, \]

2.2.7. Sorptive capacity measurements

The experiments on SO_2 sorption were carried out in a laboratory setup. Its schematic is displayed in Fig. 1.

A SO_2-containing mixture (1% SO_2 in air) from vessel (1) was delivered to the reactor through stopcock (2) and gas batching unit BDG-85 (3). The adsorbent sample (1.1 g for dry samples with a humidity of 14-18%; 1.5 g for wet samples with a humidity of 43-50%), a 0.5-1.0 mm fraction, was placed in glass reactor (4) with the inner diameter of 6 mm. The SO_2-containing mixture was fed at a rate of 60 ml/min. SO_2 sorption was performed at room temperature, and then at 50, 90, 150 and 350 ºC.

The concentration of SO_2 in the gas mixture after the reactor was determined by a Kristall-2000M gas chromatograph equipped with a FID detector. Argon was used as a carrier gas. The gas mixture was separated on a chromatographic column HayeSep C + 0.5% H_3PO_4, Ø 3 mm, L = 3 m at a column temperature of 100 ºC. The method of analysis was described elsewhere [22].

![Fig. 1. Schematic of the setup for investigation of SO_2 sorption: 1 – a vessel with SO_2-containing mixture (1%SO_2 in air); 2 – stopcock; 3 – gas batching unit BDG-85; 4 – reactor; 5 – Kristall-2000M gas chromatograph.](image)

2.2.8. Calculation of the sorptive Capacity for SO_2

Sorptive capacity for SO_2 before a breakthrough of sulfur dioxide from the reactor was expressed in grams of adsorbed SO_2 per gram of the moisture-free adsorbent and calculated as

\[ \xi = \frac{V \cdot C_{SO_2} \cdot M_{SO_2} \cdot \tau_o}{22400 \cdot m_c}, \]

\( \xi \) – sorptive capacity, g_{SO2}/g_{dry adsorbent}
\( V \) – feed rate of SO_2-containing mixture to the reactor, cm^3/min
\( C_{SO_2} \) – concentration of SO_2 in the mixture, volume fractions
\( M_{SO_2} = 64 \text{ g/mol – SO}_2 \text{ molar mass} \)
\( \tau_o \) – time before the start of SO_2 outlet from the reactor, min
\( m_c \) – weight of the moisture-free adsorbent, g
3. Results and Discussion

3.1. Investigation of physicochemical parameters of the starting material

The FMNs as raw materials exist as humid pellets and dry powder. Dependence of properties of pellet and powder on temperature treatment was studied.

The average sample was taken from a batch of FMN ore with the pellet size of 7-12 mm. This sample was used to measure the total humidity ($\Sigma_{\text{hum}}$), specific surface area ($S_{\text{BET}}$), moisture capacity ($V_{\Sigma}$ with respect to $H_2O$), and mechanical crushing strength ($P$). These parameters were studied in dependence on the calcination temperature. Chemical analysis for the main elements was made with a sample calcined at 550 ºC. The results are listed in Table 1.

The average sample from a batch of powdered FMN was used to find the total humidity ($\Sigma_{\text{hum}}$) and specific surface area ($S_{\text{BET}}$); chemical analysis for the main elements was performed with a sample calcined at 550 ºC (Table 2).

### Table 1

| No. | Drying and calcination temperature, ºC | $\Sigma_{\text{hum}}$, % | $S_{\text{BET}}$, m$^2$/g | $V_{\Sigma}$ with respect to $H_2O$, cm$^3$/g | $P$, kg/cm$^2$ | Chemical composition (550 ºC), wt. % |
|-----|--------------------------------------|---------------------------|-----------------|---------------------------------|----------------|---------------------------------|
| 1.  | Starting humid sample                | 50.3                      | not determ.     | not determ.                     | not determ.    | Fe = 25.21                      |
| 2.  | 25/24 h                              | 19.3                      | 120             | not determ.                     | not determ.    | Mn = 23.29                      |
| 3.  | 25/24 h                              | 15.9                      | 150             | 0.65                            | 6.0            | Al = 5.26                       |
| 4.  | 200/3 h                              | 9.5                       | 154             | 0.76                            | 6.0            | Si = 9.86                       |
| 5.  | 300/3 h                              | 1.9                       | 1488            | 0.79                            | 5.5            | P = 4.84                        |
| 6.  | 400/3 h                              | 1.5                       | 152             | 0.89                            | 5.2            | S = 1.13                        |
| 7.  | 550/3 h                              | 0                         | 78              | 0.66                            | 5.5            | K = 3.33                        |

### Table 2

| No. | Drying and calcination temperature, ºC | $\Sigma_{\text{hum}}$, % | $S_{\text{BET}}$, m$^2$/g | Phase composition | Chemical composition (550 ºC), wt. % |
|-----|--------------------------------------|---------------------------|-----------------|-------------------|---------------------------------|
| 1.  | Starting sample                      | 20.5                      | not determ.     | - X-ray amorphous phase | Fe = 25.13 |
|     |                                       |                           |                 | - phases not identified from radiographic database | Mn = 23.39 |
| 2.  | 25/24 h                              | 17.8                      | 113             | The same as No.1    | Al = 4.91 |
| 3.  | 100/3 h                              | 14.3                      | 153             | The same as No.1    | Si = 10.75 |
| 4.  | 200/3 h                              | 9.0                       | 155             | The same as No.1    | P = 4.98  |
| 5.  | 300/3 h                              | 1.6                       | 158             | - $\alpha$-SiO$_2$  | S = 1.01  |
|     |                                       |                           |                 | - magnetite traces  | K = 3.31  |
|     |                                       |                           |                 | - unidentified phase | Mg < 2   |
| 6.  | 400/3 h                              | 1.0                       | 150             | The same as No.5    | Na < 3   |
| 7.  | 550/3 h                              | 0                         | 65              | - $\alpha$-SiO$_2$  | Ca = 1-2 |
|     |                                       |                           |                 | - magnetite traces  |                   |
|     |                                       |                           |                 | - X-ray amorphous phase |                   |
|     |                                       |                           |                 | - phase not identified from radiographic database, close to KFeSi$_3$O$_8$ |                   |
| 8.  | 700/3 h                              | 0                         | 12              | - hematite Fe$_2$O$_3$ |                   |
|     |                                       |                           |                 | - magnetite Fe$_3$O$_4$ |                   |
|     |                                       |                           |                 | - $\alpha$-SiO$_2$  |                   |
|     |                                       |                           |                 | - unidentified phase |                   |
| 9.  | 1000/3 h                             | 0                         | 1.0             | - hematite Fe$_2$O$_3$ |                   |
|     |                                       |                           |                 | - magnetite Fe$_3$O$_4$ |                   |
|     |                                       |                           |                 | - $\alpha$-SiO$_2$  |                   |
|     |                                       |                           |                 | - two unidentified phases |               |
The comparison of Table 1 and Table 2 data shows that the FMNs in pellet and powder forms have close characteristics of the total humidity and specific surface area. The FMN dry sample (25 ºC) and samples calcined at 100-400 ºC have quite high values of specific surface area, 150-158 m²/g, which is determined by mesoporous structure and the number of fine pores. The N₂ adsorption isotherm of samples and the differential pore size distribution are presented in Figs. 2 and 3 correspondingly.

Fig. 2. N₂ adsorption-desorption isotherm of FMNs samples treated at 25, 100 and 400 ºC.

Fig. 3. The differential pore size distribution in FMN samples treated at 25, 100, and 400 ºC.

Investigation of the pore structure of FMN samples by nitrogen adsorption and determination of the pore volume with respect to water showed that an increase in the calcination temperature from 100 to 400 ºC decreases the number of fine pores with the size of 30-40 Å (Fig. 3) and increases the total pore volume from 0.65 to 0.89 cm³/g (Table 1, Samples No.3-6). This is caused not only by transformation of the skeleton of FMN samples due to sintering of the fine pores but mostly by burnout of organic inclusions and formation of cavities in the pellets. As the calcination temperature is raised above 550 ºC, the mesopore volume sharply decreases (Fig. 4) and large pores gradually sinter, which is accompanied by deterioration of all parameters of the pore structure, both the specific surface area and the total pore volume (Table 1, sample 7).

Fig. 4. The differential pore size distribution in FMN samples treated at 400 and 550 ºC.

The phase composition of a dry sample of FMN ore includes the X-ray amorphous phase and phases that cannot be identified using the radiographic database. To identify such phases, the FMN sample was heat treated at 200-1000 ºC. We expected that this treatment would lead to crystallization of compounds constituting the ore, which could show up in the diffraction pattern. This experiment revealed the crystallized phases of hematite, magnetite and quartz in the samples calcined at 700 ºC, although trace amounts of the magnetite phase were present also in the low-temperature samples (Tₐₜₜ of 300 and 500 ºC). The phase composition of sample 1000 ºC is similar to that of sample 700 ºC, except for the appearance of an additional phase, which was not identified using the radiographic database. The crystallized phases of manganese compounds are not observed in the diffraction patterns of the samples calcined at 1000 ºC.

The study has demonstrated that the ore of ferromanganese nodules has a developed pore system and a high content of Mn and Fe compounds in the amorphous state; this forms a basis for possible applications of such materials as the promising sorbents of acid gases.

3.2. A study of the sorption properties of FMN

In the experiments, the sorptive capacity of FMN for SO₂ was examined in dependence on humidity of the starting FMN at the adsorption temperature of 25 ºC and on the temperature of adsorption process on dry and wet samples.
The dependence of sorption properties of FMN samples on their humidity was studied with the following FMN samples: the starting sample with a humidity of 43%; the sample that was dried at 100 °C and supplemented with water to a humidity of 43%; the sample that was dried in air to a humidity of 18%; and the sample that was dried at 100 °C to a humidity of 14%. Results of the study are listed in Table 3.

The dependence of sorption properties of the samples on the sorption temperature was examined using the FMN samples: the dry sample with a humidity of 43%; the sample dried at 100 °C to a humidity of 14%; and the sample dried at 100 °C and supplemented with water to a humidity of 43%. The results obtained are shown in Table 4 and Fig. 5.

Sorptive capacity of FMN samples for SO₂ was found to increase with humidity of the sorbents. In the case of a dry SO₂-containing gas mixture fed on dry adsorbents at 25 °C, total capacity of the samples was 0.05-0.06 g SO₂/g moisture-free adsorbent. Sorptive capacity, which was measured before a breakthrough of SO₂, of the samples of wet FMN or dry FMN moistened to a humidity of 43% was 0.21-0.22 g SO₂/g moisture-free sorbent. This suggests that at the beginning of the process SO₂ is oxidized by air oxygen to SO₃, which readily dissolves in water with the formation of sulfuric acid. After that, H₂SO₄ is adsorbed by Mn and Fe oxides to yield respective sulfates until deactivation of the samples. As the sorption temperature increases to 90 and 150 °C, which decreases humidity of the samples, the sorptive capacity for SO₂ decreases (Table 4 and Fig. 5). A further raising of the process temperature to 350 °С increases sorptive capacity of the samples, which may be related to a 1.5-fold extension of the pore volume.

Table 3
Sorptive capacity of FMN samples for SO₂ versus FMN humidity

| No. | Sample | Sample weight, g | Adsorption temperature, °C | Sorption time, min | Sorptive capacity, g SO₂/g moisture-free sorbent (0.5-1 mm pellets) |
|-----|--------|-----------------|---------------------------|--------------------|---------------------------------------------------------------|
| 1   | FMN – wet Humidity = 43% | 1.5 | 25 | 111 | 0.22 |
| 2   | FMN – 100°+H₂O Humidity = 43% | 1.5 | 25 | 107 | 0.21 |
| 3   | FMN – air dried Humidity = 18% | 1.1 | 25 | 32 | 0.06 |
| 4   | FMN – 100° Humidity = 14% | 1.1 | 25 | 26 | 0.05 |

Table 4
Sorptive capacity of FMN samples for SO₂ versus sorption temperature

| No. | Sample | Sample weight, g | Adsorption temperature, °C | Sorption time, min | Sorptive capacity, g SO₂/g moisture-free sorbent (0.5-1 mm pellets) |
|-----|--------|-----------------|---------------------------|--------------------|---------------------------------------------------------------|
| 1   | FMN – 100° Humidity = 14% | 1.1 | 25 | 25 | 0.05 |
|     |        |                 | 50 | 34 | 0.06 |
|     |        |                 | 90 | 12 | 0.02 |
|     |        |                 | 150 | 10 | 0.02 |
|     |        |                 | 350 | 46 | 0.08 |
| 2   | FMN – wet Humidity = 43% | 1.5 | 25 | 111 | 0.22 |
|     |        |                 | 90 | 47 | 0.09 |
|     |        |                 | 350 | 37 | 0.1 |
| 3   | FMN – 100°+H₂O Humidity = 43% | 1.5 | 25 | 107 | 0.21 |
|     |        |                 | 50 | 76 | 0.15 |
|     |        |                 | 90 | 33 | 0.065 |
|     |        |                 | 150 | 27 | 0.05 |
|     |        |                 | 350 | 44 | 0.09 |
4. Conclusion

Physicochemical properties of FMN samples (as pellets and powder) were studied in dependence on the calcination temperature. The FMN samples calcined over the temperature range of 100-400 °C were shown to have quite high specific surface areas, 150-158 m$^2$/g, which are determined by the number of fine pores. When the calcination temperature exceeds 550 °C, the mesopore volume sharply decreases and large pores gradually sinter, which is accompanied by deterioration of all parameters of the pore structure, both the specific surface area and the total pore volume. According to XRD data, the starting FMN sample is characterized by an X-ray amorphous phase. Well crystallized phases of iron oxides corresponding to hematite and maghemite are observed for the samples calcined at 700 °C and higher temperatures.

FMN samples were tested in the sorption of SO$_2$. It was found that the sorptive capacity of FMN samples for SO$_2$ increases with humidity of adsorbents. The FMN samples with a humidity above 40% were shown to be promising for the removal of sulfur dioxide from gases. Sorptive capacity of wet FMN can be as high as 0.22 g SO$_2$/g moisture-free sorbent, which is an appropriate value for industrial application of ferromanganese nodules in the removal of SO$_2$ from flue gases.

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