Experimental Examination of Sulfur Dioxide Dry Removal from a Mixture of Gases by Calcium Oxide, Calcium Carbonate and Dolomite

Durić, Slavko*; Brankov, Saša
Faculty of Technical Sciences, 6, Trg Dositeja Obradovića, Novi Sad, SERBIA

Stanojević, Petko
Railways of Republika Srpska, 71, Svetog Save, Doboj, BOSNIA AND HERZEGOVINA

Božičković, Ranko
Faculty of Transport and Technical Sciences, 52, Vojvode Mišića, Doboj, BOSNIA AND HERZEGOVINA

ABSTRACT: This paper refers to an experiment of SO$_2$ absorption to the three different sorbents: calcium-oxide (CaO), calcium-carbonate (CaCO$_3$) and dolomite (CaMg(CO$_3$)$_2$). In the reactor under the condition of oxidation atmosphere with the mass of sorbent (sample) of 100 g with fractional composition of 500-700µm, the gas temperature varied from 200°C and 400°C. The examinations indicated that the highest degree of sorbent utilization of 14 % was determined for dolomite (CaMg(CO$_3$)$_2$) and the binding degree of SO$_2$ from 65-80% and the lowest for calcium oxide (CaO) of 4% and the binding degree of SO$_2$ from 34-60% at the reaction temperature of 200°C. When the reaction temperature is 400°C the degree of sorbent utilization is a bit lower. The achieved results indicate the injection capability of powdered sorbents CaMg(CO$_3$)$_2$, CaCO$_3$, and CaO in the areas of lower temperatures as it is the convective part of the boiler or furnace flue of the thermo power plant.

KEY WORDS: Sorbent, Absorption degree of SO$_2$, Reaction temperature, Particle diameter.

INTRODUCTION

During dry Flue Gas Desulfurisation (FGD) processes the dry particles of the reagent are injected into the stream of gases, which absorb SO$_2$ through chemical reactions. Most often used reagents are calcium compounds CaCO$_3$, CaO or Ca(OH)$_2$. The industrial examinations show that the binding degree of SO$_2$ from flue gases to the sorbent particles is usually function of stoechiometric relation of calcium share in injected sorbent and sulfur share from coal and it is in a wide range of 30-70% utilizing CaCO$_3$, 35-45% utilizing Ca(OH)$_2$, 25-60% utilizing CaO, 60-70%, utilizing dolomite (CaMg(CO$_3$)$_2$) in stoechiometric relations Ca/S= 1-5. [1-3]. Theoretical
and experimental researches of binding of SO$_2$ to the sorbent particles on the base of calcium compound, confirmed the practical utilization of dry process of flue gases refinement from SO$_2$ [4-14].

A significant number of theoretical and experimental studies examines indirect and direct sulfatization [4,10,11]. During the indirect sulfatization researchers conclude that the process of formation CaCO$_3$ is enabled by formation of intermediate product CaSO$_3$ which depending on the reaction temperature and O$_2$ volume ratio in flue gas oxidizes in CaSO$_4$ or CaS. According to some other researchers two mechanisms of CaSO$_4$ synthesis are recommended. The first mechanism includes the formation of intermediate product CaSO$_3$, and the second mechanism the formation of SO$_3$. For the both mechanisms the reaction temperatures of around 850 °C are sufficient, but reaction temperatures over 850 °C favor the formation of CaSO$_4$ only for the second mechanism.

The complex interaction of various resistances (diffusion over film gas, diffusion in the product layer and surface reaction of SO$_2$), is the main cause of inconsistency between mechanisms and results which were given by various researchers.

**EXPERIMENTAL SECTION**

**The scheme and description of experimental facility for SO$_2$ separation from a mixture of gases**

The scheme of laboratory facility for SO$_2$ separation from a mixture of gases in dry processes with lime (CaO), limestone CaCO$_3$, dolomite CaMg(CO$_3$)$_2$ and the list of Measuring Points (M.P.) are shown in Fig. 1, and in Fig. 2 the schemes of thermal furnace and reactor container for SO$_2$ separation are shown.

During the experiment of SO$_2$ absorption process, sorbent mass (sample) in the reactor was 100 g and the sorbent particle size was 500-700 µm. The heating process of the reactor and sorbent inside the reactor is achieved by electrical heaters. For each experiment the reaction temperature was adjusted to approximately firstly 200 °C, and secondly 400 °C. When the desired temperature was reached (=200 °C i.e.=400 °C), the sorbent is mechanically injected into the reactor using the dispenser. Then the gas mixture N$_2$+O$_2$+SO$_2$ is introduced into a reactor by vacuum pump in order to separate SO$_2$. A gas analyzer was continually used in measuring the concentration of SO$_2$ and O$_2$ in the inlet and gas outlet until the measured concentration.
of SO$_2$ is stabilized in the gas outlet. The experiments last up to 50 min. The speed of the gas which flows through the reactor for SO$_2$ separation was between 0.13-0.14 m/s. The gas velocity provides constant sorbent mass in the reactor. The sorbent mass was measured before and after the process of SO$_2$ separation from the gas mixture. During the experiment the gas value flow rate through the reactor in order to separate SO$_2$ was between 1.97-2.03 m$^3$/h.

During the experimental process of SO$_2$ separation from the gas mixture, the following measuring instruments were used:

**Measuring the temperature of the sample (sorbent) layer in the reactor**

Instrument: Thermo element Ni-Cr-Ni with digital gauges Fluke 54 –N series  
Measurement range: -200-300 °C  
Measuring error: ±0.2%

**Measuring the volume share of O$_2$ and concentration of SO$_2$ at the entrance and exit of the reactor**

Instrument: Gas analyzer Testo 350 XL  
Measurement range: 0-25%  
0-5000 ppm SO$_2$  
Measuring error: for SO$_2$ je ±5% for concentration range 100-2000 ppm and ±10% for concentration range 2001-5000 ppm.  
Measuring error: for O$_2$ is ±0.8% for the range of 0-25%

**Measuring the sample (sorbent) mass**

Instrument: Platform scale EMB 500-1  
Measurement range: 0-500g  
Measuring error: ±0.001g

**Measuring the air volume flow**

Instrument: Rotametar RAGK  
Measurement range: 0.2-6300 l/h
Fig. 3: The change of $\text{SO}_2$ concentration in a mixture of gases which flows through the reactor without sorbent and adding sorbent depending on the reaction time ($m_{\text{sorbent}} = 100 \text{ g}$, $t = 200 \degree\text{C}$, $d_p = 600 \mu\text{m}$).

Measuring error: ±4%

Measuring the $\text{SO}_2$ volume flow
Instrument: Rotametar RAGK
Measurement range: 0.2-6300 l/h
Measuring error: ±4%

Measuring the gas mixture volume flow
Instrument: Rotametar RAGK
Measurement range: 0.2-6300 l/h
Measuring error: ±4%

RESULTS AND DISCUSSION

In Fig. 3 and 4 the $\text{SO}_2$ concentration change in the flue gas which flows through the reactor for $\text{SO}_2$ is shown, without the sorbent and adding the sorbent depending on the reaction time (time of flue gas flow through the reactor) and reaction temperature. As it was expected, the concentration of $\text{SO}_2$ in the outlet gas is the lowest for CaMg(CO$_3$)$_2$ sorbent particles.

Being at reaction temperature of approximately 200 $\degree\text{C}$ and mean diameter of sorbent particles = 600 $\mu\text{m}$, the degree of $\text{SO}_2$ absorption to the sample particles occurs in the interval of 65-80% with sorbent CaMg(CO$_3$)$_2$, 42-66% with sorbent CaCO$_3$ and 34-60% with sorbent CaO (Fig. 5).

Increasing the reaction temperature from 200 $\degree\text{C}$ to 400 $\degree\text{C}$ the degree of $\text{SO}_2$ absorption to sorbent particles has no greater influence and it is in the interval of 37-78% by CaMg(CO$_3$)$_2$ sorbent utilization, 45-78% by CaCO$_3$ sorbent utilization and 19-60% by CaO sorbent utilization (Fig. 6).

During the first 6 min of the process examination of $\text{SO}_2$ absorption to sorbent particles, all sorbent samples quickly absorbed $\text{SO}_2$ at the studied reaction temperatures. After 6 minutes $\text{SO}_2$ absorption decreases, starts to stabilize and takes an approximately constant value (Fig. 5 and 6).

During the research, the value of stochiometric relation of calcium in sorbent and sulfur in flue gases ($\text{Ca}/\text{S}$) varied and it was above 5.

According to the data from the literature [4,8,11] following mechanisms of direct sulfatization can be recommended:

$$\text{CaMg(CO}_3\text{)}_2 + \text{SO}_2 + \frac{1}{2} \cdot \text{O}_2 \rightarrow \text{CaSO}_4 + \text{MgO} + 2 \cdot \text{CO}_2 \ (1)$$

$$\text{CaCO}_3 + \text{SO}_2 + \frac{1}{2} \cdot \text{O}_2 \rightarrow \text{CaSO}_4 + \text{CO}_2 \ (2)$$

$$\text{CaCO}_3 + \text{SO}_2 + \frac{1}{2} \cdot \text{O}_2 \rightarrow \text{CaSO}_4 + \text{CO}_2 \ (3)$$

During the direct sulfatization of CaMg(CO$_3$)$_2$, CaCO$_3$ and CaO (reactions (1) - (3)) the degree of sorbent utilization can be determined with the expression:

$$\eta_{\text{sorbent}} = \frac{m_{\text{sorbent}}}{M_{\text{SO}_2}} \cdot \frac{m_{\text{SO}_2}}{m_{\text{sorbent}}} \cdot \eta_{\text{SO}_2}, \ (%) \ (4)$$

where:

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CONCLUSIONS

The aim of the SO$_2$ separation experiment from a mixture of flue gases, which is demonstrated in this paper, was to determine the dependability of SO$_2$ separation degree from flue gas using different sorbents: dolomite (CaMg(CO$_3$)$_2$), calcium carbonate (CaCO$_3$) and calcium oxide (CaO), the influence of reaction temperature in the reactor and degree determination of sorbent utilization. Because of the previously said the temperatures varied from 200 and 400 °C. The sample mass within the reactor was 100 g and the mean diameter of sorbent particles was ≈600 µm. During the examination the following conclusions were made:

(1) It was acknowledged that the increase of the reaction temperature from 200 to 400 °C has no greater impact on SO$_2$ absorption to CaMg(CO$_3$)$_2$ and CaO sorbent particles. When the reaction temperature is somewhere around 200 °C and the mean diameter of sorbent particles is ≈600 µm, the degree of SO$_2$ absorption is in the interval of 65-80% by CaMg(CO$_3$)$_2$ sorbent utilization, 42-66% by CaCO$_3$ sorbent utilization and 34-60% by CaO sorbent utilization. When the reaction temperature is somewhere around 400 °C and it has the same diameter of sorbent particles, the degree of SO$_2$ absorption is in the interval of 37-78% by CaMg(CO$_3$)$_2$ sorbent utilization, 42-66% by CaCO$_3$ sorbent utilization and 34-60% by CaO sorbent utilization. When the reaction temperature is somewhere around 400 °C and it has the same diameter of sorbent particles, the degree of SO$_2$ absorption is in the interval of 37-78% by CaMg(CO$_3$)$_2$ sorbent utilization, 45-78% by CaCO$_3$ sorbent utilization and 19-60% by CaO sorbent utilization.

(2) During the first 6 min of examination of SO$_2$ absorption process to the sorbent particles, all sorbent samples quickly absorbed SO$_2$ at the studied reaction temperatures. This leads to a conclusion that the first 6 min of the process is completely controlled by the chemical reaction on the solid surface of the sorbent. After 6 min, the SO$_2$ absorption starts to stabilize and takes an approximate constant value (Fig. 5 and 6) and

\[
\eta_{SO_2} = \frac{\varphi_{SO_2, in} - \varphi_{SO_2, out}}{\varphi_{SO_2, in}} \cdot 100, \text{ (})
\]

- the degree of SO$_2$ absorption to sorbent particles
- $\varphi_{SO_2, in}$ - SO$_2$ volume ratio in flue gas at the entrance of the reactor, (ppm),
- $\varphi_{SO_2, out}$ - SO$_2$ volume ratio in flue gas at the exiting of the reactor, (ppm).

Using the numerical data gained by measurements (Fig. 3-6) and using the Eq. (4), the achieved values of the degree of sorbent utilization at the reaction temperature of 200 °C are low and are up to 14 % for sorbent dolomite, 6 % for sorbent CaCO$_3$ and 4 % for sorbent CaO. At the reaction temperature of 400 °C the degree of sorbent utilization is slightly lower. The low degree of sorbent utilization is a consequence of consumption of greater mass of sorbent than the mass of SO$_2$ in flue gases entering the reactor. (Fig. 7 and 8).
Fig. 7: The degree of sorbent (sample) utilization depending on the time of flue gas flow through the reactor for SO$_2$ separation ($m$(sorbent) = 100 g, $t = 200$ °C, $d_p = 600$ µm).

Fig. 8: The degree of sorbent (sample) utilization depending on the time of flue gas flow through the reactor for SO$_2$ separation ($m$(sorbent) = 100 g, $t = 400$ °C, $d_p = 600$ µm).

leads to a conclusion that the diffusion resistance becomes significant due the thickness of product layer.

(3) Low degree of sorbent utilization is determined: 14 % by dolomite (CaMg(CO$_3$)$_2$) utilization, 6 % by calcium carbonate (CaCO$_3$) utilization and 4 % by calcium oxide (CaO) utilization at the reaction temperature somewhere around 200 °C. When the reaction temperature is somewhere around 400 °C only higher degree of sorbent utilization for calcium carbonate CaCO$_3$ is marked and it is of 7 %. Low degree of sorbent utilization is due the utilization of great mass of sorbents (samples) in relation to the SO$_2$ mass entering the reactor.

The attained results of the experiment described in this paper indicate the capability of injecting powdered sorbents CaMg(CO$_3$)$_2$, CaCO$_3$ and CaO in the areas with lower temperatures as it is the convective part of the furnace and the flue canal of thermal power plant and which aim was to reduce the SO$_2$ emission in flue gas.

The achieved results can also represent the starting point in the phase of equipment designing for FGD as well as the equipment for separation the solid particle, whose practical utilization would improve the energetic and ecological efficiency of thermal-energetic power plants.

Nomenclature

- $d_p$ Sorbent particle diameter, µm
- $m$ The mass of sorbent, g
- $t$ Reaction temperature, °C
- $\phi_{SO_2, in}$ Gas flue SO$_2$ volume ratio in flue gas at the entrance of the reactor, ppm
- $\phi_{SO_2, out}$ Gas flue SO$_2$ volume ratio in flue gas at the outlet of the reactor, ppm
- $\eta_{SO_2}$ Retention degree of SO$_2$ for particle sorbent, %
- $\eta_{sorbent}$ Exploitation degree, %

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REFERENCES


