Absorption of pollutants from exhaust gases by low-temperature heating surfaces

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Abstract. One of the most effective methods aimed to improving the environmental safety is fuel oil combustion in the form of specially prepared water-fuel emulsions. The combustion of water-fuel emulsion in internal combustion engines makes it possible to reduce a rate of low-temperature corrosion at wall temperatures below the dew point temperature of sulfuric acid vapor, to install a condensing low-temperature heating surface in the exhaust gas boiler that leads to increase the efficiency of boiler. Therefore, it is of great importance to assess the effect of the presence of condensate (water, acid) and pollution on these surfaces on the processes of NOx, SO2 absorption from exhaust gases. Investigations of SO2, NOx and particulate matter emission were carried out on the experimental installation for fuel oil and water-fuel emulsion combustion with different water content. Using condensing heating surface enables to reduce the concentration of NOx and SO2 by 65 %. Experimental studies have shown that condensing heating surface ensures the capture of up to 30 % of particulate matter from the exhaust gas flow.

1 Introduction

One of the most perspective methods aimed to enhancing the environmental safety of atmospheric and water basin is fuel oil combustion in the form of specially prepared water-fuel emulsions (WFE) [1]. The use of wastewater as additional water makes it possible to focus the thermal power plants to low-waste technology by utilizing all wastewater contaminated with fuel products. The increased efficiency of the WFE combustion process is provided through the microexplosions of its droplets [2, 3], due to which the process of mixing fuel with air is improved, as a result the process of fuel oil combustion is intensified, that leads to decrease in the emission of particulate matter (PM), NOx and SO2 oxides [4, 5].

In addition, the WFE combustion in internal combustion engines makes it possible to reduce the low-temperature corrosion (LTC) rate at wall temperatures below the dew point temperature of sulfuric acid vapor [6, 7], to install a condensing low-temperature heating surface (LTHS) in the exhaust gas boiler (EGB) and thereby increase the EGB efficiency [8, 9] and thermal power plant [10, 11].

2 Literature review

Since the use of condensing heating surfaces determines the possibility of increasing the efficiency of boiler [12, 13], it is of great importance to assess the effect of the presence of condensate (water, acid) and pollution [14-16] on the LTHS on processes of NOx conversion during the movement of gases in the gas duct and in the atmosphere, on the amount of SO2 in exhaust gases.

Exhaust gases of power boilers nitrogen oxides NOx consist of 97 % of NO and 5 % of NO2 [17]. For auxiliary boilers and industrial boilers, according to [18] the content of NOx in NOx significantly increases (the proportion of nitrogen dioxide reaches 12 %). When water vapor is injected into the combustion chamber of a gas turbine an increase of NOx content up to 38 % is observed [19].

Conversion and absorption of higher oxides of sulfur and nitrogen by water and solutions of other liquids is used in a number of technologies for cleaning exhaust gases from toxic ingredients SOx and NOx [20, 21]. The introduction of ozone into exhaust gases ensures the oxidation of lower oxides of sulfur and nitrogen (SO2 and NO) to higher oxides (SO3 and N2O5), well absorbed by water and aqueous solutions.

The passage of the nitrous mechanism of H2SO4 formation in the condensate at the LTHS (in the pollution layer) during the WFE combustion should have an impact on the emission of SOx and NOx into the atmosphere. Therefore, it is necessary to assess the environmental indicators of EGB during the WFE combustion, providing the nitrous mechanism of formation of H2SO4 passes through the layer of pollutions at wall temperatures below the dew point temperature of sulfuric acid vapor [6, 16].

When WFE was burnt with W' = 30 %, the NOx/NO ratio is closer to the equimolar ratio (0.5) than while
burning fuel oil M40 and diesel fuel. Consequently, during the WFE combustion, a better absorption of NOx, as well as SO2, should occur. The nature of the change in the SO2 content with increasing \( W_r \) is practically the same for all types of fuels and is directed towards a decrease in concentration.

It has been experimentally proved [22] that the rate of absorption of nitrogen oxides NOx by condensate \( \text{H}_2\text{SO}_4 \), consisting of an equimolecular mixture of NO and NO2 (50 % NO and 50 % NO2), is almost 2 times higher than from a gas containing only NO2 or NO. The main influence on the rate of NOx absorption is exerted by chemical processes occurring in the liquid phase [23].

Modern methods, including ANSIS [24-26], can be used for simulation [27, 28] and optimizing [29-31] the processes and regimes of operation as well as statistical treatment of experimental data [32-34]. For estimating the efficiency of such greening and fuel saving technologies during the operation in actual climatic conditions various methods of modelling [35, 36] are applied.

The aim of research is to assess the effect of the presence of condensate (water, acid) on condensing LTHS on the processes of NOx, SO2 and PM absorption from exhaust gases during WFE combustion.

### 3 Research methodology

#### 3.1 Experimental research

Investigations of the SOx and NOx emission were carried out on an experimental installation (Fig. 1, a) with fuel oil and WFE based on this fuel oil combustion [37, 38] with different water content with an almost constant excess air ratio.

![Fig. 1. General view of the experimental setup (a) fabric filter during WFE (b) and fuel oil (c) combustion.](image-url)

The analysis of the exhaust gases composition was carried out with an OXI-5M gas analyzer behind the furnace and LTHS.

In order to assess the decrease of PM emission, experimental studies of their concentration in exhaust gas flow before and after the LTHS were carried out, depending on the water content of WFE. The values of PM concentration before and after LTHS were determined by filtering the exhaust gas flow using a glass cloth filter and checked by calculation (Fig. 1, b, c).

#### 3.2 Theoretical research

The mass flow rate of soot particles (SP) in the exhaust gas flow after LTHS has been determined by the formula [39], g/h:

\[
G_{SP \text{ in flow}}^{\text{after LTHS}} = q_i \cdot 10^3 \cdot \frac{Q_F'}{10^3} \cdot \frac{Q_F'}{r_p Q_i},
\]  

(1)

where \( Q_F' = 32800 \text{ kJ/kg}, \) \( Q_F' = 39360 \text{ kJ/kg} \) - the lowest and highest heat of fuel combustion (fuel oil M-100).

The mass flow rate of PM in the exhaust gas flow before LTHS has been determined by volume concentration, g/h:

\[
G_{PM \text{ in flow}}^{\text{after LTHS}} = f(C_{PM \text{ in flow}}) = C_{PM \text{ in flow}} \cdot V_{\text{gas flow}} \cdot 10^{-3}
\]

(2)

where \( V_{\text{gas flow}} = 15 \text{ m}^3/\text{kg} \) - volume flow rate of exhaust gases through the gas duct of experimental setup.

The experimental value of the volume concentration of PM in exhaust gas flow before LTHS \( C_{PM \text{ in flow}}^{\text{before LTHS}} \) at corresponding water contents of WFE, mg/m^3.

The mass flow rate of PM deposited on the LTHS \( G_{PM \text{ settle}} \) has been determined at the number of deposits \( K_p \), g/h:

\[
G_{PM \text{ settle}} = K_p \cdot S
\]

(3)

where \( K_p \) - the estimated number of deposits on LTHS of experimental setup, g/(m^2·h) [14].

The total area of the outer surface of the experimental samples installed in the gas duct of the experimental setup:

\[
S = \pi d \ln n
\]

(4)

The diameter of the experimental sample is \( d = 25 \text{ mm} = 0.025 \text{ m}, \) its length is \( l = 80 \text{ mm} = 0.08 \text{ m}, \) the number of installed samples is \( n = 13. \) Thus, \( S = 0.082 \text{ m}^2. \)

The volume concentration of PM deposited on the LTHS has been determined by mass flow rate, mg/m^3:
The volume concentration of PM after LTHS was determined, mg/m³:

\[ C_{PM \text{ settle}} = \frac{G_{PM \text{ settle}}}{V_{\text{gas flow}}} \times 10^3 \] (5)

The volume concentration of PM after LTHS in flow was:

\[ C_{PM \text{ in flow}}^\text{after LTHS} = C_{PM \text{ in flow}}^\text{before LTHS} - C_{PM \text{ settle}} \] (6)

The "degree of capture" LTHS has been determined, %:

\[ \gamma = \frac{100 \times (C_{PM \text{ in flow}}^\text{before LTHS} - C_{PM \text{ in flow}}^\text{after LTHS})}{C_{PM \text{ in flow}}^\text{before LTHS}} = \frac{100 \times C_{PM \text{ settle}}}{C_{PM \text{ in flow}}^\text{before LTHS}} \] (7)

The total toxicity index for the maximum concentrations of ingredients has been determined

\[ \phi_t = \frac{C_{NO_x}}{ПДК_{NO_x}} + \frac{C_{SO_2}}{ПДК_{SO_2}} \] (8)

### 4 Results

Experimental studies have shown that a decrease of NOₓ and SO₂ concentration with an increase in \( W' \) in the WFE is accompanied by an increase of the aerosol H₂SO₄ content, which is observed at \( W' = 10 \% \) and above. The same process takes place in sulfuric acid production by the nitrous mechanism. The results of investigation the content of NOₓ, SO₂ in exhaust gas flow with an increase of water content in the emulsion to 30% are shown in Fig. 2.

![Fig. 2. Influence of water content of WFE on the NOₓ and SO₂ emission.](image)

The dependence of the concentration of \( C_{PM} \) of solid ash and soot particles (particulate matter) in the exhaust gas flow before and after the LTHS during WFE combustion with different \( W' \) in Fig. 3-4 have shown.

It shows a sharp decreasing of \( C_{PM} \) concentration almost 3 times with an increasing of the water content of WFE from 2 to 15 %. This is explained by an increasing of the plume turbulence due to the effect of "microexplosion" of WFE droplets, which leads to their additional fragmentation and better burnout PM of smaller size.

![Fig. 3. Dependence of the change in the PM concentration in the exhaust gases before and after LTHS during the WFE combustion.](image)

![Fig. 4. Dependence of the change in the SP concentration in the exhaust gases after LTHS during the WFE combustion.](image)

Despite the increasing moisture content of the surface with an increase of the water content \( W' \) to 10 %, due to a decrease of the size (mass) of particles (accordingly, the action of inertial forces) and the entrainment of small particles by the exhaust gas flow, the deposition intensity of these particles decreases.

With a further increase of the water content, the particle sizes continue to decrease, which should have led to a decrease of their deposition intensity. However, due to an increase of surface moisture and an increase in its adhesion effect, the deposition of such solid particles remains practically at the same level with an increase of the water content of WFE \( W' \) from 17 to 30 %.

Measurements of the particle concentration before and after the LTHS installed in the exhaust gas duct of experimental setup had shown that this heating surface ensures the capture of up to 30 % of PM from the exhaust gas flow. By increasing of the surface size with a wall temperature below 120 °C, it is possible to capture up to 50 % of particles from the flow. This is also confirmed by the decreasing values of the PM concentration, due to their deposition on the surface.

Taking into account the accepted rule of summing up the harmful effects of NOₓ and SO₂, the total toxicity index was determined for the maximum concentrations of ingredients (Fig. 5).
When WFE combustion with a water content of 30 %, the toxicity $\Phi_C$ is significantly lower and amounts to 5750 (when fuel oil combustion $\Phi_C = 15740$), i.e. the toxicity of exhaust gases during emulsion combustion is reduced by 2.7 times in comparison with fuel oil combustion.

5 Conclusions

The results of research have shown the presence of water vapor, their uniform distribution (as well as other components) over the torch volume due to "microexplosions" during the combustion of WFE, the uniform distribution and decrease in temperature in the combustion zone, the participation of $\text{H}^+$ and $\text{OH}^-$ ions in chemical reactions lead to reducing the emission of NOx and SO2 oxides.

Due to the condensation of water vapor and sulfuric acid vapor during the WFE combustion, nitrogen oxides are absorbed in the condensate, thereby the number of harmful substances in the emission is reduced.

When condensing economizer is installed, due to the absorption of a part of nitrogen oxides by condensate, the real toxicity of exhaust gases in the atmosphere will be significantly reduce (2.7 times).

6 Nomenclature

- $C_{PM \text{ in flow after LTHS}}$ – volume concentration of PM in exhaust gas flow after LTHS, [m$^3$/kg]
- $C_{PM \text{ in flow before LTHS}}$ – volume concentration of PM in exhaust gas flow before LTHS, [m$^3$/kg]
- $C_{PM \text{ settle}}$ – volume concentration of PM deposited on the LTHS, [m$^3$/kg]
- $G_{PM \text{ in flow after LTHS}}$ – mass flow rate of PM in exhaust gas flow after LTHS, [g/h]
- $G_{PM \text{ in flow before LTHS}}$ – mass flow rate of PM in exhaust gas flow before LTHS, [g/h]
- $V_{gas \text{ flow}}$ – volume flow rate of exhaust gases through the gas, [m$^3$/kg]
- $K_\ell$ – estimated amount of deposits on LTHS, [g/(m$^2$·h)]
- $Q'_p$ – highest heat of fuel combustion, [kJ/kg]
- $S$ – total area of the outer surface of the experimental samples, [m$^2$]
- $\gamma$ – degree of capture, [%]
- $\Phi_C$ – total toxicity index.

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Fig. 5. Dependence of total toxicity index $\Phi_C$ of exhaust gases from water content of WFE.
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