Composite Materials for Adsorption-Catalytic Purification of Toxic Organic Impurities

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Abstract: Design of high effective catalysts with unique properties opens good perspectives for solving environmental problems of cleaning waste gas, particularly from toxic impurities of organic solvents. A new thermostable nanostructured composite materials based on shungyte was obtained and determined their adsorption capacity on the model substance o-xylene. Xylene used as a model substance is a part of the organic solvent presents in waste industrial gases. Best xylene sorbents (449.7 mg/g)—activated shungyte containing 70% carbon and saksaul charcoal (554.3 mg/g). Then polyoxide catalysts on the base of transition metals (Ni, Mn, Cu) modified by the rare earth elements (La, Ce and Nd), supported on a granular carrier were studied in the reaction of deep oxidation of o-xylene. 100% activity showed 7.0% Cu-Mn-Ce-catalyst at $T$: 250-300 °C, $C_{o-xylene}$: 0.6 g/m$^3$, the optimum space velocity of gas—1,200 h$^{-1}$ for catalysts supported on a granular carrier. The results obtained can be used in the purification of waste gas from paint, furniture, cable, printing and other enterprises from the impurities of toxic organic solvents.

Key words: Composite material, adsorption, xylene, catalyst, sorbent, shungyte.

1. Introduction

At the moment, when the waste recycling enterprises fully yet, the main task of gas cleaning is bringing the toxic impurities in the flue gas to the minimum permissible concentrations established sanitary standards. Industrial gases are usually contaminated with harmful impurities, so more and more attention is paid to research on cleaning and disposal of traces of organic solvents in the exhaust gas [1-5]. Of thermal, catalytic and adsorption methods of neutralization adsorption-catalytic method is promising due to the high efficiency of purification from impurities and can be cleaned with large amounts of gases containing a small fraction of impurities.

The aim of the work was the development of new composite materials for directional adsorption-catalytic purification of industrial waste gas from impurities of toxic organic solvents. Adsorption-catalytic method of purifying toxic exhaust gases from organic impurities [6] based on the implementation of a cyclic treatment process in two successive stages. In the first stage of the cycle, occurs prior accumulation due to adsorption and/or chemisorption of a toxic impurity in existing gas in the catalyst bed. The second cycle step is oxidation of toxic impurities adsorbed in the catalyst layer, which occurs when passing air through the catalyst layer heated to a temperature above the characteristic temperature of the onset of deep oxidation of adsorbed impurities. This method provides a high degree of purification of exhaust gases with low initial concentrations of impurities, thus, it is substantially less energy compared with the other. Purified gas can be returned to the system if the impurity concentration is significant. The combination of these two purification steps of the organic impurities was used in this study.

2. Experiments

This article is a continuation of a series of studies [7, 8]
devoted to the development of highly efficient adsorbents-catalysts for the two-step adsorption-catalytic purification of industrial waste gas from impurities of organic solvents. Previously performed experimental studies to obtain composite materials based on carbon shungytes and saksaull coal and heat treated with subsequent activation of the vapor-gas mixture, allowing developing a large specific surface.

This paper describes following composite materials—shungyte and activated carbon based on the chips of saksaull studied for their adsorption capacity of the model substance—o-xylene. In the second step, proposed polyoxide catalysts supported on carriers for the deep oxidation of the adsorbed organic substances.

2.1 Synthesis of Composite Materials for the Adsorption of Xylene

Shungytes—specific carbonaceous rocks. Their constituent carbon with a metastable amorphous molecular structure, not prone to graphitization. The main element of the supramolecular structure is shungyte carbon globule—multilayer formation of sizes up to 10 nm with pore inside. Shungyte is a black, mineral shiny with conchoidal glass fracture, brittle. Shungyte rocks vary in chemical and mineralogical composition and, as a consequence, have different adsorption and catalytic properties. They represent a natural composite—a uniform distribution of fine crystalline silica particles in an amorphous carbon matrix. Various samples of activated shungyte with carbon content of 25.0%, 45.0% and 70.0%, and saksaull charcoal were prepared according to the developed technique, comprising the steps of froth flotation and treatment with acid. As a result, the samples prepared with the following content of carbon and the particle size:

- shungyte with 25.00% carbon, the particle size of minus 74 microns;
- concentrate with a percentage of carbon 45.00% and a particle size of minus 40 microns;
- concentrate with a percentage of 70.00% carbon and tails, a particle size of minus 40 microns.

There were studied the effects of temperature, space velocity and the concentration of o-xylene in the reaction mixture on the adsorption capacity of the composite materials on the basis of shungyte with activated carbon of different content and saksaull coal on the pilot installation consisting of a compressor (Parker Balston, Denmark), zero clean air generator (Parker Balston, Denmark), a pressure regulator (Swagelok, USA) syringe pump KDS 100 (KD Scientific, USA), 250 mL calibrated bulb for averaging the flask analyte concentration with teflon valves and sampling port (Supelco, USA) and a modified sampling vial (CTC Analytics AG, Switzerland) connected to the inlet of the calibrated bubble flowmeter. Analyzed samples were taken from 20 mL vial using Combi-PAL (CTC Analytics AG, Switzerland) autosampler mounted on a gas chromatograph with mass spectrometric detector 7890A/5975C (Agilent, USA).

Before the test, sorbents are dried at 200 °C for 3 h. Determination of the adsorption capacity of the adsorbent was carried out by passing through it a mixture of air with xylene and measuring the xylene concentration in the outlet stream after the adsorbent. The experiment was carried out over a period of time required for the saturation of the adsorbent and the equilibrium in the system.

2.2 Design of Catalysts with Different Active Phase Supported on a Granular Carrier

Design of catalysts with different active phase based on the oxides of nickel, manganese, copper and their binary mixtures, as well as the modification by the rare earth elements was carried out when supported on the granular carriers. Preparation of granular catalysts was carried out by capillary impregnation of the support for water capacity.

Oxidative conversion of model substance—o-xylene with atmospheric oxygen was investigated in the flow catalytic laboratory facility over the catalysts supported
on a granular carrier in the temperature range 350-600 °C, with space velocities 400-1,200 h⁻¹ and atmospheric pressure. Ratio xylene: air varied from 6:3 to 3:3. The volume of catalyst supported on the granular carrier was 2 cm³. Installation consists of a reactor, heating system, mixer and instrumentation. Air is supplied from a balloon, and xylene with a metering pump in a rheometer and then to the mixer, where upon the mixture of xylene with air fed to the reactor with the catalyst. The temperature was measured in the catalyst layer using a chromel-alumel thermocouple on the device KSP-3. Sampling was made before and after the catalyst. Analysis of xylene and its deep oxidation products was performed on a GC chromatograph Hromas GH-1000, issue 2012.

3. Results and Discussion

3.1 Investigation of the Adsorption Capacity of Sorbents

Figs. 1 and 2 show typical curves of the output o-xylene adsorption for sorbents. The adsorption capacity of the sorbent was calculated on the basis of observed peaks considering the sensitivity factor. Results obtained on adsorption capacities are shown in Table 1, from which it follows that as the shungyte carbon content increase from 11% to 23%, its adsorption capacity is increased from 18.0 mg to 41.0 mg of xylene per gram (mg/g) of shungyte. With increasing of carbon content in shungyte to 50 % and 70%, its adsorption capacity reaches 261.6 mg/g and 449.7 mg/g, respectively. Maximum equilibrium capacity (554.3 mg/g) was found for the saksaul coal. The size of the equilibrium adsorption capacity in all these conditions remained constant at 449.7 mg/g sorbent. At the space velocity of 5,000 h⁻¹ o-xylene adsorption, tests were carried out at elevated temperatures from 20 °C to 200 °C. It was found that increasing the temperature to 100 °C sharply reduces the adsorption capacity of activated shungyte to 44.9 mg/g. At 200 °C, xylene adsorption does not occur.

The study of the dynamics of the adsorption of o-xylene from air mixture showed that increasing the concentration of xylene in the reaction mixture from 300 mg/m³ to 500 mg/m³ reduces saturation time
shungyte by xylene on 1.7 times while maintaining the equilibrium adsorption capacity of the activated shungyte with carbon content 70%. Thus, the adsorption capacity of composite materials on xylene by shungyte with different carbon contents is defined. It has been established that the best of xylene sorbent is activated shungyte sample containing of 70% carbon (449.7 mg/g). The maximum value of this characteristic (554.3 mg/g) was obtained by using saksaul coal.

3.2 Studying of the Activity of the Catalysts

The first step was to prepare and test of mono oxide catalysts: Ni-, Cu- and Mn. Further, there were prepared and studied binary oxide catalysts: Ni-Cu, Cu-Mn and Mn-Ni. The next step of varying the composition of the active phase of the catalyst was modification of the best binary oxide catalyst by the rare earth metals—cerium, lanthanum and neodymium.

Following are the results of the test of the catalysts as the complexity of their composition in the complete oxidation of a model substance—o-xylene by air oxygen. To determine the effect of the active phase on the direction of the oxidation reaction of o-xylene were first tested mono oxide catalysts: Ni-, Cu- and Mn-, supported on a granular carrier. Their activity was determined in the complete oxidation of o-xylene under the conditions: $T=300 \, ^\circ C$, the space velocity of oxygen—1,200 h$^{-1}$, the catalyst volume is 2 mL.

As can be seen from Table 2, among the investigated monoxides of nickel, copper and manganese, the best activity showed the copper catalyst whose activity in complete oxidation of o-xylene was 83%, while for the nickel catalyst 69% and manganese 78%. Next, under the same conditions were studied binary oxide catalysts: Ni-Cu, Cu-Mn and Mn-Ni. Among them, the highest activity (98%) is typical for Cu-Mn. Activity of Ni-Cu 87%, and the Mn-Ni 91%. Catalyst Cu-Mn, which showed the highest activity was modified with oxides of REE (rare earth elements): cerium, lanthanum and neodymium. One hundred percent of xylene oxidation is observed in all studied (Cu-Mn-Ce, Cu-Mn-La and Cu-Mn-Nd) catalysts at 300 °C (Table 2).

However, when 250 °C oxidation of o-xylene 5.6% (Fig. 3) above for Cu-Mn-Ce-catalyst than Cu-Mn-La and Cu-Mn-Nd.

Thus, among the new synthesized oxide catalysts based on transition and rare earth metals Cu-Mn-La, Cu-Mn-Ce and Cu-Mn-Nd, the highest activity showed Cu-Mn-Ce-catalyst.

3.3 Investigation of the Influence of Process Parameters on the Activity of Deep Oxidation of Xylene by Atmospheric Oxygen

Activity of prepared Cu-Mn-La, Cu-Mn-Ce and Cu-Mn-Nd—supported on a granular carriers catalyst were studied in the reaction of deep oxidation of o-xylene by atmospheric oxygen to CO$_2$ and H$_2$O with varying of process parameters: temperature, space velocity and ratio of the reaction medium. Temperature of the process being studied at the range from 100 °C to 300 °C, and was as follows: 100 °C, 150 °C, 200 °C, 250 °C, 300 °C. Fig. 3 shows the activity of Cu-Mn-La, Cu-Mn-Ce and Cu-Mn-Nd catalysts supported on granular carrier on the temperature of the reaction.

Most active at all temperatures studied characteristic for the Cu-Mn-Ce-catalyst and at 150 °C, it is 41%, with an increase in temperature to 50 °C activity increased to 83% and at 250 °C and 300 °C activity of Cu-Mn-Ce-catalyst equal to 100%, while over the Cu-Mn-La and Cu-Mn-Nd catalysts oxidation at 250 °C reaches 94% and 95%, respectively.

Table 2 Comparative activity of supported catalysts to a granular carrier of complete oxidation of o-xylene.

| Catalysts | Activity (%) |
|-----------|--------------|
| Ni        | 69           |
| Cu        | 83           |
| Mn        | 78           |
| Ni-Cu     | 87           |
| Cu-Mn     | 98           |
| Mn-Ni     | 91           |
| Cu-Mn-Ce  | 100          |
| Cu-Mn-La  | 100          |
| Cu-Mn-Nd  | 100          |
| Cu-Mn-Ce  | 100          |
| Cu-Mn-La  | 100          |
| Cu-Mn-Nd  | 100          |
| Cu-Mn-Ce  | 100          |
| Cu-Mn-La  | 100          |
| Cu-Mn-Nd  | 100          |

As can be seen from Table 2, among the investigated monoxides of nickel, copper and manganese, the best activity showed the copper catalyst whose activity in complete oxidation of o-xylene was 83%, while for the nickel catalyst 69% and manganese 78%. Next, under the same conditions were studied binary oxide catalysts: Ni-Cu, Cu-Mn and Mn-Ni. Among them, the highest activity (98%) is typical for Cu-Mn. Activity of Ni-Cu 87%, and the Mn-Ni 91%. Catalyst Cu-Mn, which showed the highest activity was modified with oxides of REE (rare earth elements): cerium, lanthanum and neodymium. One hundred percent of xylene oxidation is observed in all studied (Cu-Mn-Ce, Cu-Mn-La and Cu-Mn-Nd) catalysts at 300 °C (Table 2).

However, when 250 °C oxidation of o-xylene 5.6% (Fig. 3) above for Cu-Mn-Ce-catalyst than Cu-Mn-La and Cu-Mn-Nd.

Thus, among the new synthesized oxide catalysts based on transition and rare earth metals Cu-Mn-La, Cu-Mn-Ce and Cu-Mn-Nd, the highest activity showed Cu-Mn-Ce-catalyst.
Thus, the most active among the three studied catalysts is a Cu-Mn-Ce-catalyst. Optimum temperature at which all the studied catalysts are the most active is 250-300 °C. The most active (100%) Cu-Mn—modified by cerium catalyst at 250-300 °C. Further detailed investigation of the effect of the concentration of xylene and a space velocity of the gas-air mixture on the conversion of xylene was carried out on Cu-Mn-Ce-catalyst.

It was studied the effect of the concentration of o-xylene: 0.1, 0.3 and 0.6 g/m³ at gas space velocities of 1,000, 5,000 and 10,000 h⁻¹ at a temperature of 300 °C on the activity of Cu-Mn-Ce-catalysts (Fig. 4).

Most activity of Cu-Mn-Ce-catalyst is observed in xylene concentration of 0.6 g/m³ at all gas space velocities as compared to the concentrations of xylene 0.3 g/m³ and 0.1 g/m³. However, at concentrations of xylene 0.1 g/m³ and 0.3 g/m³ at gas space velocity 1,000 h⁻¹ activity of Cu-Mn-Ce-catalyst is high and is 99% to 100%.

Thus, the optimal concentration of xylene—0.6 g/m³ in which Cu-Mn-Ce is the most activity. Investigation of activity of the catalyst depends on a space velocity of the feed gas was carried out in the following ranges: 1,200, 3,000, 4,000, 5,000, 10,000 h⁻¹. It is shown that, the optimal space velocity of the gas which is manifested in the maximum activity of Cu-Mn-Ce-catalyst for the whole studied temperature range (100-350 °C) is 1,200 h⁻¹.

4. Conclusions

Adsorption of o-xylene by shungytes with different carbon contents was studied and it showed that the best of xylene sorbent among the activated shungytes is sample (449.7 mg/g) with 70% carbon. The maximum adsorption capacity of xylene (554.3 mg/g) is observed at saksaul charcoal.

New oxide catalysts are developed and prepared on granular carrier with desired properties based on transition and rare earth metals (Ni, Mn, Cu, La, Ce and Nd). Investigation of the influence of the active phase of the catalyst on the direction of oxidation of o-xylene showed that the most active among the investigated oxides of nickel, copper and manganese is the copper catalyst (83%), among the two-component catalyst (98%)—Cu-Mn. Modifying of bioxide catalysts by the rare earth elements enhances their activity.

The maximal activity (100%) in the deep oxidation of o-xylene with atmospheric oxygen showed Cu-Mn-catalyst modified by cerium, containing sum of metals —7.0%.
The influence of process parameters on the activity of optimal Cu-Mn-Ce-catalyst was defined. Maximum activity of all the catalysts examined is shown at 250-300 °C, the optimal concentration of xylene—0.6 g/m³ and the optimum gas space velocity in the entire investigated temperature range (100-350 °C)—1,200 h⁻¹ for catalysts supported on a granular carrier.

The results obtained can be used in the adsorption-catalytic cleaning of industrial waste gas from impurities of toxic organic solvents.

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