**Abstract.** In this work the synthesis technology of zinccontaining polymer - inorganic composite on the basis of products of secondary raw material processing at joint precipitating with carbamide and formaldehyde (ZnCFO) is described. The structure and properties of ZnCFO are investigated by the differential-thermal analysis, electronic microscopy, and IR-spectroscopy. The action of ZnCFO as vulcanization active component of elastomeric compositions on the basis of rubbers of general and special purpose with various vulcanization systems is investigated. The ZnCFO influence on the character of the formed morphological structure of rubbers is determined by the method of percalation analysis.

**Key words:** zinccontaining polymer-inorganic composite, rubber, vulcanization, elastomeric composition, vulcanization activator, physico-mechanical properties, morphological structure.

**1. Introduction**

In spite of 150-year history of vulcanization process, it is impossible to consider that fundamental and applied researches in the direction of vulcanization systems improvement are complete. For today one of the ways of rubbers properties improvement is the synthesis and application of new chemicals-additives, including the vulcanization active ones. This is connected, first of all, with reduction of global zinc ores stock as basic raw material for production of the traditional activator – zinc oxide. Besides, modern increase of industrial potential which results in the accumulation of a large quantity of wastes cause a number of ecological problems that require emergency measures. Therefore creation of resource saving technologies of the new compounds reception from products of secondary raw material processing has gained paramount importance.

**2. Experimental**

In view of the above said this work offers the first investigation of the effectiveness of zinccontaining carbamide-formaldehyde composite (ZnCFO) as vulcanization active component of various vulcanization systems for rubbers of general and special purpose. Structure and properties of the composite have been studied by means of instrumental analysis methods.

The electronic microscopy method using the EM-125 for definition of ZnCFO particles size and surface characteristic was applied. Known zinc oxides were chosen as the comparison object.

The definition of thermal stability of inorganic zinccontaining (Zn(OH)$_2$), organic carbamide-formaldehyde (CFO) components, and ZnCFO composite was carried out by the method of differential-thermal analysis on the derivatograph “Q-1500 D” of F. Paulik, J. Paulik, L. Erdey system in the temperature range of 293–1273 K at the heating rate of 283 K/min and the samples mass of 200 mg.

The study of ZnCFO was also carried out by the method of IR-spectroscopy on spectrometer UR-20. The researches of ZnCFO compatibility with the matrix of isoprene rubber were carried out in plasticorder “Brabender” PLE 651.

An estimation of ZnCFO effectiveness as vulcanization active component was carried out in modelling unfilled elastomeric compositions on the basis of isoprene, butadiene-nitrile, chloroprene, and butyl rubbers of sulphur, thiuram, peroxide, metaloxide, and resin vulcanization systems.

The rubber compounds were mixed in a laboratory internal mixer. The tests of elastomeric compositions were carried out according to working techniques and requirements of state standard.
3. Results and Discussion

The technological circuit of ZnCFO synthesis has been developed in the Ukrainian State Chemical-Technological University. ZnCFO is the product of reaction of carbamide and formaldehyde polycondensation in zinc salts solution occurring at recycling of metal containing wastes of chemical manufactures of Ukraine. The material base of ZnCFO manufacture are dead catalysts, zincing solutions and other zinc-containing wastes of various origin.

The maximal degree of zinc extraction (ρ Zn$^{2+}$) = 95–98 % and mass ratio of zinc hydroxide to carbamide-formaldehyde component in the composite (m$_{Zn(OH)_{2}}$ : m$_{CF}$O) = 1:0.7–1 is achieved at the observance of the following technological parameters: temperature (Т) = 298–308 K; concentration of zinc salts in solution (c$_{Zn^{2+}}$) = 150–170 g/l; pH of zinc salts solution = 2; pH of the reactionary mix at complete precipitating 170 g/l; pH of the reactionary mix at polycondensation = 3–5; mole ratio of carbamide to formaldehyde (n$_{C}$ : n$_{F}$) = 1:1 [1,2].

The electronic photos of powders (Fig. 1) show that the new composite and zinc oxide have external similarity by the form of particles, differ by dispersiveness (0.4–6.0 microns for zinc oxide, Fig. 1a; 0.3–6.0 microns for ZnCFO, Fig. 1b) and contain both crystal and amorphous phases in their structure.

DTA and TG curves of the researched products are presented in Fig. 2. As one can see, endothermic peak on DTA curve of Zn(OH)$_2$ and the greatest loss of weight at Т = 423–433 K are caused by transition of hydroxide into oxide. DTA curves of CFO and ZnCFO have external similarity by the character of the thermal effects, however, for ZnCFO the shift of maximums into the range of lower temperatures is observed. Thus, peaks at Т = 363–373 K are caused by the loss of atmospheric moisture contained in the samples. Endothermic maximum and greatest loss of weight (=35 %) in the temperature range 533–543 K characterize the beginning of CFO thermodestruction, and secondary exothermic peak at Т = 833 K is caused by the subsequent decomposition of the sample. Unlike CFO on DTA curve of ZnCFO the first endothermic maximum is observed at Т = 443–453 K, and secondary exothermic maximum – at Т = 723–733 K, which is characteristic of organic component decomposition. Appearance of a new endothermic peak at approximately 1083 K indicates expansion of the temperature range of ZnCFO destruction process in comparison with CFO, while the temperature interval of weight loss for CFO is 533–833 K, and for ZnCFO – 443–1083 K, and ZnCFO has extremely greater mass residue. It is necessary to note that ZnCFO at thermal action is decomposed not reaching a melting point, which was confirmed by the method of capillary melting temperature definition [3, 4]. During the experiment an intensive gas release and change in sample color from white to grey at the temperatures 443 K and 493 K that correspond to the first endothermic peak on DTA curve, were visually observed.

For detection of possible chemical bonds between inorganic and organic components in ZnCFO the spectra of Zn(OH)$_2$, CFO and ZnCFO were studied (Fig. 3). As can be seen, IR-spectrum of ZnCFO the characteristic CFO absorption bands repeat at 3350, 1640, and 1550 cm$^{-1}$, caused by the presence of a secondary amide group [5]; at the same time the intensity of absorption band in the range 2340–2360 cm$^{-1}$, identifying $\text{R}_2\text{C} \equiv \text{N}^{+} \text{H}$ group, is considerably decreased [6]. In comparison with IR-spectrum of Zn(OH)$_2$, ZnCFO spectrum the intensity of characteristic absorption band at 1384 cm$^{-1}$ decreased. The distinctive feature of ZnCFO spectrum is the presence of absorption band at 1120 cm$^{-1}$, which probably corresponds to bonds (e.g. coordination ones) between inorganic and organic components of the composite. It can be assumed, that the coordination bonds in the composite arise between nitrogen atom of $\text{R}_2\text{C} \equiv \text{N}^{+} \text{H}$ group and zinc atom in Zn(OH)$_2$.

The estimation of ZnCFO compatibility with the rubber matrix have shown that the composite dispersion process is accompanied by lower power consumption and
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best compatibility in comparison with zinc oxide (Fig. 4). The absence of ZnCFO particles as extraneous impurities in rubber mix was visually observed, while the zinc oxide particles were clearly visible [7].

Thus, the given results of experimental researches have confirmed compositional and chemically bound structure of ZnCFO with the presence of functionally active groups, which due to its organic-inorganic nature has determined relationship with rubber matrix, is easily dispersible and combinable with it.

The influence of ZnCFO concentration (3.0; 5.0 and 7.0 phr) on formation of properties complex of the unfilled rubber mixes and their vulcanizates on the basis of isoprene rubber at 428 K with ZnCFO as the activator:

- ZnO – 5.0 phr, \( k_2 = 0.47 \text{ min}^{-1} \) (1);
- ZnCFO – 3.0 phr, \( k_2 = 0.51 \text{ min}^{-1} \) (2);
- ZnO – 5.0 phr, \( k_2 = 0.58 \text{ min}^{-1} \) (3) and
- ZnO – 7.0 phr, \( k_2 = 0.69 \text{ min}^{-1} \) (4)

The absence of ZnCFO particles as extraneous impurities in rubber mix was visually observed, while the zinc oxide particles were clearly visible [7]. Thus, the given results of experimental researches have confirmed compositional and chemically bound structure of ZnCFO with the presence of functionally active groups, which due to its organic-inorganic nature has determined relationship with rubber matrix, is easily dispersible and combinable with it.

The influence of ZnCFO concentration (3.0; 5.0 and 7.0 phr) on formation of properties complex of the unfilled rubber mixes and their vulcanizates on the basis of isoprene rubber of the following composition, phr:

- isoprene rubber – 100.0;
- sulfur – 1.0;
- di-(2-benzothiazolyl)disulfide – 0.6;
- N, N'-diphenylguanidine – 3.0;
- stearic acid – 1.0

was carried out in comparison with the known activator – zinc oxide (5.0 phr). The analysis of Rheometer data of sulfur vulcanization process of elastomeric compositions on the basis of isoprene rubber at 428 K with ZnCFO as the activator:

- ZnO – 5.0 phr, \( k_2 = 0.47 \text{ min}^{-1} \) (1);
- ZnCFO – 3.0 phr, \( k_2 = 0.51 \text{ min}^{-1} \) (2);
- ZnO – 5.0 phr, \( k_2 = 0.58 \text{ min}^{-1} \) (3) and
- ZnO – 7.0 phr, \( k_2 = 0.69 \text{ min}^{-1} \) (4)
compositions at 428 K (Fig. 5) shows that by crosslink density and cure rate, which is proved by the constants of speed in the main period ($k$), they surpass the control composition with 5.0 phr of zinc oxide. Improvement of the complex of elasticity-strength parameters of rubbers with ZnCF$\text{O}$ both at normal test conditions and after thermal air aging (Table 1) is probably caused by the influence of the activator under investigation on vulcanization network character. Thus, the percent part of polysulfide bonds (C–S$_x$–C) and amount of sulfur atoms corresponding to one crosslink (S atoms/crosslink) in vulcanizates with ZnCF$\text{O}$ decreases and the percent part of disulfide bonds (C–S$_2$–C) increases (Fig. 6).

Thus, the analysis of the received results, has indicated a possibility of equal-mass replacement of the traditional activator – zinc oxide by the polymer under investigation-inorganic composite (5.0 phr) at maintenance of a high activation level of sulfur vulcanization process of rubber mixes on the basis of diene isoprene rubber and improvement of the physical-mechanical properties complex of their vulcanizates.

As zinc oxide can act not only as an activator of sulfur vulcanization of diene rubbers, but also is a required component of other vulcanization systems, the further research of the ZnCF$\text{O}$ effectiveness was carried out in elastomeric compositions on the basis of polar butadiene nitrile rubber of general and special purpose of sulfur, thiuram, and peroxide vulcanization.

The influence of the ZnCF$\text{O}$ contents (3.0; 5.0 and 7.0 phr) on crosslink kinetics of the modelling unfilled rubber mixes of NBR-26 sulfur, thiuram, and peroxide vulcanization systems of the following composition, phr: NBR-26 – 100.0; sulfur – 1.5; 2-mercaptopentazole – 0.8; stearic acid – 1.5; tetramethylthiuramdisulfide – 3.0; peroximon F-40 – 3.0, is possible to estimate using the data from Fig. 7. As can be seen, the increase of ZnCF$\text{O}$ concentration results in increase of the maximum torque and, accordingly, crosslink degree of elastomeric compositions, decrease of optimum cure time, which, in turn, causes increase of cure rate, confirmed by the calculated values of relative constant of speed in the main period ($k$). The analysis of vulcanizates physical-mechanical properties proves that with the increase of ZnCF$\text{O}$ contents the increase of the relative tensile strength, hardness, and resilience and the decrease of the relative elongation at break and residual deformation at compression by 20 % are observed. Thus, ZnCF$\text{O}$ is an effective component of the mentioned vulcanization systems, as at equal-mass replacement of the known zinc oxides (5.0 phr) the cure rate and the concentration of crosslink bonds are increased and general properties complex of rubber mixes and their vulcanizates are improved.

The comparative estimation of the effectiveness of similar concentrations of zinc oxides and ZnCF$\text{O}$ (3.0; 5.0 and 7.0 phr) as the agents of metaloxide vulcanization system was carried out using simulated unfilled elastomeric compositions of chloroprene rubber of the following composition, phr: chloroprene rubber – 100.0; magnesium

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ZnO 5.0 phr</th>
<th>ZnO 3.0 phr</th>
<th>ZnCF$\text{O}$ 5.0 phr</th>
<th>ZnCF$\text{O}$ 7.0 phr</th>
</tr>
</thead>
<tbody>
<tr>
<td>300% Modulus, MPa</td>
<td>10.0/9.0</td>
<td>10.0/9.0</td>
<td>11.0/10.0</td>
<td>10.0/9.0</td>
</tr>
<tr>
<td>Tensile strength, MPa</td>
<td>27.0/20.5</td>
<td>26.0/22.4</td>
<td>28.5/22.8</td>
<td>20.6/19.4</td>
</tr>
<tr>
<td>Elongation at break, %</td>
<td>890/735</td>
<td>830/750</td>
<td>835/750</td>
<td>835/730</td>
</tr>
<tr>
<td>Tear strength, kN/m</td>
<td>39/28</td>
<td>41/38</td>
<td>43/39</td>
<td>40/32</td>
</tr>
<tr>
<td>Shore A hardness</td>
<td>38/35</td>
<td>40/36</td>
<td>42/38</td>
<td>42/38</td>
</tr>
<tr>
<td>Elasticity, %</td>
<td>62/60</td>
<td>66/64</td>
<td>69/65</td>
<td>68/64</td>
</tr>
</tbody>
</table>

* in the denominator the parameter value after thermal aging of the sample at 373 K and 24 h is given
The negative influence of ZnCFO on the properties of elastomeric compositions of resin vulcanization is caused by interaction between the vulcanization agent (phenolformaldehyde resin on the basis of p-octylphenol – ST-137) and ZnCFO, which results in desactivation of resin as crosslinking agent of resin vulcanization system.

Thus, from the analysis of the results of experimental researches on estimation of ZnCFO vulcanization activity in comparison with zinc oxide in the structure of various vulcanization systems (VS) it follows, that its effectiveness decreases in the series (Fig. 10):

sulfur VS > thiuram VS > metaloxide VS > peroxide VS > resin VS

Taking into account the positive influence of ZnCFO on the formation of the complex of the physical-mechanical characteristics of vulcanizates on the basis of various rubbers and theoretical preconditions of the “composition-structure-properties” interrelation, it was of interest to define the morphology features of elastomeric compositions. With this purpose the percolation method of the analysis by the rheometer data of rubber mixes was used [9].

According to the theory of “composition-structure-properties” interrelation the maximum level of the physical-mechanical characteristics of elastomeric compositions is reached under condition of heterogeneous structure formation with the minimal particles size of heterophase (r – value, opposite to tangent of the corner of kinetics curve inclination) and its optimum contents. The analysis of the data presented in Fig. 11 shows, that replacement of zinc oxide by ZnCFO composite in various vulcanization systems for rubbers of general and special purpose influences morphological structure of compositions, reducing the particles size of heterophase and thereby providing thus high complex of the elasticity-strength properties of vulcanizates. It is necessary to note, that at the increase of the ZnCFO contents from 5.0 up to 7.0 phr parameter r is not essentially changed and retains the minimal value, i.e. the best complex of properties have the compositions with ZnCFO, in which the morphology with the minimal particles size of heterophase is formed at its contents ≈5.0 phr. The given statement corresponds with the experimentally obtained results of physical-mechanical tests of vulcanizates.

The decrease of ZnCFO effectiveness as component of various vulcanization systems for rubbers of general and special purpose in the above presented series (Fig. 10) can be also explained by the character of the formed morphology of compositions. Thus, at use of ZnCFO as the activator of sulfur vulcanization the structure of rubbers with the minimal value of parameter r is formed, and at transition from sulfur to peroxide

is observed, which explains the unsatisfactory level of physical-mechanical parameters of the rubber.
Fig. 7. Cure curves of vulcanization process of modeling elastomeric compositions on the basis of nitrile-butadiene rubber at 428 K with various vulcanization systems.

**Sulfur vulcanization system**
- ZnO 5.0 phr; $k_t=0.29$ min\(^{-1}\)
- ZnCFO 3.0 phr; $k_t=0.22$ min\(^{-1}\)
- ZnCFO 5.0 phr; $k_t=0.25$ min\(^{-1}\)
- ZnCFO 7.0 phr; $k_t=0.29$ min\(^{-1}\)

**Thiuram vulcanization system**
- ZnO 3.0 phr; $k_t=0.33$ min\(^{-1}\)
- ZnCFO 3.0 phr; $k_t=0.41$ min\(^{-1}\)
- ZnCFO 5.0 phr; $k_t=0.56$ min\(^{-1}\)
- ZnCFO 7.0 phr; $k_t=0.58$ min\(^{-1}\)

**Peroxide vulcanization system**
- ZnO 3.0 phr; $k_t=0.11$ min\(^{-1}\)
- ZnCFO 3.0 phr; $k_t=0.08$ min\(^{-1}\)
- ZnCFO 5.0 phr; $k_t=0.12$ min\(^{-1}\)
- ZnCFO 7.0 phr; $k_t=0.13$ min\(^{-1}\)

Fig. 8. Cure curves of metaloxide vulcanization process of modeling elastomeric compositions on the basis of chloroprene rubber at 428K with various type and contents of vulcanization agents.

**Fig. 9.** Vulcanization kinetics of modeling elastomeric compositions on the basis of butyl rubber of resin vulcanization at 433 K with ZnO or ZnCFO.

Fig. 10. Change of elastomeric compositions properties (%) with various vulcanization systems (VS) at ZnCFO presence (in comparison with ZnO): sulfur VS (1); thiuram VS (2); metaloxide VS (3); peroxide VS (4) and resin VS (5).
4. Conclusions

Based on the results of the researches of the ZnCFO effectiveness as the component of various vulcanization systems for rubbers of general and special purpose, it is possible to make the following conclusions:

1. ZnCFO is a chemically connected composite with functionally active groups, which due to its organic-inorganic nature is easily dispersed and combined with rubber matrix.

2. ZnCFO is an effective component of sulfur, thiuram, peroxide and metaloxide vulcanization systems for isoprene, nitrile-butadiene and chloroprene rubbers; at the same time it is not effective in resin vulcanization system for butyl rubber. By the positive influence degree on the properties of elastomeric compositions vulcanization systems with ZnCFO are arranged in a series: sulfur VS > thiuram VS > metaloxide VS > peroxide VS

3. ZnCFO at the contents H° 5.0 phr promotes the formation of morphological structure of compositions with the minimal particles size of heterophase, which results in the improvement of physical-mechanical properties of rubbers.

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The authors express their thanks to professor E. Djagarova (University of Chemical Technology and Metallurgy, Sofia, Bulgaria) for assistance in organization and conducting of experimental researches of estimation of rubber mixes rheological properties, and to professor W. M. Rzymski (Technical University, Lodz, Poland) for determination of sulfidity of vulcanizates crosslinks.

Fig. 11. Dependence of particles size of heterophase of modeling compositions on the basis of various rubbers with different vulcanization systems from the ZnO or ZnCFO contents: nitrile-butadiene rubber of sulfur vulcanization (a); nitrile-butadiene rubber of thiuram vulcanization (b); nitrile-butadiene rubber of peroxide vulcanization (c) and chloroprene rubber of metaloxide vulcanization (d)
References