



PROPERTIES OF THE COMPOSITION BASED ON MODIFIED POLYETHYLENES

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Abstract

The subject of this paper is to study the properties of compositions based on a modified polyethylene and a thermoplastic polyolefin structural features. As a result of the modification of polyethylene, the high elasticity of the material grows, which outwardly makes this polymer similar in complex to the physico-mechanical properties of rubber. The calculations made allowed us to estimate the parameters of the observed thermal transitions and to assign them to the mobility of specific elements of the structure

Keywords: Polyethylene, oil, gasoline, strength, temperature, swelling, fuel, rheology.

1.Introduction

1.1.Structural features of polyolefin thermoplastic elastomers

Polyolefin thermoplastic elastomers were obtained on the basis of ethylene-propylene elastomers and high-density polyethylene by two methods, in the first case sulfur-containing compounds (Altax , sulfur) were used as the modifying agent, in the second - MPC.[1-4]

“A” series thermoplastic elastomers were obtained by combining the initial polymer components in the melt on the rollers at a roll temperature of 433 ÷ 4530 K with a gap of ~ 0.2 mm between them. After melting the crystalline thermoplastic, EPDM and structuring agents were added. To achieve a good homogenization, the rolling was carried out for 25 minutes.

The composition formulation had the following composition (parts by weight): EPDM - 100; high strength polyethylene HDPE - 10 ÷ 80; decimal peroxide 1-2.[5-7].

The thermoplastic elastomers of the “B” series were obtained in the same way as the “A”-type thermoplastic elastomers, according to the following formulation (wt.): EPDM-100; HDPE - 10 ÷ 80; altax 1.8; sulfur - 2.0.

2.Method

2.1.The effect of aggressive liquids on the basic properties of polyethylene

When operating polymeric materials, it is necessary consider the influence of aggressive liquids on their basic properties. The stability of HDPE and its stabilized samples for various types of oils and petroleum products has been studied. In this

connection, it was important to clarify the effect of stabilizers on swelling processes in petroleum products [8-9].

Four types of oil have been researched: Balkan's, Shirvan's, Turkmen's, and coal oils, as well as A-72, A-93, diesel fuel "L" as petroleum products. The samples of HDPE were immersed in the form of plates 1 mm thick in hermetic glassware, where they were kept for a long time at room temperature -200°C . Significant deterioration of strength properties is observed in samples after 12 months of contact with Balkan's oil; comparatively worse properties are shown by HDPE samples containing TU and Thioalkofen-MBP, both in its deformation and strength properties. The original HDPE retains its properties for a long period. The samples containing stabilizers Diafen-NN (N, N'-Di-2-naphthalenyl-1,4-benzenediamine) exhibit high stability [10].

3. Result

Table 1 shows the results of the swelling of HDPE and its stabilized compositions in various petroleum and petroleum products. Samples in a relatively greater degree swell Table 1. The degree of swelling of HDPE and its stabilized samples in oil and fuels at a temperature of $+200^{\circ}\text{C}$, mass%

in petroleum products such as A-72, A-93 gasoline. As the contact time increases, the degree of swelling increases. Significant swelling is observed in the first months, then the swelling rate slows down, and in some cases does not change. Analyzing the data, it can be seen that the HDPE samples stabilized by Diafen-NN and Benzon-00 and Thioalkofen-MBP swell slightly. This, probably, is related to the spherulitic structure of these HDPE samples, which exerts a resistance to the diffusion of petroleum products into the polymer volume.

In oils, swelling occurs to a lesser extent. The exception is coal oil, in which the swelling of the initial HDPE in 12 months reaches 8.4%. In the presence of the Diafen-NN stabilizer, the swelling for this period does not exceed 2.16%. And in this case, the presence of the stabilizer gives a positive effect, which consists in reducing the degree of swelling in various oils and petroleum products. Important in this case is the change in the physico-mechanical properties of the samples after prolonged exposure to the above media.

№	Medium	HDPE				HDPE+TU				HDPE+Benzon-00				HDPE + Thioalkofen-MBP				HDPE+diphene NN			
		Observation time, month																			
		1	4	8	12	1	4	8	12	1	4	8	12	1	4	8	12	1	4	8	12
1	Balkan's oil	1,69	1,82	2,16	2,22	1,73	1,91	2,08	2,15	1,72	1,86	1,88	1,95	1,41	1,54	1,66	1,62	1,70	1,80	1,88	1,92
2	Shirvan's oil	1,67	1,97	2,23	2,52	1,71	2,38	2,41	2,34	1,70	2,18	2,45	2,54	1,38	1,65	1,94	2,01	1,68	2,14	2,39	2,48
3	Coal oil	4,94	6,16	7,36	8,41	5,80	7,05	7,45	7,96	1,50	1,69	2,08	2,21	6,52	6,68	6,84	6,78	1,44	1,65	2,04	2,26
4	Turkmenian oil	1,34	1,72	2,01	2,42	2,31	1,08	1,50	1,64	1,26	1,33	1,37	1,44	1,01	1,24	1,36	1,34	1,23	1,28	1,32	1,40
5	Auto-gasoline A-72	4,33	4,62	4,50	4,90	3,62	3,85	4,12	4,10	3,52	3,59	3,95	4,03	3,75	3,96	3,84	4,02	3,50	3,55	3,81	4,00
6	Auto-gasoline A-93	4,30	4,72	5,03	4,81	3,72	3,74	4,20	4,30	3,59	3,68	3,92	4,01	3,77	3,86	3,95	4,18	3,61	3,74	3,89	4,05
7	Diesel fuel	1,85	2,02	2,60	3,80	1,12	1,43	1,78	1,80	1,11	1,20	1,47	1,71	1,15	1,52	1,84	1,87	1,08	1,16	1,45	1,68

Depending on the field, oil has a tangible effect on the physical and mechanical properties (Table 2) HDPE.

Table 2. Influence of oils from various fields on the physico-chemical properties of HDPE and its stabilized compositions after 12 months.

№	Oil	HDPE		HDPE+ Benzon -00		HDPE+Dep hene-NN		HDPE+TU		HDPE+Thia lcophene MBP	
		σ_p	$\varepsilon, \%$	σ_p	$\varepsilon, \%$	σ_p	$\varepsilon, \%$	σ_p	$\varepsilon, \%$	σ_p	$\varepsilon, \%$
1.	Initial properties	370	300	375	320	394	420	367	300	454	370
2.	Shirvan's oil	382	350	405	850	405	350	350	210	370	-
3.	Turkmenian oil	395	350	320	800	370	870	380	-	206	150
4.	Coal oil	368	400	390	730	350	300	340	250	352	-
5.	Balakhan's oil	342	650	338	750	340	900	318	195	295	200

3.1. Dielectric properties of stabilized polyethylenes.

The use of stabilizers in modern polymer processing technology plays an important role: they are not only inhibitors, but also in some cases structure-forming agents and substantially improve the performance properties of polymers [3-10].

When choosing stabilizers for polyethylene, we were guided by the results

of studies on the inhibitory activity of a number of compounds in the thermal oxidation of HDPE, and we selected: a mixture of amines (diaphone + benzoin), thioalkofen MBP.

The choice of the optimum amount of stabilizer was made by comparing the dielectric properties of HDPE measured at different frequencies (Table 3)

Table 3 The effect of stabilizers on the dielectric properties of HDPE (SD)

№ of the samples	Melting index, g/10 min, J	Ash content, % Z	Stabilizer	stabilizer content weight. %	Dielectric properties			
					ε'		$\text{tg}\delta \times 10^4$	
					f=10 ⁶ Hz	f=10 ¹ Hz	f=10 ⁶ Hz	f=10 ¹⁰ Hz
1	6,3	0,08	Thioalkofen MBP	0,1	2,37	2,32	2,1	2,2
2	-<<-	-<<-	-<<-	0,2	2,34	2,33	3,1	3,2
3	-<<-	-<<-	-<<-	0,3	2,44	2,35	4,5	4,7
4	-<<-	-<<-	-<<-	0,4	2,45	2,35	6,0	6,0
5	-<<-	-<<-	-<<-	0,5	2,47	2,37	7,14	7,4
6	-<<-	-<<-	-<<-	1,0	2,49	2,37	8,9	8,9

7	5,3	0,07	Amines' mixture	0,2	2,37	2,31	3,4	3,4
8	-<<-	-<<-	-<<-	0,1	2,31	2,31	2,7	2,7
9	-<<-	-<<-	-<<-	0,3	2,37	2,27	4,9	4,9
10	6,3	-<<-	-<<-	0,4	2,30	2,30	6,0	6,0
11	5,3	-<<-	-<<-	0,5	2,30	2,30	8,8	8,8

An analysis of the results shows that thioalkofen MBP and a mixture of amines can be introduced into HDPE in an amount of up to 0.2%. At the same time, dielectric parameters remain within the limits of technical requirements. An additional assessment of the stabilizing effect of these stabilizers on HDPE was carried out in the study of atmospheric and artificial aging.

The aging process was carried out on an IP-1-3 wester meter and under atmospheric conditions. The periodic part of the samples was removed and monitored for a change in their appearance, color, melt index, strength dielectric parameters. Atmospheric aging was carried out in the climatic zone of Absheron in the period June-September, which is the richest in solar radiation.

Despite the saturation, there are abnormal structural groups in the HDPE chain, which facilitates the oxidation process under the influence of ultraviolet irradiation. HDPE contains carbonyl groups formed as a result of direct oxidation during aging. In addition, the number of methyl groups in the polymer was 7-15 per 1000 carbon atoms.

In HDPE, there are also at least three types of carbon-carbon bonds. These are terminal ($R-CH=CH_2$), Internal ($R-CH=CH-R$) and lateral

R''

R' >C=CH bonds. The total amount of unsaturation of HDPE subjected to that atmospheric aging was from 1 to 3 g/l.

The formation of carbonyl groups, i.e. the appearance of structural damage in HDPE affects the increase in dielectric losses. Under these conditions, the permittivity practically does not change. As can be seen, in the initial aging period, the melt index decreases from 5.3g / 10min to 2.6, and then increases almost linearly, slightly exceeding the original value by the end of the second month.

The obtained data indicate the simultaneity of the processes of destruction and structuring, with structuring dominating at first, accompanied by intermolecular cross-linking, which is in full accordance with a decrease in the melt index and a slight increase in strength. The increase in the melt index indicates destructive processes, accompanied by a decrease in MW polyethylene. The latter circumstance easily explains the total loss of strength indicators. Thus, the tensile strength in the first 20 days of exposure changes insignificantly, and in the next 10 days it decreases to 25.0 MPa, and in the first week of the second month HDPE becomes practically brittle, as indicated by a significant loss of elongation, from 800 % before aging, up to 250% (Figure 1).

In the process of aging for 90 days, samples of unsterilized HDPE changed their color from white to matte white.

The change in color indicates once again the structural changes that occur during the aging of the polymer.

The aging of HDPE, stabilized with a 0.2% mixture of amines and thioalkofen, is shown in Figure 1 ÷ 4.

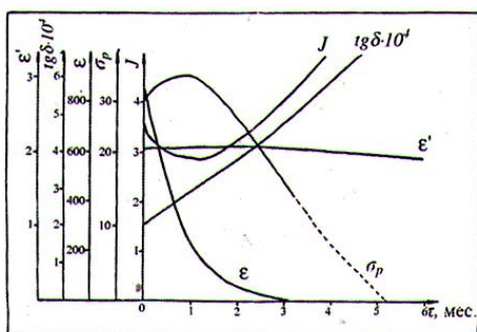


Figure.1. Change in the properties of HDPE in the process of atmospheric aging.

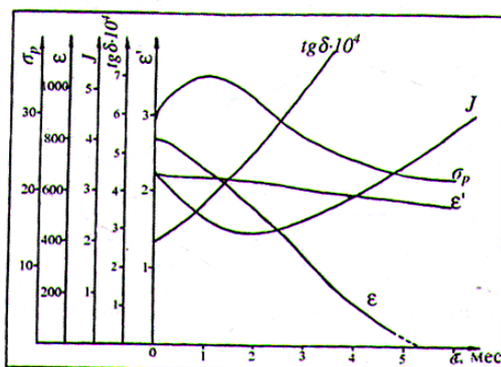


Figure.2. Atmospheric aging of HDPE, stabilized with a mixture of amines.

The tensile strength, as in the case of unstabilized samples, first increases, reaching 32 MPa, and by the end of the first month of exposure in atmospheric conditions, and in the case of artificial aging, after 375 hours, significantly decreases.

With increasing exposure time, an increase in the MM mass of the polymer, controlled by the value of the melt index, was observed. In the process of aging, the tangent of the dielectric loss angle increases linearly, the permeability value drops somewhat, and the color changes from matte to dark yellow. All this testifies to the complexity of the oxidative processes taking place in the polymer under the influence of sunlight [11-12].

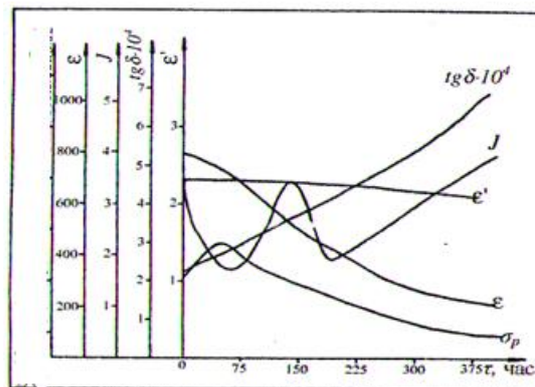


Figure.3. Artificial aging of HDPE, stabilized by thioalkophene.

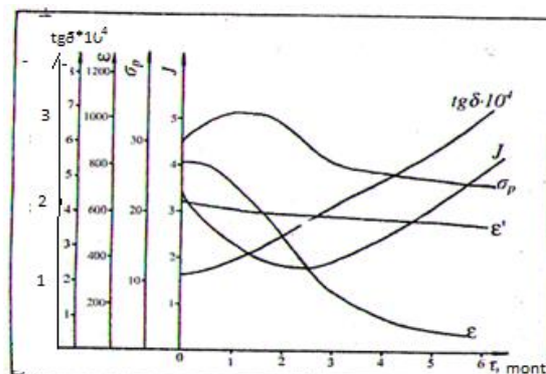


Figure 4. Atmospheric aging of HDPE, stabilized by thioalkofen-MBP.

Simultaneously, the effectiveness of using a mixture of amines to prevent the aging of HDPE was investigated. The results are shown in Figure5.

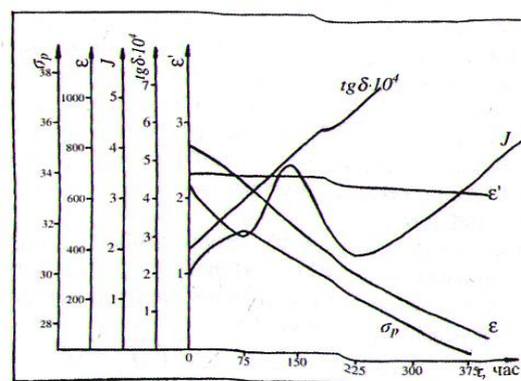


Figure5. Artificial aging of HDPE, stabilized with a mixture of amines.

The data obtained indicate a long-term preservation of dielectric parameters in the aging process.

When using a mixture of amines and thioalkofen MBP as stabilizers HDPE there is an effective stabilizing effect. The use of these stabilizers increases the stability of HDPE properties by 1.5-2 times in atmospheric and in severe artificial aging conditions. However, a lower toxicity, a lower degree of polymer staining, and a lower value of $\text{tg}\delta$ when the thioalkofen MBE is introduced into HDPE makes its use preferable to the use of a mixture of amines. Therefore, further in the development of composite materials, high density polyethylene was used, stabilized with thioalkofen MBE in an amount of 0.15% by weight.

HDPE after 100 hours exposure under UV irradiation completely loses its strength, elasticity and becomes brittle.

This is explained by destruction of the polymer chain and a decrease in MW, the appearance of the gel fraction, $[\eta]$ soluble fraction of polyethylene after 200 hours of irradiation decreases from 1.6 to 0.6. Relatively high resistance to UV irradiation is shown by HDPE containing stabilizing modifiers-diafen-HH (0.9), benzon-00 (1.25) and TU (1.25 mass%), which is explained by the appearance of micro-oriented regions around the stabilizer particles and amplification of amorphous phase.

The introduction of stabilizers-diafen-HH, thioalkofen-MBP and TU raises T_{melting} and $T_{\text{decomposition}}$ HDPE, which is explained by the formation of a less defective fine-crystalline structure in the presence of stabilizers.

The destruction of high-molecular ($M_n \geq 2 \times 10^5$) fractions in polyethylenes after 200 ÷ 500 hours exposure under UV radiation disappears, which is explained by the intensive destruction of the polymer, but the content of the gel fraction also increases up to 40%.

The processes of destruction and *struktuvirovaniya* occur not only in amorphous (20 ÷ 25%), but also in the crystalline phase of polyethylenes.

The presence of the gel fraction is explained by the formation of a network structure due to the recombination of macroradicals.

In the presence of stabilizers, in particular, diphen-NN, as a result of dissociation of light energy by energy, which is not capable of causing chemical degradation of the polymer, partial destruction occurs.

In the melt mode in the process of step cooling, the rate of compaction at the association of small structures into more perfect and large ones occurs according to the established mechanism, regardless of the type of stabilizer.

Stabilizers diphen-NN, thioalkofen-MBP and TU promote a sharp increase in specific volume HDPE and decrease in the density of the melt. HDPE containing these stabilizers over the entire temperature range have higher values of V_c . Stabilizers increase the rate of formation of crystallization centers ($T_{cr} - 124 \div 128$ °C), which is 90 minutes, initial HDPE-150, in the presence of benzene-00-130, TU-130 and thioalkofena-120 minutes. In the presence of stabilizers, the induction period of crystallization is reduced to 6 ÷ 12 minutes, the initial polyethylene is 12 minutes. The obtained results are of practical importance, since stabilizers, in particular, diphen-NN, increase the service life of polyethylene products, reduce the induction period of crystallization and the time of complete crystallization, accelerate the process of polyethylene processing, and significantly shorten the sample holding time in the mold.

The rheological characteristics of stabilized HDPE, in contrast to the initial HDPE in the presence of stabilizers, the viscosity of the melts decreases significantly, depending on the shear stress

and decreases with increasing τ . This is explained by the shape of the associates formed around the particles of the stabilizers and, in connection with this, some increase in the activation energy of the viscous flow of stabilized polyethylene indicates a cooperative displacement of the macromolecules.

HDPE stabilized by diaphene-NN, benzon-00 and tialkofen-MBP in motor gasoline swells slightly. Oil swelling occurs to a lesser degree, except for coal-coal oil, in which the swelling of the initial HDPE in 12 months reaches 8.4% in the presence of diaphene-HH, during this period, the swelling is 2.16%.

The use of a mixture of amines and tialcophene MBE increases the stability of HDPE properties in the atmospheric and artificial aging conditions by a factor of 1.5 to 2, which is explained by the phase reactivity of these stabilizers with the active macroradicals of the systems and their associative action. A lower degree of polymer colouring and a lower value of $g\delta$ is achieved when tialkofen-MBP is added to HDPE in an amount of 0.15% by weight.

The thermoplastic elastomers of the "B" series were prepared by mechanically mixing the calculated quantities of HDPE and EPDM in a melt by rolling. In contrast to the existing methods for the production of thermoplastic elastomers of the "D" series (dielectric), they were obtained on the basis of HDPE and EPDM-60, synthesized under specific conditions: copolymerization of ethylene with propylene and vinylcyclohexene(VCH) was carried out in the presence of catalysts of iso- C_4H_9 AlCl₃ chloride and vanadium triacetyl acetone catalysts (AC₃V) at a temperature of 100 °C.

The resulting powdered crosslinked product had the following characteristics: propylene content,% mol-47; content of IGCC,% mol-1.1; cross-linking density, mol / l- 10. The composition based on cross-linked VCH of ethylene-propylene elastomer (EPDM + VCH) and

polyethylene was prepared similarly to the A and B thermoplastic elastomers.

The exclusion from the composition of the polar activators in this case, such as sulfur-containing and peroxide compounds, leads to an improvement in the dielectric characteristics of polyolefin thermoplastic elastomers.

Figures 6 and 7 show IR spectra of high-density polyethylene, ethylene-propylene terpolymer (3rd component - ethylidene-norbornene) and thermoplastic elastomers based on them, containing various amounts of HDPE in the composition.

The carbon-carbon bond of the polymer chain represents an absorption band at 1100 cm^{-1} . Information on the lateral methyl branches carries an absorption band at 1378 cm^{-1} and the total number of branches per 1000 carbon atoms calculated for HDPE was only 4-7, while for the EPDM they are about 200-300, i.e. many times more. In the initial HDPE, along with the alkyl branching, there is a small amount of unsaturated vinyl (910 cm^{-1}) and vinylidene (3020-3010 cm^{-1}) bonds. The presence of the crystalline phase was fixed by the doublet of the bands 720-731 cm^{-1} . The determination of the ratio of the integral absorption values of the two components of the doublet of the given polymer for the estimation of the degree of crystallinity was not carried out by us because of the large measurement error in comparison with the x-ray phase method.

In the ethylene-propylene elastomer of EPDM-60 there is a band of pendulum vibration of the CH₂ group at 720 cm^{-1} , while the absence of the doublet 720-731 cm^{-1} indicates its amorphousness. The amount of propylene in the chain was characterized by the intensity of the band 1378 cm^{-1} . The presence of a diene in the polymer chain is indicated by a band in the 3045 cm^{-1} region.

For TEP, a decrease in the intensity of the absorption bands inherent in the

initial elastomer is noted, but a disproportionate change in these characteristics allows us to note some important results.

With an increase in the elastomer content in TEP, its characteristic absorption bands at 720, 620, 870, 930, 1150, 1380, 1460, 70, 2725, 3180, 4250 and 4310 cm^{-1} increase in intensity, however, the reverse estimated calculations of the EPDM content in TEP's intensity of the absorption bands of the propylene unit, does not correspond to its practical content. In addition, the observed weakening of the band at 690-900 cm^{-1} , which refers to the end vinylidene groups $>\text{C}=\text{CH}_2$ in EPDM, combined with a disproportionate change in the band at 1150 cm^{-1} indicates chemical transformations during the preparation of the TEP.

The very small fraction of the uncertainty in all TEP series is evidenced by the absorption in the region of 1580 \div 1800 cm^{-1} , 3020 \div 3010 cm^{-1} . 1000 cm^{-1} . The IR spectra of the T-series of the 'B' series show that in this case a simple superposition of the spectra characteristic of the initial polymer components i.e. the presence of a mechanical mixture is observed, while the TEPs of the 'B' and 'D' series are observed to convert the 980/960 cm^{-1} singlet at 980 cm^{-1} , which indicates the interaction of the elastomeric phase over the double bonds during the modification process. For the TEP of series 'A', a strong absorption of the 1378 cm^{-1} band characterizing the number of CH_3 -groups is observed.

The appearance of alcohol bands of CH groups at 3370, 1040, 1120 cm^{-1} is also observed, which are probably the result of interaction of polymer components with MPC and indicate that partial interaction between the elastomer and the polyethylene phase occurs in the TEP of series 'A'. To confirm this assumption, TICs have been treated with boiling xylene, which is the selective solvent of the crystalline phase of the polyolefin.

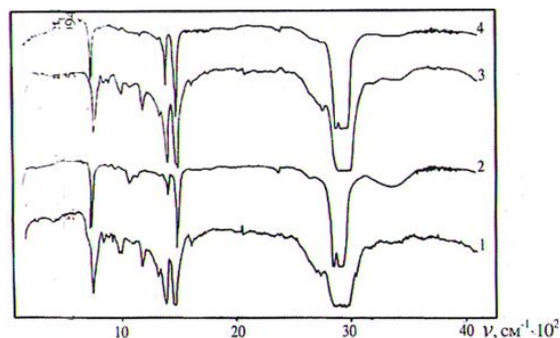


Figure 6. IR spectra of HDPE (2), EPDM (1), and thermoplastic elastomers of series 'A' (3) and 'D' (4).

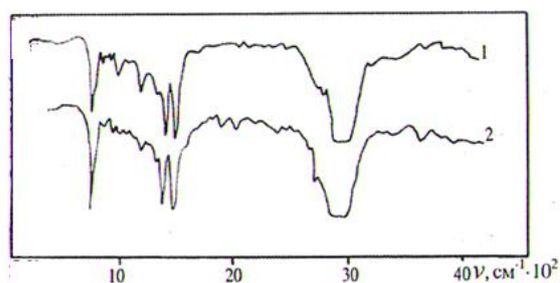


Figure 7. IR spectra of TEP series 'B' (1) and 'D' (2), subjected to boiling xylene treatment.

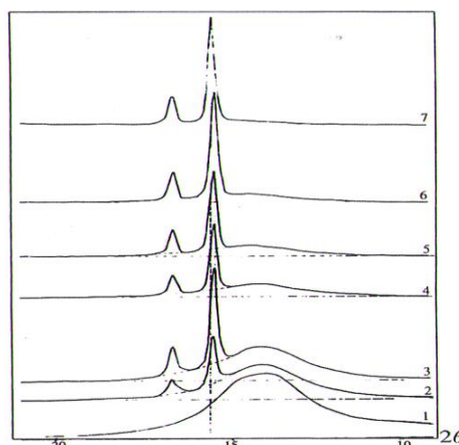


Figure 8. Diffraction curves of the initial EPDM-E (1) and HDPE (7) and thermoplastic elastomers 'A'-series containing 10, 20, 30, 40 and 50% HDPE (respectively 2 \div 6); 1 \div 3 at 200 imp / s, 4 \div 7 at 400 imp / s.

For TEPs of series B and D after aging in boiling xylene, almost complete removal of the polyethylene crystalline phase from the elastomer matrix is observed, which indicates that there is no chemical

interaction between the polymer components. In the TEP series A, even after prolonged exposure to boiling xylene, it was not possible to completely remove the polyethylene phase, which indicates partial interaction of the initial polymer

components during the modification process, increasing the MPC concentration leads to a decrease in the fraction of the extracted amount of PE in the boiling xylene (Table 4.).

Table 4. The amount of extracted HDPE after holding TEP in boiling xylene for after 72 hours.

Name	TEP series A amount of peroxide			'B' series TEP	'D' series TEP
	0,2	0,8	1,0		
The amount of PE extracted from TEP, % by mass.	97	90,7	81,6	100	100

Thus, the IR spectroscopic data allow us to conclude that, in contrast to the mechanical mixture, TEP, under the action of modifiers, chemical transformations occur in the elastomeric matrix, and in TEPs of the 'B' and 'D' series, these transformations do not affect the polyethylene phase. In the TEP of the 'A' series, both polymeric components enter into the interaction with the modifier (dicumyl peroxide), which allows one to conclude that partial PE grafting to the elastomer.

The crystalline phase in the TEP was studied by X-ray diffraction. The degree of crystallinity decreases monotonically with increasing volume fraction of the elastomeric component for all series of TICs. On the diffraction curves (Fig. 8) of the A, B and D series samples, in contrast to the 'B'-series (mechanical mixture) samples, a certain shift of the diffraction reflections of the polyethylene component

toward smaller angles occurs. In addition, the diffraction reflexes broaden, which in combination with the observed shift allows us to conclude that in the synthesis of TICs of series A, B, and D, an elementary crystal cell of PE is distorted. The performed calculation of the parameters "a", "b" and "c" of the crystal cell of these TEPs (Table 5) confirms the above assumption.

It should also be noted that with the increase in the content of the elastomeric phase, the overall background of the amorphous halo grows on the diffractogram, however, there is a tendency for it to narrow and increase its maximum position along the intensity scale, that is, in the arrangement of disordered HDPE molecules and elastomer, "Short-range order," perhaps, due to the formation of sub segmental microorganisms. This, in turn, indicates a greater degree of structural disorder of the TIC in comparison with the mechanical mixture-TEP of the 'B' series.

Table 5

The effective sizes of crystallites and the parameters of a crystalline cell of polyolefin TEP

TEP series	Degree of crystallinity, %	The effective size of the crystallites L, Å	The unit cell parameters, Å		
			a	b	c
A	40	202	7,41	4,943	2,53
B	42	202	7,42	4,943	2,53
D	41	200	7,42	4,942	2,53
C	40	170	7,42	4,972	2,53

Direct studies of the morphological features of the TEPs studied were carried out by electron microscopy. This allowed us to visually assess the degree of heterogeneity and reveal the nature of supramolecular formations in TEP.

From the above-mentioned electron micrographs, for all the investigated series

of TEPs, a characteristic formation of a typical heterophase system is evident. An increase in the degree of crystallinity in the TEP leads to the formation of spherulite structures with diffusion boundaries. It should be noted that the TEP series A and the mechanical mixture ('B' series) are more homogeneous systems than the other types of polymers studied (Figure.9)

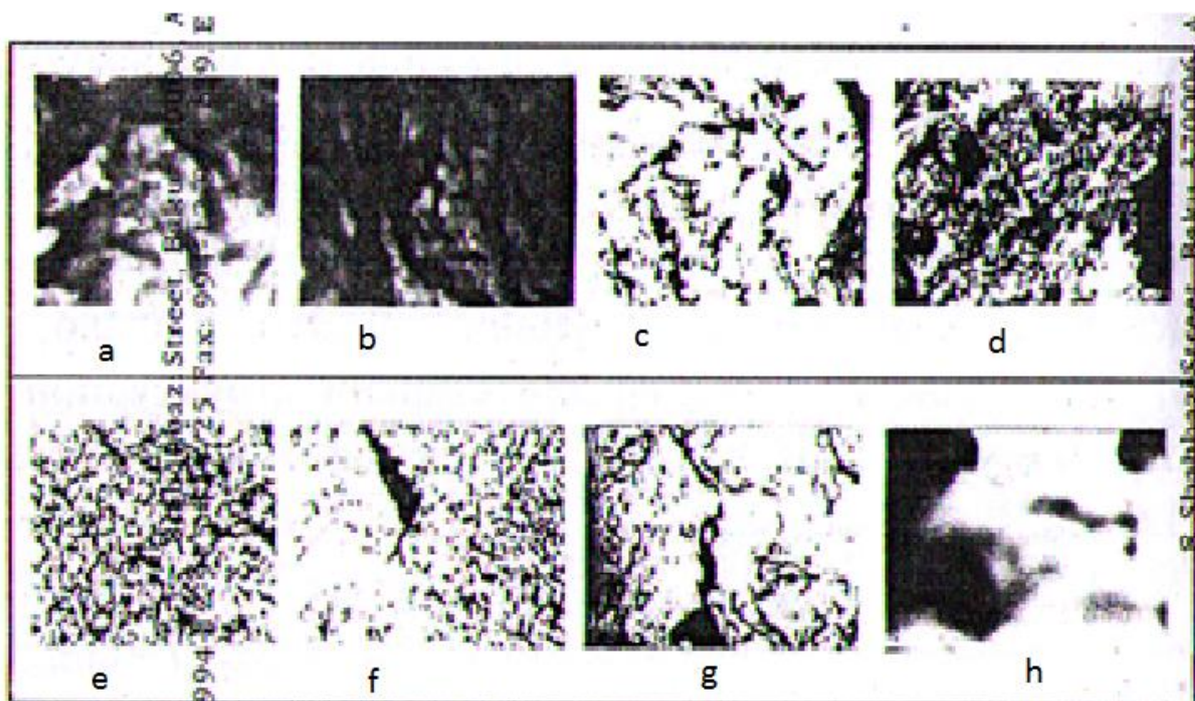


Figure9. Electronic photomicrographs of TEP series A with SREPT-60 content, mass%: 40 (a) and 60 (b); Series B SREPT-60 HDPE 40 (c) and 60 (d); Series D SREPT-60 20 (e), 40 (f), 60 (g) and B series SREPT-60 40 (h).

Electron microscopic data show that in the region of SREPT-60 concentrations of about 60%, the formation of polymer mixtures of a static nature is observed, which, unlike the mixtures of the matrix chaotactor, have an uneven distribution of one phase into the other with larger domain switches. Table 6 gives the dimensions of the domain inclusions in the TEP, determined by electron microscopy sampling. These data show that in the

region of critical concentrations of the polyethylene phase (50-60% w / w), a considerable increase in the size of the domain inclusions is observed, which leads to loosening of the interphase layer of the elastomeric matrix. A further increase in the concentration of HDPE leads to the reversal of polymer phases, where the continuous phase-matrix is already a crystalline component, which leads to an increase in the density of molecular packagings

Table 6. Effect of the composition of the thermoplastic elastomer series 'D' on the degree of crystallinity and the dimensions of heterogeneous microregions

HDPE content in TEP, % wt.	Degree of crystallinity, %	The effective size of the crystallites L, A'	The sizes of heterogeneous inclusions, A'		
			Minimum	Medium	Maximum
20	18	150,3	30-40	200	350
30	26	161,5	35-40	420	750
40	40	169,0	35-45	500	900
50	47	180,2	45-50	900	1700
60	54	192,0	45-50	1200	2700

The DTA results of the TEPs under study are in good agreement with the above data. Demi-HDPE according to DTA is 136 °C. Ethylene-propylene elastomers do not have a definite value due to the amorphous structure. According to DTA, as the content of the amorphous phase of the elastomer in the TEP increases, the areas of melting peaks decrease. The melting of crystalline domains in the matrix of elastomers in TEPs of series 'B' and 'D' occurs at a temperature of 2 ÷ 30 °C lower than that of the initial SREPT-60, which is characteristic for mechanical mixtures and indicates the absence of interaction between the polymer components. The TEP of series A recorded a sharper fall in the T_{cl} of crystalline domains in the elastomer matrix (T_m = 126°C); which indicates a partial interaction between the initial polymer components. A certain increase in the oxidation effect is also observed with the formation of hydroperoxide groups and a decrease in the exo-effect, which prevents the destruction of the sample. For the TEP of the 'D' series, the oxidation process is characterized by a low intensity with a maximum of the exoeffect at 295 °C. A more intense exoeffect with a maximum at 280°C is observed for the mechanical mixture, and the exothermic thermo-oxidative degradation occupies an intermediate position between the initial polymer components and has a maximum at 420°C.

Mass losses change in the following sequence: TEP series D < TEP series B < TEP series A. DTA data in combination with the results of IRS show that the presence of by-products of the decomposition of modifiers leads to a decrease in the resistance to thermal oxidative destruction of TEP, for example a sample of the A series. incompatibility of HDPE and EPDM, which is obviously caused by differences in the formation of final TEP structures, under conditions of their production. (Figure 10)

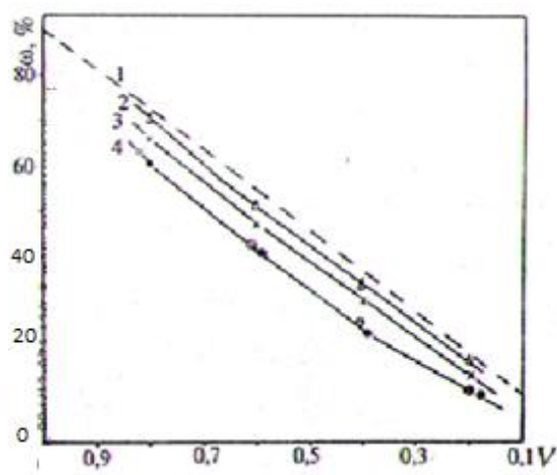


Figure 10. Comparison of the experimental dependences of the degree of crystallinity ω on the volume fraction of HDPE in the TEP of the series A (4), B (3), C (4) and D (2) with a relatively limiting theoretical dependence (1).

To estimate the degree of inhomogeneity, a technique was used that takes into account that ω for extremely inhomogeneous compositions is proportional to the volume fraction V of the crystalline component $\omega = \omega_v = 1 (V)$. Extremely homogeneous mixtures at $V = \text{const}$ have a lower degree of crystallinity, and the more consistently the more homogeneous the composition, the more complex the process of crystallization.

Comparing the values of the experimental curve $\omega = \omega(V)$ with respect to the limiting theoretical values, one can judge the degree of compatibility of the compositions under study. From the degree of heterogeneity, polyolefin TEPs can be arranged in the following series: $D > B > A$.

It follows from the above studies of the TEP structure that this polymer system is characterized by a heterogeneity of a higher order than simple mechanical mixtures. Depending on the method of production, the TEPs differ somewhat, the D and B samples being the most heterogeneous. Samples of the A series occupy an intermediate position between them and the mechanical mixture.

The different degree of heterogeneity of the above polymer systems necessitates a more detailed consideration of the compatibility of the initial components on the basis of which TEPs are formed. Thus, in the process of studying the properties of a large series of mixtures of polymers, it was shown that the compatibility factor has a significant effect on the phase state of the system under investigation

From the point of view of thermodynamics, a stable single-phase system is characterized by a decrease in the isobaric-isothermal potential ($\Delta G < 0$) and is described by the well-known equation:

$$\Delta G = \Delta H - T\Delta S \quad (.1)$$

where ΔH is the enthalpy change, ΔS is the entropy change. In view of the smallness of the $T\Delta S$ value, the evaluation of the state of the structure of polymer systems is characterized by ΔH , where the negative values ($\Delta H < 0$) correspond to the compatibility of the polymers.

It is practically very difficult to determine ΔG by direct methods in the process of polymer displacement, and therefore the degree of compatibility is determined indirectly by the interaction parameter h , which is defined as:

$$h = V_s/RT(\delta_1 - \delta_2)^2 \quad (.2)$$

where, h is the Hildenbrand interaction parameter, δ_1 and δ_2 are the solubility parameters of the components being mixed, and V_s is the volume of the mixtures

Calculation of solubility parameters for HDPE and SREPT-60 compositions leads to the following values:

$$\Delta_{HDPE} = \delta_{EPDM} = 0,81 \text{ (cal/sm}^3)^{0,5}$$

Despite the equality of the solubility parameters for these polymers, due to the specificity of their structure, thermodynamic incompatibility can be observed, since the gain of entropy as a result of mixing of two dissimilar molecules compensates for the energy expenditure on the destruction of supramolecular structures and intermolecular bonds in the original components. Thus, the complexity of the molecular structure of polymers makes it difficult, in this case, to apply thermodynamic criteria in assessing the degree of compatibility, as a result of which an attempt was made to apply relaxation spectrometry.

4. Discussion

4. 1. Relaxation processes in thermoplastic elastomers.

When combined, intensive processes of thermal mobility occur, causing the greatest

dissipation of energy. In particular, the vitrification process is more intensive, i. e. α -process of segmental mobility. The manifestation of relaxation regions associated with segmental mobility, makes it possible to uniquely determine whether these polymer systems are homogeneous or heterogeneous. The presence in the complex binary polymer mixtures of the two main relaxation regions, vitrification, indicates the heterogeneity of the system, while the existence of one main glass transition region indicates its homogeneity

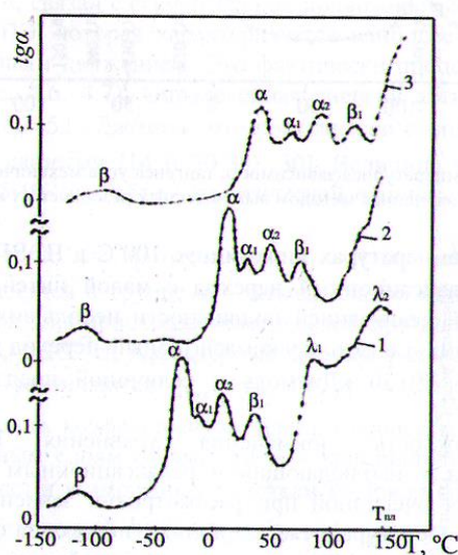


Figure 11. Temperature dependence of the tangent of the angle of mechanical losses in SREPT-60 at a frequency of 2 Hz (1), 10 Hz (2) and 1000 Hz (3).

We have obtained the temperature dependences of the tangent of the angle of mechanical losses for several frequencies by the forced oscillation method. These spectra for SREPT-60 and a similar picture of losses in the regime of forced resonance oscillations are shown in Figure 11-12

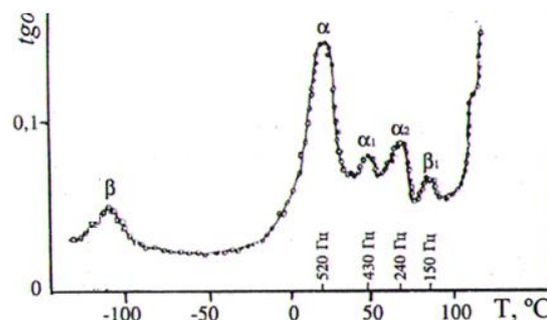


Figure 12. The temperature dependence of the tangent of the angle of mechanical losses of SREPT-60, obtained by the method of forced resonance oscillations.

At temperatures below minus 100 °C, a β -relaxation transition with a low intensity due to the mobility of small structural units is observed in SREPT-60. this is a small-scale transition with an activation energy of $29 \div 30$ kJ / mol and a pre-exponential value of $B_i = 1.6 \times 10^{-12}$ s.

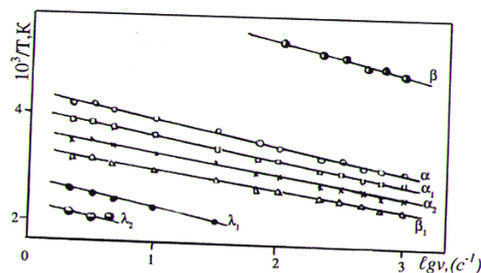


Figure 13. Dependence between the inverse temperature of the relaxation transitions in SREPT-60 and the frequency

The next temperature relaxation α -transition is associated with segmental mobility in the amorphous phase of PE, which is characterized by the most intense thermal motion. This is actually the process of vitrification (Figure 11, 12,13). The specific activation energy is $U_\alpha = 52$ kJ / mol, which practically coincides with the known data. The value of the pre-exponential is $B_i = 1.6 \times 10^{-12}$ s. and the volume of the kinetic unit is $v = 10^{-21}$ cm³ (Table 7).

The next two processes, i.e. α_1 and α_2 - processes in SREPT-60 belong to the α -segmental mobility group, since for all these processes the kinetic unit is the segment / $B_i = 5 \times 10^{-12}$ s. The characteristic relaxation times of these processes differ ($\tau_\alpha < \tau_{\alpha_1} < \tau_{\alpha_2}$) due to the difference in the temperature coefficients. These processes refer to transition layers from the amorphous and crystalline phase (α_1 -process) and to amorphous regions of spherulite fibrils, i.e. crystalline phase (α_2 - process), the reason for the multiplicity of the α -transition in polyethylene is, first of

all, the complexity of the morphological structure that causes differences in energy conditions and changes in the freedom of segments.

B_i - processes small-scale mobility in the crystalline phase. Three slow relaxation λ_i transitions (Figure 14, 15) with an activation energy of 49 kJ / mol, i.e., were detected by the voltage relaxation method in HDPE. close to the activation energy of the viscous flow of SREPT-60. Dimensions of kinetic units are presented in (Table 7)

Table 7. The characteristics of the relaxation transitions in SREPT-60, determined from the totality of the oscillations:

Relaxation period	Activation energy, kJ/mol			coefficient B_i^{**} , s	Kinetic unit		
	a	b	c		Volume, V_k , sm^3	Average linear dimension, A^*	Structural representations
β	30	29	-	$1,6 \times 10^{-13}$	2×10^{-22}	5-7	Several CH_2 groups
β_1	68	67	-	$1,6 \times 10^{-13}$	2×10^{-22}	5-7	-«-
α	52	51	-	5×10^{-12}	10^{-21}	30-40	20-25 groups CH_2 (segment)
α_1 of glassification	54	54	-	5×10^{-12}	10^{-21}	30-40	Segment
α_2	59	60	-	5×10^{-12}	10^{-21}	30-40	Segment
λ_1	48	48	49	$5,2 \times 10^{-8}$			Physical node (microblock)
λ_2	48	-	49	$4,9 \times 10^{-7}$			-«-
λ_3	-	-	49	$5,1 \times 10^{-6}$			-«-

a-stimulated oscillations, b-resonant oscillations, c-relaxation of voltage ** точность определения U ($\pm 0,5+0,8$ kJ/mol); B_i ($\pm 20\%$).

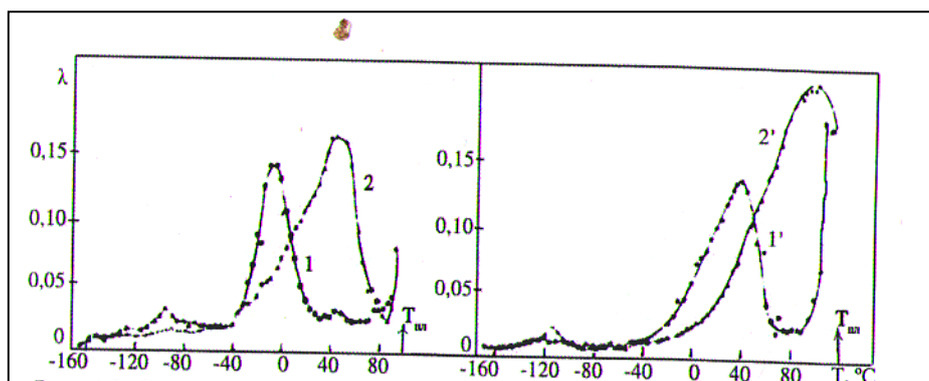


Figure14. Spectra of internal friction for SREPT-60 with different degrees of crystallinity: 1- 0.62; 2-0.56; 1'- 0.50; 2'-0.44

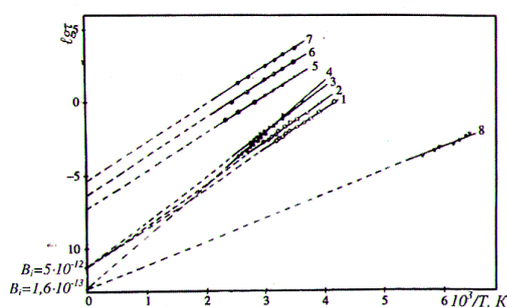


Figure15. Dependence of the relaxation times of slow (λ_i) and fast relaxation (α) transitions in HDPE on the inverse absolute temperature (according to Table 7).

The internal friction spectra of the SREPT-60 epylene-propylene elastomer, weakly cross-linked to retain the MPC form (a), sulfur-containing components (b) and vinylcyclohexene (c) (Figure16.) were obtained.

Area of manifestation and mechanism β - process in the elastomer is similar to that studied in HDPE. The activation energy (α) of transitions for ethylene-propylene elastomers, regardless of the type of cross-linking, is 47-49 kJ / mol (Table 8). The main structural unit of this process is a free segment with average sizes of $30 \div 40 \text{ \AA}^*$ and a volume in the order of 10^{-21} cm^3 , which indicates the generality of segmental mobility in EPDM with other elastomers.

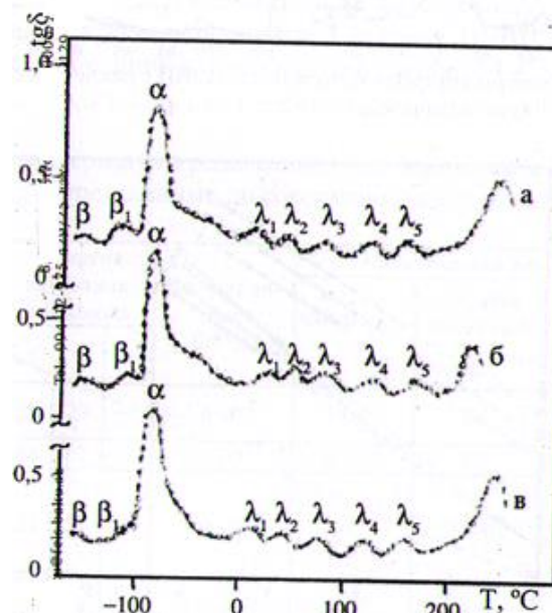


Figure16. Spectra of internal friction of ethylene-propylene elastomer crosslinked with dicumyl peroxide (a), sulfur-containing components (b) and vinylcyclohexene (c) 1 Hz.

In the region of higher temperatures, δ_s and δ_c processes are observed in the elastomeric matrix, depending on the type of cross-links. Process δ_s - is characteristic of sulfide cross-links and has an activation energy of 130 kJ / mol. (EPDM, cross-linked with sulfur-containing compounds). The process of δ_c -relaxation refers to the chemical relaxation of transverse bonds of the "C-C" type, characterized by an activation energy of 150-152 kJ / mol, observed in the EPDM, crosslinked MPC and IGCC.

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Table 8.Characteristic of relaxation transitions in the EPDM, crosslinked by MPC (a), sulfur-containing agents (b) and VCH (c).

Relaxation transition	Activation energy $U_i, \text{kJ/mol}$			Coefficient B_i, C			The size of the kinetic unit, A^*		
	a	b	B	a	b	B	a	B	B
β	28	29	28	$1,6 \times 10^{-13}$	$1,6 \times 10^{-13}$	$1,6 \times 10^{-13}$	5	+	-
β_1	40	42	41	$1,6 \times 10^{-13}$	$1,6 \times 10^{-13}$	$1,6 \times 10^{-13}$	5	+	-
α	45	46	45	5×10^{-12}	5×10^{-12}	5×10^{-13}	30	+	40
λ_1	46	46	45	$1,1 \times 10^{-7}$	$9,8 \times 10^{-6}$	$1,0 \times 10^{-7}$	500	600	550
λ_2	46	46	45	$9,3 \times 10^{-7}$	$5,4 \times 10^{-6}$	$8,5 \times 10^{-7}$	1000	1300	900
λ_3	52	53	54	$8,4 \times 10^{-6}$	$1,2 \times 10^{-6}$	$1,7 \times 10^{-6}$	1200	1000	1200
λ_4	59	59	60	$1,0 \times 10^{-7}$	$9,3 \times 10^{-7}$	$9,0 \times 10^{-7}$	900	1200	1100
λ_5	59	59	60	$6,8 \times 10^{-7}$	$6,3 \times 10^{-7}$	$6,7 \times 10^{-7}$	2500	2200	2000
δ_s	-	132	-	-	$8,2 \times 10^{-14}$	-	-	1-2	-
δ_c	152	-	150	$3,3 \times 10^{-14}$	-	$3,3 \times 10^{-14}$	2-3	-	2-3

5. Conclusion

The specificity of the structure that arises when highly crystalline polymers of the high-strength polyethylene (HSPE) type are added to the ethylene-propylene matrix and, as already noted, is characterized by the presence of inhomogeneity, leads to a sharp change in the relaxation properties of the polymer system as a whole.

Since the presence of a crystalline phase in the mixture significantly affects the entire complex of physico-mechanical properties of similar polymer systems, thermoplastic elastomers were taken for comparison, differing only in methods of preparation, i.e. thermoplastic elastomers (TPE) of various series containing 40% by weight of polyethylene. were accepted.

The temperature dependences of the mechanical loss tangent of B-series TPE at different frequencies (from $5 \times 10^{-3} \text{s}^{-1}$ to 10s^{-1}) are presented. As the frequency increases, an uneven shift of the peaks of the manifested internal losses to the right on the temperature scale is observed.

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