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MEMBRANOTROPIC ACTION OF MACROHETEROCYCLIC CROWN-ETHERS

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One of the actual scientific problems nowadays should be the investigation of newly obtained xenobiotics, with the goal of establishment of their biological mechanism and potential danger on the environment. Crown-ethers could be regarded as quite widely-spread industrial chemical pollutants of biosphere, especially of water ecosystems. These compounds are derivatives of polyesters and represented by macroheterocyclic systems with 9-60 atoms in the cycle from which one third belongs to the atoms of etherized oxygen divided by ethane groups. Crown-ethers are characterized by high volume of production, for they are applied in different fields of electrochemistry, pharmacy, medicine, because of their unique properties of solubility in many non-aqueous solvents, high stability, selectivity of oxido-reductive reactions, ability to form complexes with metals, etc. The insufficient effectiveness of water-cleansing constructions and methods from the given substances may result in the crown-ethers invasion to the human organism in the composition of drinkable water.

Previously we showed that in the processes of hydrolytic and thermal aqueous destruction and of biological organism transformation, crown-ethers heterocyclic rings break down to the enormous spectrum of biologically active low-molecular compounds. The majority of these compounds were proved to be far more toxic than their precursors and to cause membranotropic, radiomimetic, gonadotoxic, and other negative effects upon the organism of warm-blooded animals [1; 2]. Besides, crown-ethers themselves, which are quite lipophilic and extremely cumulative substances [2] with complex-forming ionophoric properties, may cause membranotropic action.

Objective. Investigation of membranotropic action which could be a feature of crown-ethers biological activity by establishment of negative influence of 12-crown-4, aza-12-crown-4 and thia-12-crown-4 upon erythrocytes and hepatocytes membrane phospholipid composition, as well as serotonin receptors state, and cyclic nucleotides system (cAMP and cGMP concentrations, cyclic nucleotides metabolism enzymes activities) state in white rats brain.

The investigation involved the usage of 32 male rats of Vistar line (body mass 200-220 g). The animals were divided into three experimental and one control group. The experimental groups of rats were administered with emulsion of the investigated crown-ethers in 1/100 LD50 (0.0117; 0.022; 0.0365 g/body mass kg, for 12-crown-4, aza-12-crown-4 and thia-12-crown-4 respectively [2]) daily within 30 days perorally. The animals of the control group were given water at the same conditions. On the 30th day of the experiment the rats of all groups were anesthetized by sodium thiopental (50 mg/body mass kg [3]) and slaughtered by decapitation with the Guillotine knife. The rat brain neocortex was isolated in the cold and was frozen in liquid nitrogen subsequently for investigation of receptors and membrane-bound enzymes.

Investigation of phospholipid composition of hepatocytes and erythrocytes was performed using liver and blood of the same animals. For this, erythrocytes were washed out thoroughly by isotonic NaCl solution using triple centrifuging, whereas hepatocytes were obtained by homogenizing rats liver in the Potter's glass homogenizer. Membranes were isolated by general methods using recommendations [4]. Lipid extraction was performed by [5]. The ratio of phospholipid factions was calculated as a phosphorus percentage of each faction phospholipids to the total phosphorus sum of all phospholipids taken as 100 %. For the investigation of erythrocyte phospholipid composition we determined the contents of phosphatidyl choline (PC),

sphingomyeline (SM), phosphatidylserine (PS), lysophosphatidylcholine (LPC) and phosphatidylethanolamine (PEA). In the liver, we additionally investigated the contents of lysophosphatidylethanolamine (LPEA), phosphatidylinositol (PI), phosphatidic acid (PA) and cardiolipin (CL).

The binding parameters of selective ligands with the first and second type serotonin receptors were established by aid of determination of ^3H -serotonin (for 5-HT-1 receptors) and ^3H -piperone (for 5-HT-2 receptors) with rat neocortex synaptosomes membranes using the method [6]. The synaptosome fraction was obtained with the method [7]. The protein contents were determined by the Lowry method [8]. The protein contents were estimated at 300-500 mkg per sample. The result calculations were performed by using Skatchard graphs of IBM program "Ligand". The cyclic nucleotides system state was evaluated by cAMP and cGMP contents determination in rough membrane fraction of rat brain neocortex with the usage of cyclic nucleotides "AMERSHAM" standards. Adenylate cyclase (E.C. 4.6.1.1) activity was determined with the method described by [9] with insignificant modifications; guanylate cyclase (E.C. 4.6.1.2) activity – with the method [10]. The basal level of the enzymes activity was considered. Phosphodiesterase (E.C. 3.1.14.17) activity was determined using the method [11].

The experimental results of phospholipids composition investigation show crown-ethers as the agents which significantly alter the ratio of rat erythrocytes and hepatocytes membrane phospholipid fractions, and the alterations directions were typical for 12-crown-4 and aza-12crown-4. Particularly, in hepatocytes, the consequences of the xenobiotics action displayed as the increase in PC and CL contents and decrease in PI (authentically only for 12-crown-4) and SM. The percentage ratio of PS and PEA remained at the control level. The percentage of PEA and PC lysoforms should be noted to get authentically increased in erythrocytes and hepatocytes of animals having been toxified by 12-crown-4 and aza-12-crown-4. The ratio of erythrocytes and hepatocytes membrane phospholipid fractions in the organism of rats toxified by thia-12-crown-4 had only the tendency to alter mostly in the similar way with the two previous experimental groups. The increase in phospholipids lysoforms contents percentage may be explained by induction of lipid peroxidation. This is verified by the earlier obtained data signifying malonic dialdehyde and dienic conjugates accumulation in liver and blood, blood biochemiluminescence induction, decrease in reduced glutathione contents, and prooxidant protection enzymes activity changes in the organism of rats, intoxicated by crown-ethers [2]. The cause of lipid peroxidation enhance is the increased generation of oxygen active forms by microsomal monooxygenase system and formation of xenobiotics biotransformation products (aldehydes, ketones, alcohols), which possess prooxidant effects. In spite of increased lysoform percentage, the percentage of PEA did not change and the one of PC even grew up, which could have been connected with the increase in metabolism rate of the mentioned phospholipids fractions in erythrocytes and hepatocytes of rats of the experimental groups. As far as CL are the main lipid components of mitochondrial membranes their contents alterations, and, as a consequence, alterations of phospholipid microsurrroundings of mitochondrial membrane enzymes may be one of the reasons of bionenergetics processes impairment which is proved by our previously reported experimental results about the decrease in activities of succinate dehydrogenase, monoamine oxidase, and ATPases rat hepatocytes [12]. The decrease in liver PI at the influence of 12-crown-4 may be caused, on one side, by activation of free radical processes, on the other side, by induction of prostaglandins synthesis which is also proved by our experimental results [2]. 5-HT-1- and 5-HT-2-receptors selective ligands binding parameters in the brain of rats toxified by the xenobiotics were different from the same indexes of the control animal group. The differences in the indexes of all the three experimental rat groups were similarly directed. The investigated crown-ethers action resulted in the untypical alterations of selective ligands binding character by both of the receptors types. The influence of the experimental substances manifested as the increase in receptors affinity and decrease in binding sites quantity. The influencing force of the three individual xenobiotics was not quite different between them. 5-HT-1-receptors affinity increased by 24-36 %, their binding sites quantity decreased by 16-21 %. Kddifferences of 5-HT-2-receptors high affinity pool of experimental animals from the control magnitudes were in the range

of 16-32 %, of 5-HT₂-receptors low affinity pool - in 12-25 %. ³H-spiperone binding sites quantity of toxified rats were lower from the control magnitudes by 10-25 % (high affinity pool), and by 17-32 % (low affinity pool).

The crown-ethers action resulted in reduction of adenylate cyclase activity in rats' neocortex. 12-crown-4 displayed 43 % fall of this enzyme activity whereas aza-12-crown-4 and thia-12-crown-4 showed only 28 and 18 % falls respectively. All results are authenticable compared to the control magnitude. The reduction in the adenylate cyclase activity correlated with decrease in cAMP contents: - 47 %, - 25 % and - 17 % for 12-crown-4, aza-12-crown-4, and thia-12-crown-4 respectively. The opposite character of the xenobiotic influence was found for the system "guanylate cyclase - cGMP". 12-crown-4, aza-12-crown-4, and thia-12-crown-4 action led to the induction of guanylate cyclase activity by 122, 81, and 54 %, and to increase in cGMP contents by 105, 77, and 48 %, respectively, and authentically, compared to the control magnitude. The activity of phosphodiesterase - a catabolic enzyme of cyclic nucleotides metabolism manifested its induction in the organism of rats toxified with 12-crown-4, aza-12-crown-4, and thia-12-crown-4 by 108, 77 and 15 % respectively compared to the control magnitude (authenticable only for the first two substances). Thus, the obtained results display the pronounced influence of crown-ethers upon the system of cyclic nucleotides in experimental rats' neocortex. This influence shows the alterations of cyclic nucleotides catabolic and anabolic enzymes activity as well as the changes in cAMP and cGMP contents. The experimental crown-ethers do not have a similarity in chemical structure with endogenic bioregulation molecules - hormones and neurotransmitters, which realize their specific influence upon cells, particularly, via cyclic nucleotides, as second messengers. Thereby, we cannot predict a selective influence of crown-ethers upon receptor and post-receptor links of intercellular information realization whatsoever. The obtained data may be explained by non-specific modulatory character of these xenobiotics influence on membrane receptor, enzymic, channel-forming protein complexes. This specific action could be a consequence of crown-ethers ability to evoke conformational rearrangements of the mentioned protein complexes, to stimulate membrane phospholipid peroxidation, to alter lipid microsurroundings of membrane receptors and enzymes. Besides, the investigated crown-ethers may lead to cellular ionic imbalance via their negative action on channel-forming proteins, as well as via the ability of these substances to form complexes with biogenic elements. The ionic imbalance could be one of the reasons of observed alterations in the cyclic nucleotides system of rats toxified by crown-ethers. The alterations, which are developed by influence of crown-ethers, are one of the reasons, and a reflection of metabolic processes impairment inherent to organism cells at conditions of xenobiotics toxic action.

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