Physicochemical and biochemical properties of the Keplerate-type nanocluster polyoxomolybdates as promising components for biomedical use

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The paper discusses the results of a research on physicochemical and biochemical properties of the Keplerate-type molybdenum-based nanocluster polyoxometalates (POMs), which show promise in the field of biomedicine as a means of targeted drug delivery, including the transport to immune privileged organs. POMs can be considered as components of releasing systems, including the long-acting ones with feedback (for controlling the drug active component release rate). POMs are promising drugs for the treatment of anemia. Also, the paper deals with the results of studies of POM effect on living systems at the molecular and cellular levels, at that of individual organs, and on the organism as a whole. The mechanism and kinetics of POM destruction and possibilities of stabilization, the oscillatory phenomena manifestation, the formation of POM conjugates with bioactive substances which can be released during the destruction of POM, with polymer components, and with indicator fluorescent dyes, as well as forecasts for further research, are considered.

Keywords: Keplerate-type nanocluster polyoxomolybdates, physicochemical and biochemical properties, impact on living systems, means of targeted delivery, releasing systems, prolonged action.

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1. Introduction

1.1. Nanoscale components of targeted drug delivery systems

The targeted delivery of drugs can employ nanostructured systems that differ in terms of the nature of components and design, to which, for instance, liposomal systems belong [1–13]. Some natural aspects can impede the use of liposomal agents. These aspects include the active capturing of liposomes by macrophages, which results in the limited possibility of drug delivery to some organs and tissues. The capturing also significantly reduces the time of drugs circulation in the organism, thus leading to the liposome membrane degradation under the influence of lipoproteins. In addition, the manufacturing of liposomal products employs rather sophisticated technologies. Various inorganic nanoscale objects are considered as components of targeted drug delivery systems; these include metal, oxide or complex oxide objects, and some other types of nanoparticles [14–25], as well as carbon clusters (fullerenes, nanotubes, graphene, and nanodiamonds) [26–59]. These objects can be used, for instance, as shells or cores of nanocapsules, providing the release of drugs, for example, under local electromagnetic influence. They can play the
role of carriers of bioactive substances in surface functionalization [44–46, 48, 49, 51–56], or when the porous structure of nanoparticles and clusters is used, including non-magnetic and magnetic systems for delivery, therapy [57–83] and diagnostics [60–68]. The use of carbon clusters and inorganic nanoparticles is associated with the problem of removing such materials from the body after they have performed their function, in order to reduce the xenobiotics load.

Nanocluster polyoxometalates (POMs) can be produced and characterized in terms of composition and structure using fairly simple and proven techniques. Representatives of this class of compounds, in particular molybdenum-containing ones, do not pose a danger to a living organism and are practically non-toxic. They do not accumulate in the body for long, because, for instance, they are capable of gradual decomposition into decay products, which are assimilated as microelements and further evacuated in the course of natural metabolism. For these reasons, POMs are considered as promising materials for targeted delivery of drugs, of long-acting therapeutic agents, including those with feedback for controlling the drug release rate. Also, the advantages of POMs include their solubility in aqueous media and the charge of polyanions, which provides the possibility of iontophoretic delivery. The present review briefly discusses the biochemical properties of the most promising POMs, their ability to associate with biologically active substances, as well as issues related to the processes of POM destruction and stabilization as a key aspects in terms of biomedical application.

1.2. Nanocluster polyoxometalates (POMs)

Variously composed and structured POMs, which include both hetero- and isopoly compounds, have found a wide application as catalytic systems, components of hybrid structures, supramolecular assemblies, etc. [84–87]. New types of molybdenum-containing nanocluster polyoxometalates with toroidal [88–91] and fullerene-like structures (Keplerates) containing subnanometer-sized windows and pores [92, 93] (Fig. 1) became known at the turn of the past and present centuries due to studies carried out at the Bielefeld University (Bielefeld, Germany) under the guidance of Prof. A. Müller, whose laboratory synthesized these compounds and confirmed their structure. In comparison with other nanosized particles, such water-soluble clusters have a high ionic charge and are capable of gradual degradation. The discovery of such POMs was, apparently, one of those cases, which are far from rare in the history of science, when such POMs had been most likely obtained in earlier experiments, but the researchers were unaware of their unique structure, which was later deciphered and purposefully reproduced. The POM structure was primarily characterized by single crystal X-ray structure analysis. The everyday scientific practice employs for this purpose reliable and more rapid spectroscopic methods (e.g., IR, Raman, NMR spectroscopy) [94]. The obtaining of giant POM clusters was also confirmed by mass spectrometry (MALDI method) [95]. Among the merits of Prof. Müller’s group is the creation of fairly simple and reproducible methods for obtaining POM [93] with a specified structure and composition. In addition to the above-mentioned, basket-like molybdenum-containing POMs [96], with a structure derived from Keplerates, as well as giant clusters named by their creators “nano hedgehog”, “blue lemon” [97, 98] were also obtained.

The structure of these POMs is based on such structural blocks (coordination oxygen polyhedra surrounding multicharged metal cations) such as octahedra, pentagonal bipyramids, which are connected to each other by vertices or edges. In particular, Mo$_2$ dimolybdenum bridges, Mo$_6$ and Mo$_{11}$ blocks should be attributed to basic structural elements. The symmetry of the latter (C$_5$ or C$_s$ point group) determines the shape of the resulting nanocluster (Keplerate or toroid, respectively). The structure of a POM is stabilized, as a rule, by the organic or inorganic acid residues and water molecules. POMs contain a significant number of molecules of both constitutional and intrACLuster water. The latter also occupy the internal cavity of Keplerates in the normal state and can exchange with molecules of other substances through the pores. POM synthesis proceeds in the self-assembly mode under conditions of a specified medium acidity and in the presence of a reducing agent, since, for instance, the basic structure of Mo$_{132}$ Keplerate (Fig. 1) includes both hexaylad molybdenum and Mo(V):

$$(\text{NH}_4)_{12}[\text{Mo}_{72}^{4+}\text{Mo}_{180}^{6+}\text{O}_{372}^{2-}(\text{CH}_3\text{COO})_{30}(\text{H}_2\text{O})_{72}]\cdot 300\text{H}_2\text{O} (~10\text{CH}_3\text{COOHNH}_4).$$

There is a possibility to use the basic structure of a POM and substitute ions in it. For instance, the isostructural substitution of Mo(V) with ferric ions leads to the formation of a Mo$_{72}$Fe$_{30}$ Keplerate [99] (Fig. 2):

$$[\text{Mo}_{72}\text{Fe}_{30}\text{O}_{252}(\text{CH}_3\text{COO})_{12}][\text{Mo}_2\text{O}_7(\text{H}_2\text{O})]_{2}^2[\text{H}_2\text{Mo}_2\text{O}_8(\text{H}_2\text{O})](\text{H}_2\text{O})_{91}]~\sim 150\text{H}_2\text{O}.$$

The POMs isolated from the concentrated stock solution are mainly crystalline substances, which may include a fraction of the material in an amorphous state [94]. Upon subsequent dissolution in water, POMs dissociate into Keplerate polyanions and cations (ammonium, sodium and potassium), or do it as a weak acid (Mo$_{72}$Fe$_{30}$). The crystallographic size of a Mo$_{132}$ ion is 2.9 nm (Mo$_{72}$Fe$_{30}$ is 2.5 nm), while the hydrated Mo$_{132}$ ion in a solution has a size of 3.2 nm. The charge of polyanions depends on the acidity and environment, and can reach $-42$ and $-22$ [93, 100] for Mo$_{132}$ and Mo$_{72}$Fe$_{30}$, respectively.

The POMs capable, in principle, of changing the energy state of molecules included in their cavity, are considered promising catalytic materials in reactions of fine organic synthesis, membrane technologies, and sensors [101–113].
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Fig. 1. Structure of Mo$_{132}$ polyoxometalate. The red polyhedron (No. 1) marks the pentagonal MoO$_7$ bipyramid, linked by common edges with five MoO$_6$ octahedra (blue, No. 2) – this is the so-called Mo$_6$ structural unit; Highlighted in green (No. 3) are MoV$_6$O$_6$ polyhedra, which are part of Mo$_2$ dimolybdenum bridges.

Fig. 2. Structure of Mo$_{72}$Fe$_{30}$ polyoxometalate. Iron-coordinated oxygen polyhedra are marked in yellow (No. 1 is MoO$_7$; 2 is MoO$_6$; 3 is FeO$_6$).

At the same time, such compounds are also of interest for the biomedical industry (Scheme 1). On the one hand, they can be interesting as individual drugs, because molybdenum and iron are widely known to be used in vitamin and mineral supplements as microelements involved in a number of enzymatic processes of importance for the organism [114]. In addition, it was shown that simpler molybdenum-containing POMs can be used in chemotherapy [115,116]. On the other hand, the initially predicted possibility of the formation of POM conjugates with bioactive substances, drugs, and metal cations using intra- and extraspHERE interactions, the possibility of their transport (including iontophoretic) in the body, adds relevance to the studies of POM in the field of targeted drug delivery and diagnostics [117–131]. These prospects may relate, inter alia, to immunomodulatory and antiviral agents. Important aspects of iontophoretic transport include the potential ability to deliver drugs to lesion areas and body tissues with low degree of vascularization. The very first experimental results showed that these ideas are promising. The low toxicity of Keplerate POMs for the warm-blooded animals, in particular of the iron-containing compounds, has been established on the basis of a set of data [132–137]. When studying the acute and subacute effects of Mo$_{72}$Fe$_{30}$ on rats, no significant negative changes in internal organs were revealed. The animal organism is able to adapt to the action of the named POM, which is confirmed by the absence of a pronounced autoimmune reaction of lymphocytes to hepatocytes during the intramuscular administration of a Mo$_{72}$Fe$_{30}$ preparation in the form of an aqueous solution. There is no long-term and substantial accumulation of POMs and their components in the body (various organs and tissues), as they are excreted by metabolism. This occurs, inter alia, due to the POM degradation into simpler components [106,137]. The issues of POM toxicity are discussed in more detail below. At the same time, in the case of iontophoretic administration, it is possible to create a temporary Mo$_{72}$Fe$_{30}$ depot localized, for example, in the skin and partly in the subcutaneous fatty tissue, which ensures a prolonged action of POM and its conjugates. In general, the present review is devoted to the
2. Investigation of the effect of Keplerate-type POMs on living systems at the level of the organism, of individual organs, and at the molecular-cellular level

2.1. Study of POMs accumulation characteristics and toxicity

The chosen and studied POMs Mo$_{132}$ and Mo$_{72}$Fe$_{30}$ exhibited both similar and different behavior due to their different composition. While the first of them contains Mo(V) in addition to Mo(VI), the second one includes both molybdenum and iron in the highest oxidation states, which makes it promising in terms of low toxicity (no lethality found in animals). The accumulation of Mo$_{132}$ [136] was studied by injecting it into the gastrocnemius muscle of rats at a concentration of 1.5·10$^{-4}$ mol/L daily for a month. A single dose of POM amounting to 0.15 mg/100 g of the body weight, corresponded to the upper limit of the normal daily intake of molybdenum and to the supposed therapeutic dose. Intact animals served as the control. The absence of the drug and its components accumulation in the liver and bones was established. A sharp decrease in the content of molybdenum in the kidneys was found, the cause of which is, apparently, the excretion of the excessive amount of this element via the urinary tract. Experiments with the Mo$_{72}$Fe$_{30}$ polyoxometalate [132, 133, 137] administered at the same dosage, made it possible to establish for this POM the absence of accumulation of molybdenum in the liver, kidneys, bones, and skin of animals, including its hairy part and subcutaneous fatty tissue, since the average values of molybdenum content in the experimental samples after the termination of the procedures did not differ significantly from those in the intact animals.

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The histological analysis [132, 137] of animal organs [138] and the results of hematologic studies [134] indicated a relatively higher toxicity of Mo$_{1:12}$ as compared to Mo$_{72}$Fe$_{30}$. Signs of hepatocyte damage were observed for Mo$_{1:12}$. However, the characteristic signs of intensified liver regeneration processes were obviously observed. The histologic analysis of kidney sections from the experimental group of animals after administration of Mo$_{72}$Fe$_{30}$ showed that the structural changes in the organ were minimal. It should be stressed that a weaker toxic effect of iron-molybdenum POMs [133, 139] was confirmed by the analysis of such biochemical parameters [140] characterizing the state of various organs, as aspartate aminotransferase (AST), alanine aminotransferase (ALT), alkaline phosphatase, alpha amylase, urea, and the total protein.

An analysis of peripheral blood parameters [138, 140] of animals allowed one to draw the conclusion that there was no anemia [141] in the group of animals that got Mo$_{1:12}$ injections, since no significant difference was found in the number of erythrocytes, concentration of hemoglobin, the content and concentration of hemoglobin in erythrocytes in the experimental and intact rats. The total leukocytes in animals that got Mo$_{1:12}$ injections also did not increase, which indicates the absence of a systemic inflammatory process. The total leukocytes did not significantly increase in the animals within 30 days after Mo$_{72}$Fe$_{30}$ injections, and the leukocyte formula remained almost unchanged, which indicates the absence of inflammation at the organismal level.

An analysis of red blood parameters in rats injected with Mo$_{72}$Fe$_{30}$ yielded a conclusion that there was no anemia – a sign characteristic of many chronic intoxications, since the content of hemoglobin in erythrocytes of experimental rats did not decrease, but increased, on the contrary [108, 139, 142]. Moreover, the experimentally revealed effect of rapid recovery of the hemoglobin level in animals after modeling the post-hemorrhagic anemia [141] is interesting from the point of view of the possible use of POMs in treatment and prevention of iron deficiency anemia, which is confirmed by a patent [142]. The presence of molybdenum in the preparation does not prevent an increase in its effectiveness; besides, molybdenum is effectively removed from the body. Given its lower toxicity, Mo$_{72}$Fe$_{30}$ was subjected to additional studies, in particular, of acute and subacute toxicity [133]. In order to further study the pathways of possible toxic action of POM [133], three methods of Keplerate administration were chosen, that is, the intramuscular, intraperitoneal, and intragastric. The intragastric (1 day) method of administration allowed a significant increase in the POM dose by using not only a POM solution, as in other methods, but also a suspension (concentration of 60 mg/ml). The introduction of the maximum possible dose of POM by the intragastric method was not accompanied by lethality, therefore, the conclusion about low toxicity of the tested compound was confirmed, especially since the amount of POM was 200 times higher than the estimated therapeutic dose of 0.15 mg/100 g.

The blood plasma of rats in all experimental groups [133] showed no increase in the activity of alkaline phosphatase, which is one of the indicators of cholestasis (impaired bile secretion). This fact indicated that the bile-producing function of hepatocytes was preserved after both short and long-term administration of POM. Alkaline phosphatase is also a specific enzyme of osteoblasts, it is involved in mineralization, and the activity of this enzyme is an indicator of changes in the rate of mineral components remodeling in the bone tissue. Molybdates are known of being able to be capable of reducing the activity of phosphatases, probably through the interaction of molybdenum with the cysteine thiol groups and histidine imidazole groups located in the active sites of enzymes [138, 140]. The groups of animals with single intramuscular, intraperitoneal, and intragastric administration of Mo$_{72}$Fe$_{30}$ have shown a relatively small (within 20–25%), but significant decrease in alkaline phosphatase activity [133], possibly due to the inhibitory effect of molybdenum. Just a short-term, 1 day-long, decrease in the activity of alkaline phosphatase eliminates the risk of osteoporosis development after a single administration of Keplerates.

When comparing the average values of other indicators in groups of animals with acute administration of POM (alpha-amylase activity, total protein and creatinine content in blood plasma), no significant differences with those of intact rats were found, which evidences the absence of pronounced changes in the kidneys, liver, and pancreas of animals in these groups [133]. The group with subacute POM administration showed no significant deviations of all the studied biochemical indicators from the norm, which suggested the absence of cytolysis and damage to the liver, kidneys, myocardium, pancreas, and bone tissue after a long (30 days) daily POM administration.

It is important to note that after POM administration by all the studied methods, a significant increase in the content of molybdenum in the blood was recorded [133] after a single POM load. The amount of molybdenum found in the blood 1 day after the administration of Keplerate POMs was proportional to the amount of the injected nanomaterial. A significant decrease in the level of molybdenum in the blood after 1 day compared to the content of the microelement one hour after intramuscular injection, as well as the normalization of the indicator after daily injections of POM within 30 days, was consistent with previously obtained data on the absence of molybdenum accumulation [132, 137, 141]. The absence of molybdenum accumulation [133] in the liver, kidneys, skin, and bone tissue [138] is probably associated with the decomposition and accelerated excretion of the elements constituting POM, which contributes to the normalization of the studied parameters (glucose, aminotransferase, and alkaline phosphatase values) [138] by day 30, a change of which was recorded upon a single injection of POM.
2.2. POMs impact at the cellular-molecular level

Further investigations of Mo$_{72}$Fe$_{30}$ were aimed at studying its effect at the cellular and molecular level, since the use of Mo$_{72}$Fe$_{30}$ as a means of targeted delivery requires knowledge of potential risks and side effects that manifest themselves not only at the macrolevel (in organs and tissues) [138, 140]. For instance, the effect of POM on the content of histone and heat shock proteins in the blood and liver of rats was investigated upon intramuscular injection of this compound into the body [143], since the changes, undetectable by means of assessing traditional indicators, can be identified using other methods. One of the possible mechanisms of the damaging effect of nanoparticles at the molecular level can be their effect on the tertiary structure of proteins. It may be accompanied by immune reactions towards denatured proteins that acquire the properties of antigens alien to the body, by increased proteolysis, apoptosis, necrosis, inflammation, and membrane permeability [144, 145]. Fractons of histone proteins and heat-shock proteins are informative indicators of changes in cells at an early stage of nanoparticles influence before the appearance of the "macrotoxicity" signs. The synthesis of inflammatory factors by lymphocytes depends on the modifications and the amount of histone proteins, the release of which from DNA is essential for the expression of proinflammatory factor genes [146–148]. The disruption of the structure of various body proteins is counteracted by heat shock proteins (Hsp), a class of conservative proteins, whose expression increases with the increasing temperature, the action of toxic substances [150], or under other conditions stressing the cell [149]. Heat shock proteins can act as chaperones, restoring native conformation in other proteins. They can enhance proteolysis, reduce apoptosis, and participate in the transport of other proteins and gene expression [149–152]. The liver is the main organ of detoxication, it is saturated with a large number of specific cells presenting antigens, and it normally deposits iron. Therefore, this organ was chosen as the object of experimental research. The action of most toxicants leads to liver infiltration by lymphocytes coming from the blood. The toxic effect of Mo$_{72}$Fe$_{30}$ in cases of single and repeated administration at the cellular, subcellular and organismal levels was studied as regards the content of histone proteins, heat shock proteins and cytolytic enzymes. The experiment showed no expression of heat shock proteins Hsp70 in the liver cells of intact animals [153], and the expression of Hsp60 was detected on average in about 30±7 cells per unit area. When determining the content of histone proteins in blood lymphocytes [143], during all periods of POM load, no significant changes were found in the fractions of histone proteins, which, according to the literature, regulate the expression of genes of proinflammatory factors [148, 154]. A study of heat shock proteins [143] has detected no Hsp70 in the liver in the groups of both intact and control animals, while Hsp60 were recorded. It indicated the inducible nature of Hsp70 and the constitutive of Hsp60. The normalization of biochemical parameters (characterizing the state of the liver) within 30 days of POM administration [132, 143], gives evidence for the activation of defense systems in the organism. Heat shock proteins restore the native conformation of denatured proteins, which can also segregate proteases, and new proteins are synthesized instead of the segregated ones.

Thus, the single and multiple administration of iron-molybdenum POMs manifested itself at the molecular level only in changes in the content of heat shock proteins [143] involved in the adaptation and maintenance of homeostasis. The studies carried out by a number of authors yielded a suggestion that the absence of an increase in heat shock proteins in response to the action of a stress factor is a violation of adaptation [155, 156]. The action of POM (single and multiple) did not cause a decrease in the amount of histone proteins in blood lymphocytes, which allows an assumption about the absence of autoimmune aggression of these cells against liver tissue. An increase in the amount of Hsp60 and Hsp70 heat shock proteins in rat liver after single and multiple administration of POM can be considered as a way of adaptation, which is confirmed by a study of indicators of cytolytic and synthetic liver function [157, 158].

The content of heat shock proteins in the thymus and spleen was also studied, and it was shown that POM administration is accompanied by an increase in the number of cells containing Hsp60 and Hsp70 heat shock proteins in both organs [108, 132, 159]. The increased expression of heat shock proteins in the thymus [160], despite the presence of a hematothymic barrier, gives reasons to suppose the ability of POM to overcome the histo-hematic barriers of immune privileged organs [161]. Taking into account the results of the ongoing studies [161], it has been shown that POM can also exert a similar effect on other organs with special histo-hematic barriers, in particular, hematocerephalic, hematopophalmic, hemato-testicular, etc. According to the results of histological studies, the effect of POM on immune privileged organs did not lead to a change in their morphology and structure. There are prospects concerning the transport of medicinal substances with the help of POM across the histo-hematic barriers.

In the course of further investigation of the action of POMs at the cellular level, the effect of Mo$_{72}$Fe$_{30}$ and Mo$_{132}$ on normal and transformed fibroblasts was assessed [162]. The effect of polyoxometalates on cell survival in culture and their cytotoxic action have been studied [163]. Human dermal fibroblasts (HDF) are more resistant to Mo$_{72}$Fe$_{30}$. On the contrary, Mo$_{132}$ causes extensive destruction of the membrane organelles of dermal fibroblasts in the first hours after exposure. Thus, it can be concluded that Mo$_{72}$Fe$_{30}$ is not toxic for normal fibroblasts (it causes only weak changes in cells), while Mo$_{132}$ is toxic for both normal and transformed fibroblasts, and its damaging effect
is more pronounced on transformed fibroblasts. A decrease in the proliferative activity of cells (cell multiplication by division) was recorded. An assessment of the nanocluster toxicity using the MTT test demonstrated a change in the mitochondrial activity after the impact of Mo$_{132}$. Probably, this type of nanocluster inhibits cellular respiration and causes disruption of energy metabolism in cells. According to the obtained data, the effect of polyoxometalates on cells is selective. The transformed cells accumulated inorganic elements included in the composition of Mo$_{72}$Fe$_{30}$ polyoxometalate (e.g., molybdenum) more intensively than of Mo$_{132}$. At the same time, iron and molybdenum were absorbed from simpler compounds (iron and molybdenum salts) in smaller amounts, which confirms the selectivity of the action. For instance, 24 hours after the exposure to Mo$_{72}$Fe$_{30}$, the number of transformed cells decreased, and cells with an uncountoured cell membrane (indicated by an arrow in Fig. 3) and an increased number of microvilli appeared. This increases the membrane surface, as was recorded by means of optical, electron, and scanning probe microscopy (a comparison of images obtained by different methods is of interest), and can lead, in turn, to an additional increase in its ability to interact with drugs. The obtained results confirm the prospects for the application of POMs in the field of biomedicine, including the use of Mo$_{132}$ in oncological practice [163].

![Fig. 3. Transformed fibroblasts after exposure to Mo$_{72}$Fe$_{30}$ POM: a – optical microscopy (×1000 magnification); b – atomic force microscopy; c – scanning electron microscopy (×15550 magnification) (equipment of the Ural Center of Shared Use “Modern Nanotechnologies” of the Ural Federal University). The increase in the number of microvilli is indicated by arrows](image-url)

3. Association of POMs with bioactive substances and polymers

Polyoxometalates that have been selected in the course of research (Scheme 1) as having characteristics that make their practical use in biomedicine promising, are studied for the compliance of their properties to requirements to the materials for targeted drug delivery systems (Scheme 2).

The Keplerate-type POMs and alike, being hydrophilic particles, are able to associate (form conjugates) in their external sphere with diverse substances, which include surfactants (SAs) of various nature [164–171], dyes, photosensitizers [109, 110, 172–174], and polymers [174–179], mainly water-soluble ones. POMs functionalization makes it possible to selectively influence their hydrophilic and hydrophobic properties [110], create catalytic and photocatalytic
systems with adjustable characteristics, as well as compositions for targeted delivery in biomedicine, controlled by an electric [179] or magnetic field [180, 181]. The hydrophobized POMs acquire an ability to more actively interact with molecules of nonpolar substances and adsorb them [110]. There appears a possibility of formation of ionic associates based on POM and cations of metals (alkaline earths, rare earths, transition metals, including luminescent labels and magnetic resonance imaging (MRI) tags [19, 121–123]), and of cationic surfactants. Weaker, e.g., hydrogen, bonds also actively participate in the interaction of organic compounds with POMs [182]. The formation of POM associates with water-soluble nonionic polymers [174–179, 183–185], including the biocompatible ones, makes it possible to create protective shells on the POM surface, similar to polyethylene glycol (PEG) layers, or to diblock copolymers on nanocapsules [186–194]. In a number of cases, the energy factor (interaction enthalpy) contributes to the formation of these associates. For instance, the enthalpy of POM and polymer interaction is negative over the entire concentration range of the components in the Mo$_{72}$Fe$_{30}$–polyvinylpyrrolidone (PVP) system [177]. Similar local concentration regions were found for the systems containing polyethylene glycol (PEG) and polyvinyl alcohol (PVA) [183]. In some of the studied systems, the formation of associates is facilitated by the entropy factor [183].

As is noted above, the use of ionic associates of POMs with rare earths as components of diagnostic systems in biomedicine has been proposed [19, 121–123, 195, 196]. In such associates, POM recharging occurs, so the resulting complex can be positively charged [197], unlike the initial POM, and be transferred in an electric field by electrophoresis to the cathode instead of the anode. The structure of a POM is retained in associates [197], which was shown by IR and Raman spectroscopy. A number of drugs, including antibiotics, form bioactive anions in solutions, the association of which directly with POM is prevented by the Coulomb repulsion. The competitive interaction of drug cations with the existing complexing anions also complicates the formation of the cation-POM complexes [197]. On the other hand, there is a way to obtain transportable associates of anionic bioactive substances with POM, for example, through the bridging calcium cations [119, 198]. The discovered POM stabilization in associates with calcium will be discussed below. It is important to note the ability of the Keplerate and compositionally similar toroidal [88–91, 110] POMs to self-associate in solutions, i.e., to self-assembly into hollow single-walled globules (Fig. 4) [91, 199], the diameter of which can vary from tens to hundreds of nanometers, depending on concentration, acidity, and dielectric permittivity of the medium [199]. These formations can also be considered as transport units for the targeted delivery of medicinal substances, and the formation of associates, e.g., with surfactants, makes it possible to control the properties of POMs not only at the level of single molecules, but also at the level of globules and aggregates [111, 180, 181].

In the course of investigations, the ratio of components in the formed POM-polymer associates (Fig. 5) was established, which in some cases, is close to the calculated value of the monomolecular coating of POM with polymer chains [175, 178, 184]. Such systems include, for example, associates based on polyvinyl alcohol, PVP with Mo$_{112}$, and on PVP with Mo$_{72}$Fe$_{30}$. As mentioned above, spontaneous energetically favorable formation of associates occurs in some systems, e.g., of PVP with Mo$_{72}$Fe$_{30}$ (in the entire concentration range of components), as well as in certain concentration ranges - of polyvinyl alcohol and PEG with Mo$_{72}$Fe$_{30}$ [177, 183]. The interaction of POM with polymer macromolecules has specific features, and one of the main ones is the nonlinearity of the associates physicochemical properties dependence on concentration [175, 176, 200, 201]. The oscillating concentration dependencies (Fig. 6–8) are
associated with the non-statistical nature of the mutual distribution of the interacting nanoscale components [200,201]. They are observed for such independent characteristics as optical density, solution viscosity, electrical properties, particle size, etc. Obviously, this fact should be taken into account when choosing the ratio of components in associates used for the targeted drug delivery.

It has been established that Mo$_{72}$Fe$_{30}$ POM forms associates with the drugs that are most promising in terms of targeted delivery. Thiamine chloride (vitamin B1) was considered as a model substance [161, 197]. Thiamine is present in a solution in the form of cations, which get associated with the POM polyanion. It was confirmed by UV/Vis spectroscopy, by measuring the pH of solutions, zeta potential, and the size of the particles in them. In concentrated solutions, colloidal particles get formed along with the solid phase of the resulting conjugates, in which the preservation of the original POM structure and the presence of thiamine are confirmed. To this end, IR and Raman spectroscopy were used. The thiamine-POM ratio in the associates was estimated as 4:1, with the retention of the total negative charge of the conjugate. Mo$_{72}$Fe$_{30}$ associates with substances belonging to the class of immunomodulators, e.g., with 2-morpholino-5-phenyl-6H-1,3,4-thiadiazine bromide [197–202].

Mo$_{72}$Fe$_{30}$ forms conjugates with such an antibiotic as kanamycin sulfate, without reducing the activity of this drug when affecting colibacilli cultures [197]. Also, it was possible to establish that Mo$_{72}$Fe$_{30}$ can associate with insulin in aqueous solutions [197], as well as with amphiphilic protein molecules, such as albumins [161, 197]. This result is of great importance in terms of the possibility of obtaining complexes with POM and with homing peptides for the implementation of bioactive substances targeted delivery to specific cells. Protein molecules, the so-called protein
Fig. 6. Dependence of the solutions dynamic viscosity (η) [201] on the ratio of the polyvinylpyrrolidone macromolecular units number per 1 ion of Mo$_{132}$ polyoxometalate (1); 2 – viscosity of the specified POM aqueous solution; the dotted line indicates the viscosity of a 1% pure polymer aqueous solution. ©Pleiades Publishing, Ltd. 2015

Fig. 7. Optical density (A) (λ = 450 nm) [201] of Mo$_{132}$ POM and polymer solutions, depending on the molar ratios of the components: a – of polyvinylpyrrolidone ($C_{POM} = 1.5 \times 10^{-6}$ mol/L); b – of polyvinyl alcohol ($C_{POM} = 1.5 \times 10^{-7}$ mol/L). ©Pleiades Publishing, Ltd. 2015

Fig. 8. Concentration dependencies of the dielectric loss factor [175] at the 25 Hz frequency for the PVA-based composites, including (1) Mo$_{132}$, (2) Mo$_{138}$, and (3) Fe$_7$Fe$_{30}$. C is the POM content (wt %). ©Pleiades Publishing, Ltd. 2016
corona, significantly influence the biological properties of nanoparticles [203–207]. It should be noted that the blood serum proteins stabilize Mo$_{72}$Fe$_{30}$ [161,177], in spite of the fact that the serum is an alkaline medium, where the POM is unstable, in principle, which is illustrated by its rapid decomposition in buffer solutions. Stabilization by the natural protein corona opens up prospects for the prolonged action of the POM-based drugs. While in buffer solutions with the corresponding pH 8 Mo$_{72}$Fe$_{30}$ decomposes rapidly, and the effective constant of the POM destruction rate has a value of $2.6 \times 10^{-2}$ h$^{-1}$ even for the self-aggregated ones, its value is only $1.8 \times 10^{-3}$ h$^{-1}$ in the composition of conjugates with proteins [96]. The initial experiments aimed at establishing the possibility of obtaining conjugates of Mo$_{72}$Fe$_{30}$ with doxorubicine yielded positive results. Doxorubicine is an anthracycline antibiotic [208–212] that is actively used in medicine in the treatment of various types of solid tumors. Two main mechanisms of doxorubicine’s antitumor activity are proposed in the literature, namely DNA intercalation and disruption of the viral DNA repair, and the generation of free radicals that damage the cell membrane, DNA, and proteins [213]. Along with high efficiency, doxorubicine has high toxicity associated with low selectivity for its action. In addition to high cardiotoxicity, a decrease in bone marrow activity (myelosuppression) is often observed in the course of doxorubicine administration [214]. To date, a significant amount of research has been conducted with the purpose of reducing the toxic effect of doxorubicine on healthy cells by creating means of its targeted delivery. Various types of different systems for the targeted delivery have been considered, in particular, based on liposomes [215–222] (similar forms have already found application in medical practice [223–227]); proteins [228–233]; polymers [234–243]; fullerenes and CNT [35,244–251]; metal and oxide nanoparticles [52,76] (gold [250–255], silver [255], and iron oxide [256–260]). The further study of the properties of POM-doxorubicine conjugates should be recognized as promising, given the corresponding properties of POM.

An important aspect of the possible use of POM in the field of biomedicine is the possibility of iontophoretic, including percutaneous, transport of Keplers and their conjugates into the body. These processes were considered in our previous work [179]. As is noted above, these processes are important, in particular, for the delivery of drugs, e.g., of doxorubicine, to the regions with a low degree of vascularization.

Additional opportunities for the use of Mo$_{72}$Fe$_{30}$ in biomedicine are opened by the preparation of its new coordination compound [261] that can serve as a transportable donor of nitric oxide – a mediator of some important processes in the body [262–267], which include the activation of antiviral and antimicrobial activity, and the cardiotropic action. It is important that the compound [POM-(NO$_2$)$_x$]$_y$Fe$_{30}$ (where $x = 6$, $y = 14 \pm 3$), obtained by the interaction of nitrogen oxides with POM and characterized by a complex of methods, belongs to the group of water-soluble compounds. Iron compounds are used as nitric oxide donor and transport vehicles for its delivery [268–275]. Examples of obtaining other POMs containing nitric oxide can be found in the literature [276–278].

4. POMs destruction process and their stabilization

One of the main above-mentioned properties of POMs is their ability to degrade in diluted solutions [279], which depends on both the POM initial concentration and the pH of the medium. This property can be useful for POM removal from the body after the function of bioactive substances delivery has been performed. The results of studies of thermal stability and destruction of the Keplerate and toroidal type POMs in the solid state using spectroscopic methods are given in [280–286]. At the same time, it was shown that a thorough spectroscopic analysis and a comprehensive interpretation of the results are necessary [280,286] in order to avoid unreliable conclusions [284] about the POMs’ thermal stability. The features of the processes of POMs destruction in solutions have been considered in the earlier performed works [135,198]. For each of the considered POM, there is a region of stable state in aqueous solutions in terms of POM concentration and medium pH. For Mo$_{132}$, this region can be clearly illustrated by a diagram (Fig. 9). The region of Mo$_{72}$Fe$_{30}$ stability in sufficiently concentrated solutions is also limited by pH within the 3.7–5.9 range [100,106]. Under the conditions that exist beyond the range of POM stability, they undergo gradual destruction into simpler components.

UVVis spectroscopy was used for studying the kinetics of POM degradation in solution. The time dependencies ($\tau$) of the optical density $A$ (directly proportional to the current POM concentration) obtained [135] for different initial concentrations of Mo$_{72}$Fe$_{30}$ POM were approximated by equations describing reactions of different orders, in order to establish which of them the experimental data obey. A satisfactory agreement with the linear dependence was observed in the ln($\frac{A_0}{A_\tau}$) = $f(\tau)$ coordinates for Mo$_{72}$Fe$_{30}$ solutions in the domain of relatively low concentrations. The experimental data correlated best with the first order reaction equation. For relatively high Mo$_{72}$Fe$_{30}$ concentrations, the experimental data were approximated by two lines with different slopes (for different time periods). Each of them was linearized in the ln($\frac{A_0}{A_\tau}$) = $f(\tau)$ coordinates. In this case, the constant of POM decomposition rate in the
first time interval coincided with the constant of Mo$_{72}$Fe$_{30}$ destruction rate in low concentration solutions. Such a situation can be explained by a change in the mechanism of the destruction process depending on the concentration and a change in the limiting stage of the general process where inflection between two lines was observed. An example of the experimental data processing is shown in Fig. 10. For comparison, the table also shows the data on the destruction kinetics for a toroid-structured Mo$_{138}$ POM, for which the one-step destruction process in a solution with a relatively high ($2.4 \cdot 10^{-6}$ mol/L) initial POM concentration was satisfactorily described by the first-order reaction equation.

In [135], a supposition was made concerning the possible differences in the process of Mo$_{72}$Fe$_{30}$ POM destruction at different concentrations. The mentioned POMs tend to form aggregates or hollow globules in solutions [89,199,286–292]. In solutions with high concentrations, aggregation proceeds in such a way that a significant part of the available POM molecules (ions) happens to be bound into aggregates within a relatively short time. In diluted solutions, the relative amount of free POM molecules is more noticeable. Therefore, it can be assumed that the free POM destruction rate constant is higher than that of those bound into globules, in which they are protected due to the presence of bonds between molecules (ions) and a lesser contact with the environment. When the experimental dependencies demonstrate

![Hypothetical diagram of Mo$_{132}$ POM stability](image)
TABLE 1. Polyoxometalates (k_i) destruction rate constants at the naturally formed acidity of solutions, h^{-1} [135]

<table>
<thead>
<tr>
<th>Solution concentration, mol/L</th>
<th>Polyoxometalate composition</th>
</tr>
</thead>
<tbody>
<tr>
<td>1·10^{-7}–6·10^{-7}</td>
<td>Mo_{72}Fe_{30} , k_1 = 0.02</td>
</tr>
<tr>
<td>1·10^{-6}–1·10^{-5}</td>
<td>Mo_{138} , k_1 = 0.02, k = 0.05</td>
</tr>
<tr>
<td></td>
<td>k_2 = 0.002</td>
</tr>
</tbody>
</table>

an inflection within the same initial concentration, a change in the limiting stage occurs, apparently, after the majority of the available free POM molecules have been spent.

The proposed hypothesis is supported by the fact that the constant of the aggregates formation rate [289] (calculated also for the first-order reaction) is lower than the POM decomposition rate constant, and equals 5.4–7·10^{-5} h^{-1} for Mo_{72}Fe_{30}, in particular. Therefore, the initially obtained solution should contain at the first stage a sufficiently large number of unbound POM molecules, which decompose faster than the aggregated ones. A relatively faster process of decomposition of free Mo_{72}Fe_{30} molecules, which are initially present in a significant amount, can also explain a steeper initial section in the kinetic dependencies of aggregate formation [289]. The rate constant in this section can be estimated as 2.5–3.9·10^{-4} h^{-1}. In this case, it can be assumed that the appearance of iron ions due to the POM destruction leads to acceleration of the aggregation process. The literature indicates a similar effect of cations on the formation of aggregates [291]. At the same time, the existence of singlet POMs in a solution is not typical for the toroid-like Mo_{138} [286]; therefore, its destruction is a one-step process in a certain time interval.

In [289], the ratio between the number of aggregates and singlet nanoclusters was determined, from which the degree of aggregation was calculated. The authors apparently did not take into account a possibility of this ratio increase due to a higher rate of Mo_{72}Fe_{30} POM free ions destruction, as well as the change of the total concentration of POM, which could lead to an overestimation of the globule formation rate constant.

Another question that arises is about the mechanism of POM decomposition s included in the aggregates. Do the POM molecules get destroyed directly in the globules, and then the latter disintegrate, or does the preliminary disintegration of the globules occur when the concentration of free POM ions decreases? The destruction of POM in globules could have possibly led to a change in their size; however, according to the data from [289], the size of the existing globules remains practically unchanged. Therefore, it can be assumed that the destruction of POM molecules (ions) into simpler compounds occurs after the preliminary destruction of these globules, which goes along with the decreasing concentration of the studied POM in the course of its decomposition in the solution. Some previously found differences in the kinetics of Mo_{138} POM and Mo_{72}Fe_{30} destruction can be associated with both the globules destruction rate, and with the stability of the polyoxometalates themselves. Thus, it was shown in [135] that, depending on the concentration of Mo_{72}Fe_{30} POM solutions, the disintegration of single nanoclusters, or the “dissolution” of the existing globules consisting of nanoclusters, can be the limiting stage of the overall destruction process.

The destruction of Mo_{138} has also been investigated [198]. An analysis of the spectrophotometric data from experiments on the destruction of Mo_{138} POM showed that the time dependencies of the optical density and, accordingly, of the POM concentration, have an induction period when the concentration decreases rather slowly. Then the decomposition accelerates, and after passing the interval with the maximum rate, it noticeably slows down again. For this reason, the standard linearization of the data using kinetic reaction equations of various orders (including those at the stages after the induction period) was a failure. Considering the shape of the obtained dependences, it was logical to assume that the destruction proceeds in the autocatalytic reaction mode. A typical picture of the kinetic dependence is shown in Fig. 11a.

The experimental kinetic data obtained at the initial concentration of the studied POM of 10^{-5} mol/L, were processed using an equation that describes the autocatalytic process [293–295]:

\[ C = C_0 (K_0 + K_C C_0)/(K_C C_0 + K_0 \exp((K_0 + K_C C_0) t)) \]

where \( C_0 \) is the nanocluster initial concentration, \( K_0 \) is the non-catalytic process rate constant, \( K_C \) is the catalytic process rate constant, \( C \) is the nanocluster current concentration, and \( t \) is the time. The values of \( K_0 \) were estimated from the slope of the dependence during the induction period, and it was assumed that the catalytic process rate is negligible during this period (of the non-catalytic reaction), which is a standard assumption in such cases. The experimental data for the interval, corresponding to the rapid course of the catalytic process with further deceleration
due to the consumption of autocatalytic particles, were used for determining $K_c$ values for each experimental point, and then plotting the time dependence of this parameter.

It is impossible to deduce the $K_c$ value from the above equation by using the analytical method, therefore a special algorithm was developed for determining this value. It consists in the following:

1. Since the equation is transcendental, it was rewritten as:
   \[ C(K_cC_0+K_0 \exp((K_0+K_cC_0)t)) - C_0(K_0+K_cC_0) = 0 \] or \( F(C,t,K_c)=0 \).

2. The available data array $C=C(t)$ was used to construct the function \( F=F(C,t,K_c) \), according to which, the time-dependent $K_c$ values were reconstructed.

The performed calculations showed that the value of the catalytic process rate constant practically did not change in the corresponding interval in time and averaged about $4.8 \times 10^2$ L/mol/min at the initial POM concentration of $10^{-5}$ mol/L, and $6 \times 10^2$ L/mol/min at $5 \times 10^{-6}$ mol/L, respectively. The half-transformation time estimated on the basis of such rate constant values, correlates well with the experimental data (about 4-5 hours). Thus, it can be considered that the destruction process proceeds as an autocatalytic reaction, and the used assumptions are correct, since the shape of kinetic dependence could be approximated successfully by a corresponding equation. The possible reasons for the differences in the obtained $K_c$ values at different initial concentrations will be discussed below. The value of $K_0$ in this case was about $1.8 \times 10^{-3}$ min$^{-1}$, regardless of the initial POM concentration in the range of 1-10 µM.

The features of Mo$_{132}$ destruction in the presence of calcium ions, associated with the POM polyanion, are also considered. The hydrated calcium ions are capable of forming differently configured associates with POMs [119]. Calcium can stabilize POM in solutions, thus increasing the effective value of its lifetime in solutions. In this case, as in that of POM association with polymers, the non-additive properties are observed, for instance, the nonlinear concentration-dependent ability of calcium to stabilize POM. In the processes of calcium cations and POM interaction, apparently, 20 pores in the Keplerate structure are involved as active centers, i.e., the places where the association of the hydrated ions of divalent metals occurs [296]. It cannot be excluded that the filling of these positions leads to the associate’s stabilization. In addition, the next level of interactions is realized along with the increase in the associated ions concentration. Naturally, ion associates are found in dynamic equilibrium with free ions in a solution, and a certain excess of associated ions is required to ensure the filling of active positions. The change in time of the solutions optical density at the POM characteristic wavelength indicated that the almost completely destroyed pure Mo$_{132}$ persisted for several days in the presence of calcium, especially at the Ca-POM ratios of 8:1, 24:1, and 60:1. The presence of POM with the original structure was confirmed by Raman spectroscopy.

This is another case when the POM destruction process is described as an autocatalytic reaction. Such parameters as the induction period duration, and the rate constants in this interval and during the catalytic process were determined for the destruction reaction. When the Ca content increases, the induction period lengthens, and the rate constant of the noncatalytic stage decreases (Fig. 12), but nonmonotonically depends on the Ca content in solutions, and has minima for some compositions. The autocatalytic process rate constant increased along with an increase in the Ca content. It
Physicochemical and biochemical properties of the Keplerate-type nanocluster...

It was noted during the research that the pH of freshly prepared POM solutions decreased depending on the amount of Ca (Fig. 13) at POM concentrations of 1 and 10 µM, and the nature of the kinetic parameters dependence (in absolute value) on the Ca concentration was generally similar to the pH concentration curve, although the pure Mo\textsubscript{132} solution did not fit this series. On the one hand, this shows that the destruction mechanism and rate are determined not only by the solution pH. Along with that, it can be stated that the maximum stability of some compositions is determined by a combination of a sufficiently long induction period and a relatively low non-catalytic stage rate constant.

When considering the pure Mo\textsubscript{132} destruction process again, a monotonic decrease in the pH of solutions was observed at a concentration of 10 µM. Notable is that this decrease was more rapid during the induction period. From the point of view of the direction of change in acidity, this regularity is quite logical taking into account the possible simultaneous process of the available pentavalent molybdenum oxidation with atmospheric oxygen, which can be conventionally described as follows:

\[7\text{Mo}^{VI} + \text{Mo}^{V} + 10\text{O}_2 + 354\text{H}_2\text{O} \rightarrow 132\text{Mo}^{VI} + 708\text{H}^+ + \text{H}_2\text{O}^+\]

Due to the presence of the above-described change in acidity, the hydroxonium ions H\textsubscript{3}O\textsuperscript{+} were considered as autocatalytic particles, which is often the case. However, it was taken into account that along with the decreasing pH of calcium-containing solutions and increasing amount of Ca\textsuperscript{2+} in them, the rate constant of the non-catalytic stage of the process decreases, and its duration increases. Therefore, non-condensed molybdate (or dimolybdate) ions that appear during the destruction of Mo\textsubscript{132}, can act as the autocatalytic particles. Further on, in addition, the diagram from [184] (Fig. 9) was taken into account, as it showed the regions of stability of various molybdate forms (including Mo\textsubscript{132} ions) depending on the total molybdenum in solutions, and on the pH of the medium. The point corresponding to a POM solution with initial concentration of 10 µM in a more detailed fragment of the diagram (Fig. 14), is found in the region of heptamolybdate ions stable existence. It should be noted that the earlier studies of POM decomposition by photon correlation spectroscopy [184] discovered the appearance of POM particles with an effective size of about 1.3 nm, which is close to the estimated size of the heptamolybdate ion. If, in addition, fragments containing less than 7 molybdenum atoms also appear during the POM destruction, then they must be completed up to a heptamolybdate...
form that is stable in this region. Then, it cannot be ruled out that these particles “attack” a POM to obtain the required material. An accelerated destruction in this case is achieved when weakly condensed compounds or monomolybdate forms get accumulated in the solution. The slowing down of the destruction process corresponds to the consumption of the mentioned autocatalytic particles. At the initial $\text{Mo}_{132}$ concentration of 1 $\mu$M, the resulting monomolybdate particles do not need to build up to heptamolybdate.

**Fig. 13.** Dependencies of the $\text{Mo}_{132}$ solutions pH on the calcium-POM ratio (a) at the concentration of $10^{-6}$ mol/L, as well as on the experimental time at concentrations of $10^{-5}$ (b) and $10^{-6}$ mol/L (c) [198]. ©Pleiades Publishing, Ltd. 2016

**Fig. 14.** Fragment of the phase diagram of forms of ions in a solution: A - heptamolybdate ions stability region; B - monomolybdate existence region. The arrows indicate the direction of change in the acidity of the solution during the $\text{Mo}_{132}$ destruction [198]. ©Pleiades Publishing, Ltd. 2016
The conclusions made above about the nature of autocatalytic particles are consistent with the data on increase in POMs stability in acidic buffer solutions [279]. These data indirectly indicate that hydronium ions do not catalyze destruction. The catalytic action of monomolybdate ions can be confirmed by a direct experiment. The addition of sodium molybdate into the Mo\textsubscript{132} solution (with the initial concentration of 5 \( \mu \text{M} \)) practically did not affect the pH of the solution, and led to the exclusion of the induction stage from the destruction process (Fig. 11b). In this case, the molar content of molybdenum in the form of monomolybdate was set equal to the molar content of molybdenum contained in the form of Mo\textsubscript{132}. Changes in the solutions acidity during the destruction of more diluted (e.g., \( \mu \text{M} \)) Mo\textsubscript{132} solutions without additives did not have a monotonic character (Fig. 11). It is also related to the thermodynamic factor (diagram in Fig. 14). The point corresponding to the indicated concentration is found in another region of the diagram, where simpler ions are stable. However, the pH decreases in the course of destruction, and the composition transits across the boundary between two regions (Fig. 14). At the initial Mo\textsubscript{132} concentration of \( \mu \text{M} \), the appearing monomolybdate particles do not need to build up to heptamolybdate before crossing the boundary of the indicated regions. Such a flow of processes is even more complicated than in the first case, so it becomes difficult to unambiguously interpret the kinetics and describe it by the used autocatalytic equation. The course of reaction can be additionally complicated by the fact that products of destruction, including pentavalent molybdenum, have different degree of stability in different regions of phase diagram. They have a higher stability [200] in the condensed forms in region A (Fig. 14). As for the Keplerate structures destruction, it was noted that the Keplerate decomposition stage (with the solution turning blue) [104] can be followed by the Mo(V) compounds oxidation, accompanied by discoloration. The change in the medium acidity during POM destruction in the presence of Ca\textsuperscript{2+} is also not monotonic in all cases. However, the above-noted general trend of an increase in the induction period duration along with an increase in the amount of Ca\textsuperscript{2+} in the solution, may be associated with the fact that calcium ions are able to form sufficiently stable associates with molybdate, which leads to at least partial binding of autocatalytic particles. However, calcium molybdate (CaMoO\textsubscript{4}) solubility product was not obtained at the studied concentrations of solutions, and no precipitation in noticeable amounts was observed. To summarize, let us note once again that POM destruction is a process, the rate of which is determined by both POM concentration and medium acidity. The destruction mechanism can vary depending on these conditions. There are possibilities for temporal stabilization of POMs and a decrease in the value of the process rate constant – by regulating either the acidity of the medium and POM self-aggregation, or POM interaction with stabilizing agents, including those of natural origin.

The effect of light on POM solutions increases the rate of their destruction [297, 298] and can change the mechanism of this process (to be discussed in more detail in the next section). Mutual stabilization of POMs and water-soluble nonionic polymers (including biocompatible ones) during the formation of their associates, which manifested itself under sufficiently severe photothermal effect on systems, was studied in [198, 297–299]. This aspect is important not only in itself, but also when using POM as, e.g., part of ointment compositions, or sterilized materials. The stabilization effect was studied by means of the EPR spectroscopy for determining the number of paramagnetic centers (spins) as an indicator of the composition components destruction processes. Quite interesting are the results of studies, which indicate the above-noted significant stabilization of Mo\textsubscript{72}Fe\textsubscript{30} POM due to the interaction with proteins of the blood serum [161, 300], which has an alkaline pH. In buffer solutions with a similar pH, POM destruction proceeded at a very high rate, while in the serum they persisted for up to dozens of hours. The destruction rate constant of POM in the resulting conjugates with serum proteins was at least an order of magnitude lower than in aqueous solutions of the POM itself. Thus, there exists a possibility to regulate POMs stability, their lifetime, and the period of the conjugated drugs release. Preliminary stabilization of POM in solutions can be carried out outside the body.

5. Prospects for the creation of POM-based releasing systems with feedback

An integrated approach to the study of the physicochemical properties of the Mo\textsubscript{132} and Mo\textsubscript{72}Fe\textsubscript{30} Keplerate POMs [138, 161, 301], as well as the possibility of their association with bioactive molecules and fluorescent dyes, made it possible to formulate the concepts [302–304] of creating long-time-drug-releasing systems (LDRS) with a function LDRS will be based on the release of molecules of bioactive (medicinal) substances, initially bound with POM, and the intensification of fluorescent labels luminescence. The release occurs at the normal pH (7.2-7.4) of the body environment, due to the POM destruction. The main principle LDRS are based on is the discovered effect [302–304] of a fluorescent label (xanthene dye rhodamine-B, RhB) inactivation on the POM surface due to charge transfer processes. This makes it possible to monitor the drugs release due to the intensification of the signal from the fluorescent labels [305, 306] incorporated in the applied LDRS. In [307], it was found that RhB, chosen from a number of xanthene dyes, is characterized by the concentration quenching of luminescence due to a static mechanism, i.e., formation of non-luminescent H-aggregates. It was shown in [304] that the reason for the RhB fluorescence quenching in an associate with POM is the photoinduced charge transfer from the level of the lowest unoccupied molecular orbital (LUMO) of the donor (RhB) to the LUMO level of the acceptor (POM). According to the cyclic
voltammetry data [308,309], the LUMO level of RhB is 0.65 eV higher than the LUMO level of POM, which leads to the spontaneous electron transfer from RhB to POM in an excited state and formation of the RhB+/Mo\(_{132}\) ionic pair.

It should be noted that the transfer of electrons between the HOMO/LUMO levels of dye molecules and POM (Fig. 15) also underlies the promising use of such photosensitized systems in catalysis [310]. The experiments performed to date have yielded such components of the above systems (Fig. 16) as the NHS ester of rhodamine-B - a potential fluorescent label [302, 303], which can be covalently fixed on the macromolecular backbone of biocompatible polymers (chitosan, polyallylamine) by an amide bond. It was shown that the Mo\(_{132}\) nanocluster, which had been chosen as a model POM, can be connected with matrix macromolecules of biocompatible polymers, such as polyallylamines, aminopropyltrimethoxysilane, and chitosan, resulting in microgels based on similar matrices. In principle, the proposed LDRS can be used as a subcutaneous implant with the function of on-line pharmacokinetic monitoring.

Such nanoscale systems can, in principle, be created on the basis of hybrid supramolecular assemblies that act autonomously, like liposomes with highly specific ligands on their surface [311]. Such systems can also be components of a microscopic implant [312] or of a biocompatible polymer matrix capable of carrying a large number of bioactive molecules (BAM) for the long-time release [313, 314]. As regards the future, the further steps under consideration relate to both realization of the target properties of the proposed LDRS systems [302], and to optimization of their chemical design, and \textit{in vitro} and \textit{in vivo} study of their biological properties. Such a task can be solved, for instance, by studying a possibility of pairwise assembly of components of a composition that includes a polymer, a POM, a xanthene dye (rhodamine) covalently bound in the form of NHS-ester, and a drug.

\textbf{FIG. 15.} Photochemical processes in a POM-dye system on the example of a toroidal compound Mo\(_{138}\)

\textbf{FIG. 16.} Diagram of a promising POM-based releasing system. BAM - bioactive molecules; GLUA - glutaric dialdehyde

The oscillatory nature of the dependence of the physicochemical characteristics of POM-based associates (Section 3) and the possibility of influencing the compositions by ultraviolet radiation have been pointed out above. It
is important to understand the light-influenced behavior of nanostructured objects when developing photoinduced controlled-release systems (Section 1.1) and feedback systems (Section 5). It is impossible to miss the fact that in some cases, oscillatory phenomena are observed during the impact on the systems that contain gradually destructing POMs, [297, 303]. In this case, fluctuations depend not only on the composition; temporal oscillations of fundamental physical and chemical properties, such as optical density, photovoltaic potential, etc. also occur. Oscillations of physical properties are most often characteristic of the systems containing macromolecular components, e.g., polymers or POMs [175, 201, 315]. For POMs, the oscillatory phenomena prerequisites include the large size of polyanions, as well as the nonlinearity of characteristics, including the optical properties [316] of molybdenum-containing complexes (associates) with organic substances, e.g., with citric acid [317]. Due to the reversibility of redox reactions of d-metal ions (electron transfer in associates [297]), POMs are able to participate in oscillatory processes. The oscillatory phenomena mechanism can have significant differences in the differently composed systems and in varying conditions in which these phenomena are observed. Each case requires a detailed consideration, which we did below for the above-described systems containing Mo$_{132}$, rhodamine-B, and also showing signs of properties oscillation.

The experiments aimed at studying Mo$_{132}$ destruction in aqueous solutions with a concentration of 5.4 $\mu$M and in contact with argon showed that this process is stimulated by the influence of light [303]. For irradiation, two LEDs were used together, the first of which emitted light mainly at 630 and 570 nm wavelengths, and the second at 466 nm. This combination made it possible to reproduce the main spectral components of daylight. Before irradiation of the solutions, they were preliminarily purged with argon. In the course of the experiments (Fig. 17), there was a general decrease in the optical density ($A$) at 455 nm wavelength, associated with POM destruction, and the measured value $A$ was found to fluctuate exceeding the measurement error. The previously performed experiments [318] demonstrated similar oscillations related to the potential difference $U$ between the frontally UV-irradiated Mo$_{132}$ solution, which also contained polyvinyl alcohol (PVA), and a likewise solution, though light-protected (Fig. 18). The potential difference had a negative value and increased with time. It occurred due to a decrease in the Mo(V) concentration in the irradiated solution during the photochemical destruction of POM, and the oxidation to Mo(VI) during Mo$_{132}$ destruction despite the presence of PVA. On the contrary, when other POMs were present in solutions instead of Mo$_{132}$, partial reduction of molybdenum ions occurred during the interaction with PVA and was accompanied by an increase in positive values of the potential difference. The presence of a photoinduced potential difference accompanying the photochromic effect manifestation in molybdenum-containing systems, as well as signs of nonlinear and oscillatory processes, are also described in [317, 319]. Oscillatory processes could occur in this case due to the partial reversibility of photochemical processes.

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![Graph](image.png)

**Fig. 17.** Relative optical density of Mo$_{132}$ solutions as a function of the irradiation time (min) in contact with argon (1) and air (2). The initial optical density is designated as 1.

It should be noted that, in contrast to the systems containing, e.g., ammonium heptamolybdate, where redox reactions are reversible [297] and the occurrence of fluctuations in the potential difference is generally understandable, Mo$_{132}$ destruction proceeds irreversibly [198]. The products of destruction do not absorb light intensely at the 455 nm wavelength characteristic of Mo$_{132}$. In this relation, there arises a question about the causes of the fluctuating optical density of Mo$_{132}$ solutions and of the concentration-induced potential difference. If the value of $U$ could fluctuate due to a reversible change in the content of Mo(V), including its non-Keplerate forms, upon their interaction with PVA, then why did the optical density fluctuate? Note that the LEDs irradiated the studied solutions from above [303], i.e.,
from where the gaseous medium was in contact with the solutions. In this case, the harder part of the LEDs radiation at 466 nm was absorbed by the solution itself, the characteristic wavelength of which is in the near region. Therefore, it can be assumed that the photoinitiated reaction of Mo$_{132}$ destruction, accompanied by the oxidation of molybdenum to Mo(VI), was localized precisely at the solution surface in a relatively thin layer, the thickness of which did not exceed 0.5 mm [320]. In this case, the described above potential difference appeared between the liquid-gas interface and bulk solution. So, the solution surface can be considered formally as a positive electrode (cathode). The reaction of POM oxidative destruction, which occurs due to the interaction with residual oxygen in argon (no more than 1 vol.%), can be written for this case, in principle, as the sum of two half-reactions, e.g., the following ones:

$$7\text{Mo}^{VI}_{12}/\text{Mo}^{V}_{60}\text{O}_{372}^{12} + 564\text{H}_2\text{O} - 252e \Rightarrow 132\text{Mo}^{VI}_{7}/\text{O}_{6}^{6-} + 1128\text{H}^{+}$$

$$\text{O}_2 + 4\text{H}^{+} + 4e \Rightarrow 2\text{H}_2\text{O}.$$  

In total: $$7\text{Mo}^{VI}_{12}/\text{Mo}^{V}_{60}\text{O}_{372}^{12} + 438\text{H}_2\text{O} + 63\text{O}_2 = 132\text{Mo}^{VI}_{7}/\text{O}_{6}^{6-} + 876\text{H}^{+}.$$  

The oxidation-reduction potentials of the components of the considered system make the above reaction feasible. At least for the second half-reaction, the standard potential is +1.229 V.

During Mo$_{132}$ decomposition in accordance with the proposed reaction scheme, the acidity of the medium is supposed to increase, which was confirmed experimentally [198]. The potential difference between the solution surface layer and its lower part should accelerate the migration of protons (hydronium ions H$_3$O$^+$) deeper into the solution and the supply of anions to the cathode. Taking into account the circumstance that the spectrophotometer beam aperture did not cover the entire measuring cell and the beam was passing through its middle part, it can be assumed that the relative increase in optical density in the observed oscillations was associated with an increase in the Mo$_{132}$ concentration in the aperture due to the accelerated diffusion from the lower part of the cell. The diffusion coefficients of hydronium ions and polyanions are sufficiently different; therefore, an oppositely directed diffusion potential could occur in the system under study, which, on the contrary, slowed down the flow of ions, thus leading to a stepwise decrease in the value of A (Fig. 17), since POM destruction continued. The interaction of these factors could lead to the observed fluctuations.

Another factor that could potentially lead to a temporary partial blocking of the reaction of POM destruction is the oxygen deficiency in the solution, because the rate of oxygen dissolution in a liquid medium is limited. The value of D, the coefficient of oxygen diffusion in an aqueous medium given in [320], is around 2·10$^{-5}$ cm$^2$/s. A comparison of this value with the effective D value of Mo$_{132}$ polyanions under electrodiffusion conditions (1.8·10$^{-7}$ cm$^2$/s) [107] (POM concentration of 4·10$^{-4}$ mol/L), shows that the latter are much less mobile. For this reason, the supply of polyanions to the solution surface is more likely the limiting stage of the oxidative photodecomposition process. In this case, the diffusion restrictions, as well as fluctuations of A values, can at least be partially removed due to intensive vertical mixing of the solution. However, this is difficult to implement with the used measurement scheme. On the other hand, in order to obtain additional information, it is possible to increase the oxygen content in the medium contacting the solution, as it was done in the next experiment (Fig. 17) by replacing argon with air. With an increase in the oxygen content in the gaseous medium, the process of POM destruction accelerated in general, but the optical density fluctuations did not disappear, which indirectly confirms the hypothesized mechanism of the occurrence of oscillations.

![Fig. 18. Potential difference ($\Delta V$) between the illuminated and light-protected Mo$_{132}$ solution (0.015 wt. %) - PVA (10 wt. %) [318]. ©Pleiades Publishing, Ltd. 2009](image-url)
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of A. In addition, attention should be paid to the amount of oxygen required in accordance with the reaction scheme for the oxidation of $\text{Mo}_{132}$ in the solution: it takes about $1.5 \times 10^{-6}$ g of oxygen per 1 ml of a solution in the used concentration. The solubility of oxygen at room temperature in water and diluted solutions is about $6.6 \times 10^{-3}$ g/L. If 1 ml of the initial solution could contain about $6.6 \times 10^{-6}$ g of dissolved oxygen, then its amount was about 4.4 times higher than that required for POM oxidation.

The photochemical nature of POM oxidative destruction can be confirmed by the stabilization of $\text{Mo}_{132}$ upon formation of its associates with rhodamine-B, a dye of the xanthene series [303] (the main absorption maximum at a wavelength of 558 nm, near one of the maxima of the used LED). Stabilization manifests itself, for instance, during irradiation of such solutions ($\text{Mo}_{132}$ - RhB molar ratio of 1:8) in contact with air (Fig. 19). In another case, when a substance that is more easily oxidized than POM is in contact with the above associates, the $\text{Mo}_{132}$ - rhodamine-B system is capable of playing the role of a photoactivated catalyst. The oxidation of compounds such as toluene and isopropyl alcohol proceeds under quite mild conditions. Toluene is selectively oxidized to benzyl alcohol and benzaldehyde [321], and isopropyl alcohol is oxidized to acetone. Photoactivated compounds of the nanocluster type, including d-metals in different oxidation states, can be practically useful as catalysts for the oxidation of organic compounds [322, 323], for instance in processes of organic synthesis, and of alternative energetics (hydrogen production) for reducing the potential of electrochemical decomposition of aqueous solutions [324]. Examples of the use of molybdenum and tungsten compounds in the photolysis of water [316, 325] are known from the literature.

So, it has been shown that diffusion processes can be the cause of the oscillatory phenomena occurrence, in particular, of oscillations of the measured optical density of $\text{Mo}_{132}$ POM aqueous solutions under local exposure to light. In the studied system, the processes of POM oxidative photodestruction occurring in the surface part of the solution, are limited by the mobility of $\text{Mo}_{132}$ polyanions, the speed of which periodically changes depending on the ratio of the values of the concentration and diffusion potentials arising between the surface of the solution and its deep part. Further studies of oscillatory phenomena involving POM are of interest, especially taking into account the possibility of using these compounds in catalysis. It should be assumed that similar phenomena can occur in the body’s environment, and it should be taken into account, in particular, when monitoring drug release.

6. Conclusion

Thus, there are a number of factors that make it possible to consider molybdenum-based nanocluster POMs as compounds for use in biomedicine which show promise. These physical and biochemical properties include: the ability of POMs to undergo decomposition in vivo; the absence of accumulation of POM and their components in organs and tissues due to elimination in the course of natural metabolism; the possibility of obtaining conjugates of drugs and bioactive substances (including protein), biocompatible polymers, indicator dyes; targeted transport (for instance, ionophoresis) of POM and their conjugates into the body. $\text{Mo}_{72}\text{Fe}_{30}$ is able to penetrate into immuno-privileged organs (e.g., brain, eyes, thymus, spleen, etc.) and overcome blood-tissue barriers. At the same time, $\text{Mo}_{72}\text{Fe}_{30}$ has low toxicity for warm-blooded organisms; its administration by various methods does not lead to disruption of organs

![Fig. 19. Relative optical density of the irradiated solutions of $\text{Mo}_{132}$ (1) and $\text{Mo}_{132}$ - rhodamine-B (2) in contact with air (as a function of time, in min)](image-url)
and systems, being reflected only in small reversible changes at the metabolic level, e.g., in the aminotransferase activity. In addition, it was noted that the organism of animals (rats) is capable of adapting to the influence of POM, which, among other things, provides the normalization of the above-mentioned parameter. The effect of \( \text{Mo}_{72}\text{Fe}_{30} \) at the molecular level revealed itself only in a change in the content of heat shock proteins, which are involved in the adaptation of the organism and maintenance of homeostasis. The single and multiple action of POM did not cause a decrease in the amount of histone proteins in blood lymphocytes, which makes a suggestion about the absence of autoimmune aggression of these cells against liver tissue. The effect of rapid recovery of the hemoglobin level and the number of erythrocytes was established upon administration of \( \text{Mo}_{72}\text{Fe}_{30} \) preparations to animals in a state of post-hemorrhagic anemia. It indicates the possibility of therapeutic use for POMs, which have a positive effect on the processes of erythropoiesis.

\( \text{Mo}_{132} \) which contains Mo(V), is more toxic than \( \text{Mo}_{72}\text{Fe}_{30} \), in general. A study of the effect of these two POMs on cell cultures of normal and transformed fibroblasts showed that \( \text{Mo}_{72}\text{Fe}_{30} \) is practically non-toxic for normal fibroblasts, while \( \text{Mo}_{132} \) is toxic for both normal and transformed fibroblasts, and its damaging effect on transformed fibroblasts is stronger: a decrease in cell proliferative activity and an accelerated differentiation of transformed rat fibroblasts were recorded. The experimental results indicate the potential use of \( \text{Mo}_{132} \) in chemotherapy.

It is important to study the aspects related to the processes of POM destruction under various conditions in their free and conjugated states, as it allows predicting the lifetime of POM-based functional transport units and regulating their stability. The studies have shown possible mechanisms of POM destruction, including photoinitiated ones, and established that stabilizing factors can include self-aggregation of POM into hollow globular associates, the interaction of POM with such biogenic ions as calcium, and association with water-soluble nonionic biocompatible polymers. One of the most important stabilizing factors influencing POM upon entering normal alkaline environments of the body (e.g., blood) is the interaction with proteins, in particular, with albumins (protein corona appearance), while in the environment of a corresponding alkaline buffer solution, POMs decompose much more quickly.

Fundamental approaches for the creation of new POM-based controlled releasing systems with feedback have been proposed. In these systems, POM conjugates with a drug and an indicator luminescent dye are initially bound with a matrix of a biocompatible polymeric organic or inorganic carrier in the microgel composition. This should ensure prolonged release of drugs and simultaneous monitoring of their residual concentration indicated by the fluorescent signal of the dye molecules.

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References

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Liu P., Zhang R. Polymer microspheres with high drug-loading capacity via dual-modal drug-loading for modulating controlled release properties.


Wang H., Li Z., Lu S., Li C., Zhao W., Zhao Y., Yu Sh., Wang T., Sun T. Nano micelles of cellulose-graft-poly (l-lactic acid) anchored with epithelial cell adhesion antibody for enhanced drug loading and anti-tumor effect.


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