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THE ROLE OF ARBUSCULAR MYCORRHIZAL SYMBIOSIS IN ¹³⁴CS UPTAKE BY CROP AND WILD PLANT SPECIES

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The role of arbuscular mycorrhizal fungus Glomus intraradices in ¹³⁴Cs isotope uptake by different plant species is studied. The impact of radiocaesium on mycorrhizal development and functioning of plant photosynthetic apparatus is considered. The possibility of mycorrhizal symbiosis application in phytoremediation of radioactively contaminated areas is analyzed. It is found that colonization of plants by arbuscular mycorrhizal fungus resulted to significant decrease of radiocaesium content in their aboveground parts, while it didn't have considerable impact on the radioclude uptake by plant root system. *Keywords:* radiocaesium, radioactive contamination of environment, arbuscular mycorrhiza, arbuscular mycorrhizal fungi, plant photosynthetic apparatus, mycorrhizal colonization, phytoremediation.

Роль арбускулярного мікоризного симбіозу в накопиченні ¹³⁴Cs дикорослими та культурними видами рослин. Сергій Валерійович Дубчак. Досліджено роль арбускулярного мікоризного гриба Glomus intratadices у накопиченні ізотопу ¹³⁴Cs різними видами рослин. Розглянуто вплив радіоцезію на розвиток мікоризи та функціонування фотосинте тичного апарату рослин. Проаналізовано можливість застосування мікоризного симбіозу у фіторемедіації радіаційно забруднених територій. Встановлено, що колонізація рослин арбускулярним мікоризним грибом призвела до суттєвого зменшення концентрації радіоцезію в їхній надземній частині й водночас не мала значного впливу на надходження радіонукліда до кореневої системи рослин. *Ключові слова:* радіоцезій, радіоактивне забруднення довкілля, арбускулярна мікориза, арбускулярні мікоризні гриби, фотосинтетичний апарат рослин, мікоризна колонізація, фіторемедіація.

Роль арбускулярного микоризного симбиоза в накоплении ¹³⁴Сs дикорастущими и культурными видами растений. Сергей Валериевич Дубчак. Исследована роль арбускулярного микоризного гриба Glomus intraradices в накоплении изотопа ¹³⁴Сs различными видами растений. Рассмотрено влияние радиоцезия на развитие микоризы и функционирование фотосинтетического аппарата растений. Проанализирована возможность применения микоризного симбиоза в фиторемедиации радиоактивно загрязненных территорий. Установлено, что колонизация арбускулярным микоризным грибом привела к существенному уменьшению концентрации радиоцезия в их надземной части и одновременно не имела значительного влияния на поступления радионуклида в корневую систему растений. *Ключевые слова:* радиоцезий, радиоактивное загрязнение окружающей среды, арбускулярная микориза, арбускулярные микоризные грибы, фотосинтетический аппарат растений, микоризныя колонизация, фиторемедиация.

Statement of the problem. The radiocaesium isotopes have been introduced into the environment via various routes for last several decades. Altogether, roughly 1 EBq (10^{18} Bq) of long-lived ¹³⁷Cs was released to the Earth's biosphere in the XX – XXIth centuries that resulted to contamination of vast areas all over the

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world. About 90 % of radiocaesium was originated from atmospheric nuclear testing, approximately 4 % was released by fuel reprocessing and nuclear fuel facilities and roughly 6 % - by Chornobyl and Fukusima accidents. Nowadays the ¹³⁷Cs absorption by plants and its accumulation, therefore, represents the main source of human exposure to this radionuclide. The principal route of radiocaesium entry into biological food chain in terrestrial ecosystems is the soil-to-plant pathway. This radionuclide is expected to remain in the rooting zone of plants for decades and respectively to be involved in biological migration chains. However, the mechanisms by which radiocaesium is taken up by plant roots are not completely understood.

Analysis of recent studies and **publications**. Recently the alternative strategies, orientated towards the use of plants and micro-organisms, separately or in combination, have been proposed for removing or immobilizing radiocaesium and othe radionuclides in the soil [1.2]. Among these micro-organisms. mycorrhizal fungi received a particular attention. An estimated 90 % of terrestrial plants exist in a symbiotic association with soil fungi forming mycorrhizal associations. Among them, the obligate arbuscular mycorrhizal (AM) fungal symbionts are supposed to have a principal role [3]. These fungi are important participants in the Cs cycle in the upper layers of soils. They have strong impact on mobility of radiocaesium in the soil and result to unavailability of this radionuclide to the other components in ecosystems [4]. At the same time, it was demonstrated [5] that AM fungi can transform and immobilize radionuclides and correspondingly limit their toxicity and bioavailability to plants and spreading into the soils. Accordingly, plants

growing in contaminated soil could obtain benefit from their AM fungal symbiotic partners.

Nevertheless, the role of arbuscular mycorrhizal fungi on the acquisition of radiocaesium by plants remains poorly understood and controversial. The lack of clear results on the capacity of AM fungi to accumulate or transport Cs could be principally attributed to different and inadequate experimental systems used in previous studies. Furthermore, the various AM fungi and plants studied could also explain the controversial conclusions obtained, since AM fungi and plants have probably different capacity to accumulate and transport radiocaesium.

Objectives of research. The goals of this work were to identify the capacity of AM fungi to take up and transfer caesium isotopes to their hosts as well as to estimate the influence of arbuscular mycorrhiza on radiocaesium uptake by plants and impact of radiocaesium on development of AM fungal symbioses.

The main material of the study. Four plant species (Plantago lanceolata, Medicago truncatula, Lolium perenne and Helianthus annuus) capable to form efficient association with a broad range of AM fungi were selected for our study. The plants were cultivated in the presence or absence of AM fungus Glomus intraradices (strain BIO, obtained from BIORIZE, Dijon, France). The ¹³⁴Cs isotope (obtained from "POLATOM" Radioisotope Centre, Otwock-Świerk, Poland) in was added to sterilized substrata in pots in form of CsCl water solution. ¹³⁴Cs activity concentration was adjusted to 100 000 Bg per pot (77 000 Bq·kg⁻¹). The plants were grown in transparent Sun bags (SigmaTM Aldrich, Poznan, Poland) in a

growth chamber at 20 °C, with a photoperiod of 12 h light and 12 h darkness, at photosynthetic photon flux density 30 \pm 6 µmol·(s·m²)⁻¹ and harvested each three months.

The activity concentration of ¹³⁴Cs in roots and shoots of plants was determined using a gamma-spectrometer with semiconductor p-type coaxial high purity HP-Ge detector with a relative efficiency of 15 % and resolution of 2.5 keV at 1.33 MeV, shielded by 10 cm of lead with inner lining with 2 mm Cd and 18 mm Cu.

For the estimation of mycorrhizal colonization, the roots of plants were carefully washed with tap water, softened in 10% potassium hydroxide for 24 hours, washed in water again, acidified in 5% lactic acid in water for 12 -24 h and stained with 0.01% aniline blue in lactic acid (to visualize AMF) for 24 h at room temperature. The root fragments were mounted and squashed on the slide covered with lactoglicerole. The parameters of AM colonization were assessed according to the method developed in [6] that assumes six levels of mycorrhizal colonization (from 0 to 5). The relative mycorrhizal root length (M%), intensity of colonization within individual mycorrhizal roots (m%), relative arbuscular richness (A%) and arbuscule richness in root fragments where the arbuscules were present (a%) were evaluated using Nikon Eclipse 800 light microscope equipped with Nomarski contrast and fluorescence.

The photosynthetic activity of plants was evaluated using a Plant Efficiency Analyzer fluorimeter (Hansatech Instruments, UK) estimating Chlorophyll *a* fluorescence transients of intact plant leaves. The Chl *a* fluorescence transients (OJIP transients) were induced by a

red light pulse (peak at 650 nm) of 600 W·m⁻² intensity provided by an array of three light-emitting diodes. The transients were recorded for 1 s with 12 bit resolution, starting 10 µs after the onset of illumination. Each transient was analyzed according to the OJIP-test based on the theory of energy fluxes in biomembranes [7]. The selected original data were processed by means of their utilization for the calculation of biophysical parameters by the JIP-test equation and the number of biophysical parameters were calculated. Among them, the most parameters are the performance indexes Plabs (evaluated on the base of light absorption) and PI_{total} (total performance index). Plabs and Plto. tal comply all basic biophysical parameters and represent the photosynthetic system vitality.

Part 1. ¹³⁴Cs uptake by plant species.

P. lanceolata inoculated with G. intraradices contained 66846 ± 11029 $Bq \cdot kg^{-1}$ of ¹³⁴Cs in their shoots, that is considerably lower in comparison with the radionuclide activity concentration in nonmycorrhizal plant shoots (87500 \pm 12333 Bq·kg⁻¹). At the same time, ¹³⁴Cs activity concentration in roots of mycorrhizal and nonmycorrhizal P. lanceolata wasn't differed significantly, although the slightly higher radiocaesium concentration (18.1 ± 9.9 %) was found in roots of nonmycorrhizal plants (Fig. 1A). Due to the higher biomass of mycorrhizal P. lanceolata, the ¹³⁴Cs activity in roots and shoots of single mycorrhizal and nonmycorrhizal plant (Bq.plant⁻¹, dry weight) and correspondingly the total radiocaesium activity in a single plant

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(i.e. shoots plus roots) weren't differed substantially (p < 0.05). The root/shoot ratios of ¹³⁴Cs activity concentration in *P. lanceolata* colonized with *G. intraradices* were slightly (about 11 %) higher as compared to those of nonmycorrhizal ones. As it's known [8], the higher root/shoot ratios of caesium content in plants indicates the reduced root to shoot translocation of this element, thus the tendency of the mycorrhiza to reduce radionuclide translocation from *P. lanceolata* roots to shoots was revealed.

The colonization of *M. truncatula* with G. intraradices also caused a significant reduction of radiocaesium uptake in plant shoots. Thus, ¹³⁴Cs activity concentration in aboveground part of mycorrhizal M. truncatula was 86888 ± 20022 Bg·kg⁻¹. whereas shoots of nonmycorrhizal plants contained 132100 \pm 15505 Bq·kg⁻¹ of this radionuclide. At the same time, the mycorrhiza resulted to considerable (18.8 \pm 5.6 %) increase of radiocaesium activity concentration in M. truncatula roots in comparison with that in nonmycorrhizal plants (see Fig. 2B). The distribution of ¹³⁴Cs activity between aboveground and underground parts of mycorrhizal and nonmycorrhizal M. truncatula was differed. Thus, the radionuclide activity concentration in roots of mycorrizal M. *truncatula* was 45.5 ± 14.7 % lower than in their shoots. The opposite tendency was observed in case of nonmycorrhizal plants, where ¹³⁴Cs activity concentration in shoots was 24.2 ± 8.9 % higher than in roots (see Fig. 2B). No statistically significant differences were found between dry of mycorrhizal masses and nonmycorrhizal plants, although both roots and shoots of AM inoculated M. truncatula grown on ¹³⁴Cs spiked substrata had slightly higher weight (about 14 and

12 % correspondingly) as compared to those of nonmycorrhizal plants. The colonization of plants grown on radioactively contaminated substrata also led to moderate (about 10 %) increase of their shoot length. The evaluated ¹³⁴Cs activity in shoots of single mycorrhizal alfalfa was 9.3 ± 0.6 Bq, whereas shoots of nonmycorrhizal *M. truncatula* contained significantly higher amount of radiocaesium (12.3 \pm 0.6 Bg). On the contrary, the radionuclide activity in roots of mycorrhizal alfalfa was substantially higher $(3.1 \pm 0.1 \text{ Bg})$ as compared to that of nonmycorrhizal plants ($\hat{2}.3 \pm 0.2$ Bq). Consequently, mycorrhizal M. truncatula had significantly lower total activity of radiocaesium (12.4 \pm 0.7 Bg) when compared to that of nonmycorrhizal plants $(14.6 \pm 0.8 \text{ Bg})$. The radionuclide translocation from underground to aboveground parts of plants was more intensive in case of nonmycorrhizal alfalfa. Their root/shoot ratio of ¹³⁴Cs activity concentration was 0.81 ± 0.28 being considerably lower in comparison with that of mycorrhizal M. $truncatula (1.45 \pm 0.41).$

The harvested plants of L. perenne mycorrhizal with G. intraradices had more than two fold lower ¹³⁴Cs activity concentration both in their roots and shoots as compared to those of nonmycorrhizal plants. The ¹³⁴Cs distribution within L. perenne demonstrated that the radionuclide activity concentration in aboveground parts of both mycorrhizal and nonmycorrhizal ryegrass was about three times lower when compared to that of plant underground parts (see Fig. 1C). Dry weights of mycorrhizal and nonmycorrhizal L. perenne and their shoot length weren't differed considerably, however the biomass of plants colonized with G. intraradices and grown on substrata

spiked with ¹³⁴Cs was slightly (less than 10 %) higher when compared to that of nonmycorrhizal species. Hence, the colonization with *G. intraradices* resulted to considerable decrease of ¹³⁴Cs activity (Bq per plant) in shoots (76.3 \pm 22.8 %) and roots (53.3 \pm 15.8 %) of single mycorrhizal ryegrass as compared to that of nonmycorrhizal plants.

As opposed to plant species considered above, the colonization of *H. annuus* with *G. intraradices* resulted to significant increase of ¹³⁴Cs uptake by plants. Thus, the radiocaesium activity concentrations both in underground and aboveground parts of mycorrhizal sunflowers were nearly 10 fold greater when compared to those of nonmycorrhizal plants (see Fig. 1D). At the same time, roots of both mycorrhizal and nonmycorrhizal sunflowers had about 50 % higher ¹³⁴Cs activity concentrations when compared to plant shoots.

The presence of ¹³⁴Cs didn't have appreciable impact on *H. annuus* growth parameters, and the most distinct was the mycorrhiza influence. Thus, the mycorrhizal H. annuus grown on radioactive and clean substrata produced correspondingly 12 and 11 % longer shoots as compared to those of nonmycorrhizal plants. The shoots dry weight of mycorrhizal H. annuus grown both on radioactively contaminated and non-polluted soil exceeded substantially (about 70 and 80 % respectively) shoots dry weight of nonmycorrhizal ones. The degrees of ¹³⁴Cs translocation from roots to shoots of mycorrhizal and nonmycorrhizal H. annuus weren't differed considerably due to similar root/shoot ratios of the radionuclide activity concentration $(1.54 \pm 0.10 \text{ and})$ 1.47 ± 0.21 correspondingly).

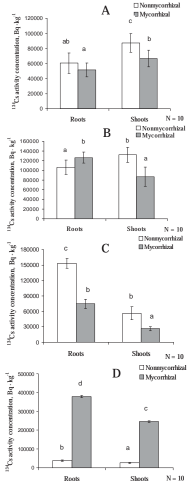


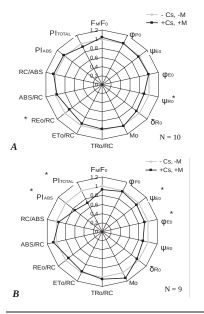
Fig. 1. ¹³⁴Cs activity concentration ($Bq \cdot kg^{-1}$) in roots and shoots of Plantago lanceolata (A), Medicago truncatula (B), Lolium perenne (C) and Helianthus annuus (D) mycorrhizal or not with Glomus intraradices and grown on substrata spiked with ¹³⁴Cs (77 000 $Bq \cdot kg^{-1}$). The results are presented as mean ± standard deviation. The different letters above bars mean statistically significant differences (p < 0.05).

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Part 2. Functioning of plant photosynthetic apparatus.

The most of photosynthesis biophysical parameters both in mycorrhizal and nonmycorrhizal plants cultivated on substrata with ¹³⁴Cs weren't varied considerably as compared to those of control ones (Fig. 2A-2D). The exception was observed only in case of M. truncatula grown on radioactively contaminated substrata which had considerably lower efficiency of trapped exciton movement into electron transport chain ($\psi_{E_0} = ET_0/TR_0$) and maximum vield of electron transport (ϕ_{Eo} = ET_0/ABS) when compared to those of control plants from clean substrata. Also, the total and absorption vitality indexes (PI_{abs} and PI_{total}) of these alfalfas were respectively 33.2 ± 12.9 and 42.0 ± 18.7 % lower than those of control plants (Fig. 2B).



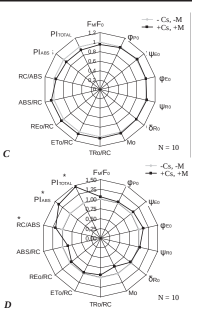


Fig. 2. Biophysical parameters of photosynthesis of Plantago lanceolata (A), Medicago truncatula (B), Lolium perenne (C) and Helianthus annuus (D): nonmycorrhizal (control) plants grown on clean soil (-Cs, -M) and plants mycorrhizal with Glomus intraradices and cultivated on substrata spiked with ¹³⁴Cs (+Cs, +M). Values on plots are presented in relative units and normalised on those of the control plants. *- means statistically significant difference (p < 0.05).

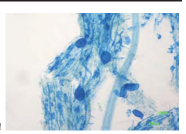
The fungal colonization of plant species grown on radioactive substrata had positive impact on functioning of *H. annuus* photosynthetic apparatus (Fig. 2D). In this case, the AM inoculation of plants cultivated on radioactive soil improved y vitality indexes of sunflowers. Thus, PI_{abs} of mycorrhizal *H. annuus* was correspondingly 41.4 \pm 11.3 and 36.3 \pm 12.5 % higher than those of nonmycorrhizal plants grown

on soil with ¹³⁴Cs and control plants. In turn, PI_{total} of mycorrhizal sunflowers exceeded considerably those of nonmycorrhizal plants cultivated on radioactive substrata as well as control plants (40.0 ± 9.7 and 44.8 ± 11.9 % respectively, Fig. 2D).

Part 3. Arbuscular mycorrhizal colonization of plants.

Both treated with radiocaesium and control plant species were characterized with high mycorrhizal frequency (F%) that exceeded 90 %. The intraradical structures of the AM fungus were morphologically typical for Arum-type mycorrhizae. The intraradical hyphea of G. intraradices propagated between cortical root cells at long distances and formed lateral branches, which penetrated cells and produced arbuscules inside them (Fig. 3). The presence of numerous intercellular vesicles was characteristic for nearly 80 % of studied root fragments. The spores of G. intraradices that have thicker walls in comparison with vesicles were found only in several root fragments.

The most of AM colonization parameters of plants cultivated on substrata spiked with ¹³⁴Cs and nonpolluted soil weren't differed significantly (Fig. 4A,C,D). Although, in case of *M. truncatula* the presence of radiocaesium resulted to considerable (about 30 %) decrease of mycorrhizal colonization intensity for all and individual mycorrhizal plant roots (M,% and m,% correspondingly) when compared to those of control plants (Fig. 4B).



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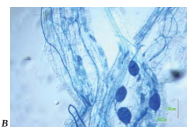
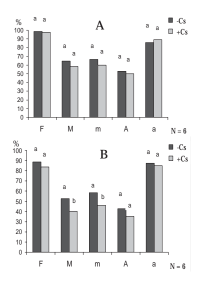


Fig. 3. Arbuscules and vesicles of Glomus intraradices within roots of Plantago lanceolata (A) and Medicago runcatula (B) cultivated on substrata treated with ^{134}Cs (77 000 $Bq\cdot kg^{-1}$).



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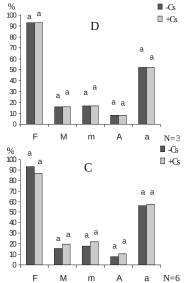


Fig. 4. Arbuscular mycorrhizal colonization parameters of Plantago lanceolata (A), Medicago runcatula (B), Lolium perenne (C) and

Helianthus annuus (D) inoculated with Glomus intraradices: F% - frequency of mycorrhiza; M% - mycorrhizal colonization intensity for all roots; m% - mycorrhizal colonization intensity within individual mycorrhizal roots; A% - arbuscular richness for all roots; a% - arbuscular richness in root fragments where the arbuscules were present, medians. Plants were cultivated on clean substrata (-Cs) and substrata spiked with ¹³⁴Cs (+ Cs). The different letters above bars mean statisti-

cally significant differences (p < 0.05).

The principal goals of the research were to compare the possible influence of mycorrhiza on various AM fungal symbionts cultivated on the same substrata under the impact of ¹³⁴Cs. Our results suggest that inoculation with AMF changed substantially the uptake of ¹³⁴Cs by studied plant species and influ-

enced the translocation of caesium isotopes within the plants. The arbucular mycorrhiza resulted to considerable decrease of ¹³⁴Cs activity concentration in shoots of *P. lanceolata, M. truncatula* and *L. perenne* when compared to nonmycorrhizal ones. The most significant (about threefold) reduction of ¹³⁴Cs activity concentration was found in shoots of mycorrhizal *L. perenne*. This result contradicts to the data obtained by [9] who found that inoculation with arbuscular mycorrhiza significantly enhanced uptake of ¹³⁷Cs by ryegrass.

The exception in our study was H. annuus where the AM colonization led to nearly tenfold increase of ¹³⁴Cs activity concentration both in plant roots and shoots. The sunflower was previously shown to be an effective hyperaccumulator of ¹³⁷Cs and ⁶⁰Co [10], although the ability of this plant to form mycorrhiza has not been studied. In our case H. annuus revealed its ability of ¹³⁴Cs hyperaccumulation only in the presence of the mycorrhiza. Also, H. annuus was only plant species in our experiment whose shoot biomass was significantly affected by the impact of AM fungus. Such contradictory findings demonstrate that basic knowledge of Cs potential uptake mechanisms are needed to facilitate the design of countermeasures to reduce or enhance the transfer of radiocaesium into plants.

Conclusions. In summary, *M. truncatula* was suggested to be the most sensitive plant species relative to the radiocaesium impact. Due to considerable reduction of caesium in their shoots this plant species as well as *P. lanceolata* and *L. perenne* couldn't be applied in phytoremediation, but they may be potentially used in phytostabilization of the radioactively polluted ecosystems.

On the other hand, the use of H. annuus with its Cs hyperaccumulation properties conditioned by mycorrhiza for the phytoremediation is also questionable. In our study, the evaluated total activity of ¹³⁴Cs accumulated in biomass of sunflowers grown in one pot (two plants) during three months was 221 Bq. This is only 2.2 % from total radiocaesium activity in the pot (100 000 Bg). Extrapolating these data for a longer term and assuming the plant active growth period is about 6 months per year, we can roughly estimate that nearly two decades are needed to remove radiocaesium completely from the soil. This assumption doesn't take into consideration the natural factors, such as the radiocaesium migration. inhomogeneous distribution in soil and possible leaching of the radionuclide below the 30 - 40 cm (i.e. outside of root zone) as well as potential impact of another AM fungi and various soil microorganisms on the radionuclide uptake by plants.

The number of authors proposed using of AM fungi in phytoremediation strategies for radiocaesium contaminated areas to enhance radionuclide re-

moval by plant biomass [1, 11, 12]. Although, the effects of AM fungi on Cs accumulation could be applied in strategies to develop crops with smaller soilto-plant transfer factors which accumulate less Cs [8, 12]. Such plant species may be potentially grown within areas with moderate radiocaesium contamination levels and further used in agricultural purposes, if the radionuclide content in their biomass doesn't exceed the prescribed permissible levels. Our results demonstrated the capacity of AM fungi to influence the acquisition and accumulation of caesium isotopes by plants by immobilizing, transporting and affecting the root-to-shoot translocation. Nevertheless the AM fungal ability to take part in phytoremediation strategies still remains questionable and needs for further researches.

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ЕКОЛОГІЯ І ВИРОБНИЦТВО

УДК 504.064; 614.835.3

ТЕХНОГЕННІ РИЗИКИ ЗАБРУДНЕННЯ ДОВКІЛЛЯ ПІД ЧАС РЕМОНТНИХ РОБІТ РЕЗЕРВУАРІВ ІЗ НАФТОПРОДУКТАМИ

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Наведено інформацію щодо поліпшення стану забезпечення екологічної безпеки навколо небезпечних об'єктів з наявністю технологічних процесів, пов'язаних з експлуатуванням та проведенням ремонтних робіт резервуарів із нафтопродуктами шляхом управління техногенними ризиками з урахуванням впливу чинників на їх значення. *Ключові слова:* техногенний ризик, забруднення довкілля, резервуар з нафтопродуктами, пробіт-функція, нафтозалишки.

Техногенные риски загрязнения окружающей среды при проведении ремонтных работ резервуаров с нефтепродуктами. Липовой В.А., Удянский Н.Н. Приведена информация об улучшении состояния обеспечения экологической безопасности вокруг опасных объектов с наличием технологических процессов, связанных с эксплуатацией и проведением ремонтных работ резервуаров с нефтепродуктами, путем управления техногенными рисками с учетом влияния факторов на их значение. Ключевые слова: техногенный риск, загрязнение окружающей среды, резервуар с нефтепродуктами, пробит-функция, нефтеостаток.

Man-caused environmental pollution during repair tanks with oil. Lipovoy V.O., Udyansky N.N. The data on the improvement of environmental safety around dangerous objects to the presence of processes associated with the operation and maintenance work tanks with oil by controlling technological risks, taking into account certain factors influence their value. Keywords: technological hazards, pollution, oil reservoir, the probit function, the residue oil.

Аварійні викиди та витоки шкідливих речовин внаслідок проведення регламентних та ремонтних робіт з очищення внутрішніх технологічних поверхонь резервуарів із нафтопродуктами можуть призвести до локального та катастрофічного рівня завдання шкоди довкіллю та життєдіяльності людей.

Статистика свідчить, що понад 20% усіх пожеж на резервуарах зберігання нафтопродуктів відбувається через порушення вимог пожежної безпеки при проведенні ремонтних