1. Introduction

The pollution of freshwater resources has presently acquired a global nature. Any changes in the composition of water have a direct impact on all the activity processes of water organisms. Current levels of anthropogenic stress on freshwater resources force to periodically evaluate water quality and the health of hydroecosystems. The water resources of Armenia have been affected by intensive anthropogenic impact. Lake Sevan, one of the largest and most exploited alpine limnosystems in the world, playing a unique role in the economic development of Armenia (Fig. 1). The lake is fed by 28 rivers that drain residential, agricultural and industrial areas. The use of water from Lake Sevan considerably improved the condition of industry and agriculture in the country. In the pollution of Lake Sevan, much role belongs to a sharp increase in the inflow of polluting and toxic substances of technogenic origin from the lake basin (Hovhannisyan, 1994; Matishov et al., 2016).

Despite the economic development, the state of the lake’s basin is deteriorated. Nevertheless, biotesting of the state of the lake ecosystem has been limited to monitoring the chemical composition of the water and biomonitoring using sensitive model test-system. (Steinkellner et al., 1999; Ma et al., 2005) The process of bioindication of genotoxic effects of complex mixtures on the water resources using plants test-objects is very appropriate and effective. Test-systems based on plant use have shown high sensitivity and are recommended for monitoring genotoxicity of xenobiotics in the air, water and soil (Grant, 1994; Ma, 1999; Aghajanyan et al., 2015). Among plants test-objects, the heterozygote clones for flower color (blue – dominant, pink – recessive) of Tradescantia (clone 02) plays a special role.

The Tradescantia stamen hair mutation (Trad-SHM) and Tradescantia-micronucleus (Trad-MCN) bioassays are the efficient and reliable biomonitoring test-systems for detecting the genotoxicity and clastogenic effects of environmental pollutants (Ma et al., 2005; Majer et al., 2005; Mizik et al., 2011; Avalyan et al., 2017). These bioassays are included in the International Program on Plant Bioassay (IPPB) under the auspices of The United Nations Environment Programme (UNEP) (Ma, 1999).

The aim of this study was to assess the genotoxicity and clastogenicity of water from different parts of the Lake Sevan basin, Armenia by Trad-SHM and Trad-MCN assays using Tradescantia (clone 02). The Trad-SHM and Trad-MCN assays were simultaneously used to determine the genetic effects of the investigated water samples.
2. Material and methods

Sampling sites
Water samples were collected at three times in different periods from seven locations of the Lake Sevan basin, Armenia: off the Artanish, Karchaghbyur, Tsapatagh, Noradus, Litchk, Masrik, and Martuni settlements (Fig. 2). The samples were collected at a depth of approximately 30 cm in three replicates per sampling site in 1 L pre-cleaned plastic bottles. Samples were stored at +4°C until delivery to the laboratory. Tests were initiated on the next day.

Tradescantia (clone 02)
Tradescantia (clone 02) used for this study was obtained from the greenhouse of Yerevan State University. It is an interspecific hybrid between Tradescantia occidentalis Briffon ex Ridb. and T. ohiensis Raf. (Fig. 3). This clone is heterozygous for flower color (blue/pink, the pink color being recessive). For Tradescantia stamen hair mutation bioassay (Trad-SHM), the appearance of pink cells of stamen hair (recessive mutation events – RME) was a marker criterion. In addition, during testing there were morphological changes in stamen hairs, dwarf (non-surviving NS) and branching hairs (BH). Using micronucleus bioassay (Trad-MCN) two main test criteria were counted: percentage of micronuclei in tetrads and percentage of tetrads with micronuclei.

Trad-SHM assay
The assay was based on counting somatic mutations and morphological changes that could be indicators of genotoxic or teratogen effects of Tradescantia (clone 02).

For testing, young inflorescences were submerged in water samples (three-five inflorescences for vessel per variant) for 24 h, and after a seven-day recovery period, flower blossoms were examined. Stamen hairs of a flower were placed on a slide and observed under a magnifying glass at 10x for counting blue to pink mutation and morphological changes. Flowers were examined daily over 21 days since their appearance. From 8000–14000 stamen hairs were obtained per sample. The number of mutations per sample was calculated per 1000 hairs according to standard protocols (Ma et al., 1994a). Tap water was used as a control, and test vessels were set in triplicate.

Trad-MCN assay
The assay was based on the scoring of the micronuclei (MCN) frequency in pollen mother cells at disturbances in the process of microsporogenesis (Ma et al., 1994b; Mizik et al., 2011). Formation of micronuclei is related to chromosomal instability and is indicative of genomic damage.

For testing, young inflorescences of Tradescantia were also submerged into the test water samples. The duration of treatment was 24 h. The inflorescences were removed and fixed in Carnoy’s solution (3:1 ethanol-glacial acetic acid). Preparation of the tetrads and the scoring of the micronuclei were carried out as described.
in Ma et al. (1994b). The following test criteria were used: the number of tetrads with micronuclei (Tetr/MCN) and the number of micronuclei per 100 tetrads (MCN/tetr). For each water sample, 3000 tetrads were analyzed.

**Chemical and statistical analysis**

Water samples were analyzed for chemical composition at a certified laboratory, following standard methods (Standard methods..., 1998). Several elements were measured (total concentrations): Al, Ni, Zn, As, Cu, Fe, Cr, Co, Mo, Cd, Sn, Pb, Mg, Mn, Na, Ca, and K (Table 1). At all sites, the pH level varied from 8.6 to 8.7. Chemical data were compared to the legislated water quality standards for aquaculture (from the Environmental Impact Monitoring Center of the Ministry of Nature Protection of Armenia (www.armmonitoring.am)).

All results were statistically processed using the Student t-test and Pearson correlation test by the computer program Statgraphics Centurion 16.2. Pearson correlation test was performed to analyze the correlation between the level of somatic mutation and the percentage of MCN as well as the concentrations of various elements and heavy metals in the studied water samples.

**3. Results and discussion**

The results of bioassay Trad-SHM have revealed a significant increase in the level of recessive mutation events (RME) frequency by water samples collected off the Noradus, Litchk, Tsapatagh, and Martuni settlements compared to the control. There is the positive correlation between the RME frequency and the concentration of some chemical elements (P, Si, Al, Mn, Fe, and Cu) in the studied water samples (Table 2).

The study of clastogenic effects in the sporogenic cells of Tradescantia has also shown an increase in the occurrence frequency of both test criteria in all studied water variants compared to the control level. The water samples collected off the Litchk, Tsapatagh, Martuni, and Masrik settlements show the maximum manifestation of these genetic effects from both investigated parameters. There is a significant positive correlation between the frequency of tetrads with MN and the concentration of Ni and Co in the studied variants.

The obtained results have shown that the significant increase in the level of recessive mutations and clastogenic effects in the studied water samples indicates the contamination of the water in the Lake Sevan basin at the tested sites near settlements with various kinds of xenobiotics (heavy metals, pesticides, etc.) as well as their influence on the processes occurring in the somatic and generative cells of Tradescantia.

A chemical mixture of metals is more toxic than individual metals if their action is synergistic. On the other hand, the antagonistic effects of some metals in a mixture diminish the uptake of individual metals e.g., by competing for the same uptake sites at the cell surface. The toxicity of a mixture, however, depends not only on the exposure concentration of each mixture constituent and its ratio but also the means of the toxicants to act jointly (Altenburger et al., 2004). Moreover, effects may be observed at concentrations far below those predicted to be safe by regulatory frameworks (Garcia et al., 2011; Liess et al., 2016). Various authors stated that it is very important to assess the genotoxicity of contaminants in the entire environmental water sample, but not for each component. The identification of toxicants in complex mixtures realized by exposure information about biological effects and approaches that deal with complex contamination typically reveals not only individual toxicants. Thus, since the lake samples used in this study are a complex mixture, we can legitimately assume the collective effect of not only individual metals even at low concentrations.

**4. Conclusion**

According to the results of our studies, the increased frequency of MCN and the elevated pink mutation (RME) rate in the Tradescantia inflorescence are indicators of clastogenicity and mutagenicity resulting from exposure to environmental pollutants. The use of both bioassays (Trad-SHM and Trad-MCN) may be effective for biotesting the water quality

|                | Na  | Mg  | Al  | P   | K   | Ca  | V   | Cr  | Mn  | Fe  | Co  | Ni  | Cu  | Zn  | As  | Mo  | Pb  |
|----------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| **Artanish**   | 47.8| 41.7| 0.05| 0.3 | 14.8| 18.4| 5   | 20  | 0.7 | 5   | 0.1 | 2   | 1   | 0.7 | 8   | 5   | 0.0 |
| **Karchaghbyur**| 72.9| 58.4| 0.1 | 0.1 | 19.4| 25.1| 8   | 30  | 1   | 460 | 0.1 | 3   | 3   | 7   | 9   | 7   | 0.1 |
| **Noradus**    | 48.6| 41.9| 0.07| 0.1 | 15.1| 18.9| 5   | 20  | 0.2 | 70  | 0.1 | 2   | 2   | 0.7 | 8   | 5   | 0.04|
| **Masrik**     | 40.6| 36.1| 0.04| 0.04| 13.6| 21.6| 5   | 20  | 0.2 | 60  | 0.1 | 3   | 1   | 0.2 | 9   | 5   | 0.03|
| **Tsapatagh**  | 61.8| 52.8| 0.09| 0.2 | 20.2| 23.9| 7   | 20  | 0.7 | 190 | 0.1 | 3   | 2   | 0.6 | 10  | 5   | 0.0 |
| **Litchk**     | 38  | 30  | 0.12| 0.08| 11.5| 16.1| 7   | 20  | 1   | 70  | 0.1 | 1   | 0.6 | 0.3 | 8   | 2   | 0.2 |
| **Martuni**    | 8.5 | 5.8 | 0.0 | 0.0 | 2.0 | 12.9| 5   | 7   | 1   | 70  | 0.1 | 1   | 1   | 0.5 | 4   | 4   | 0.5 |
| **Control**    | 8.47| 5.83| 0.0 | 0.0 | 1.96| 12.9| 23  | 7.6 | 1.1 | 8.6 | 0.0 | 1.7 | 1.1 | 64  | 3.8 | 2.3 | 0.1 |
and allows assessing the potential risk of aquatic contaminants of the natural hydroecosystem of Lake Sevan.

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Table 1. Induction of genotoxic and clastogenic effects of \textit{Tradescantia} (clone 02)

| Sampling sites | Frequency of RME/1000 | Frequency of micronuclei in sporogenic cells |
|---------------|-----------------------|---------------------------------------------|
|               | Tetrams with MCN (% ± SD) | MCN in tetrads (% ± SD)         |
| 1. Artanish   | 0.8 ± 0.31*          | 3.4 ± 0.33***                        |
| 2. Karchaghbyur | 0.7 ± 0.28*        | 5.2 ± 0.40***                        |
| 3. Noradus    | 1.8 ± 0.44**         | 5.5 ± 0.41***                        |
| 4. Masrik     | 0.7 ± 0.29*          | 8.3 ± 0.50***                        |
| 5. Tsapatagh  | 1.2 ± 0.29**         | 6.0 ± 0.43***                        |
| 6. Litchk     | 1.6 ± 0.34**         | 6.0 ± 0.43***                        |
| 7. Martuni    | 1.2 ± 0.29**         | 6.2 ± 0.44***                        |
| Control       | 0.2 ± 0.14           | 1.8 ± 0.24                           |

*- p<0.05, ** - p<0.01, *** - p<0.001