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Applying multi-criteria analysis for preliminary assessment of the properties of alginate immobilized *Myriophyllum spicatum* in lake water samples



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ARTICLE INFO

Article history: Received 13 December 2017 Received in revised form 9 May 2018 Accepted 10 May 2018 Available online 11 May 2018

Keywords: Lake water samples Immobilized aquatic weed Biosorption Water quality Multi-criteria analysis

ABSTRACT

The preliminary assessment of the properties of alginate immobilized aquatic weed Myriophyllum spicatum beads-MsAlg in a multi-element system of nine Serbian lakes water samples was done. Herein, the results obtained in the biosorption experiment with MsAlg contents of twenty-two elements analysed by inductively coupled plasma-optical emission spectrometry, biosorption capacity, element removal efficiency, total hardness (TH) and quality index of water (WQI) are presented. Scanning electron microscopy with energy dispersive X-ray spectroscopy was used for the characterization of M. spicatum and its beads. The study showed that aluminium, magnesium and strontium were adsorbed by MsAlg in the water samples from all examined lakes; barium and iron in the water samples from six lakes. The overall average efficiency of MsAlg in biosorption of elements was in the following order: Al > Ba > Sr > Fe > Mg (58.6, 51.7, 48.2, 23.9 and 17.7%, respectively). The increase of TH and WQI values after the biosorption was noticed in all studied lake water samples. The most significant correlations for pH were regarding the contents of B, Mg and Ca, whereas WQI was highly correlated to the contents of B and Mg, and pH. The complexity of the obtained data was explained by Cluster Analysis and Principal Component Analysis, which showed good discrimination capabilities between the water samples taken from different locations. Considering that the invasive M. spicatum is natural, widespread and that its immobilization is cheap and eco-friendly, presented findings could be helpful in further assessment of MsAlg beads for its potential use as biofilter.

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1. Introduction

Water is the basis for a healthy life and is directly related to human survival. So, the awareness of the limited amount of unpolluted water available to mankind is on the rise. Anthropogenic disturbances of ambient groundwater flows (ex., by agricultural, mining and industrial activities, urbanization, and/or climate change) which could cause production of different contaminants (highly toxic heavy metals or non-toxic bio-degradable materials)

greatly affect the quality of water (Kostić et al., 2016; Azizian et al., 2017). Special attention should be paid to surface water quality, as the understanding of raw water quality is equally important to humans as well as to aquatic life (Sener et al., 2017).

Various techniques (precipitation, ion—exchange, adsorption, and reverse osmosis etc.) have been utilized for the toxic elements removal from aquatic environment (Cheng et al., 2017). The methodology of the biosorption study which is based on the interdisciplinary approach of chemists, (micro)biologists and (process)engineers has emerged as an economically feasible alternative for the removal of toxic elements utilizing naturally abundant, waste biomass and immobilized biomass in a polymer matrix (Kratochvil and Volesky, 1998; Volesky, 2007).

Eurasian water milfoil Myriophyllum spicatum, grows on five

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continents and may be native and invasive plant species (Couch and Nelson, 1985). This aquatic weed can be found in various aquatic habitats such as: lakes, rivers, reservoirs, freshwater and brackish estuaries (Aiken et al., 1979). M. spicatum reproduces primarily by vegetative fragmentation and forms dense beds. Therefore, milfoil becomes a human nuisance and has adverse effects on native aquatic vegetation, decreases dissolved oxygen, reduces open areas along lake shores, creates habitats for disease-carrying insects, reduces water flow, irrigation ditches, canals, farm ponds, and irrigation equipment by clogging (Bates et al., 1985). Methods for growth control of this weed are: physical (harvesting), chemical and biological (Milojković et al., 2014). In accordance with requirements of sustainability, managing and processing of aquaticwaste need to be integrated and organized (Milojković et al., 2016a). Alginate as seaweed (brown algae) extract is a frequently used polymer for immobilization and microencapsulation technique (Taqieddin and Amiji, 2004). Calcium alginate beads are widely used as a supporting material in studies of biosorbent immobilization. They exhibit good biocompatibility and are easy to prepare. In addition, they have low prices and are easily available (Zhou et al., 2010). The alginate-immobilized Myriophyllum spicatum beads (Ms:Alginate 5:1) in biosorption tests of Pb(II) under laboratory conditions were recently described. Those M. spicatum beads achieved Pb(II) uptake of 200 mg/g (Milojković et al., 2016a).

Cluster analysis (CA) was performed to classify the samples of water. In this study, complete linkage was used and City-block (Manhattan) distance was calculated in cluster analysis. Principal Component Analysis (PCA) is a mathematical procedure used widely in chemometric data analysis, representing a multivariate technique (Tan and Lu, 2015; Mezzelani et al., 2016). PCA was performed by Eigenvalue decomposition of the correlation matrix of the obtained data (Olenycz et al., 2015; Touahri et al., 2016). PCA was applied to analyse the similarities of the samples, and also the trace element content (Pain-Devin et al., 2014; Bolotov et al., 2015; Robinson et al., 2017). The main idea for using PCA was the reduction of the number of variables, called the principal components (PCs). PCs explain the major variations within the data to make the components more interpretable (Lehtonen et al., 2016). The first two PCs were extracted and utilized in two dimensional bivariate plots; loadings were considered to evaluate correlations between variables. Prior to PCA and CA, the data sets were examined for outliers. Data were analysed by Statistica software (Data Analysis Software System, v.10.0, StatSoft, Inc, Tulsa, OK, USA).

Further investigations of *Myriophyllum spicatum* beads as a biosorbent are presented in this study with the aim to assess the performance of immobilized M. spicatum with alginate—MsAlg in a real system. For this purpose, we used water from nine accumulations: Bovan, Vlasina, Prvonek, Garaši, Ćelije, Vrutci, Sava, Gruža and Srebrno, which are located in the regions: Šumadija, Western Serbia, South and East Serbia. In this preliminary biosorption study with MsAlg beads in the multi-element system of lake water, biosorption capacity (q), element removal efficiency (R), total hardness (TH) and the quality index (WQI) of water before and after biosorption were determined. The multivariate composition of the examined samples was evaluated, and PCA and CA were applied to characterize and discriminate examined water samples. To the best of our knowledge, the current study presents the first data on the characterization and output of MsAlg in a real system (lake water).

2. Materials and methods

The studied lakes (Fig. 1) are located in the area of about 88 361 km² (US *Demographic yearbook*, 2012). Geographical coordinates and elevations of the lakes are: Vlasina—42° 42′ N/22° 20′ E, 1213 m (semi-artificial); Bovan—43° 39′ N/21° 42′ E, 250 m;

Prvonek-42° 30′ N/22° 05′ E. 580 m: Garaši-44° 17′ N/20° 28′ E. 268 m; Ćelije-43° 24′ N/21° 10′ E, 426 m; Vrutci-43° 50′ N/19° 41′ E, 620 m; Sava-44° 47′ N/20° 23′ E, 73 m; Gruža-43° 54′ N/20° 41′ E, 267 m; Srebrno -44° 45' N/21 $^{\circ}$ 25' E, 70 m (artificial lakes). All the lakes belong to freshwater lakes. Hydrological data and intended use of the lakes are presented in Table S1 (Supplementary material). Lakes Sava and Srebrno are parts of the biggest rivers in Serbia: Srebrno (located near Veliko Gradište) was formed as an oxbow lake on the right bank of the Danube, whilst Sava (in Belgrade city) was formed from the right inlet of the Sava river by building two damns. These lakes are with the lowest surface elevation amongst studied lakes and are predominantly used for recreation. Lakes Bovan, Vlasina, Prvonek, Vrutci, Gruža, Ćelije and Garaši were built for water supply purpose, and for such reason they play a fundamental role in local society as sources of drinking water. Moreover, there are additional uses of lakes for energy supply (Vlasina), flood defense (Bovan and Ćelije) and fishing (Garaši) (Anonymous, 1986, 2001). In addition, lakes Sava, Vlasina, Gruža and Srebrno are natural aquatic surroundings for Myriophyllum spicatum.

2.1. Preparation of Alginate—M. spicatum beads (MsAlg)

The aquatic weed *M. spicatum* used for the preparation of beads is from Sava Lake where it is regularly harvested with underwater mower by the public company "Ada Ciganlija", Belgrade Serbia. Since this lake area is intended for recreation and water supply the growth and spread of this water weed must be controlled. The daily amount of harvested biomass is about 30–32 m³. Fresh plant material was first washed with tap water and then 3 times with distilled water. After washing M. spicatum was dried at 35 °C. Dried plant was milled and sieved to particles <0.2 mm. Alginic acid sodium salt from brown algae-low viscosity was used (A1112, Sigma-Aldrich, St. Louis, MO, USA). M. spicatum beads were made according to the method Yan and Viraraghavan (2001). Dissolving Naalginate in distilled water a 2% polymer solution was made. After homogenization a certain amount of biomass powder (4 g in 100 ml) was added (Ms:Alginate 2:1). This slurry was then dispersed dropwise into a 0.1 M calcium chloride solution using a syringe. Therefore, soluble sodium alginate was transformed to solid calcium alginate beads. Having stayed for 24 h in 2% calcium chloride solution, beads were hardened. Those that floated on the surface were unsuitable and discarded. Then the beads were placed into the conical flasks with deionised water and mixed for 30 min. This washing process was repeated 5 times. Beads were dried at room temperature to constant weight and then used for experiments.

2.2. Surface microstructures of MsAlg

Dried *M. spicatum* and MsAlg samples coated with gold were examined on Scanning Electron Microscopy (SEM) instrument (JEOL JSM — 6610LV SEM model). Some statistical quantities were obtained by analysing the SEM figures in Gwyddion program (http://gwyddion.net/documentation/user-guide-en/statistical-analysis.html), including a few basic properties of the height value distribution, such as the mean value and median of *M. spicatum* pores.

2.3. Biosorption tests

The biosorption experiments were performed with samples taken from nine freshwater lakes. Samples were collected following ISO 5667-14 standard during autumn of 2016. For each lake, samples were collected in sterile bottles from three sampling points, and mixed together to make a composite sample before being used

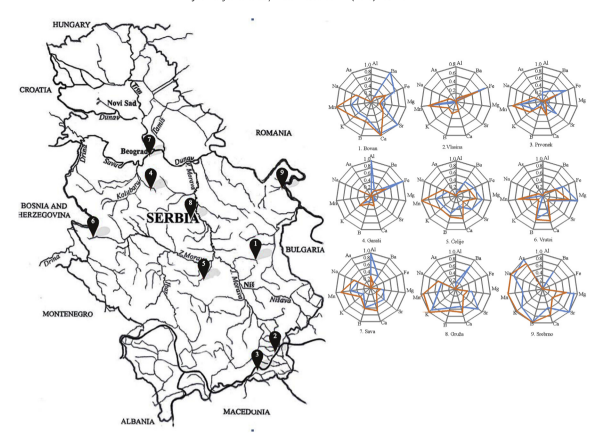


Fig. 1. Geographical locations of the lakes and the radar diagrams of the initial C_i and final element concentrations C_f (before and after biosorption) from nine freshwater lakes.

for further experimental work. Filtration of water samples was done. Subsequently, bottles with composite water samples were immediately placed in an icebox at a temperature of 4 °C, and transported to the laboratory for the following analysis on the same day of sampling. Analytical technique inductively coupled plasma—optical emission spectrometry (ICP—OES) was used for determination of twenty-two elements (ICP—OES Spectroblue equipped with Spectro Smart Analyzer data processing software—SPECTRO Analytical Instruments GmbH, Germany). US EPA (Method 200.7) was applied (1994). Method performance data are given in Table S2 (Supplementary material).

Usually in biosorption experiments with fresh tissues of *M. spicatum*, the amount of applied fresh tissues is 2 g/L calculated on dry weight. According to that, the same amount of biosorbent MsAlg was used. Beads were inserted in the samples of lake water (50 ml) in glass erlenmeyers (100 ml). The flasks were stirred at 200 rpm in a Heidolphunimax 1010 orbital shaker at room temperature (25 °C). All sorption experiments were performed in three replications. Initial and final (after specified contact time) pH values of the mixtures were measured (by a precise pH meter Sension MM340). Contact time was 24 h.

Biosorption capacity was determined from the expression:

$$q = \frac{V(C_i - C_f)}{m} \tag{1}$$

where biosorption capacity q is the amount of various elements removed by investigated biosorbent at any time (mg/g), V is the volume of solution (L), C_i and C_f are the initial and the final element concentrations (mg/L), respectively and m is the mass of the biosorbent MsAlg (g).

Element removal efficiency (*R*) was calculated from equation (2):

$$R = \frac{C_i - C_f}{C_i} \cdot 100 \tag{2}$$

Total Hardness (TH), before and after biosorption process, was calculated (Aquion, 2016) as follows:

$$TH \approx [Ca] + [Mg] \tag{3}$$

TH in °d =
$$0.1339 \times [Ca \text{ in mg/L}] + 0.2307 \times [Mg \text{ in mg/L}]$$
 (4)

The water quality index (WQI) was calculated, according to equation (5):

$$WQI = \frac{\sum_{i=1}^{n} W_i \cdot C_i}{n} \tag{5}$$

Where: *C* is element concentration, *W*—weight coefficient, i—element number, n—number of elements (Chang et al., 2001; Zhaoyong et al., 2015).

2.4. Data analysis

Cluster analysis (CA) was performed to classify the samples of water. In this study, complete linkage was used and City-block (Manhattan) distance was calculated in cluster analysis. Principal Component Analysis (PCA) is a mathematical procedure used widely in chemometric data analysis, representing a multivariate technique (Tan and Lu, 2015; Mezzelani et al., 2016). PCA was performed by Eigenvalue decomposition of the correlation matrix

of the obtained data (Olenycz et al., 2015; Touahri et al., 2016; Lehtonen et al., 2016). PCA was applied to analyse the similarities of the samples, and also the trace element content (Pain-Devin et al., 2014; Bolotov et al., 2015; Robinson et al., 2017). The first three principal components (PCs) were extracted and utilized in two dimensional bivariate plots; loadings were considered to evaluate correlations between variables. Data were analysed by Statistica software (Data Analysis Software System, v.10.0, StatSoft, Inc, Tulsa, OK, USA).

3. Results and discussion

3.1. SEM-EDX analysis

Scanning Electron Microscopy shows the surface texture and porosity of Myriophyllum spicatum and MsAlg (Fig. 2). White parts that can be seen on micrographs may be from the presence of calcium in the samples (Dibdiakova et al., 2015). The observed pore sizes varied between 0.077 and 0.563 µm, while the average and the median values were 0.316 µm and 0.324 µm. The root means square (RMS) value was calculated to explain the height irregularities within each pore (this value is computed from SEM data variance). The variations in the height of the *M. spicatum* pores were 0.121 µm. Pore-wise RMS value was determined for each pore (as a contiguous part of the inserted mask) separately, and the variance was calculated from these values. Similar to RMS, the variations in height of each pore were 0.120 µm. The skewness value was also calculated, as the measure of symmetry, and the calculated value (0.073) shows a symmetrical distribution of pore sizes. Kurtosis value was measured as comparison to normal distribution, and the evaluated value (-1.008) showed a good fit for normal distribution. The surface area was computed by simple triangulation of each pore observed in SEM figure. The surface area for *M. spicatum* pore was 0.001 mm². The observed pore sizes for biosorbent MsAlg varied between 0 and 0.796 µm, while the average and the median values were $0.225\,\mu m$ and $0.176\,\mu m$. The RMS value was $0.177\,\mu m$. The variation in height of each pore was $0.150\,\mu m$. The skewness value was 0.836, while Kurtosis value was -0.232. The surface area was 0.001 mm^2 .

The surface of samples *M. spicatum* and MsAlg was tested for the contents of sixteen elements. Elements N, F, Fe, Pb were not detected by EDX analyses on both samples. Also, on the surface of MgAlg, Na, P, Cl, K, Cu were not identified. Amounts of detected elements are presented in Tables within Fig. 2c) and d). Gold was noticed because of sample preparation. Calcium was present more than four times in MsAlg than in aquatic weed due to the process of gelation with CaCl₂.

Identified composition of elements by this instrumental technique was also confirmed by chemical composition of *M. spicatum* and calcium has the largest amount 5% (Milojković et al., 2014). Large proportion of calcium originates from the water environment in which this aquatic weed lives. For the duration of the process of photosynthesis, submerged plants are provided, not only with free CO₂, but also with one from aqueous solution of Ca(HCO₃)₂. This leads to the deposition of insoluble CaCO₃ and its accumulation on the surface of leaves of submerged plants (Stevanović and Janković, 2001). The precipitation of calcite in natural waters can be shown by the reaction:

$$Ca^{2+}+2HCO_3^- \rightleftharpoons CaCO_3\downarrow +CO_2\uparrow +H_2O$$

The moderate level of alkalinity (pH ranging from 8.2 to 8.8) in Sava Lake water favours the formation of calcium carbonate (Milojković et al., 2016b).

3.2. Element content and biosorption process in lake water samples

US EPA has prescribed National Primary Drinking water regulations with public health goal and public health secondary drinking water standards. Chemicals of health significance are described by World Health Organization guidelines (WHO) for drinking water quality in the third edition (2008) and the fourth edition (2011). WHO has also recommended other water quality parameters.

Contents of Al, As, B, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Li, Mg, Mn, Na, Ni, Pb, Sb, Se, Sr, V and Zn were examined in the water samples from studied lakes. Among the examined elements, the priority pollutants such as Cd, Cr, Cu, Ni, Pb, Se and Zn, as well as Co, Li, Sb and V

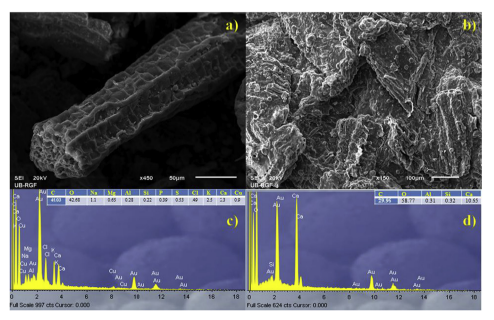


Fig. 2. SEM micrographs of a) M. spicatum b) MsAlg c) EDX analysis of M. spicatum d) EDX analysis of MsAlg.

were below the limit of detection (LOD) in all water samples. The contents of Ca, K, Mg and Na were in the range 8.87-45.44, 0.64-3.14, 1.71-23.81 and 1.96-16.22 mg/L, respectively. Concentrations of detected microelements were in the following ranges: Al $5.00-120.00~\mu g/L$, B $13.00-50.00~\mu g/L$, Ba $6.00-42.00~\mu g/L$, Fe $6.00-11.00~\mu g/L$, Mn $7.00-9.00~\mu g/L$ and Sr $21.00-170.00~\mu g/L$. Arsenic was detected only in Srebrno lake in the concentration of $7.00~\mu g/L$.

Increased amount of arsenic was detected in Srebrno lake water sample (in the concentration 7.00 μ g/L). According to US EPA (2016a) Public Health Goal for this element concentration should be zero. After application of MsAlg, concentration of arsenic decreased to 6.00 μ g/L. Secondary Drinking Water Standards for the amount of aluminium suggested by US EPA (2016b) (0.05–0.2 mg/L) and WHO (0.1 mg/L) was exceeded in samples from lakes Garaši and Sava. After biosorption aluminium content diminished, according to both standards, which approved the utilization of applied biosorption material.

Based on the obtained results, it can be noticed that the concentrations of elements in all studied lakes are within the limit values set by national and European Union (EU) regulations (EPA Ireland, 2001; Official Gazette of Serbia, 2014) (see Table S3, Supplementary material). In terms of chemical quality, water in all studied lakes was within the first category. After the treatment by appropriate methods for water purification this water may be used for drinking water supply (Official Gazette of Serbia, 2014).

Human utilization of biomass or biogenic materials as feedstock isn't new; however at this time there is a renewed interest in efficient exploitation of organic inevitable wastes, in order to decrease eco-footprint and get further stock resources that are renewable (Mohan et al., 2016). The biosorption study showed that Al, Fe, Mg, Sr, Ba and As were absorbed by MsAlg; aluminium, magnesium and strontium in the samples from all studied lakes; barium and iron in the samples from six lakes. Arsenic was absorbed in the water sample from Srebrno lake (the only sample where it was detected).

Based on the results of biosorption, radar diagrams have been plotted and they are presented in Fig. 1. Biosorption capacities (q) of MsAlg in lake water samples are presented in Fig. 3. In lakes with higher contents of certain elements such as strontium and aluminium, the highest biosorption capacity has been achieved.

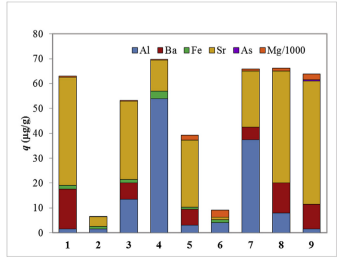


Fig. 3. Biosorption capacity (*q*) of MsAlg for Al, Ba, Fe, Sr, As and Mg in the water of the lakes: 1–Bovan 2–Vlasina 3–Prvonek 4–Garaši 5–Ćelije 6–Vrutci 7–Sava 8–Gruža 9–Srebrno.

Therefore, the maximum biosorption capacity for Al $(54\,\mu g/g)$ and for Sr $(49.5\,\mu g/g)$ was reached in Garaši and Srebrno lake, respectively. Obtained biosorption capacity values for Mg in the studied lakes were in the range from 0.13 to 2.82 mg/g (Fig. 3).

Apart from that, it was observed that boron, calcium, potassium, sodium and manganese were released (increased concentrations after biosorption were measured) in all examined lake water samples. The release of barium was noticed in samples from Vlasina and Garaši lakes.

The biosorption of aluminium was in the range of 21-90%; in five lakes it was higher than 50% (Fig. S1, Supplementary material). In all lake water samples where barium was detected, it was biosorbed in the range of 52-77% (Fig. S2, Supplementary material). The examined biosorbent has removed 25-55% of iron (Fig. S3, Supplementary material). Magnesium was biosorbed in the smallest percent (12-24%) (Fig. S4, Supplementary material) More than 50% of strontium was removed by biosorption from six lake water samples (Fig. S5, Supplementary material), while 14.3% of arsenic was removed from Srebrno lake (the only sample where it was detected). The overall average efficiency of MsAlg in the removal of elements was in the order: Al > Ba > Sr > Fe > Mg (58.6, 51.7, 48.2, 23.9 and 17.7%, respectively). The results point out that the applied biosorbent was efficient in respect to some of the detected elements.

Application *M. spicatum* as biosorbent meets all 4E requirements: i.e. it is easy to cut and collect (Engineering), does not need additional energy for drying (Energy), solves some potential ecological problems (Ecology) and it is low-cost (Economy).

Divalent cation (Ca²⁺) interacts with blocks of acid residues from alginate, resulting in the formation of a 3D network, which is generally explained as "egg-box" model (Grant et al., 1973). After biosorption experiments, MsAlg beads could easily be separated from the lake water samples (by decantation). They didn't change the shape, size and colour during the experiments nor did the mixing rate affect them. Beads were solid and easy to manage. The strength of MsAlg beads can be explained by the formation of "egg-box" model not only with calcium from calcium chloride which was added during the immobilization process, but also with calcium from *M. spicatum* water weed. The increase in the amount of calcium in the studied water samples after biosorption can be explained by probable replacement of calcium from biosorbent and metal ions from the samples according to the "egg-box" model (Mata et al., 2009). Affinity of potentially toxic elements by alginate depends on the extent of guluronic acid and other uronic acids. Carboxyl groups from these acids would be mostly included in metal biosorption (Davis et al., 2004). As suggested in literature, bidentate complexes with one carboxyl group, and ionic bonds (as presented by the "egg-box" model with Ca2+) are two possible ways of bonding between carboxyl groups and M²⁺ cation (Mata et al., 2009). Zooms into single binding zone show these types of bonds (graphical abstract). It may be assumed that elements absorbed by MsAlg in this study undergo a similar pattern, although more complex studies are necessary in order to establish a possible mechanism. According to Datta et al. (2008), the order of alginate affinity towards divalent ions is:

Pb > Cu > Cd > Ba > Sr > Ca > Co, Ni, Zn > Mn.

In our study, calcium release and biosorption of magnesium caused a change in the total hardness (TH). Generally, an increase of TH in all studied samples was noticed (Fig. 4) and the extent of the changes was in the order: Vlasina>Ćelije>Prvonek> Garaši> Sava> Gruža> Bovan> Srebrno> Vrutci. Experimental data on contents of Ca, Mg and TH in the examined water samples from lakes, before and after sorption, can be supported by literature (Mata et al., 2009; Datta et al., 2008).

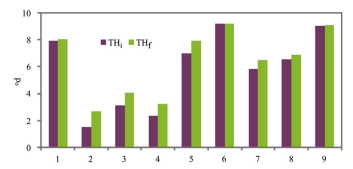


Fig. 4. Total hardness ($^{\circ}$ d): initial TH $_{i}$ and final TH $_{f}$ total hardness before and after biosorption in lake water samples: 1–Bovan 2–Vlasina 3–Prvonek 4–Garaši 5–Ćelije 6–Vrutci 7–Sava 8–Gruža 9–Srebrno.

3.3. Descriptive analysis of elements in lake water samples

The concentrations data set of eleven detected elements: Al, As, B, Ba, Ca, Fe, K, Mg, Mn, Na and Sr as well as pH values for water samples of nine examined lakes were evaluated by PCA in order to determine the average concentrations of elements and perceive their variability. The experimental data and weight coefficients are shown in Table S4 (Supplementary material).

Water quality usually pertains to the structure of the water sample. Evaluations of water quality parameters are necessary for the improvement of water resources management. Water quality index (WQI) is a mathematical tool employed to convert considerable amounts of data on water quality (based on physical, chemical and biological measurements) into a single number that represents the water quality degree (Štambuk-Giljanović, 1999; Jakovljević, 2012).

In order to evaluate changes in water after sorption process with MsAlg, the water quality index (WQI), based on the contents of elements before and after biosorption and pH values (see Table S4, Supplementary material), was accounted for each lake. The obtained results are presented in Fig. 5. It should be underlined that the general trend was observed; the final WQI values (WQI_f), were higher than the initial WQI (WQI_i) values for all studied lakes.

High correlations between observed elements were acquired using correlation analysis for detected elements in water samples. Correlation matrix of element concentrations, pH and WQI values in examined water samples is presented in Table 1. Barium is positively correlated to strontium content, statistically significant at p < 0.01 level. Iron is negatively correlated to B, K and Na content, statistically significant at p < 0.01 level, and also negatively correlated to pH value, statistically significant at p < 0.01 level.

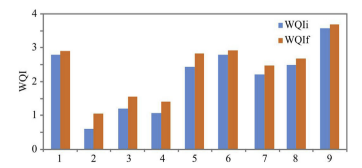


Fig. 5. WQI values of studied water samples from lakes: 1–Bovan 2–Vlasina 3–Prvonek 4–Garaši 5–Ćelije 6–Vrutci 7–Sava 8–Gruža 9–Srebrno; before and after biosorption by MsAlg evaluated by experimental results.

Magnesium is positively correlated to B content and WQI value, statistically significant at p < 0.01 level, while it is also positively correlated to As content and pH value, statistically significant at p < 0.05 level. Calcium is positively correlated to B and Mn content, statistically significant at p < 0.01 level, while it is also positively correlated to K and Na content and pH value, statistically significant at p < 0.05 level. Boron is positively correlated to Na and As content. pH and WQI value, statistically significant at p < 0.01 level, while it is also positively correlated to K content, statistically significant at p < 0.05 level. Potassium is positively correlated to Na content, statistically significant at p < 0.01 level, and also positively correlated to As content and pH value, statistically significant at p < 0.05 level. Sodium is positively correlated to As content, statistically significant at p < 0.01 level, while As content and pH value are positively correlated to WOI value, statistically significant at p < 0.05 level.

3.4. Cluster analysis

Fig. 6 shows the dendrogram of CA for the tested lake water samples. The complete linkage algorithm and City block (Manhattan) distances were used as the measure of proximity between the samples. City block distances (shown on abscissa axis) are measured as the average difference across dimensions of the tested lake water samples.

The resulting dendrogram showed three main clusters, the first cluster contained the samples from lakes Bovan, Sava and Gruža, as well as the sample from lake Ćelije (after biosorption). These samples contained a moderate amount of elements, according to experimental results. The second cluster comprised the samples from Vrutci and Srebrno lakes, as well as the sample taken from lake Ćelije. According to CA, the quality of water sample taken from Ćelije lake increased enough for the transfer to the first cluster, after biosorption. The third cluster contained the samples taken from Vlasina, Pronek and Garaši, which was characterised by the increased amount of Fe and Al, compared to all other samples.

3.5. Principal Component Analysis (PCA)

Principal component investigation was performed to examine the relationships between environmental impacts and trace element concentrations in the observed water samples. The rotation of the obtained principal components was refined by the Varimax method and Kaiser Normalization. PCA analysis of element composition found in the water samples demonstrated that the first three principal components explained 73.09% of the total variance in the original data, Fig. 7. The first principal component (PC1) contributed 45.73%, the second component (PC2) 14.96%, and the third component (PC3) explained 12.40% of the total variance. The calculated first three Eigenvalues were: 5.95, 1.94 and 1.61, respectively. By assessing the PCA graph, the Fe content (which contributed 9.1% of the total variance, based on correlations) gained the most intensive positive score for the first principal component evaluation, while the concentration of Mg (9.2%), Ca (8.7), B (14.8%), K (8.0%), Na (10.7%) and As (7.7%), and the values of pH (9.0%) and WQI (9.2%) demonstrated negative score values of the first principal component (Fig. 5). The contents of Al (which yielded 29.3% of the total variance, based on correlations), Ba (34.4%) and Sr (21.5%) showed positive influence on the second principal component evaluation, whereas the concentration of Mn (23.6%) showed negative influence on the second principal component. The contents of K (which explained 21.3% of the total variance, based on correlations) and Na (13.4%) showed positive influence on the third principal component evaluation, while the content of Mg (24.7%) and WQI value (24.5%) showed negative influence on the third principal component.

Table 1Correlation matrix of element contents, pH and WQI in the water samples from the studied lakes.

	Ва	Fe	Mg	Sr	Ca	В	K	Mn	Na	As	pН	WQI
Al	0.033	0.103	-0.245	-0.027	-0.351	-0.229	-0.263	−0.457 ^c	-0.105	-0.197	-0.056	-0.235
Ba		-0.074	0.175	0.898^{a}	0.225	0.149	0.261	-0.226	0.183	0.087	0.427 ^c	0.183
Fe			-0.364	-0.265	-0.436^{c}	-0.682^{a}	-0.625^{a}	-0.443^{c}	-0.711^{a}	-0.446°	-0.592^{b}	-0.364
Mg				0.385	0.411 ^c	0.824^{a}	0.188	0.218	0.312	0.472 ^b	0.567 ^b	1.000 ^a
Sr					0.381	0.397	0.473	0.032	0.448 ^c	0.379	0.436 ^c	0.392
Ca						0.698^{a}	0.476 ^b	0.601 ^a	0.508 ^b	0.191	0.559 ^b	0.410 ^c
В							0.507 ^b	0.423 ^c	0.701 ^a	0.622^{a}	0.710^{a}	0.824^{a}
K								0.374	0.819^{a}	0.488 ^b	0.322	0.189
Mn									0.332	0.162	0.074	0.213
Na										0.789^{a}	0.453 ^c	0.314
As											0.319	0.472 ^b
pН												0.570 ^b

- a Correlation statistically significant at p < 0.01 level.
- ^b Correlation statistically significant at p < 0.05 level.
- $^{\rm c}$ Correlation statistically significant at p < 0.10 level. Unmarked correlations are not statistically significant.

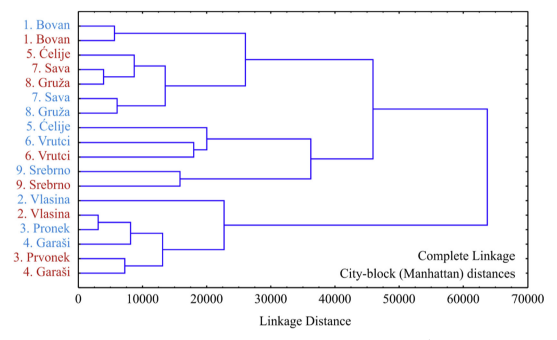


Fig. 6. Dendogram of elements in lake water samples and their locations: 1-Bovan 2-Vlasina 3-Prvonek 4-Garaši 5-Ćelije 6-Vrutci 7-Sava 8-Gruža 9-Srebrno.

As can be seen, there is good partitioning of the nine lakes according to observed elements concentrations, pH and WOI. The initial values of variables are presented by blue dots, while the final values of the observed variables are coloured as red dots in Fig. 7. The influence of different parameters that depicts the observed lake samples could be assessed from the scatter plot (Fig. 7) in which the explored sites with higher Fe content are located at the right side of the PCA scheme (lakes Vlasina, Prvonek, and Gara $\check{s}i-2$, 3 and 4, respectively), site with expanded Mg, As, Na, B, K and Ca content, pH and WQI values (lake Srebrno -9) is situated at the left side of the PCA diagram. The variations in Fe, Mg, As, Na, B, K and Ca concentration, pH and WQI values, after the biosorption with manure can be more effortlessly spotted for water of Vlasina lake, while the higher changes in Al and Mn content are observed for the lakes Prvonek and Garaši. The highest changes in Sr, Ba and Mn content are seen in the water samples from lakes Bovan, Ćelije, Vrutci, Sava, Gruža and Srebrno.

The correlation matrix of sorption properties of biosorbent MsAlg applied for biosorption process in lake water samples is introduced in Table 2. It can be noticed that TH was highly positively correlated with initial pH and biosorption process, statistically significant at p < 0.01 level. The descriptive statistics of sorption characteristics of MsAlg is depicted in Table S5 (Supplementary material).

For visual interpretation of the obtained data trends and for the appropriate discriminating efficiency of the used descriptors a scatter plot of samples using the first two principal components (PCs) from PCA of the data matrix is obtained (Fig. 8).

4. Conclusions

Preliminary assessment of the properties of immobilized *M. spicatum* with alginate—MsAlg in water samples from nine freshwater lakes was done.

• For the first time, the beads of aquatic weed and alginate (Ms:Alginate 2:1 ratio) were applied for biosorption in a real system. The experiment showed that MsAlg absorbed

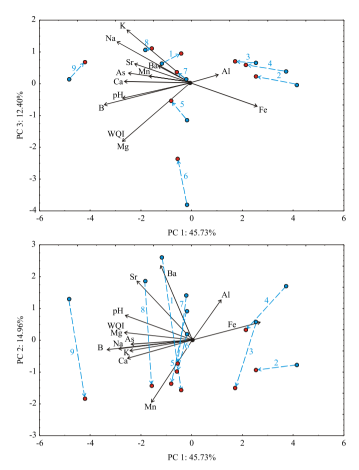


Fig. 7. Biplot of correlated elements in lake water samples and their locations: 1-Bovan 2-Vlasina 3-Prvonek 4-Garaši 5-Ćelije 6-Vrutci 7-Sava 8-Gruža 9-Srebrno.

Table 2 Correlation matrix of sorption characteristics of MsAlg applied for biosorption in lake water samples.

	Biosorption	Leakage	TH
Initial pH Biosorption Leakage	0.528	0.394 -0.497	0.861 ^a 0.830 ^a -0.025

^a Correlation statistically significant at p < 0.01 level; Unmarked correlations are not statistically significant.

aluminium, magnesium and strontium in the water samples from all studied lakes; barium and iron in the water samples from six lakes.

- The average efficacy of MsAlg biosorption ranged from 17.7% for Mg to 58.6% for Al.
- The highest content of Al was observed in water samples from lakes Garaši and Prvonek. After the biosorption with MsAlg its content was decreased. In addition, content of As was reduced in the water sample from Srebrno lake.
- · According to the results of the cluster analysis, the quality of water after biosorption process increased for the sample from lake Ćelije and the purified sample was displaced to a group with better water quality.
- The most noticeable variations in the studied parameters were observed for the sample taken from the lake Vlasina, as shown on PCA diagram, while the highest changes in Al and Mn content after biosorption were observed for samples from lakes Prvonek

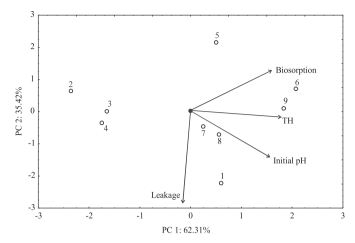


Fig. 8. Biplot of sorption characteristics of MsAlg applied for biosorption in water samples from lakes: 1-Bovan 2-Vlasina 3-Prvonek 4-Garaši 5-Ćelije 6-Vrutci 7-Sava 8-Gruža 9-Srebrno.

and Garaši. The most intensive changes in Sr, Ba and Mn content were observed in the samples from lakes Bovan, Celije, Vrutci, Sava. Gruža and Srebrno.

It can be noticed that TH was highly positively correlated to initial pH and biosorption process. The presented data contribute to a more complete evaluation of MsAlg beads' biosorption characteristics. Additional experiments are required for its potential use as a biofilter (in the treatment or pre-treatment of drinking water with the aim to gain improved quality).

Acknowledgment

Authors acknowledge the Ministry of Education, Science and Technological Development of the Republic of Serbia for financial support (Grant Nos. TR 31003, III 46009 and TR 31055). Jelena Milojković is grateful to the public company "Ada Ciganlija" (Belgrade, Serbia) for providing samples of aquatic weed M. spicatum. Authors also thank D. Popović-Beogračić for the design and processing of the graphical abstract.

Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.watres.2018.05.014.

Conflicts of interest

The authors report no declarations of interest.

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