Using ICP-OES and SEM-EDX in biosorption studies

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Abstract We have compared the analytical results obtained by inductively coupled plasma optical emission spectroscopy (ICP-OES) and by scanning electron microscopy with an energy dispersive X-ray analytical system (SEM-EDX) in order to explore the mechanism of metal ions biosorption by biomass using two independent methods. The marine macroalga Enteromorpha sp. was enriched with Cu(II), Mn (II), Zn(II), and Co(II) ions via biosorption, and the biosorption capacity of alga determined from the solution and biomass composition before and after biosorption process was compared. The first technique was used to analyze the composition of the natural and metal-loaded biomass, and additionally the composition of the solution before and after biosorption. The second technique was used to obtain a picture of the surface of natural and metal ion-loaded macroalgae, to map the elements on the cell wall of dry biomass, and to determine their concentration before and after biosorption. ICP-OES showed a better precision and lower detection limit than EDX, but SEM-EDX gave more information regarding the sample composition of Enteromorpha sp. Both techniques confirmed that biosorption is a surface phenomenon, in which alkali and alkaline earth metal ions were exchanged by metal ions from aqueous solution.

Keywords Biosorption · Microelement ions · Macroalga Enteromorpha sp. · ICP-OES · SEM-EDX

Introduction

Biosorption is a surface phenomenon. This term describes the passive binding of metal ions from aqueous solutions by non-living biomass, which implies that the process is not metabolically controlled [1]. In the literature, biosorption is analyzed with the use of techniques, which determine not the concentration of metal ions bound on the biomass surface, but either the total concentration in the biomass (by analysis of elemental composition of the biomass digested with concentrated mineral acids) or most frequently—from the mass balance, by subtraction of the equilibrium concentration of metal ions in the solution ($C_{eq}$) from their initial concentration ($C_0$) and dividing per the concentration of the biosorbent ($C_X$), using the following formula: $q^S = (C_0 - C_{eq})/C_X$ [2–5].

Denkhaus et al. (2007) in the review entitled ‘Chemical and physical methods for characterisation of biofilms’ presented techniques, which are commonly used for characterization of surfaces and interfaces, e.g. microscopic, spectroscopic and microsensoric [6]. Among them, to the most popular methods, which are used to analyze metal ions both in the biomass and in the solution are as follows: AAS (Atomic Absorption Spectroscopy) [7–9], ICP-OES (Inductively Coupled Plasma with Optical Emission Spectroscopy) [10–12], ICP-MS (Inductively Coupled Plasma
Mass Spectrometry) [13]. Although all the techniques are precise and sensitive, it is impossible to distinguish between metal ions bound by the biomass surface and those accumulated inside the cell. The application of analytical techniques such as: TEM (Transmission Electron Microscopy) [12, 14], Energy-Dispersive X-ray Spectroscopy (EDX) [12, 14] or SEM (Scanning Electron Microscopy) [14–16] enables to determine the elemental composition of the cell wall and to trace the distribution of metal ions on the surface of the cell. This technique was also used to examine the samples of macroalgae: Sargassum vulgaris [17], Sargassum vulgaris [17], Durvillaea potatorum [18].

In the present paper, two techniques were applied: ICP-OES and SEM-EDX in order to understand the mechanism of biosorption of microelement ions (Cu(II), Zn(II), Co(II) and Mn(II)) by marine macroalga Enteromorpha sp. The advantages and disadvantages of each technique are discussed in Fig. 1. Generally, an ICP-OES analysis can detect an average concentration, the SEM-EDX system can analyze an area of the sample as big as the size of the electron beam. In SEM-EDX method, sample preparation is fast and easy compared to the dissolution process necessary for conventional atomic spectroscopy. However, ICP-OES has the advantage of lower detection limits with a higher accuracy, even for the light elements, compared to the SEM-EDX [19].

In the literature it is reported, that biosorption of metal ions by the biomass could occur according to ion-exchange mechanism, whereby acidic functional groups in the biomass exchange protons and/or cations of alkaline earth metals (Mg, Ca, Be, Ba) and alkali metals (Na, K, Li) with metal ions from aqueous solutions [1, 11]. The postulated cation exchange mechanism is probably due to the presence of anionic functional groups on the surface of the cells. The cell wall of Enteromorpha contains a high level of polysaccharides—up to 63% of cell wall dry weight, 13.8% of lipids (total), 9.2% of protein and a small quantity (0.3%) of glucoseamine. These macromolecules offer host of functional groups such as: carboxyl, hydroxyl, imidazole, amino, N-acetyl, phosphate etc. [20].

The aim of the present work was to prove that elements, which are naturally bound by macroalga (Enteromorpha sp.) are exchanged with metal ions from aqueous solution in the biosorption process. For this purpose, ICP-OES method together with SEM-EDX technique were used to evaluate the concentration of elements before and after biosorption in the dry biomass of Enteromorpha sp., to determine the elemental content in the solution before and after process by ICP-OES method and additionally to map elements on the surface of macroagal cell wall by using Scanning Electron Microscope. These two techniques were also useful in identification of the mechanism of biosorption. In this paper, biosorption capacity of macroalga Enteromorpha sp. was determined not only from the mass balance, but also from the analysis of the digested biomass by ICP-OES and also from SEM with mapping. Additionally, the aim of this paper was also to indicate, which value is the most appropriate. Biosorption experiments were
carried out on microelement cations: Cu(II), Zn(II), Co(II) and Mn(II). In order to fully understand this process, it was necessary to carry out mentioned investigations.

**Experimental**

**Organism**

The macroalga *Enteromorpha* sp. was collected from the Baltic Sea (Gdańsk - Brzeźno - Poland) in April 2007. The biomass was washed with tap water several times to remove foreign matter and afterwards with deionized water three times. Then, the biomass was dried at 60 °C until the constant mass was reached.

**Biosorption experiments**

The biosorption experiments were performed in Erlenmeyer flasks, containing 500 mL of microelement ions (Mn(II), Cu(II), Co(II) and Zn(II)) in thermostated water bath shaker at 150 rpm. Initial concentration of each metal ion was 150 mg L\(^{-1}\). The maximum biosorption capacity of the examined metal ions was determined in our previous work [21]. The biomass concentration in the experiments was 1.0 g L\(^{-1}\). The solutions of metal ions were prepared in deionized water (by dissolving appropriate amounts of MnSO\(_4\), CuSO\(_4\), CoSO\(_4\), ZnSO\(_4\) from POCh S.A. Gliwice, Poland, www.english.poch.com.pl). The contact time was evaluated from previous kinetic experiments as 4 h for *Enteromorpha* sp. [22]. pH of the solutions was adjusted to 5 with 0.1 mol L\(^{-1}\) standardized solution NaOH/HCl (from POCh S.A. Gliwice, Poland). pH measurements were conducted with pH-meter Mettler-Toledo—Seven Multi (Greifensee, Switzerland, www.mt.com) equipped with an electrode InLab413 with compensation of temperature. After biosorption process, the suspension of macroalga was filtered through the filter paper and the separated enriched biomass was dried and underwent ICP-OES multielemental analysis and next SEM (EDX).

**Analytical methods**

**Multielemental analysis by ICP-OES**

The samples of natural and the biomass of *Enteromorpha* sp. loaded with microelements (c.a. 0.5 g) were digested with 6 mL of concentrated—69% HNO\(_3\) supra-pure grade from Merck (Darmstadt, Germany) in Teflon vessels with the use of microwave oven Milestone MLS-1200 (Bergamo, Italy). After mineralization, the samples were diluted to 50 g. The mineral content of macroalgae as well as the concentration of elements in the solutions before and after biosorption process were analyzed by Inductively Coupled Plasma-Optical Emission Spectrometer—Varian VISTA-MPX ICP-OES (Victoria, Australia) with ultrasonic nebulizer in the Chemical Laboratory of Multielemental Analyses at Wroclaw University of Technology, which is accredited by ILAC-MRA and Polish Centre for Accreditation (No. AB 696) according to EN-ISO 17025 [23].

**Validation of the method**

For the calibration of the apparatus, the multielemental standard (100 mg L\(^{-1}\) Astasol®, Czech Republic, www.analytika.net) was used. In order to prepare the calibration curve, the following working dilutions of the analytical standard were prepared: 1.0, 10, 50 mg L\(^{-1}\). As a “check standard”, the standard solution—10 mg L\(^{-1}\) was used after each series of 20 samples. The acceptable result was assessed as 10%.

The analytical process was controlled by the use of Certified Reference Material Hard Drinking Water (UK)—metals from LGC Standards (www.lgcstandards.com) (LGC6010) for analysis of solutions and Polish Certified Reference Material for multielement trace analysis Oriental Tobacco Leaves (CTAOTL-1) from Institute of Nuclear Chemistry and Technology (Poland, www.ichtj.waw.pl) for the analysis of digested samples of the biomass. Values of the measurements of the CRMs were within the certified range.

The examined samples were measured in three repeats. The final result was an arithmetic mean, which differed less than 5%.

**Scanning electron microscopy**

Natural and loaded with microelements biomass of *Enteromorpha* sp. was also examined by Scanning Electron Microscopy (SEM). The elemental analysis and mapping was performed at Wroclaw University of Environmental and Life Sciences (Electron Microscope Laboratory). Samples of macroalga were fixed in 2.5% of glutaraldehyde (Sigma, www.sigmaaldrich.com). Then all the samples were dehydrated by ethanol (from 30% till 100% concentration). In the next step macroalga was prepared in two planes for the observation of cross-section and its surface. Samples of the macroalga were mounted on appropriate stub, and thereafter gold-sputtered (using Scan-Coat six equipment—Oxford) and were observed and photographed with a Scanning Electron Microscope—Leo Zeiss 435 VP SEM (Oberkochen, Germany), operating at 20 kV. The microscope was equipped with a RONTEC energy dispersive X-ray system in order to obtain information on elemental composition of the surface of macroagal...
cell wall. The X-ray spectrum of each macroalga loaded with a given microelement was obtained.

**Results and discussion**

Multielemental analysis by ICP-OES

ICP-OES method was used to determine the mineral composition of the natural and loaded with microelements macroalgal biomass in a single-metal system (MA-Mn, MA-Zn, MA-Cu, MA-Co, where MA means macroalga) and the concentration of elements in the solution before and after biosorption process. This technique was used to identify the mechanism of biosorption process and also to indicate differences in the calculation of biosorption capacity of the biosorbent towards metal ions from the mass balance and from the natural and loaded biomass composition.

**Analysis of the solution before and after biosorption process**

On the basis of the composition of the solution it was observed that in the case of the biomass: MA-Cu and MA-Zn, light metal ions were released in the following order: Na(I)>Ca(II)>Mg(II)>K(I). During biosorption of Mn(II) and Co(II) ions, Ca(II) ions were bound by the biomass, and the remaining light metal ions were released as follows: Na(I)>Ca(II)>Mg(II)>K(I) (16,283 mg kg\(^{-1}\)). It is also important to mention that high concentration of Na(I) ion was the main alkali metal, which was exchanged during biosorption process with all the examined microelement ions. These data are in accordance with the results obtained in the analysis of the composition of the solution. In Table 2, the balance sheet of microelement ions in the solution and in the biomass is presented. The average recovery for microelement ions was 84%.

**Calculation of the biosorption capacity**

On the basis of the composition of the solution, it was possible to evaluate the amount of released metal ions and simultaneously ions bound by the biomass (expressed in molar units (meq g\(^{-1}\)), which considers the charge of the ion). The multielemental analysis of the solution after the process revealed the presence of cations that were not detected in the solution in high concentrations before the process. It was found that the higher amount of bound microelements—\(q(b)\), the higher amount of released light metals ions—\(q(r): Na(I), K(I), Mg(II) and Ca(II)\). The equation, which shows this relationship is as follows: \(q(r) = 0.510 \cdot q(b) + 0.980(R 0.918)\).

These data confirmed the hypothesis that ion exchange plays a major role in the binding of metal ions by algal

| Element | Natural biomass | MA-Mn | MA-Zn | MA-Cu | MA-Co |
|---------|----------------|-------|-------|-------|-------|
| K       | 3,920±784      | 285±42.7 | 246±36.9 | 213±31.9 | 280±42.0 |
| Mg      | 8,814±1 763    | 1,630±326 | 1,345±269 | 857±129 | 1,506±301 |
| Ca      | 9,550±1 910    | 2,789±558 | 3,077±615 | 1,556±311 | 3,077±615 |
| Na      | 16,283±3 257   | 590±88.5 | 505±75.7 | 417±62.6 | 1,107±221 |

Table 1 presents the mineral composition of the natural and loaded biomass. All types of the biomass after biosorption contained in the highest amounts Ca(II) ions, then Mg(II) ions, Na(I) ions and finally K(I) ions. After the comparison of the composition of loaded and the natural biomass, it can be concluded that the biomass released light metal ions during biosorption process in the following order: Na(I)>K(I)>Mg(II)>Ca(II) (the average content of Na(I) in the loaded biomass was 25 times lower than in the natural biomass, the content of K(I) 15 times lower, Mg(II) 7 times lower and Ca(II) 4 times). This allows to suppose, that Na(I) ion was the main alkali metal, which was exchanged during biosorption process with all the examined microelement ions. These data are in accordance with the results obtained in the analysis of the composition of the solution. In Table 2, the balance sheet of microelement ions in the solution and in the biomass is presented. The average recovery for microelement ions was 84%.

**Analysis of the natural and loaded biomass**

Table 1 presents the mineral composition of the natural and loaded biomass. All types of the biomass after biosorption contained in the highest amounts Ca(II) ions, then Mg(II) ions, Na(I) ions and finally K(I) ions. After the comparison of the composition of loaded and the natural biomass, it can be concluded that the biomass released light metal ions during biosorption process in the following order: Na(I)>K(I)>Mg(II)>Ca(II) (the average content of Na(I) in the loaded biomass was 25 times lower than in the natural biomass, the content of K(I) 15 times lower, Mg(II) 7 times lower and Ca(II) 4 times). This allows to suppose, that Na(I) ion was the main alkali metal, which was exchanged during biosorption process with all the examined microelement ions. These data are in accordance with the results obtained in the analysis of the composition of the solution. In Table 2, the balance sheet of microelement ions in the solution and in the biomass is presented. The average recovery for microelement ions was 84%.

**Calculation of the biosorption capacity**

On the basis of the composition of the solution, it was possible to evaluate the amount of released metal ions and simultaneously ions bound by the biomass (expressed in molar units (meq g\(^{-1}\)), which considers the charge of the ion). The multielemental analysis of the solution after the process revealed the presence of cations that were not detected in the solution in high concentrations before the process. It was found that the higher amount of bound microelements—\(q(b)\), the higher amount of released light metals ions—\(q(r): Na(I), K(I), Mg(II) and Ca(II)\). The equation, which shows this relationship is as follows: \(q(r) = 0.510 \cdot q(b) + 0.980(R 0.918)\).

These data confirmed the hypothesis that ion exchange plays a major role in the binding of metal ions by algal

| Element | The composition of the natural and loaded biomass (mg kg\(^{-1}\) of dry mass) |
|---------|---------------------------------------------------------------------|
| Natural biomass | MA-Mn | MA-Zn | MA-Cu | MA-Co |
| K       | 3,920±784      | 285±42.7 | 246±36.9 | 213±31.9 | 280±42.0 |
| Mg      | 8,814±1 763    | 1,630±326 | 1,345±269 | 857±129 | 1,506±301 |
| Ca      | 9,550±1 910    | 2,789±558 | 3,077±615 | 1,556±311 | 3,077±615 |
| Na      | 16,283±3 257   | 590±88.5 | 505±75.7 | 417±62.6 | 1,107±221 |

Table 1 The concentration of light metal ions in the natural and loaded biomass by ICP-OES in digested samples of the biomass.
Table 2  The balance of microelement ions in 1.0 L of the solution before and after biosorption process (for biomass concentration 1.0 g L\(^{-1}\))

| Element | Before biosorption process | | | After biosorption process | | | Recovery (%) |
|---------|---------------------------|-----------------|-----------------|---------------------------|-----------------|-----------------|---------------|
|         | Solution                  | Biomass         | ∑               | Solution                  | Biomass         | ∑               |               |
| Mn      | 104±16 mg                 | 0.0302±0.0076 mg| 104±16          | 79.1±11.9 mg              | 18.2±2.7 mg     | 97.3±12.2 mg   | 93.6%         |
| Zn      | 163±25 mg                 | 0.0680±0.0170 mg| 163±25          | 94.1±14.1 mg              | 31.0±4.7 mg     | 125±15          | 76.7%         |
| Cu      | 215±32 mg                 | 0.0200±0.0050 mg| 215±32          | 132±20 mg                 | 34.5±5.2 mg     | 167±21          | 77.4%         |
| Co      | 118±18 mg                 | 0.0010±0.0002 mg| 118±18          | 76.8±11.5 mg              | 27.9±4.2 mg     | 105±12          | 88.7%         |

Fig. 2  SEM images of: A natural biomass; B MA-Mn; C MA-Zn; D MA-Cu; E MA-Co (SEM, Leo Zeiss 435), where: \(WD\) working distance, \(EHT\) extra high tension, \(SE1\) secondary electrons (Signal A), \(Mag\) magnification. Mag = 1.70KX (1700 magnification)
biomass [3]. On the basis of the composition of the solution it can be concluded that 51% more light metal ions were released than bound microelement ions. This relationship was compared with the data obtained from the analysis of the digested biomass. It was found that Enteromorpha sp. released during biosorption 24% more light metal ions than bound microelement ions: 
\[ q(r) = 0.244 \cdot q(b) + 1.51 \] 
These values confirmed, that there is a difference in the approach to calculation of biosorption capacity by the two methods. Higher correlation coefficient was obtained in the case of the analysis of the solution composition (\( R = 0.918 \)) than in the case of the biomass composition (\( R = 0.608 \)). In the next step, biosorption capacity determined from the solution (\( q_{(ICP)}^{S} \), mg g\(^{-1}\) or meq g\(^{-1}\)) and from the composition of the digested biomass (\( q_{(ICP)}^{X} \), mg g\(^{-1}\) or meq g\(^{-1}\)) was compared. The equation, which describes this relationship is as follows: 
\[ q_{(ICP)}^{X} = 0.250 \cdot q_{(ICP)}^{S} + 14.2 \] 
The values of \( q_{(ICP)}^{S} \) for Cu(II), Mn(II), Zn(II) and Co(II) cations were higher (averagely 1.87±0.52) than the values of \( q_{(ICP)}^{X} \). However, the same tendency in the increase of \( q \) for microelement cations was obtained \( q_{Cu(II)} > q_{Zn(II)} > q_{Co(II)} > q_{Mn(II)} \).

Scanning electron microscopy

Scanning Electron Microscopy was used to show macro- and ultrastructural changes of the surface of dry biomass of

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**Fig. 3** X-ray spectrum, mineral trace analysis, RONTEC (SEM, Leo Zeiss 435)—A natural biomass; B MA-Mn; C MA-Zn; D MA-Cu; E MA-Co
Enteromorpha sp. before and after biosorption process (Fig. 2). General morphological changes of the shape were observed in the case of MA-Mn, MA-Zn, MA-Cu and MA-Co when compared to the natural biomass. SEM electronographs showed detail changes of ultrastructure of macroalgae loaded with microelement ions. All tested macroalgae, except of the natural biomass, were characterized with considerable shrink and possess a tendency to wrinkle the cell wall. Similar changes were observed by Raize et al. (2004) in the cell wall matrix of Sargassum biomass. Furthermore, breaking of the continuity of the morphological structure and free area between intracellular spaces were observed [17]. It could suggest that different microelement ions were responsible for larger or smaller destruction of the structure by binding ions on the surface.

RONTEC energy dispersive X-ray system (Fig. 3) gave a visible evidence of binding microelement ions on the cell wall of macroalgae. Mapping images clearly showed that Mn(II), Zn(II), Cu(II), Co(II) ions were sorbed on the surface of Enteromorpha sp. after biosorption (Fig. 4). Specific localization of microelement ions on the surface of macroalgae was observed. All the tested macroalgae showed qualitative and quantitative changes in ions binding to the surface. Different concentration and localization of ions were indicated by appropriate colors. In the case of the natural biomass, the distribution of ions on the surface was uniform (Fig. 4A). On the surface of macroalga loaded with

Fig. 4 X-ray mapping of SEM images, mineral trace analysis, RONTEC (SEM, Leo Zeiss 435)—A natural biomass; B MA-Mn; C MA-Zn; D MA-Cu; E MA-Co. Where: C—blue, S—orange, Ca—green, Al—yellow, Si—brown, A natural biomass—Mg—purple, Cl—red; B MA-Mn—Mn—violet; C MA-Zn—Zn—green, Mg—bright green; D MA-Cu—Cu—red, P—bright green; E MA-Co—Co—red
Mn(II), Zn(II), Cu(II) and Co(II) cations (Fig. 4B–E, respectively), overbalance of these ions was found. It is visible as dense clusters giving glaring and intensive color.

Table 3 shows the atomic concentrations of Mn(II), Zn(II), Cu(II), Co(II) in the different metal-laden Enteromorpha sp. samples, according to the analysis of X-ray spectrum. The changes in algal biomass observed after metal uptake included generally increases in carbon concentrations (with exception of MA-Cu) and decreases in sulfur (with exception of MA-Zn), oxygen (with exception of MA-Cu), calcium and magnesium (with exception of MA-Zn). These data are in accordance with the results obtained by Raize et al. 2004, who observed similar changes in C, S, O, Ca and Mg concentration after biosorption of heavy metals (Cd, Ni, Pb) by Sargassum [17]. In the case of Enteromorpha sp. the biggest changes concerned the concentration of oxygen, since, as it was previously shown [21], carboxyl groups play the dominant role in biosorption process (oxygen was probably covered by microelement ions).

For the natural biomass of Enteromorpha sp.—Fig. 3A showed that C, O, S, Cl and Ca constituted the major elements of the surface of macroalgal cell, and Mn, Zn, Cu and Co signals were not detected in the X-ray spectrum. After biosorption of Mn(II), Zn(II), Cu(II), Co(II) ions signals were clearly observed in Fig. 3B–E, respectively. This means that a remarkable amount of microelements was adsorbed by the cell surface of Enteromorpha sp.

Table 4 Correlation matrix between the results obtained by ICP-OES and SEM-EDX from the biosorption of Mn(II), Co(II), Zn(II) and Cu(II) ions, (N=4)

| Element | C² LAD(ICP) | C² LAD(SEM) | q⁸ (ICP) | q² X (ICP) |
|---------|-------------|-------------|----------|------------|
| MA-natural | 1.00 | 0.681 | 0.941** | 0.874 |
| MA-Mn | | 1.00 | | |
| MA-Zn | | 0.941** | 1.00 | |
| MA-Cu | | 0.874 | | |
| MA-Co | | | | 1.00* |

Correlation coefficient is significant: *at p<0.05; ** at p<0.1
Where:

- C² LAD(ICP) the concentration of elements in the metal-laden biomass, mg kg⁻¹
- C² LAD(SEM) the atomic concentration of elements on the surface of the metal-laden biomass, %

The comparison of biosorption performance by ICP-OES and SEM

The aim of this section was to compare data obtained by ICP-OES and SEM-EDX, in order to point out the best method of evaluation of biosorption performance. In order to find the correlations between the results obtained by both methods, a correlation matrix was prepared (Table 4) with the STATISTICA (v.8) software. The application of the statistical analysis revealed statistically significant correlation:

Table 3 Atomic concentration of elements (%) on the surface of natural and metal-laden biomass of Enteromorpha sp. (according to the analysis of X-ray spectrum)

| Element | Atomic concentration of elements (%) of all detected ions |
|---------|--------------------------------------------------------|
| MA-natural | MA-Mn | MA-Zn | MA-Cu | MA-Co |
| Macromolecules | | | | | |
| C | 50.4±8.27 | 53.7±9.02 | 54.3±9.64 | 35.4±6.71 | 61.0±9.97 |
| O | 40.9±6.82 | 33.5±5.95 | 30.1±6.00 | 47.6±8.61 | 31.7±5.53 |
| P | 0.63±0.06 | <LLD | <LLD | 0.46±0.08 | 0.87±0.08 |
| S | 2.12±0.10 | 1.98±0.12 | 5.31±0.30 | 0.50±0.08 | 1.44±0.10 |
| Cl | 1.58±0.10 | <LLD | <LLD | <LLD | <LLD |
| Microelements | | | | | |
| Fe | <LLD | 0.07±0.08 | 0.80±0.21 | <LLD | 0.04±0.06 |
| Cu | <LLD | 0.05±0.08 | <LLD | 5.27±0.51 | <LLD |
| Mn | <LLD | 0.11±0.10 | <LLD | <LLD | <LLD |
| Zn | <LLD | <LLD | 0.64±0.27 | <LLD | <LLD |
| Co | <LLD | <LLD | <LLD | <LLD | 0.09±0.08 |
| Alkali and alkaline earth metals | | | | | |
| K | 0.30±0.07 | <LLD | <LLD | <LLD | 0.31±0.08 |
| Ca | 1.16±0.10 | 0.48±0.08 | 1.54±0.16 | 0.23±0.06 | 0.25±0.06 |
| Mg | 0.62±0.05 | <LLD | 1.56±0.12 | 0.11±0.07 | 0.19±0.05 |
| Na | 0.746±0.06 | <LLD | <LLD | <LLD | 0.29±0.05 |

<LLD below low limit of detection

Italic below/or above limit of detection (%): C (0.277–100); O (0.525–100); P (0.001–2.013); S (0.001–2.307); Cl (0.001–2.621); Fe (0.060–6.398); Cu (0.083–8.040); Mn (0.063–5.894); Zn (0.001–8.630); Co (0.076–6.924); K (0.001–3.312); Ca (0.341–3.690); Mg (0.001–1.553); Na (0.001–1.041)
at $p<0.1$ $q^S_{(ICP)} - C_{LAD,(ICP)}^X$ ($R = 0.941$), $q^X_{(ICP)} - q^S_{(ICP)}$ ($R = 0.940$) and at $p<0.05$ $q^X_{(ICP)} - C_{LAD,(ICP)}^X$ ($R = 1.00$).

On the basis of the conducted experiments, it can be concluded that traditionally used method of evaluation of biosorption capacity (from mass balance) is more appropriate than from the composition of the natural and loaded biomass. Relatively low value of correlation coefficient for the pair: $C_{LAD,(SEM)}^X - C_{LAD,(ICP)}^X$ ($R = 0.681$) can be explained by the fact, that $C_{LAD,(ICP)}^X$ concerned the content of the microelements both—inside and outside the biomass, whereas $C_{LAD,(SEM)}^X$ only content of the elements on the cell surface after biosorption process. Moreover, EDX in conjunction with SEM is a surface analysis method and furthermore, for small concentrations this system is not very precise and the detection limit is dependent on the matrix and its surface [24]. The strong variation of the EDX results in different sample areas is the consequence of a typical behaviour of the SEM-EDX, it is surface analysis, and shows therefore only a part of the whole analytical information, this is in total contrast to conventional ICP-OES, which shows average results for every sample [19].

**Conclusion**

It is necessary to combine different methods to explore the mechanisms of metal biosorption. ICP method is especially valuable in determination of the mineral content of the whole (outside and inside) biological samples, whereas SEM technique is useful in detection of elements on the surface of samples. Moreover, it is possible to map the elements on the cell wall of the biomass and to observe morphological changes in the cell wall matrix.

Both techniques confirmed that biosorption is a surface phenomenon, in which alkali and alkaline earth metals were exchanged with microelement ions from the aqueous solution. ICP method revealed that Na(I) was the main alkali metal, which was exchanged during the biosorption process. It was released in the highest quantities from the biomass to the solution, which can be connected with the highest concentration of this element in the natural biomass from among other examined elements (Ca(II), Mg(II), K (I)). It was also found that the higher quantity of released light metals ions - $q(I)$: Na(I), K(I), Mg(II) and Ca(II), the higher quantity of bound microelement ion - $q(b)$.

The experiments showed that there is a difference in the value of biosorption capacity calculated from the mass balance and from the biomass composition. The $q$ values determined from the first method were higher than from the second—averagely $1.87\pm0.52$. However, the same tendency in the increase of $q$ for microelement ions was obtained $q_{Cu(II)} > q_{Zn(II)} > q_{Co(II)} > q_{Mn(II)}$. The correlation coefficient was $0.940$.

The analysis of the biomass by SEM technique showed that all the tested macroalgae, except of the natural biomass, were characterized by considerable shrink and writhe of the cell wall. It could suggest that different microelement ions were responsible for larger or smaller destruction of the structure by binding them on the surface. RONTEC energy dispersive X-ray system gave a visible evidence of binding microelement ions on the cell wall of macroalgae. Mapping images clearly showed that Mn(II), Zn(II), Cu(II), Co(II) ions were sorbed on the surface of *Enteromorpha* sp. after the process of enrichment.

The final conclusion of our work is that the most appropriate way is to analyze the samples with both systems: first qualitative and quantitative overview by SEM-EDX and then a precise quantification of the results by ICP-OES.

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