Influence of hydrogen cations on kinetics and equilibria of heavy-metal sorption by algae—sorption of copper cations by the alga *Palmaria palmata* (Linnaeus) Weber & Mohr (Rhodophyta)

Andrzej Kłos · Malgorzata Rajfur

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**Abstract** The influence of hydrogen cations on kinetics and equilibria of sorption of copper cations by the marine alga *Palmaria palmata* (Linnaeus) Weber & Mohr was studied under static conditions. The competitive effect of the $H^+$ cations is described, which influenced the uncertainty of evaluation of the alga sorption capacity. Under static conditions, the variation of the $Cu^{2+}/H^+$ concentration ratio during sorption was found nonmonotonic. The Langmuir isotherm model was used to determine the sorption capacity of the alga, namely 12.4 mg g$^{-1}$ of dry algae mass. A similar value was determined from the kinetic parameters of the ionic exchange which is considered a pseudo-second-order chemical reaction. The consistent results indicated that the mathematical models used correctly described the equilibria and kinetics of the ionic exchange between algae and solutions.

**Keywords** Algae · Ionic exchange · Sorption kinetics and equilibria · Copper · pH

**Introduction**

There have been many scientific reports in recent years on equilibria and kinetics of heavy-metal sorption in various kinds of biomass, including algae (Bhat et al. 2008; Feng and Aldrich 2004; Gupta et al. 2010; Gupta and Rostogi 2008a; b; Rajfur et al. 2010), lichens (Hauck and Huneck 2007), mosses (Balarama Krishna et al. 2004) and microorganisms (Vásquez et al. 2007). The main aim of these studies was to evaluate the possibility of using the living organisms or their dead biomass in the remediation processes (Monteiro et al. 2009). In addition, these studies were intended to reveal the influence of the competitive sorption of macroelements on traces of pollutants, such as heavy metals, contained in the organisms used for the biomonitoring of environmental pollution (Kłos et al. 2005; 2007). Similar evaluation has been applied to natural and modified mineral sorbents (Genc-Fuhrman et al. 2007; Wu and Zhou 2009) and to physico-chemically modified biological materials (Wong et al. 2003). The authors generally agree that sorption of heavy metals in biomass is mostly an ionic-exchange process. The algal cell wall consists mainly of polysaccharides, lipids, proteins and alginic acids. In selected species of algae, it is enriched with compounds such as calcium carbonate, chitin (N-acetylglucosamine polymer), silica and alginates (alginic acid salts). It contains 3–5-nm-sized pores, which makes it permeable to small molecule substances, such as water, metal ions and gas substances. The function groups responsible for the addition of heavy metal cations include amino, acetyl, hydroxyl, sulfo, carbonyl and carboxyl groups (Davis et al. 2003).

Mathematical models used to describe this process at equilibrium include the Freundlich isotherm model (Bulgariu and Bulgariu 2012; Kumar et al. 2008; Zhang 2011), the Langmuir model (Bulgariu and Bulgariu 2012; Kumar et al. 2008; Lee and Chang 2011) and the Redlich-Peterson and the Koble-Corrigan models (El-Sikaily et al. 2007). Kinetics of the process is usually described with the first-order reaction model (Lagergren model), the second-order reaction model and the Weber and Morris model (Apiratikul and Pavanant 2008; Pavanant et al. 2006; Quaiser et al. 2009).

An important part of the sorption studies is evaluation of the influence of solution acidity (pH) on the process. Numerous
reports indicate that the maximal sorption of metals takes place in solutions of pH ranging from 4 to 6, depending on the metal sorbed and the sorbent used. In acidic solutions, the sorption of heavy metals can be decreased by the competitive sorption of hydrogen cations (Singh et al. 2008), while in alkaline solutions, by the formation of hydrocomplexes (Uluozlu et al. 2008) and insoluble metal compounds, mainly hydroxides (Tuzen et al. 2009), as well as by the changes of the ionic forms of the metals (Herrero et al. 2005). Laboratory studies of equilibria and kinetics of heavy-metal sorption in algae under static conditions are usually carried out at pH lower than the pH of water in which the algae live naturally. In such experiments, metal cations are sorbed along with hydrogen cations—a fact already recognised (Chen et al. 2008) but often disregarded in discussions. On the other hand, the influence of the initial pH of solutions on the kinetic (Vilar et al. 2005) and equilibrium (Pavasant et al. 2006) parameters of sorption processes has also been indicated. The influence of temperature on sorption also has been studied (Gupta and Rostogi 2008a; Sari and Tuzen 2008).

This study aimed to evaluate: (1) the influence of hydrogen cations on the equilibria and kinetics of sorption of copper cations in the marine alga Palmaria palmata (Linnaeus) Weber & Mohr; (2) the influence thereof on the uncertainty of determination of the alga sorption capacity; and (3) the interdependent parameters describing the equilibria and kinetics of the process.

Materials and methods

The marine alga Palmaria palmata, purchased at Bogutyn Młyn from Radzyń Podlaski (Poland), was used for the research. The thalli of these algae are multicellular, flattened and forked, approximately 20–50 cm long, dark-red (Graham et al. 2009). The species was selected for the research due to its availability and easy preparation. The purchased biomass was in dried form. In order to remove salts released as a result of the destruction of cell membranes (Rajfur et al. 2012), the algae were rinsed with demineralised water (conductivity \( \kappa = 0.5 \, \text{µS cm}^{-1} \)) and dried for 24 h at 50 °C. The algae prepared in this way were stored in tightly closed polyethylene containers. The concentration of copper naturally accumulated in the alga equalled to 0.0033 ±0.0008 mg g\(^{-1}\) d.m. Copper was selected due to a good repeatability of the results with the AAS method.

Study of the sorption equilibria and kinetics

Directly before each experiment, alga samples were immersed in demineralised water for 30 min to remove mineral salts released through broken cell membranes (Rajfur et al. 2012). All experiments were carried out at room temperature. Each sample (1.0 g dry algae mass (DM)) was placed in a perforated container of 30 mL volume and immersed in 0.4000 L CuSO\(_4\) solution. The solution was intensively mixed with a magnetic stirrer.

The frequently reported influence of pH on the sorption of heavy metals was confirmed by placing the prepared alga samples in 0.5 mmol L\(^{-1}\) solutions of copper sulphate of various acidities ranging from pH3 to pH9. The acidity of solutions was stabilised with 0.01 mol L\(^{-1}\) solutions of hydrochloric acid or sodium hydroxide. The sorption was carried out for 70 min, until the equilibrium was reached. After sorption, alga samples were mineralised and analysed for copper content.

Kinetics of sorption was assessed in similar experiments carried out for 70 min until the equilibrium was reached. Samples of the solution were taken periodically and analysed for copper content using AAS. The solution volume decreased upon sampling for about 0.020 L (5 % of the initial volume). Changes of the solution pH were also recorded. The equilibrium parameters obtained were used to determine the Langmuir isotherm (Langmuir 1918).

Analytical equipment and chemicals

Heavy metals were determined with an atomic absorption spectrometer SOLAAAR 969 UNICAM (Thermo Electron Corporation, USA). The spectrometer was calibrated against standards obtained from ANALYTIKA Ltd., Czech Republic. The largest concentration of the calibration solution, namely 5 mg L\(^{-1}\) copper, was assumed an upper limit of linear response of the apparatus. The pH of solutions in which alga samples were immersed was measured with a CP551 pH-meter from Elmetron Zabrze, Poland. The absolute error of readings was \( \Delta \text{pH} = 0.02 \). Alga samples were mineralised with a microwave mineraliser MARS-X from CEM. All chemicals used were purchased from MERCK.

Quality control

The detection and quantification limits for copper equalled 0.0045 and 0.041 mg L\(^{-1}\), respectively. The quality control of measurements was assured by test analyses of the BCR 414 plankton and BCR-482 lichen reference materials from the Institute for Reference Materials and Measurements in Belgium. The obtained results are summarised in Table 1.

The results presented in Table 1 confirm the relevance of the choice of copper as the element applied for the research.

Statistical evaluation of the results

The uncertainty of measurement results was evaluated using: standard errors (SE) of the parameters of straight lines defined by Eqs. (1) and (2); coefficients of
determination $R^2$ for these straight lines; and direct comparison of $c^*K_t(a,1)$ values determined independently within the kinetic and equilibrium analyses. The statistical evaluation was done both in regard to copper cations and to the sum of $H^+$ and $Cu^{2+}$ cations. For statistical computations, the R language (R Development Core Team 2009) was used.

**Results**

For the purpose of detailed comparison of the equilibrium and kinetic parameters of sorption of divalent copper cations and monovalent hydrogen cations, the concentrations thereof were expressed as $c^* = z \cdot c$, where: $z$ is the cation valency (dimensionless) and $c$ is the concentration of cations in mmol L$^{-1}$ or mmol g$^{-1}$.

The transformation made the comparison of gram equivalents of cations (former val unit) possible without violating the rules of the SI system (mol units).

Table 2 shows the concentrations of copper cations $c^*$ determined in solutions (s) and in alga samples (a), before (0) and after (1) sorption, initial and final values of solution pH and the gram equivalent fractions of copper in the initial solutions $x^*_{Cu(s,0)} = c^*_{Cu(s,0)} / (c^*_{Cu(s,0)} + c^*_{H(s,0)})^{-1}$. The initial concentrations of hydrogen and copper ions in alga samples were assumed null.

Preliminary evaluation of results showed that increasing of the initial gram equivalent fraction of copper in solutions of equal initial pH reduced the competitive sorption of hydrogen cations. Thus, the sum of sorbed hydrogen and copper cations should be used for determination of sorption capacity of algae under assumption that $c^*_{Cu(s,0)} \rightarrow \infty$.

**Competitive sorption of hydrogen ions**

Plots in Fig. 1 show how the quotients of copper and hydrogen concentrations in algae and in solution changed in time during sorption (the measurement series 6, Table 2).

The variation of the concentration quotient of copper and hydrogen in solutions was clearly nonmonotonic. In the initial period of sorption, the quotient $c^*_{Cu(s,t)}(c^*_{H(s,t)})^{-1}$ in the solution increased and so did the quotient in algae, $c^*_{Cu(a,t)}(c^*_{H(a,t)})^{-1}$. It was possible if the increase of copper concentration in algae exceeded that of hydrogen concentration: $dc_{Cu(a,t)}(dt) > dc_{H(a,t)}(dt)$; and if the decrease of copper concentration in solution relative to the initial concentration of copper in solution was smaller than the similar decrease of hydrogen concentration in solution: $-dc_{Cu(s,t)}(c_{Cu(s,0)} dt) < -dc_{H(s,t)}(c_{H(s,0)} dt)$. After the maximum value of $c^*_{Cu(s,t)}(c^*_{H(s,t)})^{-1}$ was reached (point A in Fig. 1), the conditions changed to: $dc_{Cu(s,t)}(dt) > dc_{H(s,t)}(dt)$ and $-dc_{Cu(s,t)}(c_{Cu(s,0)} dt) > -dc_{H(s,t)}(c_{H(s,0)} dt)$. Thus, the sorption of copper ions in algae had two phases—the first one which lasted until the relative equilibrium of chemical potentials of $H^+$ and $Cu^{2+}$ in

| Measurement series | $c^*_{Cu(a,0)}$ mmol L$^{-1}$ | $c^*_{Cu(s,1)}$ mmol L$^{-1}$ | $c^*_{Cu(s,1)}$ mmol g$^{-1}$ | $pH_0$ | $pH_1$ | $x^*_{Cu(s,0)}$ |
|-------------------|-------------------------------|-------------------------------|-------------------------------|-------|-------|---------------|
| 1                 | 0.0346                        | 0.0016                        | 0.0132                        | 4.73 ±0.05 | 6.04 | 0.65 |
| 2                 | 0.0463                        | 0.0025                        | 0.0175                        | 5.96 | 0.71 |
| 3                 | 0.0724                        | 0.0031                        | 0.0277                        | 5.96 | 0.80 |
| 4                 | 0.0998                        | 0.0047                        | 0.0380                        | 5.73 | 0.84 |
| 5                 | 0.1309                        | 0.0057                        | 0.0501                        | 5.68 | 0.88 |
| 6                 | 0.1520                        | 0.0076                        | 0.0578                        | 5.60 | 0.89 |
| 7                 | 0.0803                        | 0.0038                        | 0.0306                        | 4.0 | 5.36 | 0.45 |
| 8                 | 0.0815                        | 0.0046                        | 0.0188                        | 3.0 | 3.42 | 0.08 |
solution was reached (point A), and the second one in which the sorption of both cations was mutually proportional and governed by thermodynamic conditions and cation concentrations. The time in which point A was reached increased when the initial pH of solutions was decreased.

To sum up, the pH of solutions influenced not only the equilibria of copper sorption, but also the kinetics of the process, especially in the initial phase.

Figure 2 shows the influence of the gram equivalent fraction of copper in the initial solution, $x^{\text{Cu(s,0)}} = c^{\text{Cu(s,0)}} (c^{\text{H(s,0)}} + c^{\text{H(s,0)}})^{-1}$, on the magnitude of the concentration quotient at point A.

The relation in Fig. 1 is explainable. If $x^{\text{Cu(s,0)}} = 0$, there are no copper cations in the solution, so $c^{\text{Cu(s,A)}} (c^{\text{H(s,A)}})^{-1} = 0$. If $x^{\text{Cu(s,0)}} \to 1$, then $c^{\text{Cu(s,A)}} (c^{\text{H(s,0)}})^{-1} \to \infty$. However, the relation has an upper limit resulting from the dissociation of water and solubility of metal hydroxides. A metal hydroxide precipitates when the acidity of solution exceeds the value of pH = $z^{-1} \left(14 - \log c_{\text{Me}} - pK_{\text{sp}}\right)$, where $z$ is the valency of the metal cation; $c_{\text{Me}}$ is the concentration of metal cations in solution (mol/L); and $K_{\text{sp}}$ is the solubility product of metal hydroxide; $pK_{\text{sp}} = -\log(K_{\text{sp}})$.

Sorption kinetics of Cu$^{2+}$ and H$^+$ + Cu$^{2+}$ cations

Kinetics of sorption was described using the model of a pseudo-second-order chemical reaction:

$$t (c^{\text{Kt(at)}})^{-1} = 1 \left(k'' (c^{\text{Kt(a,1)}})^2\right)^{-1} + 1 (c^{\text{Kt(a,1)}})^{-1}$$

where: index Kt denotes the concentration of Cu$^{2+}$ cations or the sum of Cu$^{2+}$ and H$^+$ cations; $k''$ is the rate constant for the pseudo-second reaction. The variable $c^{\text{Kt(a,1)}}$ is the highest concentration of Cu$^{2+}$ cations or the sum of Cu$^{2+}$ and H$^+$ cations absorbed in algae, which are in equilibrium with the respective cations in the solution under given thermodynamic conditions (temperature and reactant concentrations).

Further analysis of sorption kinetics included only the variation of Cu$^{2+}$ concentration and the sum of H$^+$ and Cu$^{2+}$ concentrations in the second phases of measurements (results located right of the A point in Fig. 1). As discussed above, the variation of concentrations observed in the first phases of measurements, left of the A point, was nonlinear due to various proportions of ions sorbed and did not match model (1). The results of our own experiments are shown in Figs 3a and b.

The plots show that the slopes of straight lines describing the sorption of copper cations (Fig. 3a) are much less ordered than the slopes of lines corresponding to the sum of
H\(^+\) and Cu\(^{2+}\) cations (Fig. 3b) and clearly depend on the concentration of hydrogen ions.

Sorption capacity of the algae

Sorption capacity of the algae was determined using the Langmuir isotherm model:

\[
1 \left( \frac{c^*}{K_{t(a,1)}} \right)^{-1} = 1 \left( \frac{c^*_{Kt(a,max)}}{K} \right)^{-1} + 1 \left( \frac{c^*_{Kt(a,max)}K}{1 \left( \frac{c^*_{Kt(a,1)}}{K} \right)} \right)
\]

where: \(c^*_{Kt(a, max)}\) is the sorption capacity of algae and \(K\) is the constant.

Figure 4 compares the Langmuir isotherms (Eq. 2) determined for Cu\(^{2+}\) cations and for the sum of Cu\(^{2+}\) and H\(^+\) cations.

The Langmuir isotherms also indicate the influence of hydrogen cations on the equilibrium concentrations of copper in algae and in solutions. The trend line in Fig. 3a does not cover data collected at pH\(_0\) equal to 4 and 3.

Statistical parameters of sorption kinetics and equilibria

Table 3 lists the statistical parameters of the slopes of straight lines describing the sorption kinetics (Eq. 1) of copper cations and of the sum of H\(^+\) and Cu\(^{2+}\) cations. Numbering of measurement series corresponds to that in Table 2. Standard errors of the slopes and coefficients of determination indicate the results of measurements are well structured both in case of copper cations and in case of the sum of H\(^+\) and Cu\(^{2+}\) cations. The standard error of slopes was always smaller than 2 % of the value determined (b=1/c\(*_{Cu(a,1)}\)−1).

Table 4 shows the statistical parameters of the Langmuir isotherms determined for copper cations (Fig. 4a) and for the sum of H\(^+\) and Cu\(^{2+}\) cations (Fig. 4b). Data presented in Table 4 show that the uncertainty of determination of the sorption capacity exceeds 300 % in case of copper cations and is close to 32 % in case of the sum of H\(^+\) and Cu\(^{2+}\) cations. In order to find which parameters influenced the uncertainty of the measurement results, the component 1/c\(*_{Kt(a,1)}\)−1 contained both in the kinetic model (Eq. 1) and in the equilibrium model (Eq. 2) was analysed. The values of the parameter were determined using both models and compared. For the Cu\(^{2+}\) and H\(^+\)+Cu\(^{2+}\) experiments, the significance levels were evaluated of a hypothesis that (1/c\(*_{Kt(a,1)}\)−1)\(_{PSO}\)=(1/c\(*_{Kt(a,1)}\)−1)\(_{Ln}\), where index PSO denoted the kinetic model while index Ln, the equilibrium model. Table 5 shows the results of the analysis which include the coefficients of determination \(R^2\) and the significance levels \(\alpha\) obtained with t test for the equivalent hypothesis that the free factor a=0 and the slope b=1.

Data contained in Table 5 show that the equality (1/c\(*_{Kt(a,1)}\)−1)\(_{PSO}\)=(1/c\(*_{Kt(a,1)}\)−1)\(_{Ln}\) was statistically justified both for Cu\(^{2+}\) and for the sum of H\(^+\) and Cu\(^{2+}\), which in turn indicates that the mathematical models used fitted well the kinetics and equilibria of the ionic exchange process. Thus, the significant scattering of experimental points from the theoretical Langmuir isotherm in Fig. 4a as well as large standard error of the

![Fig. 4](image-url) Langmuir isotherms determined for: a Cu\(^{2+}\) cations; and b the sum of Cu\(^{2+}\) and H\(^+\) cations

| Measurement series | Cu\(^{2+}\) | H\(^+\) + Cu\(^{2+}\) |
|------------------|------------|------------------|
| \(b=1/c^*_{Cu(a,1)}\)±SEb | \(R^2\) | \(b=1/c^*_{HCu(a,1)}\)±SEb | \(R^2\) |
| 1 | 69.31 | 0.70 | 0.999 | 45.94 | 0.25 | 1.000 |
| 2 | 52.09 | 0.64 | 0.998 | 37.17 | 0.26 | 0.999 |
| 3 | 34.51 | 0.22 | 1.000 | 27.17 | 0.14 | 1.000 |
| 4 | 24.27 | 0.27 | 0.999 | 20.35 | 0.20 | 0.999 |
| 5 | 18.50 | 0.17 | 0.999 | 16.01 | 0.16 | 0.999 |
| 6 | 16.11 | 0.13 | 0.999 | 14.25 | 0.13 | 0.999 |
| 7 | 30.01 | 0.40 | 0.998 | 13.64 | 0.13 | 0.999 |
| 8 | 41.67 | 0.67 | 0.997 | 3.449 | 0.014 | 1.000 |
sorption capacity in Table 4 resulted probably from the competitive sorption of hydrogen cations. The largest deviation from the model was found at pH = 3 (Fig. 4a).

Discussion

The results of studies on the influence of pH on copper sorption by the alga *P. palmata* showed that the maximal sorption took place at pH 4.7 (Rajfur and Klos 2012). This is generally consistent with results obtained by other laboratories. For instance, the optimal acidity for sorption of cadmium by algae *Oedogonium* sp. and *Ceramium virgatum* Roth was pH 5.0 (Gupta and Rostogi 2008b; Saria and Tuzen 2009), while that for sorption of copper by the alga *Cladophora crispata* (Roth) Kützing was pH 4.5 (Özer et al. 2004).

The model of pseudo-second-order reaction is often used to describe the sorption kinetics in several biosorbents: algae (Senthilkumar et al. 2006); mosses (Sari and Tuzen 2009); and lichen (Uluozlu et al. 2008). It has often been compared to the model of a pseudo-first-order reaction and found to better approximate the experimental data. Impact of hydrogen cations on the kinetics of the process is rather disregarded in these research. Many authors neglected the initial phases of their experiments. For instance, they analysed data collected after first 5 min (Sari and Tuzen 2008) or even after 20 min (Chen et al. 2008). The results of the studies described in literature show that the parameters of sorption kinetics were dependent on concentration of the solution, which is confirmed by the results shown in the graphs in Fig. 3, but they are also related to the ratio of sorbent mass and solution volume (Gupta et al. 2006). The studies presented in this publication were carried out in the system where 1.0 g DM of algae was dispersed in 0.4 L of solution.

For the alga *C. crispata* (Roth) Kützing, in which copper cations were sorbed, the biggest values of $k''$ and $c_{Kt(a,1)}$ were observed in temperature of 298 K (Özer et al. 2004).

Our investigations were carried out in the temperature range 295–298 K.

Ion-exchange equilibrium, as in this article, was described usually using Langmuir isotherm model, and this model has often been compared to Freundlich, Redlich-Peterson and Koble-Corrigan models. It assumes a monomolecular layer of sorbed molecules or ions and is formed on the sorbent surface, which is justified in case of the ion-exchange processes. Also in this case, the descriptions of the sorption of heavy metal cations in algae do not include simultaneous sorption of hydrogen cations.

The sorption capacity for copper of the alga *P. palmata*, determined at comparable initial acidities of solutions (Fig. 4a), was $c^* = 0.99$ mmol g$^{-1}$ (31.5 mg g$^{-1}$ DM). On the other hand, the sorption capacity determined for the sum of H$^+$ and Cu$^{2+}$ at various conditions (Fig. 4b) was $c^* = 0.39$ mmol g$^{-1}$ (12.4 mg g$^{-1}$ DM). These results, together with scattering of data in Fig. 4a, indicate that the evaluation of sorption capacity of algae is subject to large errors. Table 6 compares sorption capacities for various algae species.

| Algae                     | $c^*_{Cu(a, max)}$ (mmol g$^{-1}$ DM) | $c_{Cu(a, max)}$ (mg g$^{-1}$ DM) | References                 |
|---------------------------|-------------------------------------|----------------------------------|-----------------------------|
| *Caulerpa lentillifera* J. Agardh | 0.18                                | 5.72                             | (Pavasant et al. 2006)      |
| *Codium vermilara* (Olivi) Delle Chiaje | 0.54                                | 17.2                             | (Romera et al. 2004)        |
| *Spirogyra insignis* (Hassall) Kützing | 0.60                                | 19.1                             |                             |
| *Sargassum fluitans* (Borgesen) | 0.80                                | 25.4                             | (Davis et al. 2003)         |
| *Cladophora crispata* (Roth) Kützing | 1.03                                | 32.7                             | (Özer et al. 2004)          |
| *Ulva* sp.                 | 1.50                                | 47.7                             | (Sheng et al. 2004)         |
| *Chaetomorpha linum* (O.F.Müller) Kützing | 2.92                                | 92.8                             | (Ahmady-Asbchin et al. 2009) |
| *Fucus serratus* Linnaeus  | 3.20                                | 102                              |                             |
| *Spirogyra* sp.            | 4.20                                | 133                              | (Gupta et al. 2006)         |

Table 6 Sorption capacity for copper of various algae species (pH of solutions: 4.5–5.5)
Data in Table 6 also indicate large differences in the sorption capacities evaluated, which not necessarily result from different sorption properties between algae species. The studies done for the algae Spirogyra sp. showed that the evaluation uncertainty of the sorption capacity for copper reached 10 %, depending on the experimental procedures used (in particular—on methods of algae preparation) (Rajfur et al. 2012). Unfortunately, the authors quoted in Table 6 had not provided any error estimates.

In conclusion, the results of the research presented indicate a significant influence of hydrogen cations on the sorption of copper cations and, probably, other heavy-metal cations by algae. The competitive sorption of hydrogen cations affects the kinetics and equilibria of ionic exchange of heavy metals. It increases the uncertainty of determination of such parameters as algae sorption capacity. At equal initial acidity (pH0) of solutions, the competitive sorption of hydrogen ions decreases when initial gram equivalent fraction of copper in the solution is increased. Thus, to determine properly the sorption capacity of algae, one has to take into account the sum of sorbed cations—hydrogen and copper. If the sum of cations is used in Eq. 2, the scattering of experimental points from theoretical model lines is reduced (Fig. 4).

The results show that the Langmuir isotherm and the pseudo-second-order reaction model properly describe the equilibria and kinetics of ionic exchange between algae and solutions.

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