CHANGES OF HAEMATOLOGICAL INDICES OF JUVENILE CARP
(Cyprinus carpio L.) UNDER THE INFLUENCE OF NATURAL
POPULATIONS OF CYANOBACTERIAL WATER BLOOMS

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Abstract

The aim of the presented paper was to evaluate the effect of cyanobacterial water blooms common in eutrophic reservoirs on blood indices of carp. A total of 180 individuals of juvenile carp (Cyprinus carpio L.) of the average body mass 36.9 g (1996) and 26.3 g (1997) were used in the experiments. The experimental fish were exposed to 4 different natural populations of cyanobacterial water blooms.

In 1996, the populations of filamentous cyanobacteria (WB I) formed by Anabaenopsis flos-aquae (90 %) and Aphanizomenon flos-aquae (10 %) at a concentration of $5.6 \times 10^3 - 3.2 \times 10^5$ cells ml$^{-1}$ without the presence of microcysts and colonial cyanobacteria Microcystis ichthyoblabe (80 %) with subdominant Microcystis aeruginosa (20 %) (WB II) at a concentration of $2.6 \times 10^4 - 3.6 \times 10^4$ cells ml$^{-1}$ with detected Microcystis LR (17.2 µg g$^{-1}$ of dry mass) were used. Exposure time was 168 h for filamentous species and 96 h for colonial species. In 1997 the monospecies population of filamentous cyanobacteria Anabaena flos-aquae (WB III) at a concentration of $3.3 \times 10^4 - 7.9 \times 10^5$ cells ml$^{-1}$ containing two different microcystins (total concentration $56.06 \mu$g g$^{-1}$ of dry mass) and water bloom (WB IV) formed by colonial species Microcystis ichthyoblabe (40 %) and Microcystis aeruginosa (30 %) with filamentous Anabaena flos-aquae (30 %) at a concentration of $1.8 \times 10^4 - 1.4 \times 10^5$ cells ml$^{-1}$ which contained three microcystins (total concentration $289.3 \mu$g g$^{-1}$ of dry mass) were used. Both populations were exposed for 48 h. Control fish in both tests were kept in treated drinking water infused 24 h before the start of the experiment.

Haematological examination showed significant changes ($p < 0.05$) in leukocrit (BC) of fish exposed to the cyanobacterial population WB I, and in haematocrit (PCV) values, total protein concentration (TP), ALT and AST activities in fish exposed to the population WB II as compared to control fish. LDH activity in blood plasma of carp exposed to cyanobacterial population WB II was increased ($p < 0.01$) as compared to control fish. TP values from cyanobacterial populations WB III a WB IV were significantly reduced ($p < 0.05$) and values of ALT activities increased ($p < 0.01$). Moreover, significant increase ($p < 0.05$) of AST activity was recorded for fish exposed to cyanobacterial water bloom WB III.

The observations confirmed adverse effects of cyanobacterial biomass on juvenile carp. The effect of toxic water bloom populations was manifested by changes of blood plasma indices. Toxins supply here water the role of catalysts enhancing the negative influence of toxic high ammonia values.

Carp, cyanotoxins, water blooms, plasma enzymes

Cyanobacteria as photosynthesizing organisms produce biologically active compounds that may affect growth and development of other water organisms and physical and chemical characteristics of water (Maršálek and Turánek 1996). Great attention has recently been paid to the impact of cyanobacterial toxins on fish. Clinical symptoms of poisoning, pathological changes and influence on blood indices have been investigated as well.
Maximum of toxins is absorbed into the fish organism through the gastrointestinal tract, whereas toxin penetration through the skin or gills is negligible (Tencalla et al. 1994). Toxic influence of Microcystin LR on carp after oral administration was manifested by torpidity and loss of reflexes, skin haemorrhages, eye chamber and in internal organs. Considerable damage were found for fish kidney and liver (Navrátil et al. 1996, 1997).

Intraperitoneal exposure to microcystins causes tissue damage in fish liver as demonstrated by significant increase of alanine aminotransferase (ALT), aspartate aminotransferase (AST) and lactate dehydrogenase (LDH) activities (Raberg et al. 1991; Navrátil et al. 1998). Toxic effect on fish exposed to media containing the dispersed microcystin is manifested in delay caused by limited penetration into the healthy fish. Toxic effect after the oral administration is approximately 10 times lower than after the intraperitoneal application (Carbis et al. 1996a).

Long-term impact of the cyanobacteria containing microcystins at lower concentrations is relatively difficult to observe in individual fish; therefore it is more suitable to check a larger number of fish. The observation of aminotransferase (ALT, AST), bile acids, bilirubin, sodium and chloride from the blood serum is recommended (Carbis et al. 1996b).

Materials and Methods

Juvenile carps were kept during 48 h before the start of the experiment in laminated basins (0.5 m³) and subsequently were relocated into the permanently aerated 100-litre aquarium (10 pieces/aquarium).

A total of 100 juvenile carp (90 experimental and 10 control fish) of mean weight 36.9 ± 7.9 g were used in experiment A (29.7 - 5.8, 1996). Fish were fed a diet Alma (20% of N-substances, 13% of fat) at the dose of 2% of the fish stock weight. During the last 48 h of the experiment no food was available.

The experimental fish were exposed to two natural cyanobacterial water bloom populations for 168 and 96 h, respectively. No microcystins were detected in cyanobacterial population WB I, Microcystin LR (17.2 μg g⁻¹ of dry mass) was found in population WB II.

During the experiment, water temperature ranged between 22.8 - 26.8 °C, dissolved oxygen concentration reached 24–98 % and pH was 7.7 - 8.4. The values of ammonia N-NH₄⁺ varied between 0.04–1.08 mg l⁻¹.

Juvenile carp (70 experimental and 10 control fish) of mean weight 26.3 ± 7.8 g were used in experiment B (12. 8. - 14. 8. 1997). The experimental fish were again exposed to two natural cyanobacterial water bloom populations for the duration of 48 h without feeding. Cyanobacterial population WB II contained 2 microcystins (total toxin concentration 52.0 μg g⁻¹ of dry mass). In population WB IV 3 microcystins were found with total concentration 193.9 μg g⁻¹ of dry mass.

During the experiment, water temperature ranged between 23.7 - 26.9 °C, dissolved oxygen concentration reached 56–122 % and pH 7.7 - 8.3. The values of ammonia N-NH₄⁺ varied between 0.02–0.36 mg l⁻¹.

Differences in values of monitored physical and chemical characteristics among particular aquarium were negligible for both of experiments, occurrence of low values of dissolved oxygen were momentary (see Fig. 1 and 2). Control fish for both of experiments were kept in the treated drinking water.

Cyanobacterial biomass was evaluated by chlorophyll a concentrations (Stepek et al. 1982), and by number of cells counted in Bürker's counting chamber. High-performance liquid chromatography (HPLC) was used for the analyses of microcystins in all chosen cyanobacterial populations. These analyses were performed at the Veterinary Research Institute in Brno.

Blood samples of 52 fish in experiment A, and of 40 fish in experiment B were taken from the head into the heparinised tubes after the termination of the exposure time. Additional processing of blood and plasma separation were carried out after Sobotová et al. (1986). Values of haemoglobin (Hb), haematocrit (PCV), leukocrit (BC) and corpuscular haemoglobin concentration (MCHC) were determined by standard methods (Sobotová et al. 1986). Commercial kit (Lachema Diagnostika, Czech Republic) was used for the detection of total protein (TP) concentration in blood plasma. Activities of aminotransferases (ALT, AST) were detected by the commercial kit Humazyme UV test (Humin, Germany), activity of lactate dehydrogenase (LDH) was detected by LDH 105 UV kit (Lachema Diagnostika, Czech Republic).

Statistical evaluation of results (Student’s t-test) was done using the software Microsoft Excel 97.

Results

Results of haematological examinations are presented in the Tables 1 and 2. Statistical evaluation of the influence of cyanobacterial population (WB I) on haematological indices of the juvenile carp showed distinct decrease of leukocrit (BC).
Table 1
Haematological indices of juvenile carp blood (experiment A)

<table>
<thead>
<tr>
<th>Experiment A (1996)</th>
<th>WB I (168 h)</th>
<th>WB II (96 h)</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>chlorophyll a (μg·l⁻¹)</td>
<td>mean 40.18</td>
<td>28.83</td>
<td></td>
</tr>
<tr>
<td>cells·ml⁻¹</td>
<td>mean 1.8·10⁵</td>
<td>1.8·10⁵</td>
<td></td>
</tr>
<tr>
<td>fish</td>
<td>n 21</td>
<td>24</td>
<td>7</td>
</tr>
<tr>
<td>N-NH₃(mg·l⁻¹)</td>
<td>min-max 0.04–1.08</td>
<td>0.06–0.42</td>
<td>0.05–0.59</td>
</tr>
<tr>
<td>Hb (g·l⁻¹)</td>
<td>mean 51.32</td>
<td>55.75</td>
<td>55.17</td>
</tr>
<tr>
<td>BC (H⁺)</td>
<td>mean 0.0078</td>
<td>0.0083</td>
<td>0.0096</td>
</tr>
<tr>
<td>PCV (H⁺)</td>
<td>mean 0.0020</td>
<td>0.0012</td>
<td>0.0015</td>
</tr>
<tr>
<td>MCHC (H⁺)</td>
<td>mean 0.25</td>
<td>0.25</td>
<td>0.29</td>
</tr>
<tr>
<td>TP (g·l⁻¹)</td>
<td>mean 0.07</td>
<td>0.04</td>
<td>0.03</td>
</tr>
<tr>
<td>ALT (μkat·l⁻¹)</td>
<td>mean 0.13</td>
<td>0.15</td>
<td>0.12</td>
</tr>
<tr>
<td>AST (μkat·l⁻¹)</td>
<td>mean 0.04</td>
<td>0.02</td>
<td>0.03</td>
</tr>
<tr>
<td>LDH (μkat·l⁻¹)</td>
<td>mean 27.72</td>
<td>23.25</td>
<td>25.76</td>
</tr>
<tr>
<td>SD</td>
<td>mean 4.25</td>
<td>3.66</td>
<td>1.31</td>
</tr>
</tbody>
</table>

**WB I** - Cyanobacterial water bloom (*Anabaena flos-aquae* 90% and *Aphanizomenon gracile* 10%). Microcystins were not detected.

**WB II** - Cyanobacterial water bloom (*Microcystis ichthyoblabe* 80% and *Microcystis aeruginosa* 20%). Microcystin LR was detected (17.2 μg·g⁻¹ of dry mass).

SD - standard deviation. One and two asterisks are used for indicating the significance of differences at the level of *p < 0.05* and *p < 0.01*, respectively.

Table 2
Haematological indices of juvenile carp blood (experiment B)

<table>
<thead>
<tr>
<th>Experiment B (1997)</th>
<th>WB III (48 h)</th>
<th>WB IV (48 h)</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>chlorophyll a (μg·l⁻¹)</td>
<td>mean 62.08</td>
<td>86.67</td>
<td></td>
</tr>
<tr>
<td>cells·ml⁻¹</td>
<td>mean 6.4·10⁴</td>
<td>7.2·10⁴</td>
<td></td>
</tr>
<tr>
<td>fish</td>
<td>n 17</td>
<td>16</td>
<td>7</td>
</tr>
<tr>
<td>N-NH₃(mg·l⁻¹)</td>
<td>min-max 0.04–0.36</td>
<td>0.02–0.35</td>
<td>0.06–0.25</td>
</tr>
<tr>
<td>Hb (g·l⁻¹)</td>
<td>mean 64.18</td>
<td>61.69</td>
<td>57.57</td>
</tr>
<tr>
<td>TP (g·l⁻¹)</td>
<td>mean <em>23.61</em></td>
<td><em>23.82</em></td>
<td>25.50</td>
</tr>
<tr>
<td>ALT (μkat·l⁻¹)</td>
<td>mean <em>2.52</em></td>
<td><em>2.39</em></td>
<td>1.51</td>
</tr>
<tr>
<td>AST (μkat·l⁻¹)</td>
<td>mean <em>0.98</em></td>
<td><em>0.47</em></td>
<td>0.55</td>
</tr>
<tr>
<td>LDH (μkat·l⁻¹)</td>
<td>mean 44.73</td>
<td>34.44</td>
<td>28.89</td>
</tr>
<tr>
<td>SD</td>
<td>mean 30.19</td>
<td>11.02</td>
<td>5.13</td>
</tr>
</tbody>
</table>

**WB III** - Cyanobacterial water bloom (*Anabaena flos-aquae* 100%). Two microcystins were detected, Microcystin LR was not found. Mcyst 1 (3.22 μg·g⁻¹ of dry mass), Mcyst 2 (48.80 μg·g⁻¹ of dry mass).

**WB IV** - Cyanobacterial water bloom (*Microcystis ichthyoblabe* 40%, *Microcystis aeruginosa* 30% and *Anabaena flos-aquae* 30%). Three microcystins were detected Mcyst 1 (34.48 μg·g⁻¹ of dry mass), Mcyst 2 (82.20 μg·g⁻¹ of dry mass) and Microcystin LR (77.20 μg·g⁻¹ of dry mass).

SD - standard deviation. One and two asterisks are used for indicating the significance of differences at the level of *p < 0.05* and *p < 0.01*, respectively.
(p < 0.05) compared to controls. Values of Hb, PCV and ALT were lower than in controls, MCHC, TP, AST and LDH were slightly and non-significantly elevated. Red blood cell indices and total protein (TP) oscillated around the lower limit of variations usual for carp (Svobodová et al. 1986) or slightly below it. Activities of blood plasma enzymes were higher compared to usual values.

The toxic effects of the cyanobacterial population (WB II) resulted in significant changes of PCV, TP, ALT, AST (p < 0.05) and LDH (p < 0.01) compared to controls. PCV and TP indices were lower and the activities of blood plasma enzymes were increased. Decrease of BC values and increase of Hb and MCHC values was non-significant. Comparison of the obtained haematological indices of the cyanobacterial population (WB II) with the range of the value variation for carp (Svobodová et al. 1986) is the same as for the previous cyanobacterial population (WB I).

Both cyanobacterial populations in experiment B contained toxins that influenced mainly blood plasma indices. Significant differences in control fish were found for TP (p < 0.05) and ALT (p < 0.01) values. Moreover, monocultural population of cyanobacteria (WB III) caused significant increase of AST activity (p < 0.05). Hb and TP values were lower as compared to values for controls and ranged around the lower limit of the value variation for carp (Svobodová et al. 1986). The AST, ALT and LDH activities in all studied groups of fish were higher and exceeded the threshold of the range of the value variation (Svobodová et al. 1986).

Discussion

Rabergh et al. (1991) reported that the values of blood plasma enzymes (ALT, AST and LDH) raise in two hours after an intraperitoneal injection of toxin as a consequence of the hepatocyte necrosis. Tencalla et al. (1994) observed already after 48 h a decrease of their activity, and interpreted this fact as a result of damage of the majority of hepatocytes that were not able to release enzymes into circulatory system. Carbis et al. (1996b) noted a delay of toxic manifestation in fish exposed to water with immersed microcystin. Falconer (1998) and Falconer et al. (1994) presented catalyzing effect of hepatotoxins from blue-green algae negatively enhancing effect of physical and chemical indices (NH₃, O₂, pH).

The ammonia values exceeded maximal feasible concentration for carp in all experiments, both of experimental and control fish were influenced. The toxic effect of ammonia caused the death of 9 fish during the experiment. The influence of these high ammonia concentrations on haematological parameters is supposed. Consequences of ammonia activity implicated histopathological changes of liver of carp from the concentration of 0.1-0.33 mg l⁻¹ NH₃ (Svobodová and Groch 1971). Since these concentrations were observed in experiments, we can suppose influence of ammonia on livers enzyme activity increase.

Nevertheless high ammonia values were measured for both experimental and control fish. Statistically significant differences of haematological parameters among experimental and control fish were ascribed to effect of microcystins operating as co-factors of toxic effect of ammonia.

Cyanobacterial toxins are secondary metabolic products. Since they are endotoxins, they cannot be actively secreted into the environment. However, after the breakdown of water bloom and decomposition of cell walls, cyanotoxins can be released into the water (Maršálek and Turánek 1996). Carp ingest cyanobacteria very rarely. The digestive tract of carp has a slightly alkaline pH, and its enzymes are not able to decompose mucilaginous envelopes of cyanobacteria. Low pH is essential for more effective lysis. Therefore, fish are mostly endangered by cell toxins of older declining cyanobacterial populations having envelopes partially lysed (Carbis et al. 1997).

Higher temperature through the experiments was the reason of faster cyanobacterial biomass decomposition that was followed by faster toxin release into the water. Chorus and
Bartram 1999, showed that for young populations 100% of toxins is located in cells whereas for decaying cells toxin concentration raise in water on values of 70-80%.

Distant toxin concentration was found in particular parts of fish body. If the carp ingests toxin with food 55% of toxins is stored in musculature, 38% in digestive tract and the rest is excluded with excrements. In case of toxin presence in water 50% of toxin was found in skin, 30% in gills, 18% in intestines and 2% in musculature (Maršálek 1996). Ingestion of cyanobacteria by carp during the experiments was minimal, the gills and skin were identified as a main penetration system into organism.

The values of haematological indices correspond well with the results of other authors (Raberg et al. 1991; Tencalla et al. 1994; Carbis et al. 1996ab; Navrátil et al. 1996, 1998; Vajcová et al. 1998). Certain differences are caused mainly by different ways of toxin penetration and by different physiological state of cyanobacterial water bloom populations.

Blood plasma indices appeared as better indicators. Liver enzymes (ALT, AST and LDH) are the most frequently tested enzymes in fish. Their values increase markedly as a consequence of necrosis (Raberg et al. 1991; Tencalla et al. 1994; Navrátil et al. 1998). These results correspond with conclusions of our experiments. The influence of microcystins in cyanobacterial populations (WB II, III, IV) was manifested mainly by enzyme activity increase. These results were also confirmed by statistical analysis.

We suppose that significant changes of haematological parameters were caused by toxic effect of cyanobacteria together with toxic effect of ammonia. We found these differences already in microcystin concentrations of 17.2 µg·g⁻¹ of dry mass (cyanobacterial population WB II).

The control of physical and chemical parameters, mainly ammonia and dissolved oxygen concentrations are essential for confirmation of presented results and subsequent experiments.

The rapid increase of metabolic ammonia could be eliminated by prolongation of starvation time of fish and by elimination of feeding during the experiments. Aquaria with higher capacity as well as lower number of experimental fish will decrease these values. Application of semistatic tests with use of several aquaria with the same cyanobacterial concentration and progressive fish transfer to the others after an increase of toxic ammonia concentration will be suitable.

### Změny vybraných hematologických ukazatelů krve pládka kapra

**(Cyprinus carpio L.)** pod vlivem přírodních populací vodních květů sínice

Cílem práce bylo zjistit vliv přírodních populací vodních květů sínice, běžně se vyskytujících v eutrofních vodách, na krevní ukazatele kapra obecného.

K experimentům byl použit pládek kapra o průměrné individuální hmotnosti 36.9 g a 26.3 g v celkovém počtu 180 ks. Pokusné ryby byly chovány v prostředí 4 různých druhů vodních květů sínice. V roce 1996 byla použita populace vlákňitéch sinic WB II (Anabaena flos-aquae 90% a Aphanizomenon gracile 10%) v koncentracích 5.6 × 10⁻⁴ až 3.2 × 10⁻⁵ buněk/ml⁻¹ bez zjištěné přítomnosti microcystinů a populace kokálních sinic WB II (Microcystis ichthyoblabe 80% a Microcystis aeruginosa 20%) v koncentracích 2.6 × 10⁻³ až 3.6 × 10⁻⁶ buněk/ml⁻¹, u které byl detekován Microcystin LR (17.2 µg·g⁻¹ sušiny). Celková doba sledování činila 168 h (vláknité sinice), resp. 96 h (kokální sinice).

V roce 1997 byla použita přírodní monokulturní populace sinic WB III (Anabaena flos-aquae 100%) v koncentracích 5.3 × 10⁴ až 7.9 × 10⁴ buněk/ml⁻¹, u které byly zjištěny dva typy microcystinů (celková koncentrace 56.06 µg·g⁻¹ sušiny) a vodní květ sínice VKS IV (Microcystis ichthyoblabe 40%, Microcystis aeruginosa 30% a Anabaena flos-aquae 30%) v koncentracích 1.8 × 10⁶ až 1.4 × 10⁷ buněk/ml⁻¹, která obsahovala tři typy microcystinů (celková koncentrace 289.3 µg·g⁻¹ sušiny). Doba pozorování u obou
populaci činila 48 h. Kontrolní ryby u obou pokusů byly chovány v upravené vodovodní vodě, napuštěné 24 h před začátkem sledování.

Hematologické vyšetření prokázalo statisticky významné změny \( p < 0.05 \) u leukokritu BC (WB I) a u hodnot hematókritu (PCV), koncentrace celkových bílkovin (TP), aktivity alanin-aminotransferázy (ALT) a aspartáte-aminotransferázy (AST) (WB II) ve srovnání s kontrolní skupinou ryb. Výsledky statisticky významné zvýšení \( p < 0.01 \) bylo zaznamenáno v krevní plazmě kaprů u aktivity lactáty-dehydrogenázy (LDH) u populace síněc WB II ve srovnání s kontrolními rybami. Hodnoty ukažovacích TP u plůdek kapra z prošlechčené populace síněc WB III a WB IV byly významně sníženy a hodnoty aktivity ALT vysoko významně zvýšeny. U ryb chovaných v prošlechčené populace síněc WB III bylo naopak prokázáno významné snížení aktivit AST.

Zjištěné údaje svědčí o vlivu biomasy síněc na plůdek kapra, projevující se především u toxických populace vodních květů síněc změnami ukažovacích krevní plazmy. Toxiny zde vystupují v roli katalyzátorů umožňujících negativní působení vysokých hodnot toxického amoniku.

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