Low pH and calcium effects on net Na\(^+\) and K\(^+\) fluxes in two catfish species from the Amazon River (Corydoras: Callichthyidae)

Abstract

The present study analyzes Na\(^+\) and K\(^+\) disturbances caused by low pH in two catfish species from the Amazon River. *Corydoras adolfoi* inhabits ion-poor, black-stained, low pH (3.5-4.0) waters, while *C. schwartzii* is native to ion-rich waters at circumneutral pH. Fish were exposed to pH 3.5 Ca\(^{2+}\)-free, and Ca\(^{2+}\)-enriched (~500 μmol/l) water to determine the protective effects of calcium. Net Na\(^+\) and K\(^+\) fluxes were measured in the water collected from the fish experimental chambers. *C. adolfoi* was unable to control the Na\(^+\) efflux at low pH, exhibiting Na\(^+\) loss up to -594 ± 84 nmol g\(^{-1}\) h\(^{-1}\) during the first hour. After 3 and 6 h, net Na\(^+\) flux increased by 7- and 23-fold, respectively. In *C. schwartzii*, at pH 3.5, the initial high Na\(^+\) loss (-1,063 ± 73 nmol g\(^{-1}\) h\(^{-1}\)) was gradually attenuated. A K\(^+\) loss occurred in both species, but remained relatively constant throughout exposure. High [Ca\(^{2+}\)] affected ion losses in both species. *C. adolfoi* had 70% loss attenuation, indicating incapacity to control Na\(^+\) efflux. In *C. schwartzii*, elevated [Ca\(^{2+}\)] completely prevented the Na\(^+\) losses caused by exposure to low pH. Rather different patterns were seen for K\(^+\) fluxes, with *C. adolfoi* showing no K\(^+\) disruption when exposed to low pH/high [Ca\(^{2+}\)]. Thus, *C. adolfoi* loses Na\(^+\) during acid exposure, but has the ability to control K\(^+\) loss, while *C. schwartzii* controls diffusive Na\(^+\) loss but exhibits a slightly higher K\(^+\) loss. Ion balance was influenced by [Ca\(^{2+}\)] at low pH in *C. schwartzii* but not in *C. adolfoi*.

Introduction

The Amazon blackwaters are known by their singular physicochemical features, which are unusual relative to the world average values for freshwater (1). The two most noticeable parameters are low pH and low ion concentration. The Rio Negro River, for instance, presents pH levels around 4.5-6.0, which are as low as 3.0 in restricted flooded forest areas (the *igapós*) (2). The Amazon nutrient-rich waters, also called whitewaters, drain from the Andean and pre-Andean regions, and their pH is usually around circumneutral values (1). In spite of such extremely adverse features for aquatic living organisms, blackwater habitats are not as inhospitable as one would expect. Fish diversity for the Rio Negro River was estimated at more than 1000 species (3). How these specimens can thrive in such conditions and what the adaptive strategies developed by them are still poorly understood.

Acid waters with low ion concentrations...
such as found in blackwaters may induce severe ion disruption in fish, including K⁺ losses from the epithelial cells themselves due to the toxic effects triggered by high [H⁺] (4). These effects can be summarized as the inhibition of ion uptake and an elevation of ion efflux (5). Calcium has long been known to reduce the toxicity of acid waters to fish (6-10). Apparently, there is a Ca²⁺ versus H⁺ competition for sites on tight junctions that maintain epithelial integrity, as well as evidence that bound Ca²⁺ shields access to Na⁺-binding sites on the apical transporters (5). Thus, high [H⁺] displaces Ca²⁺ from the binding sites and diffusive losses take place (4).

_Corydoras adolfoi_ Burgess, 1982, is a catfish commonly inhabiting blackwaters in the flooded forests of the middle Rio Negro River, withstanding pH’s around 3.5–4.0 (11). C. *schwartzii* Rössel, 1963, was used for comparison because it is a species endemic to whitewaters of the Purus River, living at circumneutral pH. The aim of the present study was to evaluate the disturbances in the net Na⁺ and K⁺ fluxes in these catfish from the Amazon region exposed to acute low pH, in an attempt to assess how they can deal with low pH. The possible protective influence of Ca²⁺ was also analyzed.

**Material and Methods**

**Experimental fish**

Fish collected from their native environments were obtained from ornamental fish exporters (Corydoras Tetra Aquarium and Turky’s Aquarium) in the municipality of Manaus, Amazon, Brazil. They were acclimated for at least 14 days before the beginning of the experiments at the Laboratory of Ecophysiology and Molecular Evolution of INPA. Fish were held in 60-liter aquaria, provided with pump aerators, carbon-zeolite filters, and supplied with INPA ground water (average concentrations: 15 μmol/l Na⁺, 16 μmol/l Cl⁻, 9 μmol/l K⁺, 9 μmol/l Ca²⁺, and 2.05 mg/l dissolved organic carbon; see Ref. 12). Ground water pH was 6.0 ± 0.5 after two days of aeration to eliminate excess CO₂ before use. Water temperature was 27 ± 2°C. All fish weighed 1.5 to 3.5 g. The specimens were fed commercial dried pellets daily, and food was withdrawn 24 h prior to experimentation.

**Experimental protocol**

Ten fish were used in each treatment, including a control group. Aerated ground water was used as the ‘standard water’. Experimental chambers consisted of 200-ml polyethylene darkened chambers; small holes on the lid were used to supply water (recirculating system) and aeration, and for pH adjustments. The fish were placed in these chambers 12 h before starting the experiments. At the beginning of the experiment, the flow was stopped, and water in the chambers was totally replaced with fresh standard water. A 10-ml water sample was taken and a second sample was taken from each chamber 1 h later (control, pH 6.0). This procedure was performed for all fish of all treatments to confirm fish homeostasis and any possible disturbance due to the transfer to chambers. Water samples were stored in plastic vials and refrigerated until subsequent analysis for Na⁺ and K⁺ concentrations.

**Low pH treatment**

After sampling during the control period, standard water in the chamber was completely changed by emptying it manually and replacing with experimental solution. Acid solution (pH 3.5 ± 0.2) was previously prepared by the addition of 1 N H₂SO₄ to standard water, and kept aerated in a plastic box for 12 h before use. Calcium concentration in the solution was about 9 μmol/l (Ca²⁺.
from the ground water). The pH inside the chambers was periodically checked using a portable pH meter and adjusted as necessary by the addition of diluted H$_2$SO$_4$. Water samples (10 ml) were taken 1, 3, and 6 h after the water was changed. Acid solution was then replaced with standard water again, and the recovery period was started. Samples in the recovery series were taken after 1 and 3 h. Fish were weighed and then released into a holding aquarium. An identical control series was performed for each species with the same water changeovers, but using pH 6.0 throughout.

**Effects of calcium at low pH**

The experimental protocol for this treatment was similar to that described before, except that CaCl$_2$ was added to obtain a ~500 µmol/l solution. The standard water was replaced with this solution after first sampling, which in turn was replaced with standard water after 6 h. Water samples were taken as previously described.

**Net Na$^+$ and K$^+$ flux measurements**

A flame photometer (CELM FC-180 model) calibrated with known standards was used to obtain Na$^+$ and K$^+$ readings in the water. Net Na$^+$ and K$^+$ flux measurements were made according to the following equation (13):

$$J_{\text{net}} = ([\text{ion}_1] - [\text{ion}_2]) \cdot V \cdot (M \cdot t)^{-1},$$

where $[\text{ion}_1]$ and $[\text{ion}_2]$ are the ion concentrations (nmol/l) at the beginning and at the end of the flux period, respectively; $V$ is the water volume in the chamber (liters); $M$ is the fish weight (g), and $t$ is the duration of the flux period (h).

**Statistical analysis**

All data are reported as means ± SEM. Mean values of net ion fluxes in the treatments were compared to the control groups by one-way ANOVA (P<0.05). When significant differences were detected, Dunnett’s multiple comparison test was applied.

**Results**

All fish were in approximate homeostasis during the control pretreatment phase, with very small net ion losses to the water. $J_{\text{net}}$ values for Na$^+$ and K$^+$ were not significantly different between the two species at the beginning of the treatments. Net ion fluxes were about $-189 \pm 22$ (Na$^+$) and $-56 \pm 14$ nmol g$^{-1}$ h$^{-1}$ (K$^+$) for *C. adolfoi* during the pretreatment period. The control values for *C. schwartzii* net fluxes were $-252 \pm 50$ (Na$^+$) and $-264 \pm 52$ nmol g$^{-1}$ h$^{-1}$ (K$^+$). Fish had acclimated to the chambers and had recovered from handling procedures during the transfer.

**Net fluxes at pH 3.5**

*C. adolfoi*, the blackwater species, had an overall Na$^+$ loss about 10-fold higher than that of *C. schwartzii*, the whitewater species, at pH 3.5, showing large and significant differences relative to the control treatment (Figure 1). In the first hour of exposure to pH 3.5, *C. adolfoi* exhibited a 2-fold increase in Na$^+$ loss (to $-594 \pm 84$ nmol g$^{-1}$ h$^{-1}$). After 3 and 6 h, $J_{\text{net}}$ Na$^+$ increased significantly by as much as 7- and 23-fold (to $-4,670 \pm 371$ and $-14,171 \pm 1,053$ nmol g$^{-1}$ h$^{-1}$), respectively, indicating that *C. adolfoi* was unable to control Na$^+$ efflux under acute low pH. No deaths occurred during this treatment, and efflux was quickly reduced during the recovery period.

*C. schwartzii* showed the capacity to withstand pH 3.5 fairly well (Figure 2). Although there was an initially high efflux during the first hour, when $J_{\text{net}}$ Na$^+$ was about 3 times higher ($-1,063 \pm 73$ nmol g$^{-1}$ h$^{-1}$) than control, the losses were gradually attenuated after 3 and 6 h ($-561 \pm 44$ and $-364 \pm 33$ nmol...
Figure 1. Net Na\(^+\) flux of Corydoras adolfi exposed to pH 3.5 and pH 3.5 plus Ca\(^{2+}\). Values are reported as means ± SEM (N = 10). *P<0.05 compared to a control measurement performed at pH 6.0 (Dunnett test). Water was changed to circumneutral pH for recovery (r) after 6 h of treatment (t).

Figure 2. Net Na\(^+\) flux of Corydoras schwartz exposed to pH 3.5 and pH 3.5 plus Ca\(^{2+}\). Values are reported as means ± SEM (N = 10). *P<0.05 compared to a control measurement performed at pH 6.0 (Dunnett test). Water was changed to circumneutral pH for recovery (r) after 6 h of treatment (t).

Figure 3. Net K\(^+\) flux of Corydoras adolfi exposed to pH 3.5 and pH 3.5 plus Ca\(^{2+}\). Values are reported as means ± SEM (N = 10). *P<0.05 compared to a control measurement performed at pH 6.0 (Dunnett test). Water was changed to circumneutral pH for recovery (r) after 6 h of treatment (t).

Figure 4. Net K\(^+\) flux of Corydoras schwartz exposed to pH 3.5 and pH 3.5 plus Ca\(^{2+}\). Values are reported as means ± SEM (N = 10). *P<0.05 compared to a control measurement performed at pH 6.0 (Dunnett test). Water was changed to circumneutral pH for recovery (r) after 6 h of treatment (t).

g\(^{-1}\) h\(^{-1}\), respectively), exactly the opposite pattern to that seen in C. adolfi. In the recovery series, net Na\(^+\) flux decreased significantly as well (-112 ± 26 and -35 ± 13 nmol g\(^{-1}\) h\(^{-1}\), respectively).

Significant K\(^+\) loss occurred in both species (Figures 3 and 4). For C. adolfi, the average loss was about 3 times higher than the control treatment (-200 ± 22 nmol g\(^{-1}\) h\(^{-1}\)), and remained constant throughout exposure. This feature suggests that the fish were able to control diffusive efflux. C. schwartz exhibited a pronounced K\(^+\) loss in the first hour of treatment (-419 ± 11 nmol g\(^{-1}\) h\(^{-1}\)), with some later attenuation. After continued exposure, these K\(^+\) effluxes were -241 ± 13 and -201 ± 9 nmol g\(^{-1}\) h\(^{-1}\), at 3 and 6 h, respectively.

**Effect of calcium at pH 3.5**

High [Ca\(^{2+}\)] greatly affected ion losses in both species at pH 3.5. However, C. adolfi was not able to control the Na\(^+\) efflux rate even in the presence of Ca\(^{2+}\) (Figure 1). Net Na\(^+\) flux was not significantly different in the first hour of treatment for this blackwater species (-235 ± 33 nmol g\(^{-1}\) h\(^{-1}\)). During the subsequent sampling periods, Na\(^+\) loss was progressively higher, as much as 5 times and 18 times greater (-1,452 ± 176 and -4,591 ± 531 nmol g\(^{-1}\) h\(^{-1}\), respectively). These losses represent about a 70% attenuation of the losses at low [Ca\(^{2+}\)], but were still substantially above those seen in C. schwartz even at low [Ca\(^{2+}\)]. For C. schwartz, elevated [Ca\(^{2+}\)] completely prevented the Na\(^+\) losses caused by low pH exposure. Indeed, Na\(^+\) losses were reduced by about 50% even compared to the control series at pH 6.0 (average losses were -118 ± 28 nmol g\(^{-1}\) h\(^{-1}\)), indicating a positive effect of Ca\(^{2+}\) in the presence of acute low pH (Figure 2).

Rather different patterns were seen for K\(^+\) fluxes. C. adolfi did not show K\(^+\) disruption when exposed to low pH under high [Ca\(^{2+}\)] (Figure 3). The average loss in the
treatment was 91 ± 10 nmol g⁻¹ h⁻¹, representing a net K⁺ flux 38% higher than the control, although not statistically different. *C. schwartzi* exhibited a ~50% reduction in K⁺ loss when exposed to low pH under high [Ca²⁺], accounting for an average 112 ± 16 nmol g⁻¹ h⁻¹ loss (Figure 4), slightly higher than that observed for *C. adolfoi*.

**Discussion**

**Net Na⁺ and K⁺ fluxes in Corydoras**

Differences in the acid tolerance were evident between the two species tested in the present study. *C. schwartzi* showed high acid tolerance, being very adept at dealing with rapid changes in water pH. *C. adolfoi*, the blackwater species, showed a marked Na⁺ disturbance at pH 3.5, developing massive and progressive losses. Such massive Na⁺ loss demonstrated for *C. adolfoi* does not necessarily imply that the species is acid-intolerant. The sunfish (*Emeacanthus obsitus*), living in peat bogs at pH 3.7 in North America, also developed massive Na⁺ loss when submitted experimentally to pH 3.5 (14). Body Na⁺ levels declined by 30% after two weeks but remained stable over the following three weeks at pH 3.5. Moreover, the high Na⁺ disruption demonstrated for *C. adolfoi* is very similar to that found for other blackwater species, namely the tetra *Gymnocorymbus ternetzi* (15), and for the neon tetra *Paracheirodon innesi* (16). Like the neon tetra, *C. schwartzi* developed a control mechanism over subsequent periods so as to adapt to low pH. *C. adolfoi* did not show Na⁺ efflux control during the 6-h treatment period, indicating that it may take longer for *C. adolfoi* to adjust to the acid conditions of the transfer from pH 6.0.

It is still not clear why some predicted acid tolerant species that thrive and reproduce at extremely low pH do not show high tolerance under experimental conditions. Blackwater fish species inhabiting the igapó areas are exposed to natural acidity due to humic and fulvic acids, that make up the dissolved organic carbon fraction. Dissolved organic carbon concentration in the flooded forest is about 35 mg/l (2). The carboxylic and phenolic hydroxyl groups of these organic acids are the main source of blackwater acidity (17). Weak acids (such as humic and fulvic types) are not completely ionized in aqueous solution, and typically have a high pKₐ. Conversely, inorganic acids (HNO₃, H₂SO₄, and so forth) such as those used in the present study are almost completely ionized in solution. These conditions do not represent the same type of acidity that fish experience in the wild, even though they have the same [H⁺] in the water.

Given its normal circumneutral habitat in whitewater, the high tolerance of *C. schwartzi* under low pH conditions is quite remarkable. This species controlled both Na⁺ and K⁺ losses well at pH 3.5. In contrast to the characiform matrincha (*Brycon* sp) and tambaqui (*Colossoma macropomum*) (18), *C. schwartzi* seems to be much more adapted to rapid changes in water pH. The efflux rate of matrincha was reduced by 40% after 18 h, but the losses were still significant. *C. schwartzi* showed a 90% decrease in Na⁺ loss after 6 h at pH 3.5, indicating the ability to limit Na⁺ losses faster than matrincha.

In spite of Na⁺ disruption, K⁺ losses were under control for both species. *C. schwartzi* had a slightly higher K⁺ loss than *C. adolfoi*, which decreased over time for both, indicating that they were able to control the disturbance generated by low pH. Jₙₖₑᵣ K⁺ was closely similar to that found in a previous study for other Amazonian fish at pH 3.5 (19). It is worth mentioning that K⁺ efflux is an indicator of intracellular osmotic imbalance, being lost primarily from intracellular media. The fact that K⁺ loss remained under control may indicate that *C. adolfoi* was not experiencing a great intracellular ionregulatory disturbance, which may be the key to survival.
Effects of calcium

Na\(^{+}\) loss in *C. schwartzii* was strongly influenced by [Ca\(^{2+}\)], but this effect was less pronounced in *C. adolfoi*. Although the Na\(^{+}\) loss decreased significantly (about 70\%) in relation to the pH 3.5 treatment, *C. adolfoi* was not able to control Na\(^{+}\) loss, which still increased progressively throughout exposure. Gonzalez et al. (15,19) pointed out that Ca\(^{2+}\) should not be involved to a greater extent in the regulation of branchial permeability at low pH. A high [Ca\(^{2+}\)] (100-500 μmol/l) had almost no effect on ion loss in blackwater species compared to 10 μmol/l [Ca\(^{2+}\)]. It is still unclear whether the acclimation period out of the wild conditions would induce different ionoregulatory patterns in *C. adolfoi*. It is reasonable to propose that Ca\(^{2+}\) does not play any special role in blackwater fish acid tolerance since this element is almost unavailable in the ion-poor waters of the Amazon region. This supports the idea that ion regulation in blackwater systems is modulated by different mechanisms, as previously suggested by others (15,16,18,19). Gonzalez et al. (19) suggested that humic compounds might interact directly with the branchial tight junctions and influence permeability.

In contrast, ionoregulatory control in *C. schwartzii* is clearly Ca\(^{2+}\) dependent, as shown for the tambaqui (*C. macropomum*), a blackwater-whitewater migratory species (12,18, 20,21). This indicates that *C. schwartzii* has a high branchial affinity for Ca\(^{2+}\), particularly at the paracellular tight junctions, thus presenting diffusive ion losses.

The adaptive strategies involved in acid tolerance in the blackwaters are still a topic of discussion. The physiological adjustments that allow fish to inhabit low pH waters with low ion concentrations seem to be their ability to maintain an ionic balance by a combination of adjustments to enhance transport and limit permeability. Most reports seem to emphasize the idea that ion loss control is the key factor for species survival in acid waters, rather than maintenance of the ion uptake rates. Naturally adapted fish from soft ion-poor waters such as the Amazon blackwaters did not show branchial affinity for Ca\(^{2+}\), indicating modulation by other still unknown physiological mechanisms. The investigation of the effect of dissolved organic carbon compounds on ion regulation should be a fruitful field for additional research concerning the natural adaptations of fish to naturally acidic waters.

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References

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