

Ultimobranchial gland of freshwater catfish, *Heteropneustes fossilis*, in response to calcitonin administration¹

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ABSTRACT.- Srivastav A.K., Singh S., Mishra D. & Srivastav S.K. 2009. **Ultimobranchial gland of freshwater catfish, *Heteropneustes fossilis*, in response to calcitonin administration.** *Pesquisa Veterinária Brasileira* 29(12):963-968. Department of Zoology, DDU Gorakhpur University, Gorakhpur 273 009, India. E-mail: ajaisrivastav@hotmail.com

The absence of a hypocalcemic effect of calcitonin (CT) in fishes has been suggested due to exceedingly high plasma levels of CT; the fish may be saturated with respect of circulating CT and therefore unable to respond to exogenously administered CT. Earlier it has been suggested that a hypocalcemic action of injected CT may be obscured by changes in the release of endogenous CT and other calcium regulating hormones. In this study we have used artificial freshwater, calcium-deficient freshwater and calcium-rich freshwater and injected the fish with CT. The aim behind selecting these media were (i) in calcium-deficient medium there would be reduced circulating levels of CT, (ii) in calcium-rich medium there would be diminished secretion of prolactin (this hormone is hypercalcemic in fish), and (iii) by keeping the fish in calcium-rich medium we can test the antihypercalcemic action of CT. Moreover, the present study would reveal the changes in the ultimobranchial gland (UBG) after keeping the fish in all the above three media and/or injecting the fish with CT. Freshwater catfish, *Heteropneustes fossilis*, were administered intraperitoneally daily with vehicle or 0.5 U/100g body wt of salmon calcitonin (CT) and kept in artificial freshwater, calcium-rich freshwater and calcium-deficient freshwater for 10 days. Blood samples were collected on 1, 3, 5, and 10 days following the treatment and analyzed for serum calcium levels. The ultimobranchial gland (UBG) was also fixed for histological studies on these intervals. In artificial freshwater there was no change in the serum calcium levels of calcitonin-injected fish. The ultimobranchial gland of calcitonin-injected fish exhibited a progressive decrease in the nuclear volume from day 5 onwards. On day 10 vacuolization in the gland was also noticed. In vehicle-injected fish (control) kept in calcium-rich freshwater hypercalcemia has been noticed which persists till the end of the experiment. In calcitonin-treated fish maintained in calcium-rich freshwater there is no change in serum calcium level as compared to vehicle-injected fish. In vehicle-injected fish the UBG depicts decreased staining response and increased nuclear volume at day 5. On day 10 the nuclear volume is further increased and few degenerating cells have been noticed. Calcitonin fails to induce any histological change in the UBG as compared to control. In vehicle-injected fish kept in calcium-deficient freshwater the serum calcium levels decrease from day 1 to day 3. The levels exhibit hypercalcemia on day 10. CT treatment to the fish kept in calcium-deficient freshwater evokes a decrease in the calcium levels on day 1 and day 3. A significant hypercalcemia has been noticed on day 5 and day 10. In vehicle-injected fish kept in calcium-deficient freshwater the UBG reveals a decreased staining response on day 10. In CT-injected fish maintained in calcium-deficient freshwater the UBG depicts an increased nuclear volume and few exhausted cells on day 10. It can be concluded that CT can provoke hypocalcemia only when the fish is kept in medium

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which reduces the circulating levels of this hormone. The UBG of the fish kept in different calcemic media responded in a manner to indicate that it produces hypocalcemic factor - CT.

INDEX TERMS: Calcitonin, ultimobranchial gland, calcium, fish, *Heteropneustes fossilis*.

INTRODUCTION

In mammals the hypocalcemic hormone, calcitonin is secreted from the calcitonin (C) cells which are derived from the neural crest and during embryonic development these cells migrate and incorporated into the thyroid and parathyroid IV (Dacke 1979, Srivastav & Rani 1988, Miller 2006). However, these cells remain separate from the thyroid gland in non-mammals and form a discrete organ - the ultimobranchial gland (UBG). In fishes administration of calcitonin (CT) produced conflicting results - no effect, hypocalcemia and hypercalcemia (Wendelaar Bonga & Pang 1991, Chakrabarti & Mukherjee 1993, Oughterson et al. 1995, Srivastav et al. 1998a,b, Mukherjee et al. 2004). This conflict regarding the effect of CT in fishes has led some workers to suggest that CT is involved either with osmotic regulation (Yamauchi et al. 1978), sex-related phenomenon (Yamane & Yamada 1977, Suzuki 2005), skeletal protection during periods of high calcium demands (Wendelaar Bonga & Pang 1991) and to protect scales from excess degradation of calcium at vitellogenesis (Suzuki et al. 2000). CT has also been suggested to control dietary calcium uptake (Suzuki et al. 1999) and inhibit calcium transport in fish gills (Wagner et al. 1997, Mukherjee et al. 2004). As yet no definite functions of CT (Hirsch et al. 2001, Clark et al. 2002, Miller 2006, Nag et al. 2007) have been established and more work is necessary to establish its functions in fishes.

The absence of a hypocalcemic effect of calcitonin (CT) in fishes has been suggested due to exceedingly high plasma levels of CT; the fish may be saturated with respect of circulating CT and therefore unable to respond to exogenously administered CT. Wendelaar Bonga (1980) has suggested that a hypocalcemic action of injected CT may be obscured by changes in the release of endogenous CT and other calcium regulating hormones. In this study we have used artificial freshwater, calcium-deficient freshwater and calcium-rich freshwater and injected the fish with CT. The aim behind selecting these media were (i) in calcium-deficient medium there would be reduced circulating levels of CT, (ii) in calcium-rich medium there would be diminished secretion of prolactin (this hormone is hypercalcemic in fish), and (iii) by keeping the fish in calcium-rich medium we can test the antihypercalcemic action of CT. Moreover, the present study would reveal the changes in the ultimobranchial gland (UBG) after keeping the fish in all the above three media and/or injecting the fish with CT. To the best of our knowledge, there exists no study regarding the effects of CT on the UBG of fish kept in calcium-deficient or calcium-rich media.

In this study we report the effect of calcitonin on the ultimobranchial gland of the freshwater catfish, *Heterop-*

neustes fossilis which were kept in different calcium environments. *H. fossilis* was selected because it is hardy, readily available, easy to handle and can be kept alive for longer duration in the aquaria. This is a common food fish and forms an important species in many water resources mainly ponds, ditches, swamps, marshes and sometimes occurs in muddy rivers (Rainboth 1996). This species is found in India, Pakistan, Sri Lanka, Nepal, Bangladesh, Burma, Thailand and Vietnam (Berra 2007).

MATERIALS AND METHODS

The freshwater catfish, *Heteropneustes fossilis* (body wt. 50±9g and size 16±3.5cm) were collected locally from Ramgarh Lake, acclimatized under laboratory conditions in dechlorinated tap water for two weeks. After acclimation they were divided into six groups each containing 48 fish and subjected to the following treatments:

Group A. Fish were maintained in artificial freshwater and were daily injected intraperitoneally (i.p.) with vehicle (0.1ml of 0.6% NaCl/100g body wt).

Group B. Fish were kept in artificial freshwater and were daily injected i.p. with 0.5 U/100g body wt of salmon calcitonin.

Group C. Fish were kept in calcium-rich freshwater and were injected daily i.p. with vehicle (0.1 ml of 0.6% NaCl/100g body wt).

Group D. Fish were kept in calcium-rich freshwater and injected i.p. daily with 0.5 U/100g body wt of salmon calcitonin.

Group E. Fish were kept in calcium-deficient freshwater and were daily injected i.p. with vehicle (0.1ml of 0.6% NaCl/100g body wt).

Group F. Fish were kept in calcium-deficient freshwater and were daily injected i.p. with 0.5 U/100g body wt of salmon calcitonin.

Calcitonin used in groups B, D and F was dissolved in 0.6% NaCl solution. The fish were not fed 24 h before and during the experiment.

Different artificial media were prepared as follows:

(a) **Artificial freshwater.** Distilled water containing 2.10 mM NaCl, 0.45 mM Na₂SO₄, 0.06 mM KCl, 0.8 mM CaCl₂, 0.20 mM MgCl₂. pH of the solution was adjusted to 7.6 with NaHCO₃.

(b) **Calcium-rich freshwater.** 13.4 mM CaCl₂ was added to the artificial freshwater.

(c) **Calcium-deficient freshwater.** same as artificial freshwater without CaCl₂.

Twelve fishes from each group were anesthetized with MS 222 and blood samples were taken by sectioning of the caudal peduncle 2 h after the last injection on 1, 3, 5 and 10 days of the treatment. The sera were separated and analyzed for calcium according to the method of Trinder (1960). After collection of blood samples, the area adjoining the heart along with the esophagus (as UBG has been found in this region) were removed and fixed in aqueous Bouin's fluid. Tissues thus fixed were routinely processed in graded series of alcohols, cleared in xylene and embedded in paraffin. Serial

sections were cut at 6 μm and stained with hematoxylin-eosin (HE).

Nuclear indexes (maximal length and maximal width) were determined (50 nuclei were measured per specimen, thus 300 nuclei were measured from six specimens) with the aid of ocular micrometer and then the nuclear volume was calculated as:

$$\text{Volume} = 4/3 \pi ab^2$$

Where 'a' is the major semiaxis and 'b' is the minor semiaxis.

Data are reported as mean ± S.E. for six specimens (although 12 specimens were taken for each group but data from 6 specimens were taken into account) and Student's t test was used to determine statistical significance. Each experimental group was compared to its specific time control group.

RESULTS

(i) Artificial freshwater (groups A and B)

The serum calcium levels of calcitonin-treated fish (group B) depicted no significant change as compared to those of vehicle-injected specimens (group A) (Fig.1).

In vehicle-injected fish (group A) the UBG usually possessed a solid parenchyma which contained cell cords

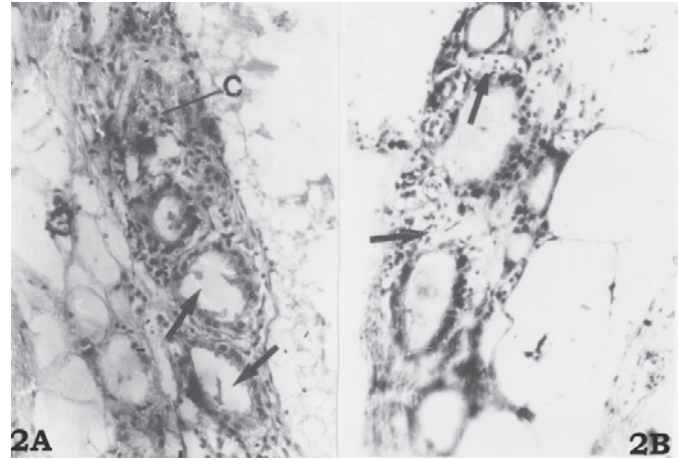


Fig.2. (A) Ultimobranchial gland of 5 days vehicle-injected (control) fish maintained in artificial freshwater. Note the presence of follicles (arrows) and cords (C). HE, 200x. (B) Ultimobranchial gland of 10 days calcitonin-treated fish kept in artificial freshwater showing vacuolization (arrows). HE, 200x.

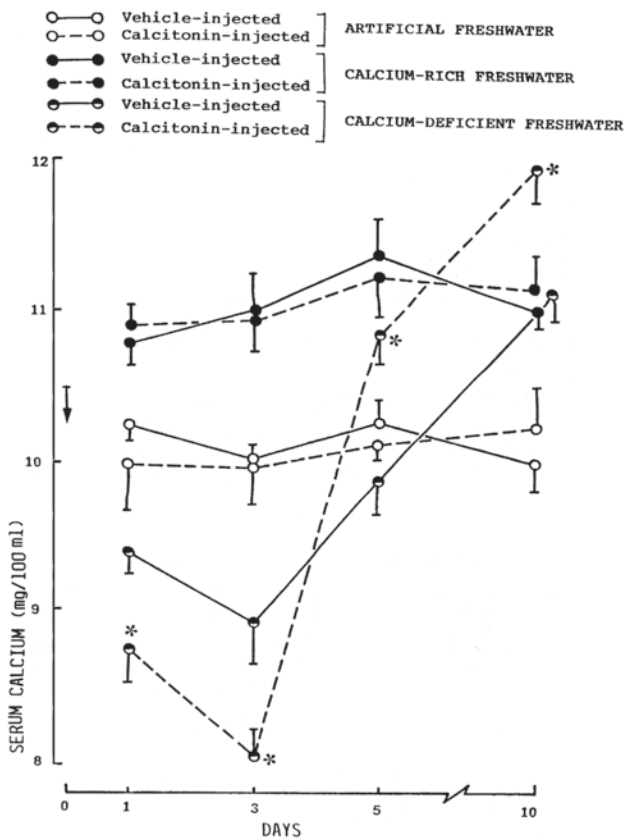


Fig.1. Changes in the serum calcium levels of vehicle (control) or calcitonin-treated (experimental) fish maintained either in artificial freshwater, calcium-rich freshwater or calcium-deficient freshwater. Each value represents mean ± S.E. of six specimens. Asterisk indicates significant differences from vehicle-injected group (P<0.05). Data of calcitonin-treated (experimental) fish were compared with vehicle-injected (control) fish in each medium. Only data which are significant are marked with an asterisk.

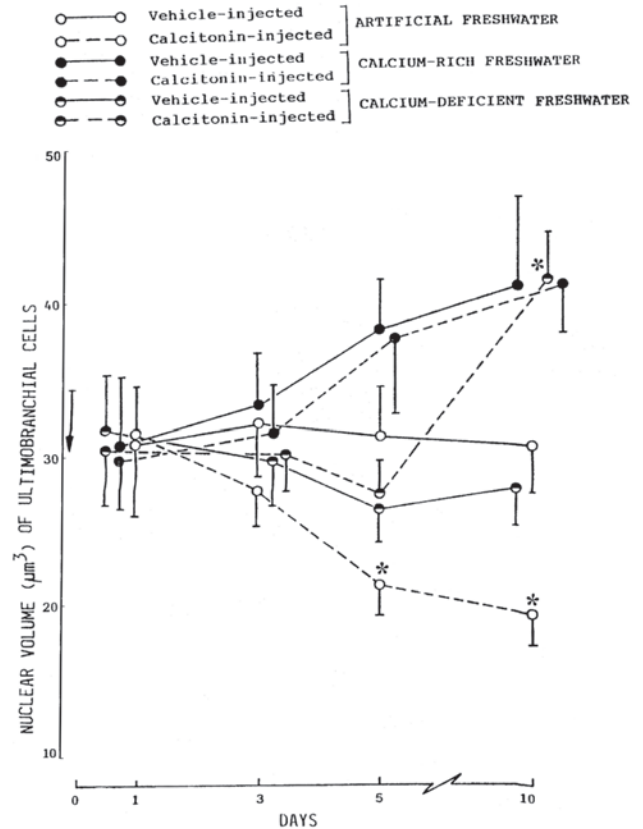


Fig.3. Nuclear volume of ultimobranchial cells of vehicle (control) or calcitonin-treated (experimental) fish maintained either in artificial freshwater, calcium-rich freshwater or calcium-deficient freshwater. Each value represents mean ± S.E. of six specimens. Asterisk indicates significant differences from vehicle-injected group (P<0.05). Data of calcitonin-treated (experimental) fish were compared with vehicle-injected (control) fish in each medium. Only data which are significant are marked with an asterisk.

and small follicles (Fig.2 A). All the cells were alike. Their cell boundaries were indistinct. When stained with HE, the cytoplasm of these cells was noticed slightly eosinophilic. The nuclei were generally rounded, oval or vesicular in shape.

Uptil day 3 following the CT treatment (group B) there was no significant histological change in the UBG. From day 5 onwards there was a progressive decrease in the nuclear volume of UBG cells (Fig.3) and degenerative changes (vacuolization of cytoplasm) were seen on day 10 (Fig.2B).

(ii) Calcium-rich freshwater (groups C and D)

Serum calcium level in the specimens of group C (vehicle-injected) showed a progressive increase up to day 5. On day 10, a slight decrease in the serum calcium level was recorded though it still remained hypercalcemic (Fig.1).

CT treatment (group D) was ineffective regarding serum calcium level which was more or less similar to that of group C (Fig.1).

Up to day 3 following vehicle-injection (group C), there was no change in the histological details of UBG. The UBG depicted a decrease in the staining response (Fig.4A) and an increase in nuclear volume on day 5 (Fig.3). On day 10, the gland became compact, the nuclear volume increases (Fig.3) and a few degenerating cells were discerned (Fig.4B).

CT failed to induce any histological change in group D as the histological structure of UBG was similar to the gland of vehicle-injected specimens.

(iii) Calcium-deficient freshwater (groups E and F)

A slight decrease in the serum calcium level of vehicle-injected fish (group E) was recorded on day 1 (Fig.1). This

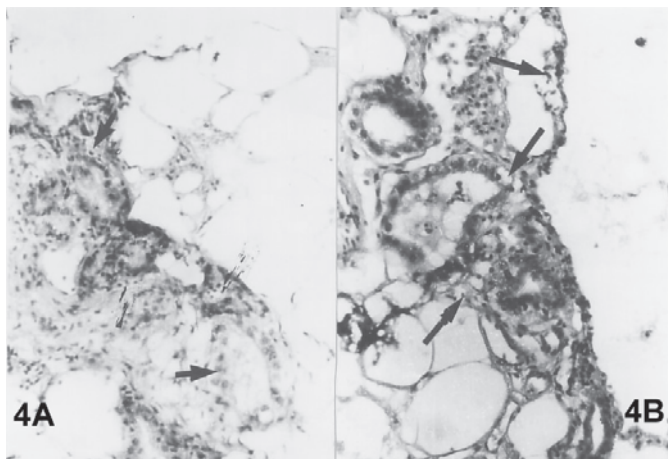


Fig.4. (A) Ultimobranchial gland of 5 days vehicle-injected (control) fish kept in calcium-rich freshwater showing decreased staining response (arrow). HE, 200x. (B) Ultimobranchial gland of 10 days vehicle-injected (control) fish kept in calcium-rich freshwater. Note certain degenerating cells (arrows). HE, 200x.

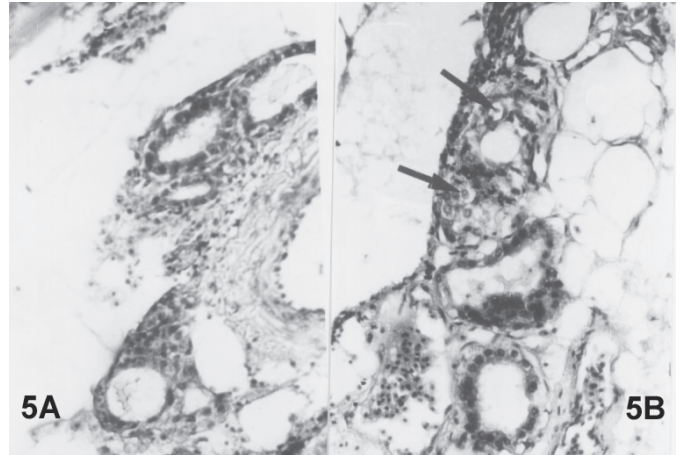


Fig.5. (A) Decreased staining response of the ultimobranchial gland of 10 days vehicle-injected (control) fish kept in calcium-deficient freshwater. HE, 200x. (B) Ultimobranchial gland of 10 days calcitonin-injected fish kept in calcium-deficient freshwater exhibiting few exhausted cells (arrows). HE, 200x.

response continued on day 3 (Fig.1). From day 5 the levels showed a trend to increase which resulted into hypercalcemia on day 10 (Fig.1).

CT treatment (Group F) evoked a decrease in the serum calcium level on day 1 which persisted till day 3. Thereafter, the value exhibited a significant hypercalcemia on day 5. This response increased progressively till day 10 (Fig.1).

No change was noticed in the UBG of vehicle-injected fish (group E) uptil day 5. However, there was a decrease in the staining response of the UBG cells on day 10 (Fig.5A). The nuclear volume of these cells displayed no significant change (Fig.3).

In CT treated fish (group F) the UBG cells remained unaffected till day 5. However, on day 10 increased nuclear volume (Fig.3) and few exhausted cells (Fig.5B) were noticed.

DISCUSSION AND CONCLUSION

In the present study, the administration of CT to *Heteropneustes fossilis* maintained in artificial freshwater causes inactivity in the UBG which is evident by decreased nuclear volume of UBG cells. This is in accord with similar findings reported by Peignoux-Deville et al. (1975), Wendelaar Bonga (1980) and Srivastav et al. (1989) as these investigators have also noticed inactivity in the gland after CT administration. The vacuolization observed in CT administered *H. fossilis* can be associated with the continuous disuse of the UBG cells in response to CT administration.

In the present study the fish subjected to calcium-deficient freshwater and/or CT treatment exhibit an increased activity of UBG on day 10. This may be due to the observed elevation of serum calcium level at this duration. No difference has been noticed in the changes in UBG of *H. fossilis* whether they are treated with either

calcium chloride or calcium chloride and CT. In both these treatments, the UBG exhibits hyperactivity which can be attributed to the increased calcium levels. To the best of our knowledge, there exists no study regarding the effects of CT on the UBG of fish kept in calcium-deficient or calcium-rich media. Hence, in this regard the findings of the present study are new and could not be compared with the observations of other investigators. Enhanced activity/secretion of CT from the fish UBG has been reported in the past by several investigators in response to experimentally induced hypercalcemia (Lopez et al. 1968, Lopez & Bagot 1971, Chan 1972, Lopez 1973, Peignoux-Deville et al. 1975, Swarup & Srivastav 1984, Fouchereau-Peron et al. 1986, Suzuki et al. 1999, Kaida & Sasayama 2003). Hyperactivity of ultimobranchial gland after induced hypercalcemia has also been reported from non-piscine vertebrates (Ziegler et al. 1970, Dubewar et al. 1978, Suzuki et al. 1982, 1984, Srivastav & Rani 1991, 1992, Srivastav et al. 1994, 2008, Srivastav & Yadav 2008, Yadav & Srivastav 2008).

It is concluded from this study that (i) CT provokes hypocalcemia in fish only when they are kept in a medium which can reduce the circulating levels of CT, (ii) the UBG of normal fish respond to CT administration by depicting inactivity, and (iii) UBG gets activated when the fish is kept in calcium-rich medium and this activity of UBG is not influenced by administration of CT.

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