Bolm. Zool., Univ. S. Paulo 8:77-91, 1984

EFFECT OF VARIATION OF SALINITY ON PROTEIN, RNA AND DNA CONTENTS OF LIVER, MUSCLE AND OVARY OF FEMALE SINGI FISH, HETEROPNEUSTES FOSSILIS (BLOCH), AT TWO PHASES OF REPRODUCTIVE CYCLE

A.K. DASMAHAPATRA AND A.K. MEDDA

Department of Animal Physiology, Bose Institute, P-1/12, CIT Sche me VII M, Kankurgachi, Calcutta 700054. (recebido em 16.IX.1983)

RESUMO - Investigou-se as mudanças causadas nos teores de proteínas e ácidos nucleicos do fígado, músculo, ovários e peso dos órgãos (HSI, GSI) pela variação de salinidade do meio (65, 135 e 225 mOsm, NaCl/litro, e salinidade zero ou a gua destilada) em fêmeas não vitelogênicas (NV) e vitelogêni cas (V) do peixe Heteropneustes fossilis (Bloch) às tempera-turas de 25°C e 30°C, 30 dias após o período de aclimação nos respectivos meios. Obteve-se valores máximos de HSI tanto nos peixes (NV) e (V) quando mantidos em água destilada e valores mínimos quando em solução de 225 mOsm. O aumento de temperatura de 25°C para 30°C causou uma redução nos valores de HSI somente nos peixes (V) Salinidade de 135 e 65 - 135mOsm produziu o valor mais alto de GSI nos dois grupos de peixes: (NV) e (V) 0 efeito estimulante da alta temperatura $(30^{\circ}C)$ no GSI foi encontrado somente nos peixes (NV) em to dos os meios salinos. O teor hepático de proteínas e RNA foi máximo nos peixes (NV) e (V) mantidos em meios de 65 mOsm de NaCl e mínimo nos de 225 mOsm de salinidade. O aumento de temperatura de 25°C para 30°C não alterou o teor proteico he pático mas aumentou e diminuiu o teor de RNA nas fases (V) e (NV). O teor muscular de proteínas e RNA foram máximos a sa-linidades zero e 65 mOsm e mínimos a 225 mOsm tanto a 25°C, como 30°C. A alta temperatura (30°C) aumentou o teor proteico, mas não alterou o teor de RNA. Salinidade de 135 mOsm causou o maior aumento nos teores de proteínas, RNA e DNA do ovário das fêmeas (NV) a 25°C ou 30°C. Salinidade zero reduziu bastante o teor desta substância no ovário. No caso do GSI o efeito estimulador da alta temperatura (30° C) no acúmu lo de proteínas, RNA e DNA se fez sentir em todas as concentrações salinas do meio. Os peixes (V) apresentaram teor má ximo de proteínas ovarianas a salinidade de 135 mOsm e mínimos em água destilada. Os ovários (V) tiveram a taxa mais al ta de DNA em 65 e 135 mOsm de salinidade do que a zero ou 225 mOsm. A influência das altas temperaturas não foi sentida nas fêmeas (V) O teor de RNA ovariano dos peixes (V) tam bém não se alterou em todas as condições experimentais.

ABSTRACT - The changes in protein and nucleic acid contents of liver, muscle and ovary and organ weight (HSI, GSI) cau sed by variation of salinity of the medium (65, 135 and 225 mOsm NaCl/liter, zero salinity or distilled water) were in vestigated on non-vitellogenic (NV) and vitellogenic (V) fe-male Singi fish (Heteropneustes fossilis Bloch) at 25° and and 300 after 30 days acclimation in the respective medium. The HSI was maximum in (NV) and (V) fish kept in distilled water and minimum in 225 mOsm NaCl solution. Rise of temperature from 25°C to 30°C caused reduction in HSI only in (V) fish, temperature but not in (NV) one. The salinity of 135 and 65-135 mOsm produced highest GSI in (NV) and (V) fish respectively. A stimu lating effect of higher temperature (30°) on GSI was found in only (NV) fish in all saline media. The amount of liver protein and RNA was maximum in (NV) and (V) fish kept in 65 mOsm NaCl medium and minimum in 225 mOsm salinity. Rise of temperature from 25°C to 30°C did not alter the liver pro tein content, but increased and decreased the RNA content at (V) and (NV) stage respectively. The muscle protein and RNA contents were maximum in zero salinity and 65 mOsm NaCl me dium, and minimum in 225 mOsm saline group at 25°C or 30°C Higher temperature (30°C) increased the muscle protein con tent, but not the RNA. The salinity of 135 mOsm caused maximum increase in protein, RNA and DNA contents of the (NV) ovary at 25°C or 30°C. The zero salinity has a very depres -sing offect on these ovarian substances. As in case of GSI , a stimulating effect of higher temperature (30° C) on the ac-cumulation of protein, RNA and DNA in (NV) ovary was noted noted in all saline media. The (V) fish showed maximum ovarian pro tein content in 135 mOsm salinity and minimum in distilled water. The (V) ovary had higher DNA content in 65 and 135 mOsm salinity than that in zero or 225 mOsm salinity. The influence of higher temperature on (V) ovary was absent. The ovarian RNA amount of (V) fish was not altered under the experimental conditions.

INTRODUCTION

In fish, an effective mechanism for osmotic regulation is necessary for maintaining the normal life processes. Sali nity of water is one of the most important environmental fac tors which control metabolism, survival and distribution $oar{f}$ fish (Holliday, 1969). Investigations have been made on the salinity tolerance and osmotic and ionic regulation in which there is a good deal of metabolic involvement of different kinds (Bashamohideen and Parvatheswararao, 1972; Venkatachari, 1974; Bhan and Mansuri, 1978; Love, 1980). Several re ports are also available on the effects of different concentrations of saline on the nitrogen metabolism in euryhaline fish (Lecal, 1958; Jones, 1959; Cowey et al., 1962; Cowey and Parry, 1963; Parvatheswararao, 1967; Huggins and Colley, 1971; Venkatachari, 1974; Mansuri and Bhan, 1978). Since the fresh water fishes are hypertonic to the surrounding water , their body organs are also likely to be affected or influen-

.

ced by the change in osmotic concentration of the medium. The salinity tolerance of Singi fish (*Heteropneustes fossilis* Bloch), a stenohaline fish, has been reported (Al Daham and Bhatti, 1977; Parwez *et al.*, 1979). But to our knowledge, the informations on the changes in different cellular constitu ents of different organs, such as liver, muscle and ovary of fresh water or stenohaline fish after adaptation in diffe rent concentrations of sodium chloride solution are inadequa te.

In view of these facts, investigations have been under taken on the effects of different concentrations of sodium chloride on different cellular components of liver, muscle and ovary of female Singi fish (*Heteropneustes fossilis* Bloch). The Singi fish of two phases of reproductive cycle, non-vitellogenic and vitellogenic, were used for the experiments in order to show the influence of reproductive stage on the changes in cellular components of different organs. In the present communication, the data on the changes in organ weight and protein, RNA and DNA contents of liver, muscle and ovary are presented.

MATERIALS AND METHODS

The female Singi fishes (Heteropneustes fossilis Bloch) of body weight 40-60 g (length 20-22 cm) were purchased from a local supplier and acclimatized in the laboratory condi-tions at 25° [±]1°C for 7 days before the experiments. The ovary of the non-vitellogenic (NV) fish contained only Stage I oocytes, whereas that of the vitellogenic (V) fish was loa ded with Stage III oocytes. The experimental non-vitelloge nic (December-January) and vitellogenic (June-July) fishes were distributed at random in different groups of 15 each for acclimatizing them to four different concentrations of sodium chloride solution, e.g., 65 mOsm, and 225 mOsm/liter. The same number of fish (15) were kept in distilled water (0 salinity) Each fish was kept in 1 liter of the respective medium in a glass jar with 12L : 12D photoperiod and fed ad libitum with Tubifex tubifex. The medium was changed at every alternate day. The fishes were acclimatized in this wav in the respective medium (sodium chloride solution of concentration or distilled water) at 25°C and 30°C for each 30 days, after which they were sacrificed for the estimation of protein, RNA and DNA contents of liver, skeletal muscle and ovary. Tissue protein was estimated by following the method of Lowry et al. (1951), RNA by the method of Mejbaum (1939) as modified by Munro and Fleck (1966) and DNA by the method of Burton (1956) as modified by Croft and Lubran (1965). The hepatosomatic index (HSI) and the gonadosomatic index (GSI) were also calculated: weight of the organ x 100/body weight.

Since the weight of the liver and ovary of Singi fish undergoes a seasonal variation (Dasmahapatra and Medda, in press) and is also affected by change of environmental tempe rature (Dasmahapatra, 1980), the results of the amounts of protein, RNA and DNA of liver and ovary were first calcula ted per 100 mg of fresh tissue, and then the final results of the amounts of these cellular substances were expressed

and in the hepatosomatic index (HSI) and gonadosomatic index (GSI) of female (non vitellogenic vitellogenic) Singi fish (*Heteropneustes fossilis* Bloch). The fishes were fed and kept the respective medium at 25°C or 30°C for 30 days. Each group consisted of 15 fishes. Table 1 - Effect of variations of salinity (0, 65, 135, 225 milliosmole NaCl per liter)

Temperature Mon-vitellogenic (fishes of December-January) vitelugenic vitelection 0^{C} Image: SiE Salinity (milliosmole per lifer) 135 225 (Distilled 55 135 0^{C} (Distilled 55 135 225 (Distilled 55 135 0^{C} (Distilled 55 1.08 bA 0.94 bBa 2.01 1.94 a 1.70 aB 25^{O} 1.49 1.25 b 1.07 bA 0.94 bBa 2.01 1.94 a 1.70 aB 25^{O} 1.49 1.25 b 1.07 bA 0.93 bBa* 1.86 * 1.70 aB 1.70 aB 25^{O} 1.32 *** 1.07 bA* 0.93 bBa* 1.86 ** 1.70 aB 1.72 bA** 25^{O} 1.32 ** 1.00 b* 0.00 2.01 b* 1.22 b** 1.70 aB 1.72 b** 25^{O} 1.35 ** 1.86 ** 1.86 ** 1.72 b** 1.12 b** 25^{O} 0.010 1.23 ** 1.25 ** 1.12 ** 1.25 ** 25^{O} 0.02	-						Keproductive pndse	t + all arrante (t)	tenes of duna-	(Alu)
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	-	Temperature	Non-vitello	genic (fishes t	of December-Jan	iuary)	~	TLETTOREUTC		
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	_					Salinity (mil.	liosmole per li	ter)		
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$		°	0 (Distilled	65	135	225	0 (Distilled water)	65	135	225
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$			Mean + S.E.		Mean ± S.E.	Mean ± S.E.	Mean <u>+</u> S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	1	250	1.0+0 +0.0		1.08 ^{bA}	0.94bBa + 0.02	2.01 <u>+</u> 0.05	1,94 ^a + 0,04	1.70 ^{aB} <u>+</u> 0.03	1.50 ^{bB8} + 0.06
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	- ISH	300	1.32*** + 0.03	* ⁵ 0.11	1.07 ^{bA*} ± 0.04	0.93 bBa*	1.86** + 0.04	1,70 ^{4***} <u>+</u> 0.05	1.52 ^{bA***} <u>+</u> 0.05	1.35 ^{bBB**} <u>+</u> 0.02
30 ⁰ 0.98 ⁴ 1.86 ^b ⁴⁴ 3.45 ^b B ⁴⁴⁴ 1.52 ^a N ^{a4} 8.71 ⁴ 13.22 ^{b^{44,4} 14.62^{bN^{46,4}} 1.52^{b^{44,4}} 14.62^{bN^{46,4}} ± 0.83 ± 0.20 ± 0.12 ± 0.19 ± 0.15 ± 0.46 ± 0.38 ± 0.83}		250	0.72 + 0.14	1,35 ⁸ + 0,15	2.01 ^{bA} + 0.28	1.22 ^{aNa} + 0.18	8.15 + 0.28	11.20 ^b	11.25 ^{bN} <u>+</u> 0.24	9,00,18 4,0,41
	E S	300	0.98* + 0.20	1.86 ^{b**} ± 0.12	3.45 ^{bB##*}	1.52 ^{aN u*} ± 0.15	8.71* <u>+</u> 0.45	13.22 ^{b***} ± 0.38	14.62 ^{bN***}	9.20 ^{nBB*} <u>+</u> 0.33

Table 2 - Effect of variations of salinity (0, 65, 135, 225 milliosmole NaCl per liter) on the protein content of liver, muscle and ovary of female (non-vitellogenic and vitellogenic) Singi fish (*Heteropneustes fossilis* Bloch). The fishes were fed and kept in the respective medium for 30 days at 25[°]C or 30[°]C temperature. Each group consisted of 15 fishes.

Tempe- Non-vitellogenic (fishes of June-July) rature Salinity (millosmole per liter) Parture Salinity (millosmole per liter) C (Distilled (Distilled Mean $\pm S.E.$ Mean $\pm S.E$					Reprodu	Reproductive phase				
rature Salinity (milliosmole per iter) salinity (milliosmole per iter) °C (0:010 uater) water) 65 135 225 135 135 °C (0:01 uater) 65 135 225 135 135 Rean ± S.E. Mean ± S.E.		Tempe-	Non-vite]	logenic (fis	thes of Decemb	er-January	Vitellog	cenic (fishes	of June-July	-
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$		rature			Salinity (mil	liosmole per l	iter)			
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$		°c	D (Distilled water)	65	135	225	0 (Distilled water)	65	135	225
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$			Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean + S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Liver protein (mg/100g of bodv	250	70.30	81.54 ⁸ <u>+</u> 3.47	56,95 ^{bB} + 2,58	45.74 bBa <u>+</u> 3.82	150.28 <u>+</u> 5.78	166.38 ⁴ <u>+</u> 2.61	131.45ªB ± 5.35	119.56 ^{bBy} + 4.68
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	weîghtĴ	300	74.38* <u>+</u> 4.70	88,54ª* <u>+</u> 3,48	53.83 ^{bB‡} <u>+</u> 4.68	35,48 ^{bB6} * <u>+</u> 2,21	157.48* <u>+</u> 2.19	171.88 ^{b*} ± 2.52	142.45 ^{bB*} <u>+</u> 3.52	124.71 ^{hB8 *} <u>+</u> 2.67
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Muscle protein (mg/100mg	25 ⁰	12.97 <u>+</u> 0.43	12.40 ⁿ <u>+</u> 0.39	10.48 ^{aB} ± 0.50	8.68 ^{bB8} <u>+</u> 0.37	14.88 <u>+</u> 0.46	14.38 ⁿ ± 0.52	12,48 ^{bA} <u>+</u> 0.44	8.82 ^{bB8} <u>+</u> 0.35
n 25 ⁰ 49.78 202.53 ^b 384.48 ^{bB} 192.28 ^{bN8} 789.00 1682.45 ^b 1888.00 ^{bA} <u>±</u> 3.87 <u>±</u> 10.15 <u>±</u> 17.40 <u>±</u> 13.19 <u>±</u> 34.38 <u>±</u> 76.63 <u>±</u> 53.83 30 ⁰ 65.45** 482.45 ^{b***} 611.44 ^b 8*** 322.36 ^b B8*** 882.45* 1792.00 ^{b*} 2064.45 ^{b8*} <u>±</u> 2.16 <u>±</u> 27.57 <u>±</u> 17.14 <u>±</u> 22.90 <u>±</u> 71.80 <u>±</u> 56.95 <u>±</u> 77.64	urresn tissue)	30 ⁰	14.08** <u>+</u> 0.28	14,44 ^{n**} <u>+</u> 0.55	10.89 ^{bA*} <u>+</u> 0.36	8.46 ^{bB8} * <u>+</u> 0.45	16.82** <u>+</u> 0.34	16.48 ^{n**} ± 0.17	12,88 ^{bB*} ±0,32	8.02 ^{bB8} * <u>+</u> 0.41
30° 65.45*** 482.45 ^{b***} 611.44 ^b B*** 322.36 ^b B8*** 882.45* 1792.00 ^{b*} 2064.45 ^b B* <u>2</u> 2.16 <u>2</u> 27.57 <u>1</u> 7.14 <u>2</u> 2.90 <u>1</u> 71.80 <u>5</u> 5.95 <u>7</u> 7.54	Ovary protein (mg/100 of body weight)	25 ⁰	.87	202.53 ^b <u>+</u> 10.15	384 48 ^{bB} <u>+</u> 17 40	192,28 ^{bN6} <u>+</u> 13,19	789.00 <u>+</u> 34.38	1682.45 ^b <u>+</u> 76.63	1888.00 ^{da} <u>+</u> 53.83	1207,55 ^{bB8} <u>+</u> 70.48
		300		482.45 ^{b***}	611.44 ^{bB***} ± 17.14	322.36 ^{bB6***} <u>+</u> 22.90	882.45* ± 71.80	1792.00 ^{b*} <u>+</u> 56.95	2064.45 ^{bB*} <u>+</u> 77.64	1282.82aBB* ± 95.48

per 100 g of body weight of fish to eliminate the error, if any, caused by change in organ weight. But, since it was dif ficult to estimate the total amount of skeletal muscle in the body, the results of protein, RNA and DNA contents of muscle were expressed per 100 mg of fresh tissue. Statisti cal analysis of the data were made using student's 't' test. In each case, the mean data were the average from 15 fishes.

RESULTS

1. Changes in weight of liver (HSI) and ovary (GSI), Table 1.

The vitellogenic (V) fish had higher HSI and GSI than non-vitellogenic (NV) fish. Increase in NaCl concentration of the medium from 65 to 135 mOsm/liter and from 135 to 225 mOsm/liter caused a reduction in HSI in each case in both(NV) and (V) fish at 25° C and 30° C. The HSI was consequently the lowest in both (NV) and (V) fish at 25°C and 30°C when the salinity was raised to 225 mOsm. The increase in temperature from 25°C to 30°C exerted no significant influence on HSI in (NV) fish kept in different concentrations of NaCl. But (V) fish showed less HSI at 30° C in each saline medium comparison to that at 25° C. Surprisingly, Singi fishes the in of both the reproductive stages kept in distilled water showed maximum liver weight, the HSI being higher at 25°C than 30°C.

Addition of NaCl or increase in osmolarity of NaCl in the medium within certain limits caused enhancement in GSI in (NV) and (V) fish. At (NV) stage, the GSI was minimum in fish kept in distilled water, maximum in fish kept in 135 mOsm NaCl solution and intermediate in 65 and 225 mOsm solutions at 25 C and 30 C. In (V) fish, GSI increased NaC1 to the maximum level when they were maintained in 65 and 135 mOsm NaCl solutions, it was the lowest in distilled water and in 225 mOsm NaCl solution. The rise of temperature from 25° C to 30° C enhanced the GSI in (NV) as well as (V) fish maintained in 65 and 135 mOsm NaCl solutions.

2 Changes in protein content of liver, muscle and ovary (Table 2)

The protein content of liver and ovary was at higher level in (V) fish in comparison to (NV) one. The muscle protein content was also higher in (V) fish, except fish in kept in 225 mOsm NaCl solution. The amount of liver protein increased to the maximum level in (NV) and (V) fish kept in 65 mOsm NaCl solution both at 25°C and 30°C, while its a - mount was minimum in fish maintained in 225 mOsm NaCl solu tion at these temperatures. The level of liver protein in (NV) and (V) fish kept in distilled water was higher in comparison to that in fish maintained in 135 and 225 mOsm NaCl solutions. Although there was a reduction in protein content of liver with the increase in concentration of NaCl solution from 65 mOsm, the rise of temperature from 25°C to 30°C had no effect on liver protein amount in fish maintained in any medium.

ter)	r Je	I S	
Table 3 - Effect of variations of salinity (0, 65, 135 and 225 milliosmole NaCl per liter)	on the RNA content of liver, muscle and ovary of female (non-vitellogenic and vitelloge -	nic) Singi fish (Hetgropneustes fossilis Bloch). The fishes were fed and kept in the res -	
Cl pe	vi t	in	
e Na(and	kept	shes
smol	genic	and	[5 fi
illic	ellog	fed	of
25 m	-vit	were	sted.
and 2	(nor	shes	consi
135	male	le fi	dno
65,	off∈	, 1	ch gr
.0)	'ary	loch	Еа
inity	vo pu	<i>lis</i> E	days
sal	ile ai	0881	r 30
lo sr	musc	tes f	C fo
atior	ver,	isnou	r 30,
vari	of li	erop	0
t of	ento	(Het	at 25
ffec	cont	fish	mni
ц - Е	RNA.	ingi	e me c
ble	the	C) S	Ctiv
Ε	<u>г</u> .	Ľ	р о

				keprodu	keprouuctive pnase				
	Tempe -	Non-vite	llogenic (fis	Non-vitellogenic (fishes of December-January)	er-January)	Vitellog	Vitellogenic (fishes of June-July)	of June-July	•
	rature			Salinit	Salinity (milliosmole per liter)	e per liter)			
	°	D (Distilled water)	តភ	135	225	0 (Distilled water)	65	135	225
		Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.
Liver RNA (mg/100 g of body	250	7.69 <u>+</u> 0.37	9, 58 ^b + 0, 34	6.55 ^{nB} <u>+</u> 0.67	3.70 ^{bB8} <u>+</u> 0.13	13.45 <u>+</u> 1.26	17.28 ^a <u>+</u> 0.48	11.48 ^{nB} + 0.45	7.45 ^{aBB} <u>+</u> 0.80
WEIGHT	30 ₀	5,448*** - 0,36	7.45b*¢ <u>+</u> 0.32	4,99nB** + 0.36	1.78 ^{bB8**} <u>+</u> 0.90	17.66** <u>+</u> 1.45	20.88 ^{2 **} <u>+</u> 1.18	16.06nA** <u>+</u> 1.75	11.28 ^{bB8 **} <u>+</u> 1.19
Muscle RNA vg/100 mg	250	84.28 <u>+</u> 4.76	86.45 ⁿ <u>+</u> 3.47	52,48 ^{bB} <u>+</u> 3.02	40.10 ^{bBa} <u>+</u> 3.42	52.44 <u>+</u> 4.28	49.28 ⁿ <u>+</u> 3.31	33,88 ^{bB} <u>+</u> 1.02	28,00 ^{bB8} ± 1.45
of fresh tissue	300	88 * + + + * *	90,46 ^{n*} + 4,46	60.48 ^{b8*} + 3.46	45.88bBa* + 4.61	51.87* <u>+</u> 3.00	46.45 ^{n*} + 3.64	36.12 ^{bA*} <u>+</u> 2.19	25.01 ^{bB6 *} <u>+</u> 1.62
Ovarian RNA (mg/100 g of body	250	12.72 <u>+</u> 0.93	15.71 ⁴ <u>+</u> 0.81	19.35 ^{bB} <u>+</u> 0.18	15.21 ^{bNa} <u>+</u> 0.88	35.19 <u>+</u> 2.81	37.84 ^л <u>+</u> 2.86	37.06 ^{nN} <u>+</u> 1.95	36.45 ^{nNY} ± 3.75
weight)	300	13.25# <u>+</u> 0.82	18.27 ^{b**} ± 0.45	22.00 ^{bB****} <u>+</u> 0.76	18.81 ^{aA6**} <u>+</u> 0.86	39.10 ⁴ <u>+</u> 1.44	44.48 ^{n*} + 3.04	42.45 ^{nN*} <u>+</u> 2.35	38.20 ^{nNY *} <u>+</u> 2.45
S.E. Stan significant temperature	Standard error. cant. Alphabets tures (25°C or 3	or. 't' test p ats denote com or 30°C) and s	probability di aparison betwe stars denote o	ifference : 'F sen different comparison bet	S.F. Standard error. 't' test probability difference : 'P' value : a,A,a, ^{4*} P<.05; b,B,B, ^{4***} = P<.01; n,N,Y, ⁴ Not significant. Alphabets denote comparison between different saline groups (including distilled water) at a particular temperatures (25°C or 30°C) and stars denote comparison between different temperature groups (25°C and 30°C) at a parti-		P<.05; b,B,B,8,*** = P<.01; n,N,Y,* ing distilled water) at a part ature groups (25°C and 30°C) at a	P<.Dl; n.N.Y.) at a par and 30°C) at	N,Y,* Not particular at a parti

Increase in saline concentration from 65 to 135 m0sm and from 135 to 225 m0sm caused reduction in muscle protein content in each case both at 25°C and 30°C in (NV) and (V) fish. The protein content of muscle was about the same in fish kept in distilled water and in 65 m0sm NaCl solution at 25° C or 30° C at these two reproductive stages, but the hi – gher temperature (30° C) increased the muscle protein content in these fishes. The muscle protein was at the lowest level in (NV) and (V) fish kept in 225 m0sm NaCl solution at 25° C and 30° C.

The (NV) fish had markedly less protein content of ovary than (V) fish kept in all media. The ovarian protein level was the lowest in 0 salinity at 25°C or 30°C in both(NV) and (V) fish. It then gradually increased when the salinity of the medium was raised to 65 mOsm and 135 mOsm, but with the rise of salinity to 225 mOsm from 135 mOsm the ovarian protein amount was reduced in both (NV) and (V) fish at $25^{\circ}C$ and $30^{\circ}C$. The increase in temperature from $25^{\circ}C$ to $30^{\circ}C$ caused enhancement of protein content of ovary in (NV)fish kept in all media, while in (V) fish such influence of higher tem perature ($30^{\circ}C$) was not found.

 Changes in RNA content of liver, muscle and ovary (Table 3)

Like protein, the RNA content of liver and ovary was found to be higher in (V) fish. But the muscle RNA content was more in (NV) fish. Addition of NaCl in the medium to а concentration of 65 mOsm caused an increase in liver RNA content in (NV) and (V) fish at 25° C or 30° C in comparison to the distilled water group. Increase in saline concentration from 65 to 135 mOsm decreased the liver RNA content in (NV) and (V) fish. The lowest amount of liver RNA was found at both (NV) and (V) stage in 225 mOsm NaCl medium at 25°C or 30°C. The rise of temperature from 25°C to 30°C caused reduc tion and enhancement of liver RNA content in (NV) and (νΣ fish respectively in all media.

The RNA content of muscle remained at about the same level in (NV) or (V) fish kept in 0 salinity and in 65 m0sm NaCl solution at 25°C or 30°C. Increase in saline concentration from 65 to 135 m0sm and from 135 to 225 m0sm decreased the muscle RNA content in each case in both (NV) and (V) fish at 25° C or 30° C. The rise of temperature from 25° C to 30° C did not cause any change in muscle RNA content in any fish in any medium under study.

The RNA content of ovary gradually increased with the increase in salinity of the medium up to 135 mOsm in (NV) fish at 25°C or 30°C. At V stage the ovarian RNA content was almost at the same level in all media at 25°C and 30°C. The influence of higher temperature (30°C) on the enhancement of ovarian RNA content was observed in (NV) fish in all NaCl me dia, but in (V) fish there was no significant change in ovarian RNA by higher temperature.

Table 4 - Effect of variations of salinity (0, 55, 135 and 225 milliosmole NaCl per liter) on the DNA content of liver, muscle and ovary of female (non-vitellogenic and vitellogenic) Sin-gi fish (*Heteropnewstes fossilis* Bloch). The fishes were fed and kept in the respective me -dium at 25^oC or 30^oC for 30 days. Each group consisted of 15 fishes.

				Reprodu	Reproductive phase				
	Tempe-	Non-vit	Non-vitellogenic (fishes	shes of Decem	of December-Jan.)	Vitel	Vitellogenic (fishes of June-July)	ies of June-J	uly)
	rature			Salini	Salinity (milliosmole per liter)	le per liter)			
	°u	0 (Distilled water)	65	135	225	0 (Distilled water)	65	135	225
		Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean ± S.E.	Mean + S.E.	Mean ± S.E.
Liver DNA (mg/100 g of	250	3.79 + 0.08	3,80 ⁿ + 0,14	3.04 ^{nN} + 0.25	3,97 ^{nNY} + 0,40	4.78 + 0.40	4,50 ⁿ + 0,46	3.98 ^{nN} + 0.42	4,00 ^{nNY}
body weight)	300	4.01# 41.0 +	3,98n* + 0,05	3.94 nN*	3.35nN Y*	4,91* + 0.24	4,28 ^{n*} + 0.60	4.57nN* + 0.48	4.25nNY# + 0.32
Muscle DNA (ug/100 mg of	250	65.32 <u>+</u> 4.48	67.1 ⁿ ± 4.15	58.45 ^{nN} ± 7.42	60.45 ^{nNY} + 4.00	48,444	42.85 ⁿ <u>+</u> 4.12	50.20 ^{nN} ± 3.47	47.77 ^{nNY} + 5.80
fresh tissue)	300	68,00* + 5,00	0 ii * 1 - * 1 8 1 * 0 9	59.44 ⁿ N* <u>+</u> 5.45	55.55 ^{nNY} * + 4.46	40.88* <u>+</u> 7.22	38.44 ^{n*} ± 5.21	42.22 ^{nN*} <u>+</u> 4.00	36,40 ^{nNY *} + 5,88
Ovarian (mg/100 g of	250	0,80 + 0,03	1,17 ^b <u>+</u> 0,06	1, 39 ^{bB} + 0,04	1.12 ^{aN a} + 0.10	7.41 <u>+</u> 0.28	9,03 ⁸ + 0.60	0,08 ^{bN}	6.75 ^{nA°} <u>+</u> 0.90
body weight)	300	1.12* + 0.08	1.52 ^{4 ***} + 0.04	1.82 ^{bB###}	1.40 ^{aNB**} <u>+</u> 0.10	6.78* <u>+</u> 0.70	10.47 ^{b*} <u>+</u> 0.65	9.51 ^{aN*} + 0.77	8.43nAoa* + 0.38
S.E. Stand significant. temperature	ard erro Alphabe (25°C or	r. 't' test p ts denote com 30°C) and st	robability di parison betre ars denote co +illed water o	fference : 'P' en different mparison betweed medium. Small	S.F. Standard error. 't' test probability difference : 'P' value : a,A,a, a = P(.05; b,B,B,B,B,B,B, P<.01, n,N,Y, ^A Not significant. Alphabets denote comparison between different salue groups (including distilled water) at a particular temperature (SSC or 30°C) and stars denote comparison between different temperature groups (23°C and 30°C) at a particular (1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,	<pre>" " " " " " " " " " " " " " " " " " "</pre>	b,B,B, ^{*±*} I stilled water) roups (25°C ar	P<.01, n,N,Y,) at a par nd 30 ^o C) at a on between dii	,N,Y, [*] Not particular at a parti- en distilled

4. Changes in DNA content of liver, muscle and ovary (Ta ble 4)

The DNA content of liver remained unchanged with the increase in saline concentration as well as in temperature from 25° C to 30° C in both (NV) and (V) fish. The same nature of results (unchanged) was obtained in case of muscle DNA in all these fishes. As expected, the DNA content of ovary was higher at (V) stage. In (NV) fish, increase in NaCl conovary centration up to 135 mOsm was associated with an increase in each case in the amount of ovarian DNA at 25°C or 30°C. When the saline concentration was raised from 135 to 225 mOsm reduction in ovarian DNA content was noticed in (NV) fish This reduced level of DNA was significantly higher in comparison to that in fish kept in distilled water at $25^{\circ}C$ or $30^{\circ}C$. In (V) fish kept in distilled water and in 225 mOsm NaCl, the DNA content of ovary was almost at the same level at 250C or 30° C, but this level of ovarian DNA was found to be less than that of fish maintained in 65 mOsm and 135 mOsm NaCl at either of these two temperatures. The rise of temperature from 25° C to 30° C had no influence on the ovarian DNA content in (V) fish kept in all media, but significantly increased the ovarian DNA amount in (NV) fish kept in all NaCl solutions.

DISCUSSION

Singi fish can survive in media which are considerably hypertonic to its natural fresh water habitat containing about 5 mOsm salinity (Al Daham and Bhatti, 1977; Parwez et al., 1979). In some representative experiments we have found that they also survive 45-50 days in distilled water but only 12-16 days in 270 mOsm NaCl medium, this osmolarity is about the same as that of its body fluid (Parwez et al. 1979). We have, therefore, choosen for experiments different NaCl concentrations having osmolarity less than that of the body fluid of Singi fish, such as 65, 135 and 225 mOsm/liter corresponding to about 75%, 50% and 17% less osmolarity than that of body fluid, and also distilled water to study the adaptive changes in the liver, muscle and ovary with res pect to protein, RNA and DNA contents and organ weight (H (HSI and GSI) at both non-vitellogenic and vitellogenic stages at 25°C and 30°C. Singi fishes show better metabolic efficien cy, particularly the ovarian activity, at a temperature ran-ge from 28.6°C to 32°C (Vasal and Sundararaj, 1976) We have also observed the aggravated activity of the ovary of Singi fish at 30° C (Dasmahapatra and Medda, 1982a) (NV)

It is evident from our results that the cellular metabolism of liver, muscle and ovary were markedly affected or influenced by changes in the concentration of NaCl in the me dium in which the Singi fishes were maintained for 30 days A number of such induced metabolic changes were temperature dependent (Dasmahapatra, 1980) The HSI was maximum in fish kept in distilled water and minimum in fish kept in 225 mOsm NaCl. But the GSI was minimum in fish kept in distilled wa -

8**6**

ter, maximum in fish kept in 135 mOsm NaCl and intermediate in fish kept in 65 mOsm and 225 mOsm NaCl at (NV) stage. The reduction in HSI in (NV) and (V) fish with the rise of sali-nity of the medium at 25° C or 30° C might be due to the deple tion or decreased synthesis of some cellular constituents of liver. The higher temperature (30°C) caused a decrease in HSI particularly in (V) fish. The concomitant increase in GSI in (NV) fish with the rise of salinity of the medium up to 135 mOsm NaCl both at 25°C and 30°C might be the result of influenced maturation of the ovary. The highest GSI was found in (V) fish maintained in 65 and 135 mOsm NaCl solu tions. But the GSI was almost the same in (V) fish kept in 0 salinity and 225 mOsm NaCl solution. Higher temperature favoured ovarian growth (more GSI) at a suitable saline concen tration. The results further indicated that higher saline concentration (225 mOsm) retarded ovarian growth in (NV) fish. But this medium appeared to be better than distilled water so far as the GSI of the (NV) fish was concerned. The higher GSI values of (V) fish in an environmental salinity of 65 mOsm and 135 mOsm may be explained as the result of better maintenance of the gravid ovary and the lower GSI va-lues in 225 mOsm NaCl and distilled water as the result of regression of the ovary. It seems, therefore, that a saline concentration ranging from about 25% to 50% of the osmolarity (65-135 mOsm per liter) of the body fluid of Singi fish may be suitable for the formation and maturation of the oocy tes in the ovary (Dasmahapatra and Medda, 1982a) possibly through some concomitant contributions of the metabolities from the liver (and muscle) leading to a decrease in weight of the latter organ (liver)

Further, suitability of NaCl medium having osmolarity from 65 to 135 mOsm for ovarian activity in preparation for the favourable conditions for the formation and maturation of eggs can be supported by the enhanced levels of protein , RNA and DNA contents of ovary of (NV) fish kept in these NaCl media (65 and 135 mOsm), the higher concentration of 135 mOsm NaCl being considered as the best medium in these respects. The combined stimulating effect of salinity and higher temperature (30°C) was evidenced from the higher va-lues of these cellular constituents of ovary at 30°C in (NV) fish kept in these saline media. Although a saline concentra tion of 225 mOsm had some depressing action on the matura tion of eggs in (NV) fish (Dasmahapatra and Medda, 1982a)and also on the cellular constituents of ovary studied, this saline medium was better than distilled water so far as the metabolic functions of this organ were concerned. This was evident from the higher levels of protein, RNA and DNA con tents of (NV) ovary in 225 mOsm NaCl group in comparison to those of distilled water group. The data further indicated that in (V) fish the salinity range from 65-135 mOsm , NaCl was favourable for the maintenance of gravid ovary. This could be supported by the enhaced levels of protein and DNA (glycogen and lipid also, unpublished) and unchanged number of mature eggs (Dasmahapatra and Medda, 1982a) in (V) ovary in such saline range. The reduction in the amounts of pro tein and DNA (glycogen and lipid also, unpublished) in the ovary of (V) fish kept in distilled water and 225 mOsm NaC1 solution in comparison to those of the fish kept in 65 and 135 mOsm NaCl solutions might be interpreted as the initial preparatory changes for the regression of mature eggs. The gradual fall of the level of protein and RNA in

in liver and muscle of (NV) and (V) fish with the rise of salinity of the medium from 65 to 135 or 225 mOsm may possibly be the result of induced breakdown and release of these subs tances for the development of maturing oocytes and/or for maintaining the osmolarity of the body fluid in a salinity higher than that of the natural habitat of Singi fish. It has been reported that some euryhaline fishes exposed to higher saline concentration than their natural habitat show so me degradation of protein in liver and muscle and the resultant increase in the amino acid level in blood (Venkatachari, 1974; Bhan and Mansuri, 1978) possibly to adjust the osmola-rity of the body fluid. Thus a constant difference in osmola rity between the body fluid and the surrounding medium is probably maintained. It may be assumed that such difference in osmolarity is maintained in Singi fish when kept in the saline of increasing osmolarity and the resultant degrada -tion of protein helps not only for supplying the metabolites for the growth and maturation of oocytes but also for adjusting the osmolarity of the body fluid possibly to maintain a constant difference. It may be noted further that the DNA content of liver and muscle was not changed by the change in salinity of the medium at 25°C or 30°C. That means the en vironmental salinity has no direct effect on DNA synthesis or degradation in liver and muscle of Singi fish.

The induced changes in different organs at the variation of salinity of the medium may be argued as the results of the changes in hormonal levels, particularly of gonadotro phin, prolactin and/or estrogen at different saline concen trations. Although the endogenous hormonal levels of Singi fishes kept in different saline media have not been studied, the ovarian changes including formation and maturation of eggs may be assumed to be due to enhacement of gonadotrophin level in different salinity. It has been reported that the gonadotrophin content of the pituitary of Mugil (Brackish water) held in fresh water is considerably species lower than that of pituitaries from sea water fish and the area of the pituitary containing the gonadotrophic cells is reduced in size in fresh water specimens (Blanc and Abraham, 1968 ; Blanc-Livni and Abraham, 1970)

The higher estrogen level in blood of (V) Singi fish than (NV) fish (Lamba et al., 1982) may be responsible for higher levels of protein, RNA and DNA contents of liver of (V) fish (Dasmahapatra and Medda, 1982b) Estrogen adminis tration in Singi fish also increased the protein, RNA, lipid and water contents and decreased the glycogen content of liver, but failed to cause any change in muscle and ovary (Med da et al., 1980; Dasmahapatra and Medda, 1982c) The reduc tion in the amount of protein and RNA contents of liver and muscle in (NV) and (V) fish with the rise of salinity of the medium may, therefore, be due to the direct effect of salini ty, rather than indirectly through the mediation of estrogen. The variation of salinity of the medium may cause an alteration in the prolactin concentration in Singi fish. It has

been reported that the transfer of sticklebacks from fresh water to sea water causes the appearance of intercellular cysts among the prolactin cells and also the diminution in number of these cells with shrunken nuclei and poorly granulated cytoplasm which indicate their decreased secretory activity (Benjamin, 1978). Prolactin is also involved in osmoregulation in fresh water fish (Brett, 1979; Gallis *et al.*, 1979). Moreover, prolactin has an antigonadal effect (de Vla ming and Vodicnik, 1977) The possibility of the involvement of prolactin in the changes of liver, muscle and ovary of Singi fish with the change in salinity of the medium is yet to be assessed.

REFERENCES

- AL-DAHAM, N.K. AND BHATTI, M.N. 1977 Salinity tolerance of Gambusia affinis (Baird & Girard) and Heteropneustes fossilis (Bloch) J.Fish Biol. 11:309-313.
- silis (Bloch) J.Fish Biol. 11:309-313. BASHAMOHIDEEN, M. AND PARVATHESWARARAO, V 1972. Adaptations to osmotic stress in the fresh water euryhaline teleost, *Tilapia mossambica*. IV Changes in blood glucose, liver glycogen and muscle glycogen levels. Mar.Biol. 16:68-74.
- BENJAMIN, M. 1978. Cytological changes in prolactin, ACTH, and growth hormone cells of the pituitary gland of Pungitius pungitius L. in response to increase environmental salinity. Gen.Comp.Endocrinol. 36:48-58.
- salinity. Gen.Comp.Endocrinol. 36:48-58.
 BHAN, S., AND MANSURI, A.P. 1978. Adaptations to osmotic stress in the marine euryhaline teleost, *Periophthalmus dipes*. III. Changes in tissue succinic dehydrogenase enzy me levels. J.Anim.Morphol.Physiol. 25:132-138.
- BLANC, N. AND ABRAHAM, M. 1968. Evaluation du pouvoir gonado trope dans l'hypophyse de Cyprinus carpio et Mugil cephalus.C.r.hebd.Seanc.Acad.Sci., Paris 267:958-961.
 BLANC-LIVNI, N. AND ABRAHAM, M. 1970. The influence of envi-
- BLANC-LIVNI, N. AND ABRAHAM, M. 1970. The influence of environmental salinity on the prolactin - and gonadotropin secreting regions in the pituitary of Mugil (Teleosti) Gen.Comp.Endocrinol. 14:184-197
- BRETT, J.R. 1979 Environmental factors and growth. In 'Fish Physiology (Hoar, W.S., Randall, D.J, and Brett, J. R. eds.) Vol. VIII, p. 599-675, Academic Press, New York.
- BURTON, K. 1956. A study of the conditions and mechanisms of the diphenylamine reaction for the colorimetric estima tion of deoxyribonucleic acid. Biochem.J. 62:315-323.
- COWEY, C.B., DAISLEY, K.W. AND PARRY, G. 1962. Amino acids, free or as components of protein, and some B vitamins in the tissues of the Atlantic salmon, during spawning migration. Comp.Biochem.Physiol. 7:29-39
- COWEY, C.B. AND PARRY, G. 1963. The non-protein nitrogenous constituents of the muscle of parr and smolt stages of the Atlantic slamon. Comp.Biochem.Physiol. 8:47-51.
- CROFT, D.N AND LUBRAN, M. 1965. The estimation of deoxyribo nucleic acid in the presence of sialic acid. Application to analysis of human gastric washings Biochem.J 95:612-620.
- DASMAHAPATRA, A.K. 1980. Physiological studies on the effect

of some physicochemical factors on liver, gonad and mus cle of fresh water fish. Ph.D. Thesis, Calcutta University.

DASMAHAPATRA, A.K. AND MEDDA, A.K. 1982a. Effect of some phy sico-chemical factors on the maturation of eggs of Singi fish, *Heteropneustes fossilis* (Bloch) Sci. & Cult. 48: 363-366.

DASMAHAPATRA, A.K. AND MEDDA, A.K. 1982b. Seasonal variation in protein and nucleic acid contents of liver, muscle and ovary of female Singi fish (*Heteropneustes fossilis*(Bloch) in relation to ovarian growth. Bangladesh J.Fish. (in press).

DASMAHAPATRA, A.K. AND MEDDA, A.K. 1982c. Effect of estradiol dipropionate and testosterone propionate on the glycogen, lipid and water contents of liver, muscle and gonad of male and female (vitellogenic and non-vitellogenic)Sin gi fish (*Heteropneustes fossilis* Bloch) Gen.Comp.Endocri nol. 48:476-484.

DE VLAMING, V.L. AND VODICNIK, M.J 1977 Diurnal variations in pituitary gonadotropin content and in gonadal response to exogenous gonadotrophin and prolactin in *Notemigonus* crysoleucas. J.Fish Biol. 10:371-383.

GALLIS, J.L., LASSERRE, P. AND BELLOC, F. 1979. Fresh water adaptation in euryhaline teleost, *Chelon labrosus*. I. Effects of adaptation, prolactin, cortisol and actinomycin D on plasma osmotic balance and (Na⁺ - K⁺) ATPase in gill and kidney. Gen.Comp.Endocrinol. 38:1-10.

HOLLIDAY, F.G.T. 1969. The effects of salinity on the eggs and larvae of teleosts. In 'Fish Physiology (Hoar, W. S. and Randall, D.J. eds), Vol.1, p.293-311, Academic Press, New York.

HUGGINS, A.K. AND COLLEY, L. 1971. The changes in the non protein nitrogenous constituents of muscle during the a daptation of the eel Anguilla anguilla L. from fresh wa ter to sea water. Comp.Biochem.Physiol. 38B:537-541.

JONES, N.R. 1959. Pyruvic acid in the skeletal muscle of fresh and chill-stored, trawled codling (*Gadus callarias*). J.Sci.Food Agr. 10:472-484.

LAMBA, V.J , GOŚWAMI, S.V AND SUNDARARAJ, B.I. 1982. Radioimmunoassay for plasma cortisol, testosterone, estradiol-17β and estrone in the cat fish, *Heteropneustes fossilis* (Bloch). Development and validation. Gen.Comp.Endocrinol. 47:170-181.

LECAL, J 1958. Influence of the salinity factor on the serum proteins of *Blennius pavo*. Compt.rend.soc.biol. 152: 1492-1494.

LOVE, R.M. 1980. The Chemical Biology of Fishes' Vol.II,Aca demic Press, New York.

LOWRY, O.H., ROSEBROUGH, N.J., FARR, A.L. AND RANDALL, R.J. 1951. Protein measurement with folin phenol reagent. J. Biol.Chem. 193:265-275.

MANSURI, A.P. AND BHAN, S. 1978. Adaptations to osmotic stress in the marine euryhaline teleost, *Periophthalmus dipes*. IV Changes in total protein levels of various tis sues. J.Anim.Morphol.Physiol. 25:139-146.

MEDDA, A.K., DASMAHAPATRA, A.K. AND RAY, A.K. 1980. Effect of estrogen and testosterone on the protein and nucleic a cid contents of liver, muscle and gonad and plasma protein content of male and female (vitellogenic and non-vitellogenic) Singi fish (*Heteropneustes fossilis* Bloch) Gen.Comp.Endocrinol. 42:427-436.

- MEJBAUM, W. 1939. Estimation of small amount of pentose especially in derivatives of adenylic acid. Z.Physiol.Chem. 258:117-120.
- MUNRO, H.N. AND FLECK, A. 1966. The determination of nucleic acids In 'Methods of Biochemical Analysis (D.Glick ed.) Vol.14, p.113-176. Wiley (Interscience), New York.
- PARVATHESWARARAO, V 1967 Some mechanism of salinity acclimation in the euryhaline teleost, *Etroplus maculatus*.Mar. Biol. 1:97-101.
- PARWEZ, I., GOSWAMI, S.V AND SUNDARARAJ, B.I. 1979. Salinity tolerance of the fresh water catfish Heteropneustes fos silis (Bloch). Ind.J.Exp.Biol. 17:810-811.
- VASAL, S. AND SUNDARARAJ, B.I. 1976. Response of the ovary in the catfish, *Heteropneustes fossilis* (Bloch), to va rious combinations of photoperiod and temperatures. J Exp.Zool. 197:243-263.
- VENKATACHARI, S.A.T. 1974. Effect of salinity adaptation on nitrogen metabolism in the fresh water fish *Tilapia mossambica*. I. Tissue proteins and amino acid levels. Mar. Biol. 24:75-63.