

## Effect of temperature on laboratory growth, reproduction and life span of *Octopus bimaculoides*

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### Abstract

Laboratory culture of 40 *Octopus bimaculoides* from April 1982 to August 1983 through the full life cycle at 18 °C vs 23 °C provided information on the growth, reproductive biology and life span of this California littoral octopus. At 18 °C, the cephalopods grew from a hatchling size of 0.07 g to a mean of 619 g in 404 d; the largest individual was 872 g. Octopuses cultured at 23 °C reached their highest mean weight of 597 g in 370 d; the largest individual grown at this temperature was 848 g after 404 d. Growth data revealed a two-phase growth pattern: a 5 mo exponential phase followed by a slower logarithmic (power function) phase until spawning. At 5 mo octopuses grown at 23 °C were over three times larger than their 18 °C siblings. However, beyond 6.5 mo, growth rates were no higher at 23 °C than at 18 °C. At 13.5 mo, the mean weight of the 18 °C group surpassed that of the 23 °C group. The slope of the length/weight (L/W) relationship was significantly different for the two temperature regimes, with the 23 °C octopuses weighing 18% less than their 18 °C siblings at a mantle length of 100 mm. Females weighed more than males at any given mantle length. Males grew slightly larger and matured before females. The L/W relationship indicated isometric body growth throughout the life cycle. Higher temperature accelerated all aspects of reproductive biology and shortened life span by as much as 20% (from approximately 16 to 13 mo). *O. bimaculoides* has one of the longest life cycles among species with large eggs and benthic hatchlings. Extrapolations to field growth are made, and the possible effects of temperature anomalies such as El Niño are discussed.

### Introduction

Cephalopods are important predators in marine ecosystems, where they compete with vertebrate fishes; this

competition has led to many forms of convergence (Packard 1972). However, cephalopods differ dramatically from fishes with respect to their high rate of growth, semelparous reproductive strategy and comparatively short life spans (see Boyle 1983, 1987). Various studies have described the effects of temperature on cephalopod growth, reproduction or life span (Van Heukelem 1979), but no study before the present report has analyzed the effect of temperature on these three aspects of octopus biology simultaneously throughout the full life cycle. The laboratory data reported herein (the first for this species) are based upon observations of two groups of *Octopus bimaculoides* cultured from eggs to sexual maturity and egg-laying in the laboratory under two temperature regimes.

*Octopus bimaculoides* is a common inter- and sub-tidal species along the southern California coast, with a known range extending from approximately Santa Barbara, California south to San Quintin, Mexico on the upper Baja peninsula. There is little published information on the biology of this species. Hochberg and Fields (1980) and Forsythe and Hanlon (1988) have reviewed the literature, and the latter authors have made laboratory and field observations on aspects of the behavior and reproductive biology of the species. The present study arose as a part of our ongoing research in evaluating various large-egged octopus species as potential laboratory-culture candidates for supplying biomedical research needs (Hanlon and Forsythe 1985).

The specific objective of this study was to evaluate the influence of temperature on growth, age at reproduction and resultant life span to see if it might be used as a tool in timing reproduction and expanding the temporal availability of eggs. By accelerating growth and reproduction of some brood-stock individuals and retarding that of others, it might be possible to achieve egg-laying throughout the year. Additionally, this type of information is useful to fisheries biologists and ecologists studying aspects of cephalopod biology in nature, where temperature can in-

fluence distributions and life-history patterns. The short-term elevation in temperature during an El Niño event is a specific example of how laboratory data can be useful in understanding the impact of such a phenomenon on rapidly growing, short-life-span predators such as *Octopus* spp.

### Materials and methods

The culture project was conducted from April 1982 to August 1983 in the Division of Biology and Marine Resources laboratory of The Marine Biomedical Institute in Galveston, Texas. All culture work was carried out in two closed, recirculating seawater systems, each consisting of a circular 2 000-liter capacity water-conditioning tank above which were supported shallow culture-trays. This system has been described in detail by Hanlon and Forsythe (1985). In the main system (SYST 1), a 1 hp seawater chilling unit was used to maintain water temperature at approximately 18 °C. The second culture system (SYST 2) was maintained at warmer ambient laboratory temperatures (mean 23 °C). Artificial sea water (Instant Ocean brand) was used exclusively throughout this study. Temperature and salinity were recorded daily and levels of ammonia (NH<sub>4</sub>-N), nitrite (NO<sub>2</sub>-N) and nitrate (NO<sub>3</sub>-N) were measured weekly. Salinity was maintained between 34 and 36‰. Periodically, 50 to 70% water changes were made when inorganic nitrogen levels increased. When pH levels fell below 7.75, additions of sodium bicarbonate increased pH. Trace elements (HW Marine Mix) were added regularly to prevent their possible depletion between water changes. The culture systems received overhead fluorescent lighting and natural, indirect sunlight from large adjacent windows. The light cycle was approximately the same as that outdoors, since the fluorescent lights were only on from 08.00 to 17.00 hrs.

Eggs were laid by a field-collected (near Los Angeles, California) female *Octopus bimaculoides* being held for physiological studies in San Antonio, Texas. The eggs were maintained at 17 °C in a 400-liter closed system using artificial sea water for the first three weeks, and were then removed from the mother and transported to Galveston in sealed plastic bags containing sea water and pure oxygen. Eggs were placed in SYST 1 and maintained at 18 °C for the remainder of their development. Hatching occurred over an 8 d period from mid- to late June 1982. The modal hatching day was considered to be Day 1 for the study. *O. bimaculoides* was cultured under two temperature regimes, beginning at hatching. The main population (initiated with 177 hatchlings) was kept in SYST 1 after hatching and cultured at 18 °C. To evaluate the tolerance of this species to higher temperatures, a second smaller group (45 hatchlings) was moved to SYST 2 and cultured at temperatures near 23 °C. During the first 6 mo, isolated individual octopuses were measured and weighed regularly. Thereafter (Day 218), most or all of the culture population

was measured. The octopuses were narcotized first in a 1.5% solution of ethyl alcohol in sea water, and then measured for mantle length (ML) and wet weight (WW) after the mantle cavity had been drained of sea water. At this time, they were also inspected visually for signs of hectocotylus development. Octopuses revived in fresh sea water without assistance within 2 to 3 min.

Linear regression analyses of the growth data were performed (Forsythe 1984). For each chronological series of length and weight measurements, a line of best fit to the data was calculated mathematically by a least-squares linear regression, and an equation describing the growth data was generated. The mean length or weight of the entire sample group at each measuring period was used in these determinations. The linear regressions were performed using standard computer curve-fitting programs. The following equations were tested to generate lines of best fit to the growth data:

$$\begin{aligned} \text{linear, } & y = a + bx; \\ \text{exponential, } & y = ae^{bx}; \\ \text{power, } & y = ax^b; \\ \text{log, } & y = a + b \ln x; \end{aligned}$$

where  $y$  = length or weight and  $x$  = age in days;  $a$  = the  $y$  intercept,  $b$  = the slope and  $e$  = the natural logarithm of 2. The square of the correlation coefficient ( $r^2$ ) was calculated for each linear regression and used to determine which equation best described a given set of data. To avoid confusion, it should be noted that, traditionally, growth described by the power equation is referred to as logarithmic growth (Brody 1945). That is what it shall be taken to mean in this paper and not growth described by the log equation as listed above.

Growth rates were determined using the instantaneous coefficient of growth,  $G$ , calculated from the equation:

$$G = \frac{\log_e Y_2 - \log_e Y_1}{t_2 - t_1}$$

where  $Y_2$  = final length or weight,  $Y_1$  = initial length or weight,  $t_2$  = age in days at  $Y_2$ , and  $t_1$  = age in days at  $Y_1$  (Brown 1957, Wells and Wells 1970, Forsythe and Van Heukelem 1987). The instantaneous relative growth rate (percent increase in body length or body weight per day; %/day) was calculated by multiplying  $G$  by 100. Multiplying any given length ( $L$ ) or weight ( $W$ ) by  $G$  gave growth rates in mm per day (for  $G \times L$ ) or grams per day (for  $G \times W$ ). Doubling time (DBL), i.e., the time required for an individual to double in weight, was calculated by dividing the natural log of 2 by  $G$ . Where necessary, data were compared using Student's  $t$ -tests after variance-ratio tests (Zar 1974) had been performed to assure means came from normally distributed populations. An analysis of covariance (Zar 1974) was used to test for differences in slopes and elevations of linear forms of calculated growth curves. This was done using log<sub>e</sub>-transformed data:  $x$  and log<sub>e</sub> $y$  for exponential functions, and log<sub>e</sub> $x$  and log<sub>e</sub> $y$  for power functions.

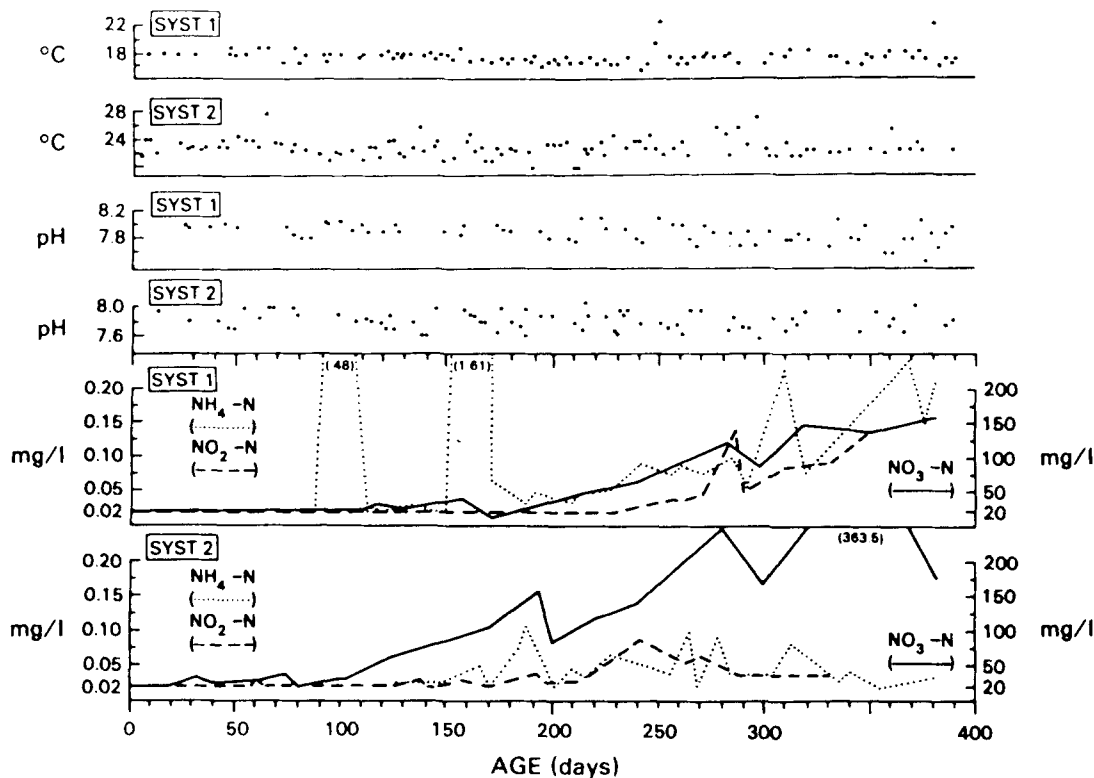


Fig. 1. Synopsis of water-chemistry data for closed systems used in this study

## Results

Forty *Octopus bimaculoides* were cultured through the life cycle from hatching to sexual maturity and egg-laying. The population was maintained in culture until 18 August 1983, when all surviving octopuses were lost during a 72 h power failure after Hurricane Alicia.

### Water quality

Since all research was conducted in closed, recirculating seawater systems, synopses of the water-chemistry data during the study are presented in Fig. 1. In SYST 1, monthly mean temperatures ranged from 17.2° to 18.6°C, with an overall mean of 17.96°C. The minimum and maximum temperatures were 16.5° and 23.0°C, respectively; on only four days were temperatures above 19°C. In SYST 2, temperatures ranged from 19.0° to 28.0°C, with monthly means ranging from 20.1° to 24.4°C. Temperatures were above 25°C on only 11 d. The overall mean temperature was 22.98°C. Temperature fluctuations were much greater in this system, since it varied with ambient laboratory temperatures. The measured levels of pH, ammonia, nitrite and nitrate (Fig. 1) were within acceptable ranges for octopuses (Hanlon and Forsythe 1985) and had no observable deleterious effects on behavior, feeding, growth or survival.

### Growth

The wet weight data are summarized in Table 1 and the mantle length data in Table 2. The calculated growth equations are listed in Table 3.

#### Growth in weight

*Octopus bimaculoides* grew from an approximate hatching size of 0.07 g to a mean of 619 g in 404 days at 18°C. Growth was exponential in form from hatching until Day 156 (Eq. 1 in Table 3), with a mean relative growth rate of 3.56% of body weight per day (%/day). The octopuses doubled in weight approximately every 22 d for a total of seven doublings during this period. Growth was slower after Day 156, becoming logarithmic in form (Forsythe 1984), as described by Eq. (2) in Table 3. Table 1 shows a decrease in mean wet weight at Day 218. Beginning on this day, most or all of the population was measured rather than a sub-sample. Growth rates decreased from nearly 3% d<sup>-1</sup> at Day 170 to less than 1% d<sup>-1</sup> at Day 404. The mean doubling time increased to about 54 d, yielding 4.5 doublings over this period.

The break in the growth phases at Day 156 coincided with the first appearance of the hectocotylus (a modification of the third right arm of males that is used in transferring spermatophores). It is noteworthy that the

**Table 1.** *Octopus bimaculoides*. Growth in weight at 18°C and 23°C. Growth rates calculated from difference between consecutive measurements. Dash indicates no growth rate calculated

N	Age (d)	Mean (g)	(SD)	Range		Growth rate		N	Age (d)	Mean (g)	(SD)	Range		Growth rate	
				min.	max.	(%/d)	(g/d)					min.	max.	(%/d)	(g/d)
Group cultured at 18°C								Group cultured at 23°C							
10	14	0.14	(0.03)	0.09	0.18	-	-	20	14	0.15	(0.04)	0.07	0.25	-	-
10	29	0.25	(0.04)	0.19	0.31	3.99	0.01	19	29	0.28	(0.09)	0.15	0.43	4.38	0.01
10	36	0.33	(0.05)	0.24	0.41	3.82	0.01	17	36	0.39	(0.12)	0.21	0.62	4.62	0.02
10	43	0.39	(0.07)	0.28	0.51	2.47	0.01	17	43	0.53	(0.16)	0.29	0.74	4.52	0.02
9	57	0.63	(0.13)	0.43	0.78	3.43	0.02	17	57	1.08	(0.30)	0.57	1.56	5.06	0.05
8	85	1.65	(0.30)	1.27	2.06	3.44	0.06	7	128	25.63	(22.07)	11.20	74.58	4.46	1.14
9	99	2.55	(0.49)	1.89	3.24	3.12	0.08	7	142	49.70	(32.09)	17.00	106.30	4.73	2.35
4	114	3.62	(0.79)	3.07	4.77	2.32	0.08	8	156	67.01	(40.24)	26.80	142.70	2.13	1.43
8	128	10.28	(2.37)	7.68	15.23	7.47	0.77	7	170	103.81	(64.39)	38.60	210.90	3.13	3.25
8	142	15.23	(3.45)	10.35	21.78	2.81	0.43	7	183	130.21	(73.42)	52.00	252.60	1.74	2.27
8	156	21.18	(4.26)	13.78	28.56	2.36	0.50	7	198	174.86	(86.64)	77.50	311.80	1.97	3.44
8	170	31.73	(6.86)	18.79	40.84	2.89	0.92	7	212	196.39	(86.29)	92.60	318.30	0.83	1.63
8	183	39.66	(9.76)	25.46	53.11	1.71	0.68	13	218	175.68	(95.79)	90.60	424.00	-	-
8	198	55.48	(11.22)	40.54	71.95	2.24	1.24	18	240	268.39	(117.46)	132.50	554.40	1.93	5.17
8	212	68.23	(12.72)	51.16	87.92	1.48	1.01	17	268	371.77	(115.77)	193.20	572.30	1.16	4.33
27	218	61.82	(19.52)	29.30	99.90	-	-	16	289	383.24	(115.39)	208.70	588.20	0.14	0.55
31	240	103.94	(32.82)	45.00	158.50	2.36	2.45	14	324	517.39	(139.75)	244.50	734.30	0.86	4.44
24	268	174.23	(53.28)	84.50	240.60	1.85	3.21	8	370	597.53	(142.19)	446.20	810.20	0.31	1.87
36	289	210.89	(79.15)	76.20	329.50	0.91	1.92	8	404	584.56	(195.02)	267.40	848.60	-0.06	-0.38
32	324	350.28	(106.55)	123.00	551.90	1.45	5.08								
30	370	505.17	(138.40)	110.90	730.10	0.80	4.02								
21	404	618.89	(140.65)	369.00	872.10	0.60	3.70								

**Table 2.** *Octopus bimaculoides*. Growth in mantle length

N	Age (d)	Mean (mm)	(SD)	Range		Growth rate (%/d)	N	Age (d)	Mean (mm)	(SD)	Range		Growth rate (%/d)
				min.	max.						min.	max.	
Group cultured at 18°C							Group cultured at 23°C						
10	14	7.76	(0.37)	7.10	8.40	-	20	14	7.16	(0.66)	6.00	8.20	-
10	29	8.79	(0.72)	7.80	9.80	0.83	20	29	8.84	(0.98)	7.10	10.70	1.40
10	36	9.81	(1.06)	8.10	11.60	1.57	17	36	10.33	(1.07)	8.50	12.20	2.23
10	43	10.40	(0.96)	8.50	11.60	0.83	17	43	11.42	(1.26)	8.80	13.30	1.43
10	57	11.01	(1.25)	8.30	12.80	0.41	17	57	13.53	(1.51)	10.90	16.10	1.21
9	85	15.73	(0.99)	14.50	17.20	1.27	17	128	40.77	(10.33)	32.10	62.80	1.55
9	99	18.81	(1.40)	17.20	21.40	1.28	8	142	51.83	(12.39)	36.60	72.00	1.71
4	114	21.45	(2.28)	19.70	24.80	0.88	7	156	59.06	(14.21)	40.50	83.00	0.93
8	128	29.56	(1.83)	27.60	31.80	2.29	7	170	67.94	(15.55)	50.60	90.70	1.00
8	142	34.85	(3.43)	29.00	39.30	1.18	7	183	69.23	(15.04)	50.00	90.50	0.14
8	156	38.64	(2.91)	33.00	41.80	0.74	7	198	78.09	(13.78)	63.30	98.50	0.80
8	170	43.50	(4.82)	35.20	49.80	0.85	6	212	79.92	(13.42)	62.50	96.80	0.17
8	183	45.66	(4.68)	39.80	51.40	0.37	13	218	78.31	(14.18)	64.00	113.00	-
8	198	51.33	(5.55)	42.70	59.80	0.78	18	240	91.00	(14.52)	74.00	120.00	0.68
8	212	52.94	(5.04)	44.50	62.30	0.22	15	268	101.87	(14.48)	84.00	130.00	0.40
27	218	54.37	(7.45)	35.00	66.00	0.45	16	289	103.75	(11.47)	90.00	130.00	0.09
31	240	63.74	(7.52)	50.00	76.00	0.72	14	324	116.29	(12.64)	85.00	140.00	0.33
24	268	75.04	(10.61)	55.00	94.00	0.58	8	370	132.38	(9.20)	122.00	147.00	0.28
36	289	75.00	(12.33)	46.00	95.00	0.00							
32	324	97.44	(18.47)	65.00	160.00	0.75							
30	370	108.40	(12.19)	80.00	140.00	0.23							

female data up to Day 198 included some males that were not yet identifiable; however, beyond Day 198 the sexes could be differentiated clearly. The equations for the male and female data are given in Table 3 (Eqs. 3 and 4, respectively). Males and females were similar in weight up

to Day 218. Thereafter, the mean weight of males was consistently higher than that of females. This is illustrated in Fig. 3. Over this time period the mean instantaneous growth rate for males was 1.8% d<sup>-1</sup>, compared to 1.5% d<sup>-1</sup> for females. A Student's *t*-test analysis revealed no statisti-

**Table 3.** *Octopus bimaculoides*. Growth equations for live wet weight, mantle length and length/weight relationship

Group	Days	Equation	Eq. No.	
Wet wt ( <i>WW</i> )				
18 °C	combined population	14–156	$WW (g) = 0.0843 e^{0.0356t}$ ; $r^2 = 0.9936$	(1)
		156–404	$WW (g) = (2.59 \times 10^{-7}) t^{3.62}$ ; $r^2 = 0.9942$	(2)
	males	156–404	$WW (g) = (1.13 \times 10^{-7}) t^{3.77}$ ; $r^2 = 0.9939$	(3)
	females	156–404	$WW (g) = (5.61 \times 10^{-7}) t^{3.47}$ ; $r^2 = 0.9935$	(4)
23 °C	combined population	14–142	$WW (g) = 0.0762 e^{0.0456t}$ ; $r^2 = 0.9999$	(5)
		142–324	$WW (g) = (5.65 \times 10^{-5}) t^{2.79}$ ; $r^2 = 0.9748$	(6)
	males	142–370	$WW (g) = (2.04 \times 10^{-3}) t^{2.16}$ ; $r^2 = 0.9799$	(7)
	females	142–324	$WW (g) = (1.12 \times 10^{-6}) t^{3.48}$ ; $r^2 = 0.9768$	(8)
Mantle length ( <i>ML</i> )				
18 °C	combined population	14–156	$ML (mm) = 6.21 e^{0.0116t}$ ; $r^2 = 0.9888$	(9)
		156–370	$ML (mm) = 0.0842 t^{1.21}$ ; $r^2 = 0.9878$	(10)
	males	156–370	$ML (mm) = 0.0699 t^{1.25}$ ; $r^2 = 0.9860$	(11)
	females	156–370	$ML (mm) = 0.111 t^{1.15}$ ; $r^2 = 0.9780$	(12)
23 °C	combined population	14–142	$ML (mm) = 5.78 e^{0.0154t}$ ; $r^2 = 0.9993$	(13)
		142–324	$ML (mm) = 0.513 t^{0.942}$ ; $r^2 = 0.9817$	(14)
	males	142–370	$ML (mm) = 1.99 t^{0.704}$ ; $r^2 = 0.9837$	(15)
	females	142–324	$ML (mm) = 0.127 t^{1.18}$ ; $r^2 = 0.9823$	(16)
Length ( <i>ML</i> ) vs wt ( <i>WW</i> ) relationship				
18 °C	combined population	14–370	$WW (g) = (3.2 \times 10^{-4}) ML (mm)^{3.06}$ ; $r^2 = 0.9984$	(17)
	males	142–370	$WW (g) = (2.95 \times 10^{-4}) ML (mm)^{3.07}$ ; $r^2 = 0.9887$	(18)
	females	142–370	$WW (g) = (4.03 \times 10^{-4}) ML (mm)^{3.01}$ ; $r^2 = 0.9881$	(19)
23 °C	combined population	14–370	$WW (g) = (4.6 \times 10^{-4}) ML (mm)^{2.94}$ ; $r^2 = 0.9989$	(20)
	males	128–324	$WW (g) = (2.1 \times 10^{-4}) ML (mm)^{3.10}$ ; $r^2 = 0.9913$	(21)
	females	128–324	$WW (g) = (3.87 \times 10^{-4}) ML (mm)^{2.98}$ ; $r^2 = 0.9877$	(22)

cally significant differences in mean wet weights of males and females at any measurement period. However, there was a significant difference ( $p < 0.05$ ) in the slopes of simple linear regressions of the log-transformed data of males and females. Thus, the slope value of Eq. (3) for male growth is significantly higher than that of Eq. (4) for females, and indicative of a slightly higher growth rate among males.

The weight data for the warm-water population are also listed in Table 1. Analysis revealed the same pattern of growth over slightly different time periods. The faster exponential growth phase lasted until Day 142 (Eq. 5) with an average growth rate of  $4.5\% d^{-1}$ . The doubling time was 15 d, producing 9.5 doublings in weight. The slower logarithmic phase lasted from Days 142 to 324 (Eq. 6). Differences in male and female growth resulted in poor curve fits to combined data beyond Day 324. Males continued to grow logarithmically until at least Day 370, with growth leveling off at Day 404. Eq. (7) for male growth was calculated through Day 370. Females es-

entially stopped growing after Day 324 (Fig. 3), and this cessation of the growth phase coincided with egg-laying in this group between Days 315 and 370. The equation for female growth (Eq. 8) was therefore only calculated through Day 324. Again, as in the main population, males had consistently higher mean weights than females. Despite this finding, analysis showed that females were growing slightly faster than males. The mean instantaneous growth rate for females from Days 128 to 324 was  $2.1\% d^{-1}$ , while males had a mean of  $1.5\% d^{-1}$ . In addition, the higher slope of the female growth equation (Eq. 8) was significantly different ( $p < 0.001$ ) from that of the males (Eq. 7). Apparently, the male octopuses were larger than female octopuses at the end of the exponential growth phase. This size difference was large enough to allow the mean male weights to stay larger than the mean female weights, although the more rapidly growing females were continually lessening this difference.

The strong influence of temperature on the growth of *Octopus bimaculoides* is seen clearly by comparing the

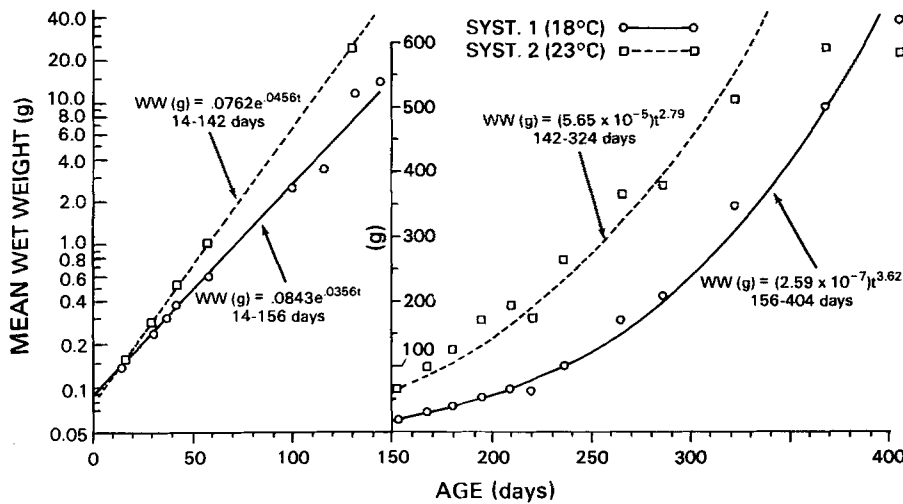


Fig. 2. *Octopus bimaculoides*. Growth curves for individuals cultured through life cycle under two temperature regimes

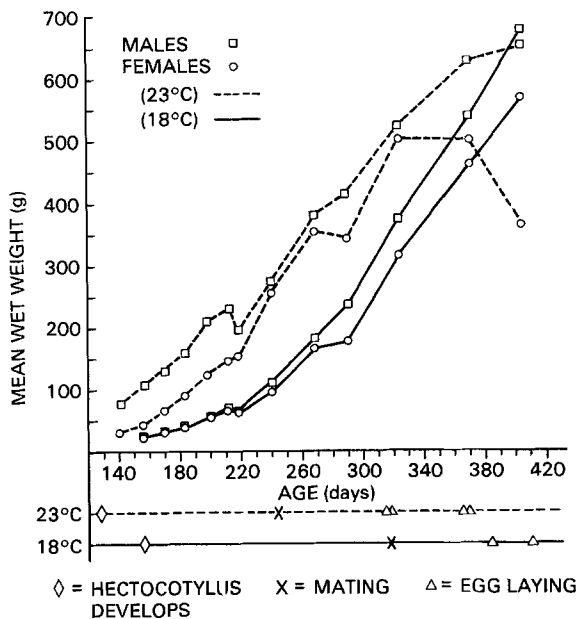


Fig. 3. *Octopus bimaculoides*. Comparative growth of males and females at 18° and 23°C. Major events in reproductive life cycle are plotted for each temperature beneath age axis

growth data of the main population (18°C) to the data of the warm-water group (23°C) in Table 1 and Figs. 2 and 3. Over the course of the exponential growth phase, the octopuses at 23°C grew on the average 1% d<sup>-1</sup> faster than the octopuses at 18°C (4.5 vs 3.5% d<sup>-1</sup>). This difference in growth rate resulted in a three-fold difference in mean size by Day 142 (49.7 vs 15 g). Over this period, the warm-water juveniles doubled in weight 9.5 times compared to 6.5 times for their cold-water siblings. This three-fold difference in weight continued up to Day 198, after which it gradually declined until Day 404, when the cold-water group had a higher mean weight for the first time. The largest individual cultured (872 g) was a male from the cold-water population. Comparing the growth curves for the two groups in Fig. 2, it is clear that the slope of Eq. (5)

(warm-water juveniles) is higher than that of Eq. (1) (cold-water juveniles); this difference is highly significant ( $p < 0.001$ ). For the curves describing the periods of logarithmic growth in Fig. 2, the slope of Eq. (2) is higher than that of Eq. (6) and this difference is highly significant ( $p < 0.001$ ). Closer examination of the data showed that growth of the warm-water group did not begin to slow until after Day 198, and only then did the cold-water group have consistently higher relative growth rates. The slowed growth of the warm-water group was essentially attributable to the males in the group. Comparing the slope of Eq. (7) to Eqs. (8), (3) and (4), it can be seen that the warm-water males had the lowest slope value. The warm-water females, on the other hand, had essentially the same slope as their cold-water counterparts, with the statistical analysis showing no significant differences in the slopes.

#### Growth in length

Mantle length is a long-standing measure in cephalopod biology, even though it is not as useful as weight. The mantle length data in Table 2 were analyzed in the same fashion as the wet weight data. The growth equations for the mantle length data are also given in Table 3. Growth in length paralleled the relationships described for growth in weight. The growth equations had lower  $r^2$  values than those of the weight data. This is due to the greater variability encountered with length measurements of the soft-bodied octopuses.

#### Length vs weight relationship

The length vs weight (L/W) relationship was examined by comparing concurrent wet weight and mantle length (ML) data. Table 3 presents the equations describing this relationship (Eqs. 17–22). Although growth in weight occurred in two phases over the life cycle, the L/W relation

maintained nearly identical slope values throughout the entire life cycle and, therefore, single equations were used to describe the combined group data from Days 14 to 370 (last-day ML measurements were taken). Eq. (17) describes the L/W relationship for the main culture population and is depicted graphically in Fig. 4. Eqs. (18) and (19) describe the male and female data, respectively. An analysis of covariance showed no statistically significant difference in the slopes or elevations of the male and female curves. Using the equations to calculate wet weights at given mantle lengths, it can be seen that females weigh more than males of the same mantle length. For example, at 100 mm ML, the equations estimate the weight of females at 422 g, the weight of males at 407 g.

Eq. (20) describes the L/W relationship for the warm-water population. The male and female L/W data for the warm-water group were curve-fit to Day 324, since this produced a better fit than to Day 370. There was no statistically significant difference in the slopes of Eqs. (21) and (22). Again, females weighed more than males at the same ML. At 100 mm ML, females are predicted to weigh 353 g and males 333 g.

The slopes of the L/W relationship for the cold-water and warm-water population (Eqs. 17 and 20, respectively) were significantly different ( $p < 0.002$ ). The slope of the L/W relationship was not found to be statistically different in comparisons between the two temperatures for just the exponential growth phase, logarithmic growth phase, males or females. At a mature size of 100 mm ML, the octopuses cultured at 23°C weighed an average of 18% less than their siblings cultured at 18°C (348 g vs 422 g). This same magnitude of weight difference was seen when comparing males or females at the two temperatures. These analyses suggest that the longer it takes an individual to reach a given ML, the more the octopus will weigh. Males reached any given mantle length before females, but females weighed more than males of the same mantle length. Likewise, the warm-water group grew faster than the cold-water group, but the latter group weighed more than the former at the same mantle length.

The slope of the L/W relationship lies typically between 2.5 and 4.0 (Brody 1945, Brown, 1957). A slope of 3.0 indicates that weight (or volume) is increasing as the cube of length (or linear size), resulting in isometric body growth and constant body proportions (Gould 1966, Ricker 1979). Slopes significantly greater than or less than 3.0 are indicative of allometric growth, which produces changes in body proportions over the life cycle. *Octopus bimaculoides* seems to display isometric body growth through its life cycle, since Eqs. (17) and (20) vary only slightly above and below a slope value of 3.0.

## Reproductive biology

### Sexual maturation

The earliest sign of sexual maturation was the formation of the hectocotylus on the third right arm of males. In the

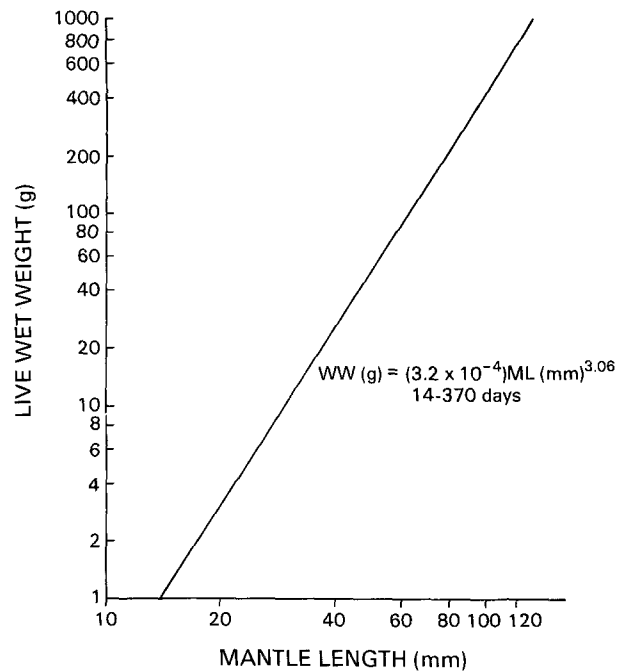


Fig. 4. *Octopus bimaculoides*. Length versus weight relationship for main-culture population grown at 18°C

main culture population (SYST 1) the first males with an identifiable hectocotylus were found on Day 156. All males could be identified by approximately Day 200. The appearance of the hectocotylus was seen approximately one month earlier (Day 128) in the smaller culture group being grown at higher temperatures (SYST 2), with all males identifiable by Day 170 (Fig. 3).

Among females, the only external visible sign of sexual maturation was the rapid enlargement of the ovary in the distal end of the mantle. This enlargement did not become evident until the month prior to egg-laying; during this period the ovary increased from less than a tenth of the internal mantle space to almost one-third to one-half the mantle space.

### Mating and egg-laying

Matings were observed on Day 317 in the main culture population and almost 2.5 mo earlier (Day 241) in the warm-water group (Fig. 3). These first matings followed first hectocotylus formation in the two groups by approximately 5.4 and 3.8 mo, respectively.

Egg-laying followed the initiation of mating behavior by two to four months in both populations. Only two broods of eggs were laid in the main culture population before it was lost. These broods were laid on Days 383 and 409, but perished before completing development. The 12 females in this group that died after the hurricane were preserved, and their ovaries examined for degree of maturation. Four females had ripe ovaries, with one apparently dying the day she would have begun egg-

laying since there were eggs at the opening of each oviduct. The eight remaining females were still immature. Four broods of eggs were laid in the warm-water population on Days 315, 316, 365 and 369. Of the four broods, two were fertile, one infertile and one brood was destroyed when the mother was killed by another larger female. A fifth female had died on Day 284 when her ovary literally burst open with fully mature eggs.

The duration of egg development varied dramatically with temperature. The brood of eggs from which the octopuses in this study hatched required 82 d to develop at a mean temperature of 17.8 °C. In contrast, the two fertile broods in the warm-water population required only 46 and 51 d at 23.4° and 22.5 °C, respectively.

**Discussion**

Temperature

Elevated temperature caused a dramatic increase in growth rate during the exponential growth phase of *Octopus bimaculoides*. This effect diminished as the octopuses entered the slower logarithmic growth phase of their life cycle. Beyond Day 200, the higher temperature (23 °C) produced no increase in relative growth rate over that at 18 °C. Van Heukelem (1976) reared *O. maya* from hatching at 20° and 30 °C. After 60 d, the 20 °C octopuses had grown in weight at a mean rate of 3.8% d<sup>-1</sup> vs 7.5% d<sup>-1</sup> for the 30 °C individuals. He also found the length of the exponential growth phase was much shorter at 30 °C (75 d) than at 20 °C (165 d). Once the 30 °C individuals ended their exponential growth phase, growth rates declined gradually until, at about Day 200, their growth rates dropped below those of their 20 °C siblings. Pascual (1978) found that the growth rates of young cuttlefish (*Sepia officinalis*) between one and three months of age were nearly twice as high at 22 °C (5.1% d<sup>-1</sup>) than at 16 °C (2.7% d<sup>-1</sup>).

All large-egged octopus species reared through the life cycle in the laboratory have displayed the same two-phase growth pattern characterized by an early, rapid, exponential growth phase followed by a somewhat slower, logarithmic (power-function) growth phase to the onset of spawning (Forsythe 1984, Forsythe and Van Heukelem 1987). Some comparisons of growth-phase duration and overall instantaneous relative growth rate for each phase are made in Table 4. *Octopus bimaculoides* and *O. briareus* show very similar growth patterns, having an extended exponential growth phase compared to the other species. *O. briareus* weighs approximately 55 g (17° to 25 °C) at the end of its exponential phase, while *O. bimaculoides* weighs 45 g (23 °C). *O. bimaculoides* has a longer logarithmic growth phase, but growth rates are similar.

This same relationship has been noted in other cephalopod species (Forsythe and Van Heukelem 1987) where, within a species, the slower-growing sex weighs more at a given ML than its counterpart. A surprising finding is that, for a given length, the cold-water *Octopus bimaculoides* weighed more than those reared at warmer temperatures. This emphasizes the potential errors of estimating biomass from length measurements, a custom followed by many invertebrate larval biologists. Overall, temperature seemed to have an impact on the L/W relationship in this species, but it did not produce individuals visibly different in morphology.

The existing L/W data for octopuses was reviewed by Forsythe (1984) and Forsythe and Van Heukelem (1987). The L/W relationship for *Octopus bimaculoides* is most similar to that reported for *O. vulgaris* (Guerra 1979) and *Eledone moschata* (Boletzky 1974). For example, *O. bimaculoides* (18 °C) and *O. vulgaris* weigh 0.36 and 0.42 g at 10 mm ML, and 421 and 410 g at 100 mm ML, respectively. The exponent of the L/W relationship is near 3.0 for all three species, indicating isometric growth throughout the life cycle. Not all octopuses grow isometrically. *O. briareus*, *O. joubini* and *O. digueti* display allometric growth during part or all of the life cycle, indicating changes in body proportions throughout the life cycle.

**Table 4.** *Octopus* spp. Comparison of growth-phase duration and growth rate for five species. (Where necessary, growth rate was calculated from data in cited papers)

Species	T (°C)	Growth-phase duration with mean growth rate, as % body wt d <sup>-1</sup>	Source
<i>O. joubini</i>	25	7.41 ————— 2.6	Forsythe (1984)
<i>O. digueti</i>	25	6.4 ————— 2.6	DeRusha et al. (1988)
<i>O. maya</i>	25	6.1 ————— 2.9	Van Heukelem (1976)
<i>O. briareus</i>	17–25	4.8 ————— 1.8	Hanlon (1983)
<i>O. bimaculoides</i>	23	4.5 ————— 1.1	(present study)
<i>O. bimaculoides</i>	18	3.5 ————— 1.4	(present study)

Age (wk)



### Reproductive biology

Van Heukelem (1979) reviewed data indicating that light, temperature and food influence sexual maturation in cephalopods. It is significant, in the present study, that temperature alone was shown to have a strong effect on all aspects of reproductive biology and eventual life span (Fig. 3). In *Octopus bimaculoides* an elevation in temperature of 5°C over the entire life span accelerated hectocotylus formation by one month, mating and egg-laying by two months, and duration of egg development by one month. It may have shortened the life cycle by as much as three to four months. As discussed previously, temperature had no effect on growth rate beyond Day 200, yet its effects on reproductive biology over this same period are clear. Richard (1966) was able to show similar effects of temperature on the sexual maturation of *Sepia officinalis*. Cuttlefish reared at 20°C reached full sexual maturity in seven months, while siblings at 15°C were showing only the first signs of gonad development and individuals at 10°C were showing no gonadal development at all. Clearly, temperature must be considered in estimating life spans of cephalopod species, particularly when temperature varies over the geographic range.

### Life span

*Octopus bimaculoides* appears to be fairly long-lived for an octopus. At 18°C, females began spawning at 12.5 mo; however, the majority of females were still immature at 14 mo. The additional three months required for egg brooding gives a total life span of at least 15 to 17 mo. At 23°C, females spawned at 10 to 12 mo, giving a complete life span with egg brooding of 12 to 14 mo. Males appear to live at least as long as females. Among large-egged octopods, only *Eledone moschata* in the Mediterranean Sea is known to have as long a life span as *O. bimaculoides* (Mangold 1983); it ranges from 12 to 24 mo, depending upon seasonal temperature. The life spans of most other large-egged species range from 6 to 12 mo (Boyle 1983). Species with planktonic hatchlings typically have slightly longer life spans (12 to 18 mo) due to their 1 to 3 mo planktonic phase (Boyle 1983). Life-span estimates have been reported for three small-egged octopus species found on the Pacific coast of the USA. Hartwick (1983) predicted a life span range of 3 to 5 yr for the giant Pacific octopus *O. dofleini*, and Dorsey (1976) estimated 2 to 3 yr for *O. rubescens*. Both estimates were for populations in the Pacific northwest (Washington, British Columbia) that have a low seasonal temperature range of 7° to 15°C.

*Octopus bimaculatus* is a sibling species of *O. bimaculoides* (Pickford and McConnaughey 1949) and its ecology has been studied by Ambrose (e.g. 1984, 1986, 1988) off Catalina Island in California. It is of similar size (at Catalina Island), appearance and habitat, except that it hatches as a "paralarva" (Young and Harman 1988) that spends several weeks to months in the plankton before

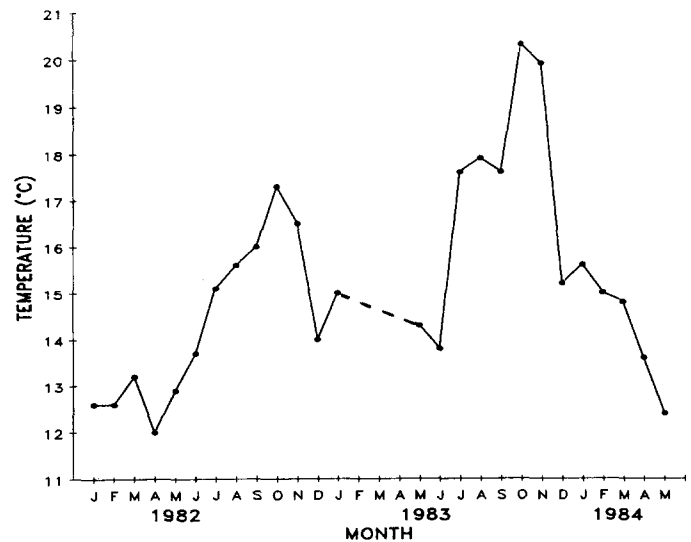


Fig. 5. Mean sea-water temperatures for first week of each month, taken from daily temperature records at University of California at Santa Barbara Marine Laboratory. Dashed portion of line indicates no data available from February through April of 1983

settling. Based upon size-frequency, abundance and reproduction data from field samples, Ambrose (1988) proposed two possible models of its life cycle: an "alternating generation" model in which octopuses that hatch early enough in spring grow sufficiently fast to spawn 13 mo after settling, and an "alternating years" model in which all octopuses take 2 yr to spawn. Although Ambrose's field data tend to support the "alternating years" model, our laboratory growth data indicate that, even at 16°C, *O. bimaculatus* could grow fast enough to spawn at 13 mo post-settlement. A major unknown factor with *O. bimaculatus* is the duration of the planktonic phase; settlement occurs throughout the year (Ambrose 1988).

However, *Octopus bimaculoides* with its direct development and very large, benthic adult-like hatchlings has a more predictable life cycle. Although few field data are available, we can estimate the following. In May 1984 we found many females brooding eggs at various stages of development (freshly-laid to posthatching) off Santa Barbara, the northernmost part of its range. With May to July temperatures about 14° to 17°C (Figs. 5), eggs would hatch about mid-summer, the time of highest temperature (17° to 19°C in most years). Even with the fast exponential growth in the first 3 to 5 mo posthatching, when temperatures are highest, it seems unlikely from our laboratory growth data at 18°C that *O. bimaculoides* could be large enough to spawn the following summer, since interim temperatures off Santa Barbara are in the range of 12° to 15°C. This scenario better fits the "alternating years" model of Ambrose (1988), in which *O. bimaculoides* would spawn after two years.

Conversely, in the southern range of *Octopus bimaculoides*, it is possible that the populations could spawn at one year of age ("alternating generation" model). Temperatures at Catalina Island (Ambrose 1988; his Fig. 2) are several degrees warmer throughout the year, and tempera-

tures nearshore can be warmer farther south in coastal areas between San Diego and San Quintin, Mexico. The surprising growth, vigor and early reproduction of this "cold-water species" cultured at a warm 23°C in our laboratory may be a reflection of its need to survive in the fluctuating environment of the intertidal zone. Experimentation in progress shows that *O. bimaculoides* can also grow well in reduced salinity (25‰), a trait very rare among cephalopods, which are generally stenohaline (the squid *Lolliguncula brevis* is the only documented exception: Hendrix et al. 1981). Furthermore, the ability of *O. bimaculoides* to grow rapidly and reproduce successfully at a temperature of 23°C suggests clearly that this species' geographic range could extend well south of its known southern limit on the upper Baja Peninsula (Forsythe and Hanlon 1988).

The strong growth and reproduction differences noted here, with a 5°C temperature difference, underscore the important effects of natural temperature anomalies such as El Niño. Off California between early 1983 and late 1984, there was a mean temperature rise of about 3.5°C, with some short-term anomalies of up to 6°C (McGowan, 1984). The temperature data from Santa Barbara reflect these increases (Fig. 5). In nature, the impact of major shifts in temperature on population dynamics and ecology is poorly understood, but would be expected to be greater on short-life-span predators such as *Octopus* spp. than on fishes that are iteroparous and longer-lived. The results of the present study combined with other published information (Forsythe and Hanlon 1988) would suggest the following effects of an El Niño event on field populations of *O. bimaculoides*: (1) the elevated temperature (up to 25°C) would have no predictable detrimental effect on the biology of this species, (2) females near maturity would spawn weeks to months sooner, (3) embryonic development in existing egg broods would be accelerated, and (4) hatchlings and juveniles still in the exponential growth phase would undergo a period of accelerated growth (assuming sufficient food availability).

This last effect would probably produce the greatest local impact ecologically. Using the temperature data in Fig. 5 for Santa Barbara, *Octopus bimaculoides* hatching in July of a "normal" year (1982) would grow for 5 mo at an average temperature of 16°C compared to 18.6°C in an El Niño year (1983). This slight difference can have a dramatic effect in an organism capable of exponential growth. Assuming a growth rate from hatching of 3.0% at 16.1°C vs 3.6% at 18.6°C, octopuses would be 2.5 times larger (16.6 g vs 6.7 g) after 5 mo in an El Niño year. The increased growth rate is produced by an increase in feeding rate due to increasing temperatures (food conversion efficiency or gross growth-efficiency remain constant). As the octopus population grows in weight it will be consuming a relatively greater biomass of prey than would be consumed in a year with normal temperatures. Such increased growth might be expected to have a measurable impact on prey species of juvenile *O. bimaculoides*, especially since juvenile octopuses are known to have feeding

rates on the order of 10 to 17% of their body weight per day (Forsythe 1984). Field data are needed to corroborate our laboratory results and to provide a baseline for ecological studies of this species.

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#### Literature cited

- Ambrose, R. F. (1984). Food preferences, prey availability, and the diet of *Octopus bimaculatus* Verrill. *J. exp. mar. Biol. Ecol.* 77: 29–44
- Ambrose, R. F. (1986). Effects of octopus predation on motile invertebrates in a rocky subtidal community. *Mar. Ecol. Prog. Ser.* 30: 261–273
- Ambrose, R. F. (1988). Population dynamics of *Octopus bimaculatus*: influence of life history patterns, synchronous reproduction and recruitment. *Malacologia* 29 (1): 23–29
- Boletzky, S. von (1974). The "larvae" of Cephalopoda: a review. *Thalassia jugosl.* 10: 45–76
- Boyle, P. R. (ed.) (1983). Cephalopod life cycles, Vol. I. Species accounts. Academic Press, London
- Boyle, P. R. (ed.) (1987). Cephalopod life cycles, Vol. II, Comparative reviews. Academic Press, London
- Brody, S. (1945). Bioenergetics and growth with special reference to the efficiency complex in domestic animals. Hafner Press, New York
- Brown, M. E. (1957). Experimental studies on growth. In: Brown, M. E. (ed.) *The physiology of fishes*, Vol. 1, Metabolism. Academic Press, New York, p. 361–400
- DeRusha, R. H., Forsythe, J. W., Hanlon, R. T. (1988). Laboratory growth, reproduction and life span of the Pacific pygmy octopus, *Octopus digueti*. *Pacif. Sci.* 41: 51–59
- Dorsey, E. M. (1976). Natural history and social behavior of *Octopus rubescens* Berry. Master's thesis, University of Washington, Friday Harbor
- Forsythe, J. W. (1984). *Octopus joubini* (Mollusca: Cephalopoda): a detailed study of growth through the full life cycle in a closed seawater system. *J. Zool., Lond.* 202: 393–417
- Forsythe, J. W., Hanlon, R. T. (1988). Behavior, body patterning and reproductive biology of *Octopus bimaculoides* from California. *Malacologia* 29 (1): 40–56
- Forsythe, J. W., Van Heukelem, W. F. (1987). Cephalopod growth. In: Boyle, P. R. (ed.) *Cephalopod life cycles, Volume II, Comparative reviews*. Academic Press, London, p. 135–155
- Gould, S. J. (1966). Allometry and size in ontogeny and phylogeny. *Biol. Rev.* 41: 587–640
- Guerra, A. (1979). Fitting a von Bertalanffy expression to *Octopus vulgaris* growth. *Investigación pesq.* 43: 319–329
- Hanlon, R. T. (1983). *Octopus briareus*. In: Boyle, P. R. (ed.) *Cephalopod life cycles, Vol. I. Species accounts*. Academic Press, London, p. 251–266
- Hanlon, R. T., Forsythe, J. W. (1985). Advances in the laboratory culture of octopuses for biomedical research. *Lab. Anim. Sci.* 35: 33–40
- Hartwick, B. (1983). *Octopus dofleini*. In: Boyle, P. R. (ed.) *Cephalopod life cycles, Vol. I, Species accounts*. Academic Press, London, p. 277–291

- Hendrix, J. P., Jr., Hulet, W. H., Greenberg, M. J. (1981). Salinity tolerance and the responses to hypoosmotic stress of the bay squid *Lolliguncula brevis*, a euryhaline cephalopod mollusc. *Comp. Biochem. Physiol.* 69A: 641–648
- Hochberg, F. G., Jr., Fields, W. G. (1980). The squids and octopuses. In: Morris, R. H., Abbott, D. P., Haderlie, E. C. (eds.) *Intertidal invertebrates of California*, Chapter 17: Cephalopoda. Stanford, Stanford University Press, p. 429–444 and P133–P136
- Mangold, K. (1983). *Octopus vulgaris*. In: Boyle, P. R. (ed.) *Cephalopod life cycles*, Vol. 1, Species accounts. Academic Press, London, p. 335–364
- McGowan, J. A. (1984). The California El Niño, 1983. *Oceanus* 27(2): 48–51
- Packard, A. (1972). Cephalopods and fish: the limits of convergence. *Biol. Rev.* 47: 241–307
- Pascual, E. (1978). Crecimiento y alimentación de tres generaciones de *Sepia officinalis* en cultivo. (Growth and feeding of three generations of lab-reared *Sepia officinalis*.) *Investigación pesq.* 42: 421–442
- Pickford, G. E., McConnaughey, B. H. (1949). The *Octopus bimaculatus* problem: a study in sibling species. *Bull. Bingham oceanogr. Coll.* 12(4): 1–66
- Richard, A. (1966). Action de la température sur l'évolution genitale de *Sepia officinalis* L. C. r. hebd. Séanc. Acad. Sci., Paris 263: 1998–2001
- Ricker, W. E. (1979). Growth rates and models. *Fish Physiol.* 8: 677–743
- Van Heukelem, W. F. (1976). Growth, bioenergetics and life-span of *Octopus cyanea* and *Octopus maya*. Ph.D. dissertation, University of Hawaii, Honolulu
- Van Heukelem, W. F. (1979). Environmental control of reproduction and life span in *Octopus*: an hypothesis. In: Stancyk, S. E. (ed.) *Reproductive ecology of marine invertebrates*. University of South Carolina Press, Columbia, p. 123–133
- Wells, M. J., Wells, J. (1970). Observations on the feeding, growth rate and habits of newly settled *Octopus cyanea*. *J. Zool., Lond.* 161: 65–74
- Young, R. E., Harman, R. F. (1988). "Larva," "paralarva" and "subadult" in cephalopod terminology. *Malacologia* 29 (1): 201–207
- Zar, J. H. (1974). *Biostatistical analysis*. Prentice-Hall Inc., Englewood Cliffs

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