# Larval Fish Recruitment and Research in the Americas 

Thirteenth Annual Larval Fish Conference<br>Mérida, México, May 1989

Robert D. Hoyt (editor)

U.S. Department of Commerce

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91. Marine flora and fauna of the northeastern United States, Echinodermata: Crinoidea, by Charles G. Messing and John H. Dearborn. August 1990, 30 p.

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# Larval Fish Recruitment and Research in the Americas 

Proceedings of the Thirteenth Annual<br>Larval Fish Conference<br>Mérida, México<br>21-26 May 1989

Robert D. Hoyt (editor)

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## U.S. DEPARTMENT OF COMMERCE

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The 13th Annual Larval Fish Conference and Annual Meeting of the American Fisheries Society Early Life History Section cohosted by Mote Marine Laboratory, United States, and the Instituto Nacional de la Pesca, Mexico, were held 21-26 May 1989, in Mérida, Yucz.tán, México. The purpose of holding the meeting in Mexico was to encourage the participation of our Latin American and Caribbean colleagues and to provide a forum for the exchange of ideas and information among researchers working in the Americas. More than 150 participants represented 24 U.S. states, the District of Columbia, and 13 foreign countries including Mexico, Canada, Puerto Rico, Costa Rico, Panama, Cuba, Columbia, Chile, Peru, Brazil, Argentina, South Africa, and West Germany.

The Conference began with registration and a social in the courtyard patio of the Mérida Holiday Inn. Fresh red grouper, the most important commercial finfish species of the State of Yucatán, was prepared and served by the hotel staff, courtesy of CPI, Itzamex, and the Terramar Trading Company.

Plenary sessions opened and closed the Conference. The first plenary session opened with a welcome from Ing. Carlos Rihani, the Federal and State Fisheries Delegate who represented Lic. Victor Manzanilla, the Governor of the State of Yucatán. Also adding a few welcoming remarks were Lic. Raúl Diego, Director of the Centro Regional de Investigación Pesquera Yucalpetén (Instituto Nacional de la Pesca), and Robert Werner, the AFS Early Life History Section President. Following the welcome, Cuban representative Biologist Maida Montolio presented a short tribute to the memory of Mar Júarez, the pioneer of ichthyoplankton research in Cuba, who died shortly before the Conference.

Four speakers delivered keynote addresses on the theme of the Conference, "Larval Fish Recruitment and Research
in the Americas'': William Richards, National Marine Fisheries Service-Miami (North America); Biologist Rosa Maria Olvera, Instituto Nacional de la Pesca, Mexico City (Mexico and Central America); Douglas Shapiro, University of Puerto Rico (Caribbean); and Juana D. de Ciechomski, Instituto Nacional de Investigación y Desarrollo Pesquero, Argentina (South America), all of whom provided a synopsis of the major areas of larval fish research being conducted in the geographic regions they represented. Juana D. de Ciechomski's paper entitled, "A Review of the Investigations on Early Developmental Stages and Larval Recruitment of Marine Fishes in South America," appears in this volume.

Following the first plenary session, four days of concurrent sessions covered the topics of migration and dispersal, taxonomy, recruitment, feeding ecology, reproductive strategies, ecology, fish culture, physiology, biochemistry and behavior, and a special scombrid session. A poster session and three major workshops were held during the Conference. William Richards conducted the Scombrid Workshop; Darrel Snyder the Taxonomy Workshop; and Stanley Warlan, Perce Powles, and J. Isley the Age and Growth Workshop. Also included were VCR presentations, exhibits by publishers and manufacturers and a special exhibit of larval fish reference literature compiled by Darrel Snyder valued at over $\$ 2,000$ and which was donated to the Instituto Nacional de la Pesca following the Conference. A total of twenty posters and eighty-nine oral papers were presented. Of the eighty-nine presentations given, twelve are published in this volume. Exclusing the keynote address of Juana D. de Ciechomski, they are organized into Reproduction and Biomass Estimation, Distribution and Abundance, Larval Transport and Migration, Microhabitat Selection, and Taxonomy and Morphology sections. Unlike other conference editions, the
topics in this symposium are diverse. As such, they are truly representative of the 13th Annual Larval Fish Conference which was in itself a break with tradition, overcoming language barriers and geographic boundaries-an example that diversity can provide the strength for unity.

There were ten candidates for the annual Sally Richardson Best Student Paper Award. The panel of judges, Grace Klein-MacPhee, University of Rhode Island, USA; Scott Holt, University of Texas, USA; Jeffrey Marliave, Vancouver Aquarium, Canada; and Yasunobu Matsuura, Instituto Oceanographico da Universidade de Sao Paulo, Brazil, presented the award to co-recipients David M. Goshorn (University of Delaware) for his paper, "The Diet of Larval Weakfish, Cynoscion regalis, in Delaware Bay and the Relationship of Prey Density to Larval Growth and Survival," and a Columbian student, Guillermo Moreno (Moss Landing Marine Laboratory, California) for the paper "Descriptions of the Early Rockfishes (Sebastes spp.) from Central California."

During the final plenary session there was an evaluation of the Conference goals and suggestions for continuing and enhancing international growth within the Early Life History Section.

A field trip was made to the Mayan ruins in Uxmal and a diving trip to the reefs off Cozumel. The Conference ended with a fiesta at a Mexican hacienda. Typical Yucatecan dishes were served while strolling mariachis serenaded banquet attendees.

As Conference Chairperson, I would like to thank Andrea Frank, my Assistant Coordinator, without whose help there would not have been a Conference. I also wish to thank other Mote Marine Laboratory personnel: Linda Franklin, Marilee Lipinski, and Robert Dixon; and NMFS personnel, William Richards and Nikki Bane, for their dedicated work on behalf of the Conference. I am indebted
to Roberto Freund who translated all the Conference abstracts and Roberto Donadi of Recursos Técnicos para Conferencias, S.C., for the excellent simultaneous translations. I also wish to thank Fred Binkowski for all his help in disseminating conference information in the Early Life History Section Newsletter. A special thank you is also in order for Conference sponsors NOAA/National Marine Fisheries Service Southeast Fisheries Center, United Nations Environment Programme, CPI, Itzamex, Terramar Trading Company, Mexicana Airlines, Collezio National de Educación Profesional Tecnica, and the Department of Tourism of the State of Yucatán. Finally, I would like to thank all those who attended and gave presentations.

All participants of the 13th Annual Larval Fish Conference were invited to submit their papers in English for this publication. Because English is a foreign language for many of the authors and international mail service can be slow and unpredictable, an extra burden was placed on many authors and especially on the reviewers. A total of thirtythree peer reviewers were involved in the review process, and their expertise and assistance in bringing this work to fruitation are sincerely appreciated. I would like to thank AFS/ELHS Editor Robert Hoyt and his Associate Editors, Dannie Hensley, Douglas Markle, John Olney, Robert Olson, Eileen Setzler-Hamilton, and William Szelistowski for their patience, perseverance, and commitment to producing this special symposium edition. For their Herculean efforts, this publication is dedicted to them.

Karen M. Burns
Conference Chairperson Mote Marine Laboratory
Sarasota, Florida

# A Review of Investigations on Early Developmental Stages and Larval Recruitment of Marine Fishes in South America 

JANINA D. de CIECHOMSKI*<br>Instituto Nacional de Investigación y Desarrollo Pesquero<br>C.C. 175, Mar del Plata, Argentina


#### Abstract

This paper presents a summary of studies on eggs, larvae, and juveniles of marine fishes in South America, with an emphasis on recruitment. A brief review is given of the history of these studies in different countries. The countries included are on the Pacific coast: Peru, Chile, Ecuador, Colombia; the Atlantic coast: Argentina, Uruguay, Brazil; and the Caribbean coast: Venezuela.


## Introduction

The purpose of this paper is to summarize research on early developmental stages of marine fishes and their recruitment recently carried out in South America (Fig. 1). This summary intends to point out various lines of research that have developed. Continued updating of our knowledge in this field will lead to improved research.

When one analyzes the evolution of these studies in South American countries, it becomes clear that South American advancements parallel advancements at a worldwide level. As in other parts of the world, scientists and managers realize that studies related to early developmental stages of fish can provide basic data for estimation both of biomass of spawning stocks and of future recruitment. This is especially evident in pelagic species that usually have a short lifespan, are directly dependent on the environment, and are subject to marked fluctuations in their populations. It has been generally accepted that fluctuations in recruitment represent one of the major causes of changes in abundance of fish populations under exploitation, and that the mechanisms determining the magnitude of recruitment depend not only on the production of eggs and larvae but also on their environmentally induced mortality (predation, starvation, advection). During the last decade in South America, oceanographers have worked with fishery biologists to improve understanding of this complex problem.

Studies related to early life history of marine fish in South America have gone from a descriptive stage, which existed in some countries in the 1950's, to a stage where investiga-

[^0]tions focus on national management of fishery resources. All of these studies began first in countries with economies heavily influenced by fishing activities, such as Chile and Peru. These countries border one of the richest current systems in the Eastern Pacific, the hydrographic characteristics of which assure high productivity (Parrish et al. 1983). This fact, together with world interest in the region, resulted in collaboration involving developed countries and international organizations that brought together scientists and funds to carry out programs that could not have been performed entirely within a single country.

An analysis of early life history and recruitment studies in each country, together with a summary of present knowledge, is presented in this paper.

## Pacific Coast

## Peru

The development of research related to early life history of fish in this country was directly related to the hydrobiological regime off Peru. Because of intense and permanent upwelling, the Peru off-shore area is one of the most productive in the world. The intense upwelling and cyclical perturbations, caused by the El Niño phenomenon, support its high productivity. Several major currents, directed towards the South Pole as well as the Equator, determine temperature, oxygen, and nutrient regimes. This habitat supports an important community of pelagic fishes that has a large biomass and few species. Marked fluctuations in stocks are observed. The most important exploited species are the anchovy Engraulis ringens, the sardine Sardinops sagax, the jack mackerel Trachurus murphi, and the mackerel Scomber japonicus.
$\qquad$


Figure 1
Location of main South American Laboratories where early life-history studies of fish are being carried out.

In the 1950's and 1960's, several papers were published in Peru on reproduction, morphology and distribution of fish eggs and larvae. In the 1960's, the research effort was directed toward the anchovy fishery which became the largest in the world, reaching 12.3 million metric tons in 1970.

Studies of ichthyoplankton and anchovy sexual maturity indicate that the spawning period for anchovy lasts 8 to 9 months, from July to March, with the highest intensity between August and September (Tsukuyama 1983). The most important area for spawning is located off-shore of the northern central Peruvian coast between lat. $06^{\circ} \mathrm{S}$ and $14^{\circ} \mathrm{S}$. This is positively correlated with phytoplankton
abundance (Rojas de Mendiola and Ochoa-Lopez 1980; Santander 1980).

Owing to intense fishing effort and environmental changes caused by "El Niño," as in 1972, anchovy biomass began to decline and other pelagic species such as the sardine, jackmackerel, and mackerel gained importance as fishery resources. The scientists of the Instituto del Mar del Peru (IMARPE) began to study other species and intensified their investigations on mechanisms which are related to fluctuations of pelagic resources and the interactions of several elements within the ecosystem.

The drastic fall of the anchovy fisheries in Peru moved many countries and international organizations, such as

FAO and UNESCO (IOC), to offer help (Dickie and Valdivia 1981). By the end of 1976, a cooperative agreement was signed between Peru and Canada and a Cooperative Research Program on the anchovy and its ecosystem (ICANE) began with the collaboration of IMARPE and several Canadian institutions. Scientists of other countries, mainly from the United States, also participated.

Later, a Peruvian-German Cooperation Program was signed (PROCOPA) involving the Technical Cooperation German Agency (GTZ) and IMARPE. Participants included W. Arntz and J. Albeit from Germany, as well as scientists from other countries and international organizations such as: A. Bakun, R. Parrish from SWFC, P. Smith, S.R. Goldberg, and B.J. Macewicz; D. Pauly from International Living Resources; and J. Csirke from FAO. All foreign groups closely collaborated with local scientists from IMARPE. Multidiscipline research efforts were the specific focus of the program. Recently, Peru has made plans to enter the yet to be implemented Sardine and Anchovy Recruitment Program (SARP).

Many papers resulting from these collaborations are included in three large volumes: 1) "Investigación cooperativa de la anchoveta y su ecosistema entre Perú y Canadá" (Bolet. Vol. extraord., Inst. del Mar del Peru, Callao, Peru, 1981; 2) 'El Niño, su impacto en la fauna marina" (W. Arntz, A. Landa, J. Tarasona, eds.), IMARPE, vol. extr., 1985 (together with G「Z); and 3) "The Peruvian anchoveta and its upwelling ecosystem: three decades of change'" (D. Pauly and I. Tsukayama, eds.), ICLARM Studies and Reviews 15. Many of these papers concern recruitment as related to parental stock, as well as to physical-oceanographic phenomena.

An important methodological paper is that of Santander et al. (1982) in which the sampling effort required for determining egg-production estimates for anchovy was developed. Santander (1987) analyzed the relationship between egg abundance and biomass of spawning stock of anchovy and found high cannibalism by adults on their own eggs. The same phenomenon was corroborated by Santander et al. (1983) and Santander and Alheit (1984), who noted that the main causes for embryonic mortality of anchovy were cannibalism and predation by sardine.

Parrish et al. (1981), working on the relationship between transport mechanisms and fish reproduction in the California Current, have provided interesting observations on anchovy reproduction in the Peruvian current system (Bakun and Parrish 1982). They pointed out that the spawning peak for anchovy occurs during winter, when the Ekman transport from the coast to off-shore areas is more intense, contradicting the currently accepted theory.

Pauly and Tsukayama (1987) and Mendelsohn and Mendo (1987) should also be noted in relation to anchovy recruitment. Pauly and Soriano (1987) analyzed anchovy monthly egg production between 1953 and 1981 and noted that production peaks did not coincide with the peaks of
the next recruitment, results similar to the observations of Mendelsohn and Mendo (1987). The latter authors suggest that it is possible to predict trends of future recruitment, information required for better management of anchovy stocks.

Santander and Flores (1983) studied egg and larval distribution of anchovy, sardine, jack mackerel, and mackerel in relation to temperature, salinity, coastal blooming, superficial currents, and "El Niño" from 1964 to 1982. Also of interest are studies by Castillo (1985) on variations of ichthyoplankton distribution and Carrasco (1988) on myctophid larval abundance in the Peruvian Sea.

Muck et al. (1987) discussed the relationships between the abundance of sardine, mackerel, and jack mackerel eggs and larvae and temperature, turbulence, and anchovy biomass. The studies of Espino and Wosnitza Mendo (1984) on the stock-recruitment relationship of the hake Merluccius gayi peruanus indicated that a Ricker-type curve may be applied, where stock and recruitment are inversely related. They found that in 1982 the number of recruits was at a low level and therefore recommended conservative management of the stock for 1983.

In 1981 the egg production method for assessing the biomass of spawning anchovy (Alheit et al. 1983; Santander et al. 1984) was first used in Peru. This method was developed by Hunter and Goldberg (1980) and Hunter and Macewicz (1980). In a related area, studies on fecundity and spawning frequency were carried out by Alarcón and Alheit (1984), Alegre and Alheit (1984), Alheit et al. (1984), Alheit (1985), and Lo et al. (1986) on sardine and Peña et al. (1986) on mackerel.

Research on anchovy larval feeding started by the end of the 1960's and has continued to the present. Some papers published in the 1980's include observations on maintenance ration, growth efficiency, and food density for the maintenance of anchovy and sardine larvae (Rojas de Mendiola and Gomez 1980, 1981; Ware et al. 1981; Villavicencio 1981; Villavicencio and Muck 1983). A study by Walsh et al. (1980) considered anchovy larval feeding in association with mortality and recruitment in the natural environment, including during an "El Niño". Villavicencio and Muck's (1985) work on minimum vital density of food for anchovy and sardine larvae based on a growth and feeding model is important. The same authors also discussed the negative influence of high temperatures on survival and development of embryos and larvae of both species. Additional studies are now being carried out on starvation in pelagic fish larvae. Recently, studies on age and daily growth of anchovy larvae have been initiated using daily rings in otoliths (Palomares et al. 1987). Another research effort presently being carried out in IMARPE is that related to Bothidae larvae by Girón.

IMARPE is the institution where the majority of early life-history studies are performed. Its scientists include S. Carrasco Barrera, O.S. de Castillo, M. Espino,
M. Girón, D. Gomes Caballero, O. Lotzano Rubio, J. Mendo, P. Muck, N. Ochoa Lopes, J. Tsukayama, Z. Villavicencio, C. Wosnitza-Mendo, and S. Zuta. In addition there is interest from some scientists from the Universities of Lima, Callao and Trujillo. Finally, Haydée Santander of IMARPE, who died in 1987, should be mentioned here because she obtained inestimable success in this field.

## Chile

The oceanographic conditions along Chile's coast correspond, as in Peru, to the eastern ocean currents. The area is characterized by high productivity, intense upwelling, and changes in oceanographic conditions caused by the cyclic phenomenon "El Niño." The hydrological characteristics are determined mainly by the influence of the Humboldt Current moving toward the Equator at a distance of 100 to 200 miles from shore. Inshore there is a poleward current called the Peru-Chile Countercurrent. Also, there are subsurface flows, with wind-driven flows predominating at the surface.

In this unstable habitat, several fish species reside, the pelagic ones being the most important. They are: the anchovy Engraulis ringens, the Spanish sardine Sardinops sagax, the jack mackerel Scomber japonicus, and the common sardine Clupea $=$ Strangomera bentincki. Interspecific interactions among the populations have subjected the species to many changes, particularly in connection with their abundance, geographic distribution, and spawning areas. The most spectacular recent events were the collapse of the anchovy and the "explosion" of the Spanish sardine. It has been determined that while the anchovy and Spanish sardine concentrate in definite spawning areas producing high egg densities, the jack mackerel spawns in larger areas with fewer eggs (Serra 1983).

Because of the importance of pelagic fisheries (approximately $80 \%$ of the total), the great majority of studies on reproduction and recruitment have been directed at these species. The first investigations were descriptive works on morphology and identification. They continue at present, the most recent including a comparative study of Clupeiformes in the Chilean coasts by Orellana and Balbontin (1983); Martínez et al. (1983) on Scomberesox saurus scombroides; Balbontin and Orellana (1983); Palma et al. (1985) and Acuña (1986, a and b) on myctophids; Herrera (1984) on blennids, Nelson et al. (1985) on Psychrolutes; Mujica (1988) on the jack mackerel; Rojas (1988) on the sardine and anchovy; Silva (1988) on flatfish P. adspersus; and Muñoz et al. (1983) on Paralichthys microps. Recent work dealing with ichthyoplankton in general include those of Palma Fuenzalida (1985), Aron (1986), Palma and Pizarro (1987), Palma (1988), Castillo et al. (1983), and of Loeb and Rojas (1988) on inter-annual variations between 1964 and 1983.

Some investigations on distribution and abundance of fish eggs with other biological parameters have been performed in order to assess the biomass of sardine and anchovy spawning stocks. After the collapse of its anchovy fishery, efforts since 1972 have been mainly concentrated on species that increased in abundance, particularly the sardine. Also, studies on other aspects of early life history of fish, particularly related to larval survival and recruitment have been undertaken.

Studies have been carried out on feeding, both at sea and in the laboratory, and include that by Herrera and Balbontin (1983) on intestinal evacuation rates and the incidence of feeding in Spanish sardine larvae. Garretón (1983) reported on the bio-energetics of yolk utilization in embryos and vitelline larvae of Hypomesus pretiosus and Balbontin et al. (1986) on food conposition, and size of prey in fish larvae of the Bransfield Strait (SIBEX) in the Antarctic. Uriarte and Balbontin (1987) studied starvation of Spanish sardine larvae, using morphometric and histological methods.

Together with studies on food and feeding habits, investigations on growth of fish larvae were undertaken using daily otolith rings on Spanish sardine (Garretón and Balbontin 1982; Castillo et al. 1985; Balbontin and Cannobio 1988; Garland 1988) and on Spanish sardine and anchovy (Herrera et al. 1985; Herrera et al., in press, a and b).

Studies have also been performed on parasitism in fish larvae by Herrera (1984, in press). At the Pontificia Universidad Católica of Chile in Talcahuano, Dr. Tarifeño is also carrying out a program related to physiologicalecological aspects of fish larvae.

In general, in all Chilean institutions where studies on early life history of fish are performed, the scientists are working on projects similar to those mentioned above, trying to enlarge the scope as far as possible.

A collaborative effort with FAO is presently underway. However, agreements with foreign institutions are rather rare. There are programs jointly developed by the Pontificia Universidad Católica of Chile in Talcahuano and Gothenburg University, through agreements with SAREC (Swedish Agency for Research Cooperation with Developing Countries) or direct joint projects between scientists of Instituto de Oceanología in Viña del Mar and of the Southwest Fisheries Center, La Jolla, California, on the influence of temperature on otolith growth in sardine larvae under experimental conditions. Chile has also made application to join the yet to be implemented SARP Project.

Research centers where studies on early life history of fish are carried out exist along the coasts of Chile as well as in Santiago. A list of the more active institutions and individuals presently involved include the following:

[^1]Universidad del Norte, Departamento de Biología Marina, Coquimbo: A. Aron, E. Acuña, H. Flores, A. Silva.
Instituto de Oceanología, Universidad de Valparaiso, Viña del Mar: F. Balbontin, M. Garreton.
Universidad Católica de Valparaiso, Facultad de Recursos Naturales, Valparaiso: S. Palma.
Instituto de Fomento Pesquero (IFOP). Santiago: A. Asencio, A. Mujica, O. Rojas.
Universidad de Concepción, Departamento de Oceanología, Concepción: T. Antezana.
Pontificia Universidad Católica de Chile, Talcahuano. BIOTECMAR: P. Bernal, G. Herrera, Gonzalez, E. Aguilera, H. Muñoz, E. Tarifeño, A. Troncoso.

Universidad Austral de Chile, Valdivia: C. Moreno, T. Rueda, G. Valenzuela.

## Ecuador

The waters of Ecuador are also under the hydrographic regime that dominates Peruvian and Chilean waters. Its coasts are not very irregular except for Guayaquil Gulf where the mouth of the Guayas River basin establishes a typical estuarine environment (García 1983). From the southeast, flows a frontal system or Frente Ecuatorial of superficial circulation, with different characteristics according to the season of the year (Enfield 1976; Pak and Zeneveld 1974). This system separates tropical superficial waters from the modified subtropical superficial ones. This is apparent during winter when the flow of the cold Humboldt Current to the northwest becomes stronger. The flow of warm waters from the north, which takes place in summer, moves it away southward. Abnormal displacements of this flow cause the "El Niño" phenomenum which produces drastic changes in biological productivity (Jiménez 1982, 1983).

French and Menz (1983) suggested that the front areas of off-shore Ecuador determine the optimum conditions for reproduction, feeding, and larval fish survival. The most intensive spawning of the chief commercial pelagic spe-cies-the mackerel Scomber japonicus, the sardine Sardinops sagax, the round sardine Etrumeus teres, Opisthonema spp., and the anchovy Cetengraulis mysticetus-takes place at times when oceanographic conditions are best for larval fish survival. Such conditions principally occur in the Gulf of Guayaquil.

Research on the early life history of fish in Ecuador is not very developed and deals mainly with distribution and abundance of eggs and larvae (Cajas and Hinostroza 1981 on clupeids and engraulids; Peribonio et al. 1981, Jiménez 1982, and García 1983 on the mackerel, sardine, round sardine, and other clupeids in the coastal waters of Ecuador and the Gulf of Guayaquil; García 1983 on mackerel; García et al. 1983 on sardine near the Galapagos Islands; and Garcia and Ochoa 1983 on the effect of "El Niño" on zooplankton and ichthyoplankton in Ecuadorian waters
between 1983 and 1985).
The few scientists who are carrying out studies in this field are M.L. García (retired at present), L. Mariduena, D. Hinostroza, L. Cajas, Y. Ochoa, and M. Luzuriaga de Cruz, mainly in Guayaquil at the Instituto Nacional de Pesca.
As far as international collaboration is concerned, it should be mentioned that many years ago there was a development Project of FAO and that Ecuador, Chile, and Peru tried to join the yet to be implemented SARP Program.

## Colombia

In Colombia, whose coasts lie mainly on the Pacific, studies on early stages of fish development have not been numerous. Greater effort has been directed to mollusks and crustaceans that have more economic importance. FAO also gave its support to such research many years ago.

In the 1970's, there were descriptive papers like those of Mercado-Silgado (1971) and Mercado-Silgado and Ciardelli (1972) on Megalops atlanticus, Artunduaga (1972) on Scomberomorus sierra, and Maldonado and Remolina (1975) on the anchovy, C. mysticetus. Mora-Lara (1983) in his paper on pelagic resources of Colombian waters commented on certain reproductive characteristics of some pelagic species.
Alvarez-León and Lesser (1986) in their paper on the recruitment of demersal resources, basically mollusks and crustaceans, also considered some fish species such as Mugil spp. and Eugerres plumieri. Their emphasis was the exploitation of this resource in coastal lagoons and estuaries.

The few scientists working on early life history of fishes are concentrated at the fisheries institute INDERENA in Bogota, or at Universidad Jorge Tadeo Lozano Bogota, Facultad de Ciencias del Mar, where several students have presented theses on ichthyoplankton.

## Caribbean Coast

## Venczuela

The coast of Venezuela lies on the Caribbean Sea. In that country very few scientists are working on ichthyoplankton and recruitment of marine fish. In the 1960's and 1970's, with the collaboration of FAO, some studies were conducted by Barrett and Howard (1961) on the anchovy, Cetengraulis mysticetus, Simpson (1965) on C. edentulus, and Simpson and Gonzalez (1967) and Lopez Rojas (1972) on the sardine, Sardinella anchovia. Recently, there has been increased interest in assessment of larval fish potential within the areas influenced by the predominant hydrological regime and its seasonal characteristics which have a noticeable influence on the fisheries in the country.

At the Fundación La Salle de Ciencias Naturales, Estación de Ciencias Marinas, Margarita, studies are carried out on some species such as clupeids and engraulids (Rodríguez et al. 1983, a and b) and mugilids and carangids (Rodríguez et al. (1983b). There also exists a collaboration at the personal level between the Centro Interdisciplinario de Ciencias Marinas (CICIMAR) in Baja California, Mexico, particularly in regard to clupeids and engraulids.

At INTECMAR at the Universidad Simon Bolivar in Caracas, studies on aspects of juvenile recruitment into fishing areas in demersal communities are being conducted (Penchaszadeh et al. 1986). In these investigations, ten fish species caught at the Triste Gulf are being studied. Among these ten species the most important are Lutjanus synagris, Sphyraena guachancho, Selene setapinnis, and Diapterus rhombeus.

The active Venezuelan early life-history institutions, including names of some scientists are the following:

Estación de Investigaciones Marinas (EDIMAR), Fundación La Salle, Margarita: C. Rodríguez.
Instituto Oceanográfico, Universidad de Oriente, Cumaná.
Centro de Investigaciones Científicas, Universidad de Oriente.
Instituto de Zoología Tropical, Universidad Central of Venezuela, Caracas.
Instituto de Tecnología del Mar (INTECMAR), Universidad Simon Bolivar, Caracas: P. Penchaszadeh.
Centro de Investigaciones Biológicas, Universidad de Julia, Maracaibo.

## Atlantic Coast

The waters at the south-western end of South America are subject to a complicated oceanographic regime that changes seasonally and depends mainly on displacements of principal marine currents: the Brazil Current of subtropical waters flowing to the South and the Malvinas Current of subantarctic waters flowing to the North. There are also local currents and the action of large rivers such as the River La Plata, the influence of which may be noted along the shores of Argentina and Uruguay. The north-western sector of the ocean between $10^{\circ} \mathrm{S}$ and $10^{\circ} \mathrm{N}$ is under the influence of the regime of warm tropical waters and the Amazon River.

The fronts of the various water masses are of great importance to the distribution and life processes of many marine organisms. This area of the Atlantic is characterized by a large continental shelf, particularly accentuated off Argentina, that is of great importance in the development and fate of early life stages of fish.

## Argentina

Studies on the early life history of fish began with the work of Ciechomski in the 1960's when the Instituto de Biologia Marina in Mar del Plata was created. This institute was later restructured and transformed into the present Instituto Nacional de Investigación y Desarrollo Pesquero (INIDEP).

At the beginning, studies were concerned with taxonomy and morphology of eggs and larvae for the purpose of identification. The research emphasis began to change with a 1967-73 FAO/Argentina Government agreement. From 1978 to 1979, agreements between Argentina, West Germany, and Japan resulted in 18 monthly cruises within Argentinan shelf waters, aboard the research vessels Walther Herwig and Shinkai Maru. Since 1981, two INIDEP-owned vessels have permitted greater continuity of data collection. It has been possible to study selectivity of plankton nets, handling of ichthyoplankton data, and sampling error (Ciechomski et al. 1983; Sánchez and Ciechomski 1984; Ciechomski and Sánchez 1986; Sánchez 1986; Ciechomski and Sánchez 1988; all in relation to anchovy (Engraulis anchoita) eggs and larvae.

Morphological and systematic studies of marine fish eggs and larvae have been published by Ciechomski and her collaborators since the beginning of the 1960's. Among the papers published in the 1980's are Ciechomski (1981) in summarizing the existing data in Atlas format; Ciechomski and Booman (1981) on Macrouronus whitsoni, Coelorhynchus fasciatus, Micromesistius australis, and Salilota australis; Ciechomski and Cassia (1982) on Cynoscion striatus; Ehrlich (1982) on Congiopodus peruvianus; Cassia (1984) on Stromateus brasilianus; Doseff and Rakitin (1987) on larvae of Trachurus lathami; Sánchez and Acha (1988) on scorpaenids, Sebastes oculatus, Helicolenus dactylopterus lahille, and Pontinus rathbuni; De Cabo (1989) on larvae of Genypterus blacodes, Trypterygion cunninghami, and Pinguipes spp.; Camina and Ciechomski (in press) on larvae of Basilichthys bonaeriersis argentinensis; and Cassia and Lasta (in press) on larvae of Sprattus fuegensis. Cassia is at present developing studies that allow for an early differentiation of the larvae of Engraulidae and Clupeidae living off the shore of Buenos Aires Province.

The following papers, published in the 1980's, deal with fish egg and larvae distribution in relation to environmental conditions and reproductive activity: Ciechomski et al. (1981); Ciechomski (1982) where only the Patagonian shelf is taken into consideration; Ciechomski and Booman (1983) on anchovy; Cassia and Booman (1985) on ichthyoplankton in general; Ciechomski et al. (1986a) on anchovy, including vertical distribution, and where it is shown that the majority of its eggs are distributed within the layers between 50 m and the surface; Ehrlich and Ciechomski (1986) on hake, Merluccius hubbsi, and its winter spawning; Ciechomski and Sánchez (1986) with a review of 20 years
of research; Lasta and Ciechomski (1988) on egg and larval distribution in Samborombon Bay, considered spawning and nursery ground for several species, and Ciechomski and Sánchez (in press) on anchovy. Also, two other papers by Hubold (1982, a and b) and Hubold and Ehrlich (1981), although published in Brazil, include work conducted in northern Argentina waters and deal with anchovy and Brevoortia spp., Lycengraulis grossidens and Anchoa martinii egg and larval distribution, respectively.

At present, comprehensive studies are being carried out by C. Lasta at Samborombon Bay in order to characterize this area as a spawning and nursery ground for Brevoortia aurea, Mugil liza, several species of the family Sciaenidae, and others. Other investigations on spawning and nursery grounds at Bahia Blanca relate to Basilichthys bonariensis, Ramnogaster arcuata, Brevoortia aurea, Parona signata and some species of Sciaenidae. Acha (in press) is studying early developmental stages of Mugil liza and other species at Mar Chiquita lagoon and Samborombon Bay. At the Instituto del Biología Marina y Pesquera "Almirante Storni" in San Antonio Oeste, L. Curtolo began studies on ichthyoplankton in San Matias Gulf (North Patagonia). At the Instituto Antartico Argentino, studies were conducted on the larval distribution of Antarctic fish species (Tomo 1981; Alder and Tomo 1987).

Studies have been performed by Ciechomski and Sánchez (1983) on the relationship between the spawning of some fish species and zooplankton abundance. Assessments of biomass of spawning adults from egg surveys and fecundity have been conducted on the hake Merluccius hubbsi (Ciechomski et al. 1983); the blue whiting Micromesistius australis (Sánchez et al. 1986); and the anchovy Engraulis anchoita (Ciechomski and Capezzani 1973; Ciechomski et al. 1983; Sánchez and Ciechomski 1984; Ciechomski et al. 1986a; Ciechomski and Sánchez 1988; in press).

In the paper by Ciechomski and Sánchez (1984) formulae were given to estimate daily egg production by relating age of embryos, temperature, and development stages, and by introducing a correction factor for embryonic mortality from the time of spawning up to collection. Studies have been conducted on the growth in weight and condition factor of anchovy larvae from different areas, months, and years (Ciechomski et al. 1986a; Sánchez et al., in press). All these studies, together with data on the incidence of food particles in the digestive tract, have allowed the characterization of different spawning areas in the sea, where different values in the nutritional condition factor have been detected and where differences in the larval mortality coefficients have been observed.

Two spawning areas for anchovy are considered of most interest, one near the mouth of the La Plata River and the other in the Patagonian region of the Valdez Peninsula, where a frontal system exists between stratified shelf waters and tide- and wind-homogenized coastal waters (Carreto et al. 1981; Glorioso 1987). This area is under the influence
of the above mentioned frontal system and is very important for the spawning of several fish species, including hake. For this reason, it is intensively studied by specialists in the different marine sciences of INIDEP.

In studies of larval mortality, aspects such as feeding, nutritional condition factor, and advection have been considered (Ciechomski and Sánchez, in press). Also, investigations have been initiated on predatory activity on fish eggs and larvae by gelatinous plankters (H. Mianzan of the Laboratory of Zooplankton of INIDEP). Studies on cannibalism in anchovy eggs are being conducted which formerly were carried out by V. Angelescu and A. Anganuzzi of INIDEP. More details on some of these aspects can be found in Sánchez (in press).

To determine the relationship between anchovy mortality and various oceanographic and biological processes which in turn affect the variability of anchovy recruitment, a team of international scientists from Argentina, Uruguay, Brazil, Germany, Sweden, and America carried out a cruise (sponsored by the IOC of UNESCO) on board the West German RV Meteor in the austral spring of 1989. Details about the objectives, plans, and participants have been sent by Sánchez to the ELHS Newsletter of the A.F.S. (1989).

In INIDEP there are ichthyoplankton samples from beyond the continental slope off Argentina obtained through an agreement with the U.S.S.R. that have yet to be analyzed.

Papers on reproduction and fecundity of marine fish have been published by Christiansen and Cousseau (1985) on the anchovy, Cassia (1986) on Cynoscion striatus, and Christiansen et al. (1986) and Olivieri and Christiansen (1987) on the hake. Research on this subject continues, particularly to estimate anchovy, hake, and mackerel spawning stocks.

The institution where the majority of studies related to early life stages of marine fish have been conducted is the Instituto Nacional de Investigacíon y Desarrollo Pesquero, Mar del Plata. The scientists at the Laboratory of Fish Biology and Ichthyoplankton are J.D. Ciechomski, R.P. Sánchez, M.D. Ehrlich, C.A. Lasta, M.C. Cassia, and M.E. Acha. Other institutions where early life-history studies are conducted are the following:

Instituto de Biología Marina y Pesquera "Almirante Storni" in San Antonio Oeste, Prov. Río Negro: L. Curtolo.

Instituto Antártico Argentino in Buenos Aires (larvae of Antarctic fish): A. Tomo (absent at present) and V.A. Alder.

## Uruguay

In this country on the Atlantic coast, where fishing activity is quite intensive, there are few studies on early life
stages of fish. Most information comes from bordering countries, mainly Argentina, where the Uruguayan waters have often been covered by cruises. The River Plata Treaty between these two countries created a common fishery zone and a common research area. This zone is markedly influenced by the La Plata River.

Uruguayans conducted some ichthyoplankton investigations from 1979 to 1983 whenever it was possible to obtain adequate materials. This research was carried out by G. Mantero at the Instituto Nacional de Pesca (INAPE) in Montevideo. She worked on the morphology of larvae and juveniles of Peprilus paru (Mantero 1981a), on the sciaenid Macrodon ancylodon (Mantero 1981b), and on the anchovy E. anchoita (Mantero 1983, 1986). For the latter, she described winter spawning from 1979 to 1982. She also studied the distribution of eggs and larvae of hake Merluccius hubbsi, Trichirus lepturus, and Stromateus sp. and assessed the spawning adult anchovy in this area from 1980 to 1981. During the last Symposium on Fisheries Investigations, held in December 1988 in Río Grande, Brazil, Nion and Ríos (1988) presented research conducted on the assessment of anchovy eggs and larvae in Uruguayan waters, together with studies on nursery grounds, mainly for the 0 group.

Other studies have included the work of Ubal et al. (1987) on the season, area, size, and structure of the spawning stock of hake, and of Grunwaldt (1986) on the fecundity of this species. Studies on early life-history stages of fish are carried out in GNAPE, Montevideo. There appears to be some interest in early life-history investigations at the Universidad de la Republica, Montevideo.

## Brazil

The first publications on early life stages of fish appeared in Brazil in the 1970's. During the period 1970-1980, many papers were published by Matsuura and his collaborators, Phonlor, Weiss and her collaborators, and others. Since 1980, this research has intensified, the number of investigators increased, and new aspects have been considered. Studies have mainly been conducted off the southeastern and mid-section coasts which support large commercial fish populations. There is little information available from the northern and northeastern zones. Brazil has a very large coastline, approximately 8000 km , from $5^{\circ} \mathrm{N}$ up to $34^{\circ} \mathrm{S}$, most of which encompasses the tropical and subtropical regions with their typical ichthyofauna.

Several species are being investigated in these studies, particularly those of economic value. One of the most important species for the Brazilian fishery is the sardine, Sardinella brasiliensis. In the southeastern area, the catch in 1973 reached 288000 metric tons (Matsuura 1987). The most intensive spawning and the highest density of larvae occurs between $22^{\circ}$ and $26^{\circ} \mathrm{C}$, and between 34.5 and $36 \%$ (Matsuura 1979). Since 1973, catches have
fluctuated from year to year and now are low, while the abundance of anchovy is high (Rossi-Wongtschowski from the Instituto Oceanografico, São Paulo, pers. commun. Dec. 1988). Consequently, sardine are being intensively studied under an Integrated Research Sardine Program, in which investigations on its early stages of development are included, and in which several institutions have participated, mainly the Instituto Oceanografico in São Paulo.

Research on this species is closely related to recruitment and the estimation of spawning stocks through the egg production method. Recent papers published on the sardine include Matsuura (1986, a and b) on its exploitation, recruitment, and assessment of spawning stocks; RossiWongtschowski et al. (1988) on spawning, reproductive cycle, and fecundity; and Campaner and Hondo (1987) on the co-occurrence of the copepod, Calanoides carinatus, and sardine larvae. Additional studies such as the estimation of the spawning stocks by Matsuura, Katsuragawa, and collaborators; larval mortality, calculated by Matsuura as $28 \%$ daily; and other studies have been carried out on the species. Also, investigations in the formation of daily increments in otoliths are being conducted.

Anchovy is another commercially important pelagic species and is shared by Brazil, Uruguay, and Argentina. Among more recent publications are Matsuura and Naka$\operatorname{tami}$ (1980) on methodological characteristics related to the collection of samples; Hubold (1982b) on reproductive ecology; Nakatami (1982), Weiss and Almeida (1983), and Phonlor (1984) on morphology and distribution of eggs and larvae; and Weiss et al. (in press) on egg mortality in different regions. Methodological aspects are being considered at present by Katsuragawa and others.

Publications on scombrids include larval abundance of five species of tunas between $23^{\circ}$ and $29^{\circ} \mathrm{S}$ by Matsuura and Sato (1981), spawning of Katsuwonus pelamis by Matsuura (1982, a and b; Matsuura 1986a), and distribution of larvae of several tuna and mackerel species by Mafalda and Weiss (1988). Yoneda, Katsuragawa, and co-workers at the Instituto Oceanografico in São Paulo are developing experiments on rearing scombrid and sciaenid larvae in the laboratory.

Papers on other species include Matsuura and Katsuragawa (1981, 1985) on larvae of Balistes capricus; Matsuura and Yoneda $(1986,1987)$ on larvae of Lophius gastrophysus; Weiss et al. (1987) on larvae of Antigonia capris and Zenopsis conchifer; Sato and Matsuura (1986) on larvae of Thyrsitops lepidopoides; Weiss and Hubold (1988) on eggs and larvae of Maurolicus muelleri. Sadowski and Almeida-Dias (1986) and Vieira (1988) have studied Mugil liza and M. platanus and discovered a long spawning period and egg and larvae transport of the former in the inverse direction of the migration of spawning adults. Sinque et al. (1983) worked on larval distribution of Sciaenidae in Paranagua Bay.

More general studies that deal with egg and larval distribution in relation to environmental conditions are Weiss (1981); Muelbert and Weiss (1991) on ichthyoplankton in the Los Patos Lagoon; Weiss and Souto (1988) on ichthyoplankton along the southern coast of Brazil; Sinque (in press) on ichthyoplankton in Paranagua Bay; and Monteiro-Ribas and Mureb (1988) on ichthyoplankton in the zone of upwelling off Cabo Frio. There are also publications on the distribution of Antarctic fish larvae in Bransfield Straits (Sinque et al. 1986, a and b). Recently Souto (1988, a and b) has investigated reproduction and extra vitelline nutrition of sharks, Mustelus schmitti and M. canis, and Ponz-Louro and Rossi-Wongtschowski (1988) the reproduction of various species of rays.

At the Universidad de Río Grande do Sul (FURG), Departamento de Oceanografía, Weiss and her group have been working with ichthyoplankton on the shelf off Río Grande and Los Patos Lagoon. At the same university, in the Laboratorio de Bioensayos, Phonlor with his coworkers have been conducting physiological experiments on fish larvae, such as time of yolk absorption, critical period, starvation, mortality and growth at various temperatures.

Sinque and his group at the Universidad Federal do Parana have continued their studies on coastal ichthyoplankton distribution, (including the estuarine area off Parana) and on the eggs and larvae of Antarctic fish in connection with the Biomass International Program.

At the Instituto de Oceanografico, São Paulo, where intensive research on sardine is performed, Matsuura, Katsuragawa, and collaborators are also working on microdistribution in spatial and temporal series of eggs and larvae of pelagic fish in the northern littoral area of São Paulo, on comparative ontogeny of genera of the family Carangidae in the southeastern region of Brazil, and on the development and distribution of some species of the genus Symphurus in the same region.

At the Universidad Federal in Río de Janeiro, A.C. Teixeira and collaborators are working on mesopelagic larvae from Guanabara Bay. They also plan to study the ichthyoplankton of Antarctic fish under an agreement between Brazil and the Federal Republic of Germany.

The principal Brazilian institutions where studies on early life stages of marine fish are carried out are

Fundaçao Universidade de Río Grande do Sul (FURG), Departamento de Oceanografía, Río Grande: G. Weiss, F.M. Souto. FURG, Laboratorio de Bioensayos: G. Phonlor.

Universidad Federal de Parana, Centro de Biologia Marina, Paranagua, Parana: Ch. Sinque, S. Koblitz, and L.M. Costa.
Instituto Oceanografíco, São Paulo: Y. Matsuura, M. Katsuragawa, N.T. Yoneda, H.L. Spach, F.W. Kurtz, E.M. Kitahara, E. Futema, E. Masami.

Universidad Federal de Río de Janeiro, Instituto de Biologia, Lab. Ictioplankton: A.C. Teixeira Bonecher, A. de Souza Días Neto.

## Conclusions

This review of studies of early life-history stages and recruitment in marine fish in South American countries shows that they began at the end of the 1950's with descriptive studies of eggs and larvae and progressed to studies of distribution in relation to the environment, to more highly developed studies in some countries today. Scientists and managers have agreed that an understanding of the factors that control recruitment is one of the most important requirements in fishery research today. Thus, studies on feeding, predation, and advection of eggs and larvae are considered in the context of the whole plankton community in which they occur.

Several international institutions recommend the study of early life-history stages of fish in relation to all environmental parameters, for predicting the success or failure of a year class. At present in South America, as in the other parts of the world, research on recruitment is being developed from various disciplines, including physics, meteorology, and solar energy, as the only way to explain and understand the causes that define the success of a year class. Spatio-temporal studies at the micro scale are also considered important, in order to obtain data on various aspects of population dynamics of fish embryos and larvae of different species. Finally, international cooperation has proved most valuable for encouragement and development of this discipline.

## Citations

Acha, E.M.
In press. Estudio anatómico-ecológico de la lisa (Mugil liza) durante su primer año de vida. Publ. Com. Téc. Mix. Fr. Mar.
Acuña, E.
1986a. El recurso mictófidos (Pisces, Myctophidae): Antecedentes en aguas chilenas y marco de referencia para su investigación. In La Pesca en Chile (P. Arana, ed.), p. 315-339. Esc. Cs. del Mar. Universidad Católica, Valparaíso, Edit. Universitaria Santiago, Chile.
1986b. Peces linterna (Familia Myctophidae) recolectados en el estrecho Bransfield (SIBEX-Fase II, Chile). Ser. Cient. INACH 35:111-124.
Alarcón, V., and J. Alheit.
1984. Frecuencia de desove y proporción sexual de la anchoveta (Engraulis ringens) y de la sardina (Sardinops sagax). (Resum.) I. Congr. Nac. Biol. Pesq. Perú, p. 23.
Alder, V.A., and A.P. Tomo.
1987. Distribución de postlarvas de peces en el sector Atlántico del Oceáno Antártico. Inst. Antartico Arg., Contrib. 342, 11 p.
Alegre, B., and J. Alheit.
1984. Estudio de fecundidad parcial de anchoveta (Engraulis ringens) y sardina (Sardinops sagax). (Resum.) 1st Congr. Nac. Biol. Pesq. Perú, p. 22.

Alheit, J.
1985. Spawning frequency of peruvian anchovies taken with a purse seine. NOAA Tech. Rep. NMFS SSRF 36:59-62.
Alheit, J., B. Alegre, V.H. Alarcón, and B. Macewicz.
1983. Batch fecundity and spawning frequency of various anchovies (Genus: Engraulis) populations from upwelling areas and their use for spawning biomass estimates. FAO Fish. Rep. 291(3): 977-985.
Alheit, J., V.H. Alarcón, and B.J. Macewicz.
1984. Spawning frequency and sex ratio in the Peruvian anchovy, Engraulis ringens. Calif. Coop. Fish. Invest. Rep. 25: 43-52.
Alvarez-León, R., and E.S. Lesser.
1986. Aspectos sobre el reclutamiento de los recursos demersales en las costas colombianas. IOC Workshop Rep. 44 (suppl.): 107-122.
Arntz, W., A. Landa, and J. Tarasona (eds.).
1985. El Niño, su impacto en la fauna marina. Inst. Maı. Perú (vol. extraord.), 221 p .
Aron, A.
1986. Taxonomía, distribución y abundancia de las larvas de pesces en el Atlántico Ecuatorial. Resultados de las investigaciones durante el Programa de Investigación, "FGGE-ECUADOR 79." Ph.D. diss., Univ. Kiel, Fed. Rep. Germany.
Artunduaga, E.
1972. La sierra Scomberomorus sierra (Jordán y Starks) del Pacífico colombiano. Divulgación Pesquera, INDERENA, Bogotá, 8(4):1-67.
Bakun, A., and R.H. Parrish.
1982. Turbulence, transport and pelagic fish in the California and Perú Current Systems. Calif. Coop. Fish. Invest. Rep. 23: 99-112.
Balbontin, F., and A. Cannobio.
1988. Growth and survival of Chilean sardine, Sardinops sagax musica, larvae reared at different densities of food. (Mimeo.) ICES, Early Life History Symp. 1988, Poster 119, 9 p.
Balbontin, F., and M.C. Orellana.
1983. Descripción de las larvas del pez linterna Hygophum bruuni del área de Valparaíso, Chile (Pisces, Myctophidae). Rev. Biol. Inst. Oceanolog. Univ. Valparaíso 19(3):205-216.
Balbontin, F., M. Garretón, and J. Neuling.
1986. Composición del alimento y tamaño de las presas en larvas de peces del Estrecho Bransfield, Antártica. Ser. Cient. INACH 33:125-144.
Barrett, I., and G.V. Howard.
1961. Studies of age, growth, sexual maturity and spawning of populations of anchoveta (Cetengraudis mysticetus) of the coast of the eastern tropical Pacific Ocean. Bull. Inter.-Am. Trop. Tuna Comm. 5:113-216.
Cajas, L., and D. Hinostroza.
1981. Huevos y larvas de clupeidos y engraúlidos en el Golfo de Guayaquil. Rev. Cienc. Mar y Limnol. Inst. Nac. Pesca 1(1): 37-47.
Camina, R.E., and J.D. de Ciechomski.
In press. Desarrollo embrionario y larval del pejerrey de Basilichthys bonaerensis argentinensis (Valenciennes, 1835) en la Bahía Blanca. Ser. Contrib. INTDEP, Mar del Plata.
Campaner, A.F., and S. Hondo.
1987. Distribution and co-occurrence of Calanoides carinatus and larvae of Sardinella brasiliensis and Engraulis anchoita over the south Brasilian Continental shelf. Bolm. Inst. Oceanogr., S. Paulo 35(1):7-16.
Carrasco, S.
1988. Abundancia larval de myctófidos en el Mar Peruano. Simposio int. rec. vivos pesq. Pacífico sudeste; 9-13 mayo 1988, Viña del Mar.

Carreto, J.J., M.L. Lasta, R. Negri, and H. Benavides.
1981. Los fenómenos de marea rojay toxicidad de moluscos bivalvos en el Mar Argentino. INIDEP, Contrib. 399, 30 p.
Cassia, M.C.
1984. Descripción y distribución de larvas del Pampanito, Stromateus brasiliensis Fowler, 1906 (Pisces, Stromateoidei), en el Mar Argentino. (Resum.) VII Jorn. Argent. Zool. Mar del Plata, 21-16 octubre 1984, 175 p.
1986. Reproducción y fecundidad de la pescadilla de red (Cynoscion striatus). Publ. Com. Téc. Mix. Fr. Mar. (Argentina-Uruguay) 1(1):191-203.
Cassia, M.C., and C.I. Booman.
1985. Distribución del ictioplancton en el Mar Argentino en los años 1981-1982. Physis (Secc. A) 43(1905):91-111.
Cassia, M.C., and C.A. Lasta.
In press. Description des larves de Sprattus fuegensis (Jenys 1842) (Pisces, Clupeiformes, Clupeidae) de l'Sud Ouest. Cybium.
Castillo, S.O. de.
1985. Variaciones en distribución del ictioplanctón en relación con las condiciones ambientales de enero a mayo de 1983. (Resum.) IX Congreso Latinoamer. Zool., Arequipa-Perú, 1985.
Castillo, G., H. Muñoz, H. González, and P. Bernal.
1983. Daily analysis of a fish larval assemblage at a fixed station in relation to environmental conditions in the Gulf of Arauco. Pontificia Universidad Catolica de Chile, Talcahuano, unpubl. manuscr.
Castillo, G., E. Aguilera, G. Herrera, P.A. Bernal, J.L. Buttler, J. Chang, H. González, C. Oyarzun, and C. Veloso.
1985. Larval growth rates of the Pacific sardine Sardinops sagax. Biología Pesquera, Chile 14:3-10.
Christiansen, H.E., and M.B. Cousseau.
1985. Aportes a la determinación de la frecuencia reproductiva de la anchoíta (Engraulis anchoita). Physis 43(104):7-17.
Christiansen, H.E., P.D. Glorioso, and C.E. Olivieri.
1986. Aplicación de la histología en la determinación de efectivos de merluza (Merluccius hubbsi). Tipificación de tejidos, cálculos de la fecundidad y vinculación con las condiciones ambientales. Publ. Com. 'Téc. Mix. Fr. Mar., Argentina-Uruguay 1(2): 567-574.
Ciechomski, J.D. de.
1981. Ictioplanctón. In Atlas de Zooplancton del Atlántico Sudoccidental (D. Boltovsky, ed.), p. 829-860. Inst. Nac. Invest. Desar. Pesq. Publicationes Especiales, Mar del Plato.
1982. Investigations on ichthyoplankton in the Patagonian Shelf off Argentina. Cybium 6(1):33-36.
Ciechomski, J.D. de, and C.I. Booman.
1981. Descripción de embriones y de áreas de reproducción de los granaderos Macrourus whitsoni y Coelorhynchus fasciatus, de la polaca Micromesistius australis y del bacalao austral Salilota australis en la zona Patagónica y Fueguina del Atlántico Sudoccidental. Physis 40(98):5-14.
1983. Distribución cuantitativa de huevos y larvas de anchoíta (Engraulis anchoita) en la plataforma continental, frente a las costas de la Argentina y Uruguay en el ciclo anual 1981/82. Ser. Contrib. INIDEP, Mar del Plata 431, 14 p.
Ciechomski, J.D. de, and D.A. Capezzani.
1973. Studies on the evaluation of the spawning stocks of the Argentinean anchovy, Engraulis anchoita, on the basis of egg surveys. Rapp. Proc. Verb. ICES 164:293-301.
Ciechomski, J.D. de, and M.C. Cassia.
1982. Observaciones sobre embriones, larvas y juveniles de pescadilla, Cynoscion striatus. Rev. Invest. Desar. Pesq. INIDEP, Mar del Plata 3:5-13.
Ciechomski, J.D. de, and R.P. Sánchez.
1983. Relationship between ichthyoplankton abundance and associated zooplankton biomass in the shelf waters off Argentina.

Biol. Oceanogr. Int. J. 3(1):77-101.
1984. Field estimates of embryonic mortality of Southwest Atlantic anchovy (Engraulis anchoita). Meeresfoorschung 30(3):172-187.
1986. Problemática del estudio de huevos y larvas de anchoíta (Engraulis anchoita) en relación con la evaluación de sus efectivos pesqueros. Reseña de veinte años de investigación. Publ. Com. Téc. Mix. Fr. Mar. 1(1):93-109.
1988. Análisis comparativo de las estimaciones de biomasa de la anchoíta Engraulis anchoita en el Atlántico Sudoccidental en diferentes años y con distintas metodologías. Publ. Com. Téc. Mix. Fr. Mar., Argentina-Uruguay No. 4:117-132.

Ciechomski, J.D. de, and R.P. Sánchez.
In press. Distribución y abundancia de huevos y larvas de Engraulis anchoita en la zona Común de Pesca Argentio-Uruguaya durante el periodo mayo 1986-abril 1987. Publ. Com. Téc. Mix. Fr. Mar. Argent., Uruguay.
Ciechomski, J.D. de, M.D. Ehrlich, C.A. Lasta, and R.P. Sánchez. 1981. Distribución de huevos y larvas de peces en el Mar Argentino y evaluación de los efectivos de desovantes de anchoíta y de merluza. Ser. Contrib. INIDEP, Mar del Plata 383: 59-79.
Ciechomski, J.D. de, R.P. Sánchez, C.A. Lasta, and M.D. Ehrlich. 1983. Distribución de huevos y larvas de anchoíta y merluza, evaluación de sus efectivos desovantes y análisis de los métodos empleados. Ser. Contrib. INIDEP, Mar del Plata 432:3-37.
Ciechomski, J.D. de, R.P. Sánchez, and C.A. Lasta.
1986a. Evaluación de la biomasa de adultos desovantes, distribución vertical y variación cuantitativa de la intensidad de los desoves de la anchoíta (Engraulis anchoita) durante la primavera de 1982. Rev. Invest. Des. Pesq. INIDEP, Mar del Plata 5:30-48.
Ciechomski, J.D. de, R.P. Sánchez, G. Alespeiti, and H. Regidor. 1986b. Estudio sobre el crecimiento en peso y factor de condición en larvas de anchoíta Engraulis anchoita Hubbs \& Marini. Variaciones regionales y anuales. Rev. Invest. Des. Pesq. INIDEP, Mar del Plata 5:183-193.
De Cabo, L.
1989. Estudio sobre el desarrollo larval de tres peces teleosteos del Mar Argentino: Genypterus blacodes, (Ophidiidae), Trypterygion cunninghami (Tripterygiidae) y Pinguipes spp. (Mugiloididae). Sem. Ocean. Biol. Univ. Buenos Aires, Inst. Nac. Invest. Des. Pesq., Mar del Plata
Dickie, L.M., and J.E. Valdivia G.
1981. Investigación cooperativa de la anchoveta y su ecosistema (ICANE) entre Perú y Canadá. Informe Sumario. Bol. Invest. Mar. Perú (vol. extraord.):I-XII, 288 p.
Doseff, A.J., and A. Rakitin.
1987. Estudio del desarrollo larval del surel, Trachurus lathami Nichols, 1920. Semin. Ocean. Biol. Univ. Buenos Aires, Invest. Nac. Invest. Des. Pesq., Mar del Plata.
Ehrlich, M.D.
1982. Desarrollo inicial de Congiopodus peruvianus (Cuvier 1829) Norman, 1937, Pisces, Congiopodidae en el Mar Argentino. Physis 41(100):21-27.
Ehrlich, M.D., and J.D. de Ciechomski.
1986. Nuevos aportes sobre el desove invernal de merluza (Merluccius $h u b b s i$ ) en aguas de la Plataforma del Atlántico Sudoccidental entre las latitudes de $34^{\circ}$ y $36^{\circ} \mathrm{S}$. Publ. Com. Téc. Mix. Fr. Mar., Argentina-Uruguay 1(2):299-300.
Enfield, D.
1976. Oceanografía de la región Norte del Frente Ecuatorial:aspectos físicos. Reunión de Trabajo sobre el fenómeno conocido como "El Niño." Quayaquil, Ecuador, 4-12 Diciembre de 1974. FAO, Inf. Pesca 195, 411 p.
Espino, M., and C. Wosnitza-Mendo.
1984. La relación entre stock y reclutamiento de la merluza peruana (Merluccius gayi peruanus). Bol. Invest. Mar. Perú 8(5):179-189.

French, S., and A. Menz.
1983. La pesqueria para peces pelágicos en el Ecuador y la distribución de las capturas en relación con factores ambientales. Rev. Com. Perm. Pacífico Sur 13:65-82.
García, M.L.
1983. Variabilidad en la distribución y abundancia de huevos y larvas de macarela (Scomber japonicus peruanus) y de algunos clupeidos en aguas ecuatorianas. In Actas de la consulta de expertos para examinar los cambios en la abundancia y composición de peces neríticos (Sharp, G.D., and J. Csirke, eds.), p. 151-178; 18 abril 1983, San Jose, Costa Rica. FAO Fish. Rep. 291, 2:151-178.
García, M.L., and Y. Ochoa.
1983. Efectos del evento El Niño sobre el zooplancton e ictioplancton en aguas ecuatorianas, 1983-1985. Unpubl. manuscr.
García, M.L., G. Larrea, and A. Vasquez.
1983. Zooplanctón, huevos y larvas de sardina en las Islas Galápagos. Unpubl. manuscr.
Garland, R.D.
1988. Determinación del momento del nacimiento de la sardina española Sardinops sagax musica (Pisces: Clupeidae) a través del recuento de lineas de crecimiento en sus otolitos sagitales. Simp. Fund. Univ., Río Grande Pesq. Pesquera, Brasil. Resum.:20.
Garretón, M.
1983. Bioenergetics of yolk utilization in embryos and yolk-sac larvae of the surf smelt Hypomesus pretiosus pretiosus (Girard 1855) under different incubation temperatures. M.S. thesis, Oregon State Univ., Corvallis, OR, 53 p.
Garretón, M., and F. Balbontín.
1982. Efecto de la temperatura en el desarrollo embrionario y crecimiento inicial de las larvas de la sardina española, Sardinops sagax musica, en condiciones de laboratorio. Rev. Biol. Mar., Valparaíso 18(1):56-71.
Glorioso, P.D.
1987. Temperature distribution related to shelf-sea fronts on the Patagonian Shelf. Cont. Shelf Res. 7(1):27-34.
Grunwaldt, P.
1986. Contribución al conocimiento de la fecundidad de la merluza (Merluccius hubbsi). Publ. Com. Téc. Mix. Fr. Mar. 1(1):66-74.
Herrera, G.
1984. Descripción de estados post-embrionales de Ophiogobius jenynsi Hoese, 1976 (Gobiidae: Blennioidei). Rev. Biol. Mar., Valparaíso 20(2):159-168.
In press. Incidence of parasitism of caligid stages on anchovy (Engraulis ringens) larvae. Talcahuano.
Herrera, G., and F. Balbontín.
1983. Tasa de evacuación intestinal e incidencia de alimentatción en larvas de Sardinops sagax musica (Pisces, Clupeiformes). Rev. Biol. Mar., Valparaíso 19(2):113-132.
Herrera, G., E. Aguilera, G. Castillo, and P.A. Bernal.
1985. Growth of anchovy larvae Engraulis ringens in Central Chile determined by daily increment counts in otoliths. Biología Pesquera, Chile 14:11-15.
Herrera, G., P. Bernal, E. Aguilera, and G. Castillo.
In press, a. Determinación de crecimiento larval de Sardinops sagax y de Engraulis ringens mediante lectura de incrementos diarios en otolitos. Memorias Segundo Congreso Latinoam. Cs. Mar., ALICMAR.
Herrera, G., E. Tarifeño, and M.C. Orellana.
In press, b. Estadíos de desarrollo larval de Strangomera bentincki y de Ethmidium maculatum. Pontificia Universidad Católica de Chile Talcahuano.
Hubold, G.
1982a. Eggs and larvae of Engraulis anchoita Hubbs and Marini, 1935, in the Southwest Atlantic between $25^{\circ} \mathrm{S}$ and $40^{\circ} \mathrm{S}$. Meeresforschung 29(4):208-219.

1982b. Zur Laichökologie der sudatlantische Sardelle Engraulis anchoita (Hubbs und Marini). Ph.D. diss., Christian-AlbrechtsUniversität zu Kiel, 161 p.
Hubold, G., and M.D. Ehrlich.
1981. Distribution of eggs and larvae of five clupeoid fish species in the Southwest Atlantic between $25^{\circ} \mathrm{S}$ and $40^{\circ} \mathrm{S}$. Meeresforschung 29(1):17-29.
Hunter, J.R., and S.R. Goldberg.
1980. Spawning incidence and batch fecundity in northern anchovy Engraulis mordax. Fish. Bull., U.S. 77:641-652.
Hunter, J.R., and B. Macewicz.
1980. Sexual maturity and fecundity, spawning frequency, and temporal pattern of spawning for the northern anchovy, Engraulis mordax, during the 1979 spawning season. Calif. Coop. Fish Invest. Rep. 21:139-149.
Jiménez, R.
1982. The 1976 El Niño: biological response off Ecuador. Paper presented at the Joint Oceanographic Assembly, August, 1982, Halifax, Canada.
1983. Variabilidad de las condiciones oceanografícas en el área del frente ecuatorial. FAO Fish. Rep. 291(2):131-150.
Lasta, C.A., and J.D. de Ciechomski.
1988. Primeros resultados sobre la distribución de huevos y larvas de peces en Bahía Samborombón en relación con temperatura y salinidad. Publ. Com. Téc. Mix. Fr. Mar., Argentina-Uruguay 4:133-141.
Lo, N.C.M., J. Alheit, and B. Alegre.
1986. Fecundidad parcíal de la sardina peruana (Sardinops sagax). Inst. Mar. Peru, Bol. 10(2):48-60.
Loeb, V.J., and O. Rojas.
1988. Interannual variation of ichthyoplankton composition and abundance relations off northern Chile, 1964-83. Fish. Bull., U.S. 86:1-24.

Lopez-Rojas, H.
1972. Distribución y abundancia estimada de huevos de la sardina (Sardinella anchovia) en la región oriental de Venezuela, 1968-1969. Proy. de Invest. y Desar. Pesq. Inf. Tec. 46:1-12.
Mafalda-Jun, P., and G. Weiss.
1988. Distribução y abundancia de larvas da suborden Scombroidei (Pisces: Perciformes) de Bom Abrigo ( $25^{\circ} \mathrm{S}$ ) a Ponta Mogotes $\left(38^{\circ} \mathrm{S}\right)$. (Resum.) Simp. Fund., Univ. Río Grande, Pesquisa Pesqueira, Brasil, Resum. p. 33.
Maldonado, J., and R. Remolina.
1975. Biología de la anchoveta colombiana Centengraulis mysticetus Gunther, 1866. Divulgación Pesquera, INDERENA, Bogotá 14(4-5), 55 p .
Mantero, G.
1981a. Descripción del juveniles de la ñata Peprilus paru (Linné, 1758). Perciformes, Stromateidae). In Res. Com. J. Cient. Nat. 2, p. 48.
1981b. Descripción del juveniles de la pescadilla de red Macrodon ancylodon (Bloch y Schneider, 1801). (Perciformes, Stromateidae.) Res. Com. J. Cient. Nat. 2, p. 60.
1983. Distribución y abundancia de huevos de anchoíta Eng:aulis anchoita Hubbs \& Marini en Marini, 1935) en la Zona Común de Pesca Argentino-Uruguaya, en los inviernos de 1979, 1980, 1981, 1982. (Resum.) VIII Simposio Latinoamer. Oceanogr. Biol. Montevideo 38 de noviembre-2 de diciembre 1982, p. 51.
1986. Análisis de los cambios morfológicos en la fase larval de la anchoíta (Engraulis anchoita). Publ. Com. Téc. Mix. Fr. Mar., 1(1):110-120.
Martínez, F.C., J. Oliva L., and C. Salazar Z.
1983. Sinopsis biológica de la agujilla Scomberesox saurus scombroides Richardson. Rev. Com. Perm. Pacífico Sur. 13:125-158.
Matsuura, Y.
1979. Distribution and abundance of eggs and larvae, Sardinella
brasiliensis, during 1974-75 and 1975-76 seasons. Bull. Jpn. Soc. Fish. Oceanogr. 34:1-12.
1982a. Distribution and abundance of eggs and larvae (Katsuwonus pelamis) larvae in eastern Brazilian waters. Bolm. Inst. Oceanogr., São Paulo 31(2):5-7.
1982b. Perspectiva de pesca de bonitos e atunes no Brasil. Ciencia e Cultura 34(3):333-339.
1986a. Distribution and abundance of skipjack larvae off the coasts of Brasil. Proc. ICAAT 1986:285-289.
1986b. Estimativa de taxa de explotação, recrutamento e biomassa do estoque de sardina-verdadeira, Sardinella brasiliensis, na região sudeste do Brasil. Ciencia e Cultura 38(5):892-904.
1987. Recursos pesqueiros. 31-Avaliaçao de recursos pesqueiros no Brasil. In Manual de pesca (M. Ogawa and J. Koike, eds.), p. 67-91. Assoc. Eng. Pesca., Fortaleza.

Matsuura, Y., and M. Katsuragawa.
1981. Larvae and juveniles of grey triggerfish, Balistes capriscus, from southern Brazil. Jpn. J. Ichthyol. 28(3):267-275.
1985. Osteological development of fins and their supports of larval grey triggerfish, Balistes capriscus. Jpn. J. Ichthyol. 31(4): 441-421.
Matsuura, Y., and K. Nakatami.
1980. Variability in quantity of zooplankton, fish eggs and larvae collected with two different mesh size Bongo nets. Atlantica, Río Grande. 4:43-52.
Matsuura, Y., and G. Sato.
1981. Distribution and abundance of scombrid larvae in southern Brasilian waters. Bull. Mar. Sci. 31(4):824-832.
Matsuura, Y., and N.T. Yoneda.
1986. Early development of the lophiid anglerfish Lophius gastrophysus. Fish. Bull., U.S. 84:429-436.
1987. Osteological development of the lophiid anglerfish, Lophius gastrophysus. Jpn. J. Ichthyol. 33(4):360-367.
Mendelsohn, R., and J. Mendo.
1987. Exploratory analysis of anchoveta recruitment off Peru and related environmental series. In The Peruvian anchoveta and its up-welling ecosystem: three decades of change (D. Pauly, and S. Tsukayama, eds.), p. 294-306. ICLARM Studies and Reviews 15.

Mercado-Silgado, J.E.
1971. Notas sobre los estados larvales del sábalo Megalops atlanticus Valenciennes, con comentarios sobre su importancia económica. Museo del Mar. Bol. 2:1-28.
Mercado-Silgado, J.E., and A. Ciardelli.
1972. Contribución a la morfología y organogénesis de los leptocéfalos del sábalo Megalops atlanticus (Pisces: Megalopidae). Bull. Mar. Sci. 22(2):153-184.
Monteiro-Ribas, W.M., and M.A. Mureb.
1988. Ictioplanctón de zona de influencia da ressurgencia de Cabo Frío (RJ) Brasil. (1986). Acta Biológica Leopoldensia, Año VIII, (1986):231-244.

Mora-Lara, O.
1983. Recursos pelágicos del Pacífico Colombiano. Rev. Com. Perm. Pacífico Sur 13:83-88.
Muck, P., O. Sandoval de Castillo, and S. Carrasco.
1987. Abundance of sardine, mackerel and horse mackerel eggs and larvae and their relationship to temperature, turbulence and anchoveta biomass off Peru. In The Peruvian anchoveta and its upwelling ecosystem: three decades of change (D. Pauly and I. Tsukayama, eds.), p. 268-275. ICLARM Studies and Reviews 15.

Muelbert, J.H., and G. Weiss.
1991. Abundance and distribution of fish larvae in the Channel Area of the Patos Lagoon Estuary, Brazil. In Larval fish recruitment and research in the Americas: proceedings of the thirteenth annual larval fish conference; 21-26 May 1989, Merida, Mexico (R.D. Hoyt, ed.), p. 43-54. Dep. Commer., NOAA Tech. Rep.

NMFS 95.
Mujica, A.
1988. Distribución espacio temporal de huevos y larvas de (T. murphyi). Simposio int. rec. vivos pesq. Pacífico sudeste; 9-13 Mayo 1988, Viña del Mar.
Muñoz, H., G. Herrera, and H. Fuentes.
1983. Desarrollo larval del lenguado de ojos chicos Paralichthys microps. Pontificia Universidad Católica de Chile, Talcahuano. Unpubl. manuscr.
Nakatami, K.
1982. Estudos sobre ovos e larvas de (Engraulis anchoita) (Hubbs \& Marini, 1935) (Teleostei, Engraulidae) colectados na regiao entre cabo Frío $\left(23^{\circ} \mathrm{S}\right)$ e Santa Marta Grande $\left(29^{\circ} \mathrm{S}\right)$. Disertacao de Mestrado, Inst. Oceanogr. São Paulo.
Nelson, J., N. Chirichigno, and F. Balbontin.
1985. New material of Psychrolutes sio (Scorpaeniformes, Psychrolutidae) from the eastern Pacific of South America and comments on the taxonomy of Psychrolutes inermis and Psychrolutes macrocephalus from the eastern Atlantic of Africa. Can. J. Zool. 63(2): 444-451.
Nion, H., and C. Rios.
1988. Los recursos pelágicos del Uruguay. (resum.) Simp. Fund., Univ. Río Grande, Pesquisa Pesqueira, Brasil, p. 37.
Olivieri, C., and H.E. Christiansen.
1987. Consideraciones preliminares sobre la frecuencia reproductiva de la merluza común (Merluccius hubbsi). Publ. Com. Téc. Mix. Fr. Mar., Argentina-Uruguay 3:67-72.

Orellana, M.C., and F. Balbontin.
1983. Estudio comparativo de las larvas de Clupeiformes de la costa de Chile. Rev. Biol. Mar. Valparaíso 19(1):1-46.
Pak, H., and J.R.V. Zeneveld.
1974. Equatorial front in the Eastern Pacific Ocean. J. Phys. Oceanogr. 4:570-578.
Palma, W.
1988. Monitoreo semanal e interdiario de huevos y larvas de sardina y anchoveta en la zona norte de Chile. Informe Final Proyecto INPESCON, Univers. Arturo Prat.
Palma, W., and R. Fuenzalida.
1985. Intensidad y extensión del desove de otoño para tres familias de peces pelágicos, Clupeidae, Engraulidae y Myctophidae en la zona norte de Chile. (Resum.) VI Jornadas Cs. Mar., Osorno, Chile.
Palma, W., and J. Pizarro.
1987. Análisis comunitario del ictioplancton en la zona norte de Chile. (Resum.) VIII Jornadas Cs. Mar., Concepción, Chile.
Palma, W., V. Fernandez, and J. Pizarro.
1985. Abundancia larval de especies de la familia Myctophidae en la zona norte de Chile. (Resum.) VI Jornadas Cs. Mar., Osorno, Chile.
Palomares, M.L., P. Muck, J. Mendo, E. Chuman, O. Gomez, and D. Pauly.
1987. Growth of the Peruvian anchoveta (Engraulis ringens) 1953 to 1982. In The Peruvian anchoveta and its upwelling ecosystem: three decades of change (D. Pauly and I. Tsukayama, eds.), p. 117-141. ICLARM Studies and Reviews 15.

Parrish, R.H., C.S. Nelson, and A. Bakun.
1981. Transport mechanisms and reproductive success of fishes in the California current. Biol. Oceanogr. 1(2):175-203.
Parrish, R.H., A. Bakun, D.M. Husby, and C.S. Nelson.
1983. Comparative climatology of selected environmental processes in relation to eastern boundary current pelagic fish reproduction. FAO Fish. Rep. 291, 3(FIRM/R291):3:731-777.
Pauly, D., and M. Soriano.
1987. Monthly spawning stock and egg production of Peruvian anchoveta (Engraulis ringens), 1953 to 1982. In The Peruvian anchoveta and its upwelling ecosystem: three decades of change
(D. Pauly and I. Tsukayama, eds.), p. 167-178. ICLARM Studies and Reviews 15.
Pauly, D., and I. Tsukayama.
1987. On the implementation of management-oriental fishery research: the case of the Peruvian anchoveta. In The Peruvian anchoveta and its upwelling ecosystem: three decades of change (D. Pauly and I. Tsukayama, eds.), p. 1-13. ICLARM Studies and Reviews 15.
Pauly, D., and I. Tsukayama (eds.).
1987. The Peruvian anchoveta and its upwelling ecosystem: three decades of change. ICLARM Studies and Reviews 15, 351 p.
Peña, N., J. Alheit, and M.E. Nakahama.
1986. Fecundidad parcial de la caballa del Perú (Scomber japonicus peruanus). Inst. Mar. Perú Biol. 10(4):93-104.
Penchaszadeh, P.E., J.J. Selaya, R. Malinet, and O. Defeo.
1986. Aspectos de reclutamiento en comunidades demersales en Golfo Triste, Venezuela. Intergovern. Oceanogr. Comm. Workshop Rep. 44(Suppl)):203-214.
Peribonio, R.G., R. Repetin, M.I. Cruz, D. Hinostroza, and M.E. Villarod.
1981. Estudio ecológico del mesoplanctón del Golfo de Guayaquil. Abundancia, ciclos nictimerales y relaciones entre el estuario del río Guayanas y el Oceáno. Bol. Cient. Téc. Inst. Nac. Pesca 4(2).
Phonlor, G.
1984. Morfología e biología de larvas de Engraulis anchoita (Hubbs \& Marini) (Osteichthyes, Engraulidae). Atlántica, Río Grande 7:85-98.
Ponz-Louro, M., and C.L.D.B. Rossi-Wongtschowski.
1988. Reprodução de Elasmobranchii do ecosistema costeiro da regiao de Ubatuba ( $23^{\circ} 30^{\prime}$ S), São Paulo, Brasil. (Resum.) Simp. Fund., Univ. Río Grande, Pesquisa Pesqueira, Brasil, p. 40.
Rodríguez, C., J. Castillo, and X. Elquetsabal.
1983. Distribución y abundancia relativa de Mugilidos y Carangidos en el Golfo de Venezuela durante el período febrero-agosto de 1986. Fundación La Salle, Margarita. Unpubl. manuscr.
Rodríguez, C., P. Núñez, and X. Elquetsabal.
1983. Distribución y abundancia relativa de Clupeidos y Engraúlidos en el Golfo de Cariaco durante el período febrero-agosto de 1986. Fundación La Salle, Margarita. Unpubl. manuscr.

Rojas, O .
1988. Variación interanual de la distributición y abundancia de huevos y larvas de sardina española $S$. sagax musica, y anchoveta E. ringens en el norte de Chile, 1983-1987. Simposio int. rec. vivos pesq. Pacífico sudeste; 9-13 mayo 1988, Viña del Mar.
Rojas de Mendiola, G., and O. Gómez.
1980. Alimento, sobrevivencia y tiempo de inactividad de las larvas de anchoveta (Engraulis ringens J.). Intergov. Ocean. Com. Workshop Rep. 28:275-286.
1981. Primera alimentación, sobrevivencia y tiempo de actividad de las larvas de anchoveta (Engraulis ringens J.). Bol. Inst. Mar Perú. Callao (vol. extraord.), p. 72-79.
Rojas de Mendiola, B., and N. Ochoa-López.
1980. Fitoplanctón y desove de la anchoveta (Engraulis ringens J.). Intergov. Ocean. Com. (FAO-UNESCO) Workshop Rep. 28: 241-254.
Rossi-Wongtschowski, C.L.D.B., M.S.M.T. Wentzel, and M.A.M. Oliveira.
1988. Programma integrado de estudios sobre sardina, Sardinella brasiliensis (Steindachner, 1879). II: Caracterizaçao macro e microscópica das gonadas, tipos de desove, ciclo reproductivo e fecundidad. (Resum.) Simp. Fund., Univ. Río Grande, Pesquisa Pesqueira, Brasil, 44 p .
Sadowski, V., E.R. Almeida-Dios de.
1986. Migração da tainha (Mugil cephalus Linnaeus, 1758 sensu latu) na costa sul do Brasil. Bolm. Inst. Pesca, São Paulo 13(1): 31-50.

Sánchez, R.P.
1986. Estudio sobre las variaciones espacio-temporales en los patrones de distribución de embriones y larvas de la anchoíta (Engraulis anchoita) en relación con la estimación de su intensidad reproductiva. Rev. Invest. Des. Pesq. INIDEP, Mar del Plata 5:92-142.
1989. Comparative study on recruitment and reproductive biology of sardine and anchovy in the South-West Atlantic Ocean. Am. Fish. Soc. Newsletter Ser. (Early Life History Sec.) 10(3):38-39.
In press. Estado actual de las investigaciones sobre ictioplancton marino en Argentina y Uruguay. Simp. Fund., Univ. Río Grande, Pesquisa-Pesqueira, Brasil.
Sánchez, R.P., and E.A. Acha.
1988. Development and occurrence of embryos, larvae and juveniles of Sebastes oculatus with reference to two Southwest Atlantic Scorpaenids: Helicolenus dactylopterus lahillei and Pontinus rathbuni. Meeresforchung 32:107-133.
Sánchez, R.P., and J.D. Ciechomski.
1984. Estimación de la biomasa de adultos desovantes de la anchoíta (Engraulis anchoita) en el área costera de la Prov. de Buenos Aires, durante la primavera de 1981 y el análisis comparativo de los recuentos de ictioplanctón recolectado con redes de diferentes mallas. Rev. Invest. Des. Pesq. INIDEP, Mar del Plata 4:49-61.
Sánchez, R.P., J.D. Ciechomski, and M.A. Acha.
1986. Estudios sobre reproducción y fecundidad de la polaca Micromesistius australis Norman 1937 en el Atlántico Sudoccidental, hasta $55^{\circ}$ S. Rev. Invest. Des. pesq. INIDEP, Mar del Plata. 6:21-43.
Sánchez, R.P., J.D. de Ciechomski, M. Pájaro, and A. Aubone.
In press. Análisis comparativo del crecimiento larval de Engraulis anchoita en diferentes áreas de cría del Mar Argentino y zona Común de Pesca Argentino-Uruguaya. Publ. Com. Téc. Mix. Fr. Mar., Argentina-Uruguay.
Santander, H.
1980. Fluctuaciones del desove de anchoveta y algunos factores relacionados. IOC Workshop Rep. 28:255-274.
1987. Relationships between anchoveta egg standing stock and parent biomass off Perú $4-14^{\circ}$ S. In The Peruvian anchovets and its upwelling ecosystem: three decades of change (D. Pauly and I. Tsukayama, eds.), p. 179-207. ICLARM Studies and Reviews 15.

Santander, H., and R. Flores.
1983. Los desoves y distribución larval de cuatro especies pelágicas y sus relaciones con las variaciones del ambiente marino frente al Perú. FAO Fish. Rep. 291 (FIRM/R291) 3:838-867.
Santander, H., and J. Alheit.
1984. Producción, mortalidad y predación de huevos de anchoveta (Engraulis ringens) y sardina (Sardinops sagax). (Resum.) I. Congr. Nac. Biol. Pesq. Perú, p. 24.
Santander, H., J. Alheit, and P.E. Smith.
1982. Determinación del esfuerzo del muestreo requerido para el estimado de producción de huevos de anchoveta, Engraulis ringens, frente al Perú. Instituto Mar. Perú 7(1):18.
Santander, H., J. Alheit, A.D. MacCall, and A. Alamo.
1983. Egg mortality of the Peruvian anchovy (Engraulis ringens) caused by cannibalism and predation by sardines (Sardinops sagax). FAO Fish. Rep. 291, 3(FIRM/R291): 3:1011-1025.
Santander, H., J. Alheit, and P.E. Smith.
1984. Estimación de la biomasa de la población desovante de anchoveta peruana (Engraulis ringens) en 1981 por aplicación del "Método de producción de huevos." Inst. Mar. Perú, Bol. 8(6): 212-250.
Sato, G., and Y. Matsuura.
1986. Early development of Thyrsitops lepidopoides (Pisces, Gempylidae). Bolm. Inst. Oceanogr., São Paulo 34:55-69.
Serra, J.R.
1983. Changes in the abundance of pelagic resources along the

Chilean Coast. In Proceedings of the expert consultation to examine changes in abundance and species composition of neritic fish resources (G.D. Sharp, J. Csirke, eds.), p. 255-284; 18-29 April 1983, San José, Costa Rica. FAO Fish. Rep. 291.
Silva, A.
1988. Consideraciones sobre el desarrollo y crecimiento larval de lenguado ( $P$. adspersus) cultivado en laboratorio. Simp. int. rec. vivos pesq. Pacífico Sudeste; 9-13 mayo 1988, Viña del mar.
Simpson, J.G.
1965. Estudio de las primeras etapas de desarrollo del rabo amarillo Centengraulis edentulus (Cuvier) en el oriente de Venezuela. Rep. Venezuela Min. Agric. Cría, Div. Invest., Centro Invest. Pesq., Ser. Biol. 1:1-24.
Simpson, J.G., and G.G. González.
1967. Algunos aspectos de las primeras etapas de vida y el medio ambiente de la sardina, Sardinella anchovia, en el oriente de Venezuela. Ser. Recursos y Explot. Pesq. 1. 2:38-93.
Sinque, $C$.
In press. Ictioplanctón do ecosistema da Bahía de Paranaguá (Paraná-Brasil).
Sinque, C., L.M. Costa, S. Koblitz, and M.J.C. Sena.
1983. Ichthyoplankton survey in the estuarine Bay of Paranagua and surrounding areas $\left(25^{\circ} 35^{\prime} \mathrm{S}\right)$ and $\left(48^{\circ} 10^{\prime} \mathrm{W}\right)$ Paraná, Brasil. Sciaenidae Teleostei. Symp. int. aquac. Coquimbo, Chile, Sept. 1983:445-465.
Sinque, C., S. Koblitz, and I.. Costa.
1986a. Ichthyoplanktón of Bransfield Strait-Antártica. Nerítica, Pontal do Sul PR1, 3:91-102.
1986b. Distribution of larval and postlarval Antartic fishes around Elephant Island and Bransfield Strait-Antártica. Nerítica, Pontal do Sul, PR1. 3:103-110.
Souto, C.M.
1988a. Nutrição extra-vitelica dos embriões de Mustelus schmitti. (Resum.) Simp. Fund. Univ. Río Grande, Pesquisa, Pesqueira, Brasil, p. 52.
1988b. Reprodução comparada dos cações Mustelus schmitii (Springer, 1939) e Mustelus canis (Mitchill 1815) no plataforma continental sulbrasileira. (Resum.) Simp. Fund., Univ. Río Grande, Pesquisa Pesqueira, Brasil, p. 53.
Tomo, A.P.
1981. Contribución al conocimiento de la fauna ictiológica del sector Antártico Argentino desde el punto de vista sistemático, ecológico y como recurso natural renovable. Dirección Nac. Antart. Inst. Antart. Argent. Publ. 14, 242 p.
Tsukayama, I.
1983. Recursos pelágicos y sus pesquerías en Perú. Rev. Com. Perm. Pacífico Sur 13:25-63.
Ubal, W., W. Norbis, B. Bosch, and D. Pagano.
1987. Estudio del stock desovante de la merluza (Merluccius hubbsi) en la Zona Común de pesca Argentino-Uruguaya. Publ. Com. Téc. Mix. Fr. Mar. 3:59-66.
Uriarte, I., and F. Balbontin.
1987. Caracterización del estado de hambruna en las larvas de sardina, Sardinops sagax musica (Pisces, Clupeiformes), mediante criterios morfométricos e histológicos. Rev. Biol. Mar. Valparaíso 23(12):77-106.
Vieira, J.P.
1988. Distribução abundancia e movimientos reproductivos da "tainha" (Mugil platanus Gunther, 1880) no sul do Brasil. (Resum.) Simp. Fund., Univ. Río Grande, Pesquisa Pesqueira, Brasil, p. 63.
Villavicencio, R.Z.
1981. Investigación preliminar de los requerimientos energéticos de anchoveta adulta (metabolismo estandar y actividad). Bol. Inst. Mar. Perú (vol. extraord):193-205.

Villavicencio, R.Z., and P. Muck.
1983. La racíon de mantenimiento, la densidad de mantenimiento y la eficiencia de crecimiento de Engraulis ringens y Sardinops sagax como una medida de su potencia ecológica. Bol. Inst. Mar. Perú. 7(4):72-107.
1985. La importancia del cambio de la temperatura y de la biomasa planctónica para la producción y mortalidad de anchoveta (Engraulis ringens) y (Sardinops sagax). In "El Niño" su impacto en la fauna marina (W. Arntz, A. Landa, J. Tarazona, eds.), p. 119-128. IMARPE (vol. extraord.)
Walsh, I.J. T.E. Whitledge, W.E. Esaias, R.L. Smith, S.A. Huntsman, H. Santander, and B. Rojas de Mendiola.
1980. The spawning habitat of the Peruvian anchovy, Engraulis ringens. Deep Sea Res. 27A:1-27.
Ware, D., B. Rojas de Mendiola, and D. Newhouse.
1981. Behaviour of the first feeding Peruvian anchovy larvae Engraulis ringens. Rapp. P.-v. Reún. Cons. Int. Explor. Mer 178:467-474.
Weiss, G.
1981. Ictioplanctón del Estuario de Lagoa Dos Patos, Brasil.

Ph.D. diss., Cs. Nat., Universidad Nacional de La Plata, 164 p. Weiss, G., and A. Almeida.
1983. Distribução sazonal quantitativa de ovos e larvas de Engraulis anchoita frente ao Estado de Río Grande do Sul, Brasil. (Resum.) VIII simp. Latinoamer. ocean. biol.; 28 nov.-2 dic. 1983, Montevideo, Uruguay, 50 p .
Weiss, G., and G. Hubold.
1988. Eggs and larvae of Maurolicus muelleri (Gmelin, 789) (Teleostei, Sternoptychidae) in the Southwest Atlantic. Meeresforschung 32(1):53.
Weiss, G., and C.M. Souto.
1988. Ictioplanctón marinho da costa sul do Brasil. (Resum.) Simp. Fund., Río Grande, Pesquisa Pesqueira, Brasil, p. 68.
Weiss, G., G. Hubold, and A.C.D. Bainy.
1987. Larval development of the zeiform fishes Antigonia capros Lowe, 1843 and Zenopsis conchifer (Lowe, 1852) from the southwest Atlantic. Cybium 2(1):79-92.
Weiss, G., G.O. Staffa, and D.A. Capezzani.
In press. Morfometría dos ovos de Engraulis anchoita para las latitudes de $26^{\circ} \mathrm{S}$ a $35^{\circ} \mathrm{S}$ do Atlántico Sul. Rio Grande, Brasil.

# Reproduction of Red Drum, Sciaenops ocellatus, in the Northcentral Gulf of Mexico: Seasonality and Spawner Biomass 

BRUCE H. COMYNS and JOANNE LYCZKOWSKI-SHULTZ<br>Gulf Coast Research Laboratory<br>P.O. Box 7000<br>Ocean Springs, MS 39564

DAVID L. NIELAND and CHARLES A. WILSON<br>Coastal Fisheries Institute<br>Center for Wetland Resources<br>Louisiana State University<br>Baton Rouge, LA 70803


#### Abstract

Identification of red drum Sciaenops ocellatus eggs, especially those in late stages of development, was found to be feasible using modal analysis of egg diameter frequency distributions. However, because of the scarcity of red drum eggs in collections, both the description of the seasonal spawning curve in 1984 and 1985 and a spawner biomass estimate in 1986 were based on larval abundance data and not egg data. Most spawning in east Louisiana (LA), Mississippi (MS), and Alabama (AL) coastal and shelf waters occurred from early September through early October when water temperatures over the shelf were decreasing rapidly from $27-29^{\circ} \mathrm{C}$ in early September to $24-25^{\circ} \mathrm{C}$ in early October. Red drum larvae used to estimate spawner biomass were collected during September 1986 over most of the east LA-MS-AL shelf with the exception of the midshelf region. The resulting first-order approximation of spawning stock in the study area, 567 metric tons ( 1.25 million pounds), is almost certainly an underestimate. Variability among catches of larvae from which egg production was estimated probably caused this underestimation. Sampling effort during a subsequent survey was tripled in an attempt to reduce sampling error and thus improve the accuracy of this estimate.


## Introduction

Concern that overfishing of adult red drum Sciaenops ocellatus in offshore federal waters was seriously reducing population levels in the northern Gulf of Mexico has resulted in emergency regulations during recent years. The main impetus for the emergency management decisions which closed down the commercial fishery and limited the sport fishery for this species was the lack of precise knowledge of the biomass of adult red drum in offshore waters. Insufficient data are available to directly estimate population size; because there is no longer a directed commercial fishery for this species, stock size estimates based on commercial catches are not available. Recently, two estimates of stock size have been produced using fishery-independent data from a mark and recapture effort in northcentral Gulf waters and a Gulf-wide aerial survey (Lohoefener et al. 1988; Nichols 1988).

Because no single method for stock size assessment is without potential biases and shortcomings (Ricker 1975; Smith and Richardson 1977), another fishery-independent method, based on the production of eggs and larvae, was used in an attempt to provide additional abundance information on that portion of the red drum stock in east Louisiana, Mississippi, and Alabama waters. This paper presents the resulting first-order estimate of red drum spawner biomass based on larval abundance, new data on batch fecundity and spawning frequency, and a description of the seasonal spawning pattern in the north-central Gulf of Mexico.

## Methods

## Collections

Ichthyoplankton collections were taken during 11 sampling


Figure 1
Occurrence of red drum eggs and larvae at nineteen locations on the east LA-MS-AL shelf during the 1986 ichthyoplanktin survey, 8-11 September. The hatched area encompasses assumed red drum spawning area. $\mathrm{H}=$ Horn Island; PB = Petit Bois Island.
periods ( 6 in 1984 and 5 in 1985) each extending 24-48 $h$ in duration at approximate biweekly intervals from late August to early November (Lyczkowski-Shultz et al. 1988a). Additional collections were taken during a shelfwide survey of 4 days duration, 8-11 September 1986 (Lyczkowski-Shultz 1987). Similar surveys were conducted in September 1987 and 1988. The sampling series in 1984 and 1985 were taken in Mississippi coastal waters in the near vicinity of a subsurface current drogue which identified the water mass to be sampled during a particular cruise (Lyczkowski-Shultz et al. 1988a). Collections from the 1986 survey were taken at nineteen predetermined locations on the east LA-MS-AL shelf (Fig. 1).

Collections were taken with a $1 \times 1.4$-m Tucker trawl
with an effective mouth opening of $1 \times 1 \mathrm{~m}$ when the net is fished at a $45^{\circ}$ angle. The opening-closing Tucker trawl consisted of three nets with $0.333-\mathrm{mm}$ mesh netting which were operated by messengers. Tow path for these collections was horizontal, and of approximately 5 -minutes duration. At each station in 1986, collections were made at nominal depth strata of 1 and 5 m . A third, deeper level, usually at $\geqslant 10 \mathrm{~m}$, was sampled in 1984 and 1985.
A conductivity/temperature/depth probe (CTD) mounted 0.5 m above the Tucker trawl on the conducting/towing cable monitored sample depth and obtained vertical profiles of temperature and salinity prior to sample collection. Digital flowmeters in each Tucker trawl measured volume filtered.

Samples were preserved at sea in $5-10 \%$ buffered formalin and later (one week to six months) transferred to $70 \%$ ethanol for final preservation. In the laboratory all fish larvae were removed from either the entire sample or from a one-half aliquot using a Motoda plankton splitter. Fish eggs were removed from either the whole sample or a known aliquot, depending on the density of eggs present. A target subsample of eggs from each sample was 200. Standard length (SL) of larvae was measured to the nearest 0.1 mm at $12 \times$; egg diameters were measured to the nearest 0.02 mm at $50 \times$ using a stereomicroscope.

## Egg Identification Methodology

Egg identifications were based on descriptions of red drum egg morphology (Holt et al. 1981; Holt et al. 1988), and comparison with modal size and size-frequency characteristics of known red drum eggs collected in the study area (Lyczkowski-Shultz et al. 1988b). The identity of these eggs was established after eggs in one collection were hatched in shipboard rearing containers and the larvae reared until positive identifications were made. Comparative information on the size and morphology of eggs of co-occurring taxa was taken largely from Moser et al. (1984).

## Biomass Estimation Procedure

The adult biomass of red drum was calculated with an equation modified from Houde (1977):

$$
B=\frac{P_{d} \cdot W}{F \cdot K \cdot S},
$$

where $B=$ biomass of adults in the stock
$P_{d}=$ total number of eggs spawned in one day (daily egg production)
$W=$ mean weight of adults in the stock (when sex ratio is $1: 1$ )
$F=$ mean batch fecundity of females
$K=$ proportion of adults that are females
$S=$ proportion of females that are spawning on a particular day (spawning fraction).

The low frequency of occurrence of red drum eggs precluded estimation of spawning biomass directly from egg densities. To calculate daily egg production $\left(P_{d}\right)$ from larval densities we estimated larval mortality rate and assumed that the mean density of individuals at time 0 approximated mean daily egg density. Mortality, the decline in numbers of fish over time, is the slope of the exponential function relating larval abundance and age:

$$
D_{t}=D_{0} \exp (-Z t) \quad(\text { Ricker 1975 })
$$

$$
\text { where } \begin{aligned}
D t= & \text { density of larvae at time } t \\
D_{0}= & \text { mean density of individuals at time } 0 \text { (i.e., } \\
& \text { density of eggs) } \\
Z= & \text { instantaneous mortality coefficient } \\
t= & \text { age of size class in days since spawning. }
\end{aligned}
$$

Size class age and densities were fitted to this exponential function with a nonlinear least squares routine using a pseudo-Gauss-Newton algorithm (Dixon and Brown 1979; Picquelle and Hewitt 1983). Ages were assigned to midpoints of the $0.5-\mathrm{mm}$ size classes using the growth equation $L=1.310 \exp (0.130 t)$, where $L=$ notochord or standard length and $t=$ larval age in days. This equation described the growth of red drum larvae in September 1985 at temperatures that were similar to those found during the 1986 survey, $27.8^{\circ}-29.0^{\circ} \mathrm{C}$ and $28.1^{\circ}-29.0^{\circ} \mathrm{C}$, respectively (Comyns et al. 1989). The density of each age class was corrected for stage duration by dividing the density estimate of each age class by their respective durations so that abundances of all size classes would be standardized with regard to time (Houde 1977).
Daily egg production was derived by multiplying the estimated mean egg (age 0 ) density by the volume of water in the spawning area. The spawning area in 1986 was defined as that portion of the survey area where red drum eggs and larvae were captured. One station (stn. 13) at which no red drum eggs or larvae were found was included in the spawning area because this station was surrounded by positive stations. The area associated with a station was taken to extend midway between that station and adjacent stations. Water volume was calculated by multiplying the area over which red drum spawned, by the depth to which collections were taken ( 5 m ).
Data on reproductive biology of red drum came from specimens collected by purse seine from Louisiana coastal waters in the vicinity of the mouth of the Mississippi River in September of 1986, 1987, and 1988, when spawning was assumed to be at or near its peak (Wilson et al. 1988). The red drum larvae used in the present study were collected just east of the Mississippi River mouth in Mississippi coastal waters. We assumed that reproductive parameters of red drum from these adjacent localities would not differ because the results of mark and recapture, electrophoretic, and morphological studies indicate that red drum in the Gulf of Mexico form a single genetic stock (Ramsey and Wakeman 1987; Nichols 1988; Poss 1988; Wakeman and Ramsey 1988).

Batch fecundity of an individual was determined from numbers of hydrated oocytes in six subsamples of ovarian tissue (Hunter et al. 1985; Wilson et al. 1988). Spawning females, i.e. fish that had spawned within 24 hours of sampling, were identified by the presence of post-ovulatory follicles in ovarian tissue (Hunter and Macewicz 1985; Fitzhugh et al. 1988; Wilson et al. 1988). Spawning fraction was the ratio of spawning females to the total number of

Table 1
Summary of collection dates, effort, frequency of occurrence, and mean density (all sampling depths and times combined) of Sciaenops ocellatus larvae captured near the subsurface current drogue in waters of the northcentral Gulf of Mexico during late summer and fall months 1984-85.

| Cruise | Date | Total volume filtered ( $\mathrm{m}^{3}$ ) | No. of collections | Occurrence |  | No. of specimens | Mean density (Number per $100 \mathrm{~m}^{3}$ ) |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  | All | Positive |
|  |  |  |  | frequency | percent |  | collections | collections |
| 84-8 | 30-31 Aug. 1984 | 1117 | 9 | 3 | 33 |  | 15 | 1.34 | 3.97 |
| 84-9-1 | 13-14 Sept. 1984 | 7427 | 29 | 28 | 97 | 5839 | 78.62 | 82.29 |
| 84-9-2 | 26-27 Sept. 1984 | 3016 | 10 | 9 | 90 | 256 | 8.49 | 9.12 |
| 84-10-1 | 10-11 Oct. 1984 | 4480 | 13 | 12 | 92 | 606 | 13.53 | 14.35 |
| 84-10-2 | 23-25 Oct. 1984 | 5016 | 19 | 1 | 5 | 2 | 0.04 | 0.63 |
| 84-11 | 07-09 Nov. 1984 | 2739 | 9 | 0 | 0 | 0 | 0 | 0 |
| 85-8 | 28 Aug. 1985 | 897 | 3 | 1 | 33 | 4 | 0.45 | 1.20 |
| 85-9-1 | 11-12 Sept. 1985 | 6864 | 27 | 21 | 78 | 447 | 6.51 | 8.44 |
| 85-9-2 | 25-26 Sept. 1985 | 2284 | 8 | 5 | 63 | 19 | 0.83 | 1.40 |
| 85-10 | 10-11 Oct. 1985 | 2445 | 9 | 7 | 78 | 54 | 2.21 | 2.90 |
| 85-11 | 4 Nov. 1985 | 1257 | 5 | 1 | 20 | 1 | 0.08 | 0.35 |

mature females in the sample. The inverse of this ratio estimates the time in days between successive spawnings and approximates spawning frequency. Mean batch fecundity and spawning fraction were determined from observations made in September of 1986, 1987, and 1988. Data from three years were combined because of the relatively small sample sizes in each individual year. Estimates of mean weight and sex ratio were calculated from observations made in September 1986.

Age in days of red drum larvae used to backcalculate spawning dates during the 1984 and 1985 seasons was determined from growth equations and/or raw length-atage data using counts of daily increments in sagittae of larvae from the same collections or from collections where water temperatures were most similar (Comyns et al. 1989). Approximate spawning dates were estimated for each cruise in 1984 and 85 by subtracting the modal age in days (or range in age when no mode was present) from the capture date.

## Results

## Occurrence of Red Drum Eggs

Examination of 38 collections from 11 sampling series conducted in late August to early November 1984 and 1985, and examination of 125 collections taken during shelfwide surveys in September of 1986, 1987, and 1988, resulted in only four occurrences of red drum eggs.

## Physical Environment

From September to early November the water column in
the sampling region was relatively well mixed (LyczkowskiShultz et al. 1988a). Temperatures generally ranged from 27 to $29^{\circ} \mathrm{C}$ in August and September, from 24 to $26^{\circ} \mathrm{C}$ in October, and from 22 to $23^{\circ} \mathrm{C}$ in early November. A seasonal trend in salinity was not as evident. Salinities within the upper 13 m generally ranged from 25 to 34 ppt in August and September, from 28 to 34 ppt in October, and from 32 to 35 ppt in early November.

## Spawning Seasonality

Seasonal Patterns in Abundance-Patterns of seasonal abundance of red drum larvae were consistent in 1984 and 1985. Red drum larvae were most numerous in midSeptember. Mean larval density (number of larvae per $100 \mathrm{~m}^{3}$, all collections) in mid-September 1984 and 1985 was 78.6 and 6.5 , respectively (Table 1). Lowest larval abundances were found in August and late OctoberNovember, when densities never exceeded 1.4. Observed spawning dates for red drum in LA-MS-AL coastal waters ranged from 21 August to 2 November (Table 2). Spawning dates backcalculated from collections during time of maximum larval abundance for each year were 7 and 8 September 1984 and 2, 6, and 9 September 1985.

## Biomass Estimate of Spawning Stock

Spawning Area-During the 1986 survey a total of 347 red drum larvae were collected at nine of the nineteen stations and probable red drum eggs were collected at two stations (Table 3). The spawning area generally enveloped the $10-\mathrm{fm}$ bottom contour and was estimated to cover 6.497 $\times 10^{9} \mathrm{~m}^{2}$ (Fig. 1). Red drum spawning does occur in

| Table 2 <br> Range in age and age of modal size class(es) (to the nearest 0.5 day) of red drum larvae collected during cruises from 1984-85. Backcalculated spawning dates were based on age of modal size class(es) except where no mode was present, then spawning date was calculated from the range in age. |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |
| Cruise | Date | Range in age (days) | Modal age (days) | Approx. spawning date |
| 84-8 | 30-31 Aug. 1984 | $3.0-7.5$ | 5.0 | 25 Aug. |
| 84-9-1 | 13-14 Sept. 1984 | $2.0-8.0$ | 5.0 | 8 Sept. |
| 84-9-2 | 26-27 Sept. 1984 | $2.0-7.5$ | 5.0 | 21 Sept. |
| 84-10-1 | 10-11 Oct. 1984 | 4.0-20.0 | $7.5,9.0$ | 3 Oct, 30 Sept. |
| 84-10-2 | 23-25 Oct. 1984 | - | 6.5 | 18 Oct. |
| 85-8 | 28 Aug. 1985 | $3.0-6.5$ | - | 25, 21 Aug. |
| 85-9-1 | 11-12 Sept. 1985 | 2.0-12.0 | 2.0,9.5 | 9, 2 Sept. |
| 85-9-2 | 25-26 Sept. 1985 | $2.0-7.0$ | 2.0 | 23 Sept. |
| 85-10 | 10-11 Oct. 1985 | $2.0-11.0$ | 6.0 | 4 Oct. |
| 85-11 | 4 Nov. 1985 | - | 2.0 | 2 Nov. |

Chandeleur Sound, but this area was not included in the 1986 survey. The volume of water used in estimating total daily egg production was approximately $3.249 \times$ $10^{10} \mathrm{~m}^{3}$.

Larval Mortality and Daily Egg Production-Mean density of red drum larvae within the spawning area (Figure 1) was 5.06 larvae per $100 \mathrm{~m}^{3}(\mathrm{SE}=2.01)$. Examination of the overall size-frequency distribution of red drum larvae from the 1986 survey indicated the presence of two size groups (Fig. 2). If combined, the less abundant group of smaller larvae would reduce the estimated mortality rate, not because mortality within this cohort was necessarily lower, but because there were fewer individuals in this cohort owing probably to either sampling variability or lower spawning intensity. Consequently, mortality and mean egg density estimates were derived only from the five older $0.5-\mathrm{mm}$ size classes ( $3-5 \mathrm{~mm}$ ).

The exponential function describing the nonlinear regression of duration-corrected age class densities (Table 4) on age for $3-5 \mathrm{~mm}$ red drum larvae was (Fig. 3):

$$
D_{t}=43.8 \exp (-0.521 t)
$$

where $43.8=$ mean density at time 0 (No. per 100 $\mathrm{m}^{3}$ ), i.e. mean daily egg density, with a standard error of 68.78
$0.521=$ instantaneous mortality rate with a standard error of 0.207 .

Daily egg production, estimated by multiplying mean daily egg density by the volume of water encompassing the 1986 spawning area, was $1.4229 \times 10^{10}$ eggs per day. The standard error of this estimate was $2.23 \times 10^{10}$ eggs per day.

Sex Ratio and Mean Weight-The sex ratio ( $K$ ) was assumed to be $1: 1$. Actual values were $51 \%$ male and $49 \%$ female ( $n=327$ ). Mean weight $(W)$ was 8.517 kg ( $n=$ $325, \mathrm{SE}=2.332$ ). Although mean weight of females is greater than mean weight of males, overall mean weight can be used in the biomass formula because the sex ratio is approximately $1: 1$.

Mean Batch Fecundity (F) and Spawning Fraction $(S)$-Mean batch fecundity was $2.128 \times 10^{6} \mathrm{eggs}(n=15$, $\mathrm{SE}=0.164 \times 10^{6}$ ). This estimate of mean batch fecundity is probably underestimated because purse seine sampling could be conducted only during daylight hours but red drum do not begin spawning until after dark (Holt et al. 1985). Consequently, eggs that would hydrate just prior to spawning would not have been included in estimates of mean batch fecundity. Mean spawning fraction, determined from the percentage of females with post-ovulatory follicles in ovarian tissue, was $0.20(n=245, \mathrm{SE}=0.045)$, indicating a spawning frequency of approximately once every five days in September.

Biomass Estimate-Incorporation of estimates of $P_{d}, W$, $F, K$, and $S$ into the biomass equation resulted in an estimated spawner biomass of 567 metric tons ( $1.25 \times$ $10^{6} \mathrm{lb}$ ).

## Discussion

Unlike most sciaenids in northcentral Gulf waters, the red drum has a restricted spawning period. Based on larval age and abundance data, red drum spawning in 1984 and 1985 extended from August until late October or early November, with peak spawning in September. During the

Table 3
Catch summary for red drum eggs and larvae collected during an ichthyoplankton survey of east LA-MS-AL shelf and coastal waters, 8-11 September 1986.

| Station | Date | Time | Volume filtered ( $\mathrm{m}^{3}$ ) | Sampling depth | Eggs |  | Larvae |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | Total no. | No. per $100 \mathrm{~m}^{3}$ | Total no. | No. per $100 \mathrm{~m}^{3}$ |
| 01 | 09/08/86 | 1807 | 504.5 | 5 | 0 | 0.00 | 0 | 0.00 |
| 01 | 09/08/86 | 1812 | 313.9 | 1 | 0 | 0.00 | 4 | 1.27 |
| 02 | 09/08/86 | 0000 | 284.8 | 1 | 0 | 0.00 | 4 | 1.40 |
| 02 | 09/08/86 | 0005 | 303.4 | 5 | 0 | 0.00 | 4 | 1.32 |
| 03 | 09/09/86 | 0325 | 267.8 | 1 | 2240 | 836.00 | 0 | 0.00 |
| 03 | 09/09/86 | 0329 | 266.1 | 5 | 0 | 0.00 | 0 | 0.00 |
| 04 | 09/09/86 | 0729 | 255.0 | 5 | 0 | 0.00 | 24 | 9.41 |
| 04 | 09/09/86 | 0734 | 309.0 | 1 | 0 | 0.00 | 4 | 1.29 |
| $05^{\text {a }}$ | 09/09/86 | 1201 | 231.6 | 5 | 0 | 0.00 | 0 | 0.00 |
| $05^{\text {a }}$ | 09/09/86 | 1206 | 221.9 | 1 | 0 | 0.00 | 0 | 0.00 |
| 06 | 09/09/86 | 1616 | 374.7 | 5 | 0 | 0.00 | 40 | 10.68 |
| 06 | 09/09/86 | 1621 | 311.9 | 1 | 0 | 0.00 | 4 | 1.28 |
| 07 | 09/09/86 | 1959 | 262.4 | 1 | 0 | 0.00 | 1 | 0.38 |
| 07 | 09/09/86 | 2004 | 165.8 | 5 | 0 | 0.00 | 0 | 0.00 |
| $08^{\text {a }}$ | 09/09/86 | 2240 | 212.5 | 1 | 0 | 0.00 | 0 | 0.00 |
| $08^{\text {a }}$ | 09/09/86 | 2245 | 192.3 | 5 | 0 | 0.00 | 0 | 0.00 |
| $09^{a}$ | 09/10/86 | 0149 | 234.8 | 1 | 0 | 0.00 | 0 | 0.00 |
| $09^{\text {a }}$ | 09/10/86 | 0154 | 262.0 | 5 | 0 | 0.00 | 0 | 0.00 |
| 10 | 09/10/86 | 0444 | 268.8 | 1 | 24 | 9.00 | 0 | 0.00 |
| 10 | 09/10/86 | 0449 | 227.3 | 5 | 0 | 0.00 | 0 | 0.00 |
| 11 | 09/10/86 | 0733 | 376.2 | 5 | 0 | 0.00 | 4 | 1.06 |
| 11 | 09/10/86 | 0739 | 222.9 | 1 | 0 | 0.00 | 4 | 1.79 |
| 12 | 09/10/86 | 1124 | 300.8 | 5 | 0 | 0.00 | 66 | 21.94 |
| 12 | 09/10/86 | 1130 | 212.9 | 1 | 0 | 0.00 | 106 | 49.79 |
| 13 | 09/10/86 | 1348 | 242.9 | 5 | 0 | 0.00 | 0 | 0.00 |
| 13 | 09/10/86 | 1353 | 215.0 | 1 | 0 | 0.00 | 0 | 0.00 |
| $14^{\text {a }}$ | 09/10/86 | 1558 | 321.3 | 5 | 0 | 0.00 | 0 | 0.00 |
| $14^{\text {a }}$ | 09/10/86 | 1603 | 253.0 | 1 | 0 | 0.00 | 0 | 0.00 |
| $15^{a}$ | 09/10/86 | 1857 | 327.3 | 1 | 0 | 0.00 | 0 | 0.00 |
| $15^{\text {a }}$ | 09/10/86 | 1901 | 404.1 | 5 | 0 | 0.00 | 0 | 0.00 |
| $16^{a}$ | 09/10/86 | 2123 | 216.7 | 1 | 0 | 0.00 | 0 | 0.00 |
| $16^{a}$ | 09/10/86 | 2129 | 213.1 | 5 | 0 | 0.00 | 0 | 0.00 |
| $17^{*}$ | 09/10/86 | 2310 | 264.0 | 1 | 0 | 0.00 | 0 | 0.00 |
| $17^{a}$ | 09/10/86 | 2315 | 222.8 | 5 | 0 | 0.00 | 0 | 0.00 |
| 18 | 09/11/86 | 0130 | 259.5 | 1 | 0 | 0.00 | 2 | 0.77 |
| 18 | 09/11/86 | 0135 | 281.6 | 5 | 0 | 0.00 | 0 | 0.00 |
| 19 | 09/11/86 | 0419 | 360.1 | 1 | 0 | 0.00 | 12 | 3.33 |
| 19 | 09/11/86 | 0424 | 274.7 | 5 | 0 | 0.00 | 68 | 24.71 |

${ }^{a}$ Stations were not considered to be in spawning area and were consequently omitted from calculations of larval density and water volume.
period of peak spawning, water temperatures over the shelf decreased from $27-29^{\circ} \mathrm{C}$ in early September to $24-25^{\circ} \mathrm{C}$ in early October 1984 and 1985. Although our collections did not extend beyond early November, water temperatures in November 1984 and 1985 were already approaching $20^{\circ} \mathrm{C}$, the lower limit for spawning in this species (Holt et al. 1981). Data on the reproductive biology of red drum from the northcentral Gulf of Mexico also indicated a restricted spawning season, with gonosomatic indices of both sexes and oocyte maturity peaking in September (Wilson et al. 1988). Peters and McMichael (1987) noted
that within-season spawning peaks of red drum in the Tampa Bay area of Florida were associated with new and full moons. Our data agree with these observations. Peak spawning in east LA-MS-AL coastal waters occurred within a week of the September full moon in 1984 and 1985.

In order to estimate egg production from egg or larva data, samples should be representative of the entire seasonal and areal extent of spawning. Temporal coverage of spawning in species with relatively short spawning seasons, like red drum, may not be as critical if the time


Figure 2
Size frequency distribution of red drum larvae captured during the 1986 survey of east LA-MS-AL shelf and coastal waters. Shaded size classes were used to estimate larval mortality and mean egg density.

Table 4
Age and duration-corrected density of size classes used to estimate mortality rate of red drum larvae.

| Size <br> class <br> $(\mathrm{mm})$ | Mean <br> age <br> (days) | Stage <br> duration <br> (days) | Density <br> $\left(\# / 100 \mathrm{~m}^{3}\right)$ | Duration <br> corrected <br> density |
| :---: | :---: | :---: | :---: | :---: |
| $3.0-3.4$ | 6.99 | 1.19 | 1.268 | 1.065 |
| $3.5-3.9$ | 8.09 | 1.03 | 0.976 | 0.948 |
| $4.0-4.4$ | 9.05 | 0.90 | 0.117 | 0.130 |
| $4.5-4.9$ | 9.91 | 0.81 | 0.291 | 0.359 |
| $5.0-5.4$ | 10.68 | 0.74 | 0.029 | 0.039 |

of most active spawning is sampled or a representative sample of spawning is obtained (Smith and Richardson 1977; Smith and Hewitt 1985). The assumption that the September 1986 survey was conducted when red drum were most actively spawning is supported by historical data on adult gonadal condition, as well as, temporal abundance patterns of larvae and postlarvae.

The spawning area represented by our 1986 collections is only a portion of red drum spawning grounds in the Gulf of Mexico, which include inner shelf and coastal waters from Mexico to the Everglades. That our survey area represents a logical unit or stratum in which to estimate spawner biomass is supported by recent Gulfwide stock assessment data from mark/recapture and aerial survey studies (Lohoefener et al. 1988; Nichols 1988). These data indicate that the biomass of red drum along the northwest Florida coast, ie. the area east of our study area, is very low compared to the area off Alabama, Mississippi, and

Louisiana. We have assumed, therefore, that spawning activity east of our study area is minimal. To the west of our study site, the Mississippi River delta restricted movement of adults into and out of our survey area during the 3 to 4 days over which spawning activity (egg and larval production) was measured in 1986. Nichol's (1988) tag recapture data support this contention in that few fish were found to have crossed the river mouth during his study.

Egg production can be calculated either directly from egg abundance or indirectly from larva abundance if larval mortality rate can be calculated (Sette and Ahlstrom 1948; Houde 1977; Parker 1980, McGowan and Richards 1986). In this study, egg production was back calculated from larval densities because of the low frequency of occurrence of red drum eggs, caused most likely by the short incubation period (1 day) (Holt et al. 1985; Holt et al. 1988) and highly contagious spatial distribution of the eggs. Use of larvae instead of eggs to calculate egg production can cause numerous problems which may bias and, most certainly, reduce precision of the resultant spawner biomass estimate. Foremost among these problems is that eggs and newly hatched larvae may experience different mortality rates, being either higher (Smith 1973; Houde 1977) or lower (Picquelle and Hewitt 1983), than older larvae. The direction and magnitude of ontogenetic differences in mortality rate of red drum is unknown.

Another serious limitation in estimating egg production from larval densities is the highly variable nature of larval catches on which the slope of the larval abundance vs. age regression, ie. instantaneous mortality rate ( $Z$ ), is based. To calculate a meaningful value of $Z$, larval density must


Figure 3
The relationship between duration corrected density and mean age of 3.0-5.0 mm size classes of red drum larvae. The solid line describes the exponential function of the data fitted to a nonlinear regression, and the dashed line describes the exponential function solved using log transformed density data.
be accurately measured, and an adequate range in size classes must be represented so that the descending limb of the size-frequency distribution approximates the exponential decline of larval abundance with age. In September 1986, $Z$ was estimated to be $0.521(\mathrm{SE}=0.207)$, which equals a daily loss of $40.6 \%$ per day ( $[1-\exp 0.521]$ $\times 100$ ). This rate is similar to $Z$ values of 0.36 and 0.64 that have been reported for another sciaenid, Cynoscion nebulosus, off Florida (Peebles and Tolley 1988). In an earlier calculation of the 1986 red drum spawner biomass (Lyczkowski-Shultz et al. 1988b), the same data were fitted to the exponential function using a logarithmic transformation, resulting in a steeper slope $(Z=0.81)$ and a spawner biomass estimate of 11.7 million pounds (Fig. 3). Examination of residuals later showed the logarithmic transformation of these data to be inappropriate.

Estimates of larval fish mortality can also be affected by two potential biases associated with sampling gear: incomplete recruitment to the sampling gear by the smallest larvae and avoidance of sampling gear by the largest lar-
vae. Using the extensive larval red drum data sets from the 1984 and 1985 seasons, we determined that these sources of bias did not affect the mortality estimate. The shape of length-frequency distributions from over 200 collections indicated that the $3.0-\mathrm{mm}$ size class, the smallest size class used in the mortality estimate, was fully recruited to the sampling gear used in this study. Comparison of daytime and nighttime collections showed no pattern of increasing catches of larvae within the larger size classes on which the 1986 mortality estimate was based (LyczkowskiShultz et al. 1988a), indicating that gear avoidance during daylight was not a factor.

Collections during the September 1986 survey were taken at 1 and 5 m because preliminary data from the 1984 and 1985 spawning seasons suggested that red drum larvae were concentrated in the upper 5 m of the water column (Lyczkowski-Shultz et al. 1988a). This premise is supported with data from 130 discrete-depth samples collected in the study area during five cruises in September of 1984, 1985, and 1987, when depths below 5 m were sampled; mean
density in the upper 5 m was 34.2 larvae per $100 \mathrm{~m}^{3}$, while mean density between 7 and 12 m (the maximum depth sampled) was only 3.7 larvae per $100 \mathrm{~m}^{3}$ (Lyczkow-ski-Shultz et al. 1988a; Lyczkowski-Shultz, unpubl. data). If the low density of larvae found between 7 and 12 m had extended throughout the entire water column (mean water depth was 17 m ), larval abundance and the subsequent biomass estimate could have been underestimated by up to $25 \%$.

Net movement of red drum larvae into or out of the survey area could potentially bias measurement of egg and larval production, ie. spawning activity. However, recent published reports (Chuang et al. 1982; Schroeder et al. 1985; Dinnel 1988), as well as our own observations on surface water circulation patterns over the LA-MS-AL shelf (Lyczkowski-Shultz et al. 1988a), clearly show that there is no consistent and predictable pattern of larval transport into or out of the survey area. Additional evidence showing that inshore displacement of larvae was negligible was the absence of an accumulation of larger and older red drum larvae at a nearshore sampling site located approximately 15 to 19 km NNW of the offshore sampling sites in 1984 and 1985.

Our estimate of red drum spawner biomass in northcentral Gulf waters, 1.25 million pounds, is lower than estimates from two other recent stock assessments. A mark/recapture study (Nichols 1988) estimated the spawning stock between Pensacola, FL and Galveston, TX to be 90 million pounds. Seventeen percent of the stock, 15 million pounds, was estimated to occur between the Mississippi River Delta and Mobile Bay (Scott Nichols, NMFS, Southeast Fisheries Center, Pascagoula, MS, pers. commun. Jan. 1989), an area covering not only the spawning grounds delimited in this study, but also the Chandeleur Sound region where red drum spawning is known to occur. Also, aerial surveys conducted between the Mississippi River Delta and Mobile Bay in the fall of 1987 estimated the biomass of near-surface schools of red drum between the shoreline and the $12-\mathrm{fm}$ contour to be approximately 11.4 million pounds (Lohoefener et al. 1988). Further indication that we underestimated the size of the spawning stock are reports from the late 1970's of individual schools of red drum in the study area that alone were estimated from aerial observations to have exceeded our biomass estimate (Overstreet 1983).

This underestimation of the spawner biomass of red drum in the study area was most likely caused by an underestimate of mean daily egg production. This estimate is not only inaccurate, it is also imprecise considering the large standard error associated with daily egg production. Although factors such as a higher mortality rate among eggs and newly hatched larvae than among older larvae (Houde 1977), or sampling only the upper 5 m , could have contributed to this underestimation of spawner biomass, the variability associated with larval catch data was most
likely the primary cause. Parker (1980) found that the estimate of egg production in the northern anchovy contributed approximately 8 times more to the coefficient of variation of the spawner biomass estimate than all other parameters combined. Sampling effort within the survey area was tripled in September 1989 using oblique tows in an attempt to increase the precision of the estimate of larval red drum abundance and thus provide a more accurate and reliable estimate of red drum spawner biomass.

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## Citations

Chuang, W., W.W. Schroeder, and W.J. Wiseman Jr. 1982. Summer current observations off the Alabama coast. Contrib. Mar. Sci., 25:121-131.
Comyns, B.H., J. Lyczkowski-Shultz, C.F. Rakocinski, and J.P. Steen Jr. 1989. Age and growth of red drum larvae in the northcentral Gulf of Mexico. Trans. Am. Fish. Soc. 118(2):159-167.
Dinnel, S.P.
1988. Circulation and sediment dispersal on the Louisiana--Missis-sippi-Alabama continental shelf. Ph.D. diss., Louisiana State Univ., Shreveport, 173 p.
Dixon, W.J., and M.B. Brown, eds. 1979. Biomedical computer programs, P-series. Univ. California Press, Berkeley.
Fitzhugh, G.R., T.G. Snider III, and B.A. Thompson. 1988. Measurement of ovarian development in red drum from off
shore stocks. Contrib. Mar. Sci., Supp. to Vol. 30:79-86.
Holt, J., A.G. Johnson, C.R. Arnold, W.A. Fable, and T.D. Williams. 1981. Description of eggs and larvae of laboratory reared red drum, Sciaenops ocellata. Copeia 1981:751-756.
Holt, S.A., C.L. Kitting, and C.R. Arnold.
1985. Diel periodicity of spawning in sciaenids. Mar. Ecol. Prog. Ser. 27:1-7.
Holt, S.A., G.J. Holt, and L. Young-Abel.
1988. A procedure for identifying sciaenid eggs. Contrib. Mar. Sci. 29 (suppl).
Houde, E.D.
1977. Abundance and potential yield of the round herring, Etrumeus teres, and aspects of its early life history in the eastern Gulf of Mexico. Fish. Bull., U.S. 75:61-89.
Hunter J.R., and B.J. Macewicz.
1985. Measurement of spawning frequency in multiple spawning fishes. In An egg production method for estimating biomass of pelagic fish: Application to the northern anchovy, Engraulis mor$\operatorname{dax}$ (R. Lasker, ed.), p. 79-94. NOAA Tech. Rep. NMFS SSRF 36.

Hunter, J.R., N.C.H. Lo, and J.H. Leong.
1985. Batch fecundity in multiple spawning fishes. In An egg production method for estimating biomass of pelagic fish: Application to the northern anchovy, Engraulis mordax (R. Lasker, ed), p. 66-77. NOAA Tech. Rep. NMFS SSRF 36.

Lohoefener, R., C. Roden, W. Hoggard, K. Mullin, and C. Rogers.
1988. Distribution, relative abundance, and behavior of nearsurface schools of large red drum (Sciaenops ocellatus) in the rorthcentral Gulf of Mexico. Technical Report, National Marine Fisheries Service, Southeast Fisheries Center, Mississippi Laboratories, Pascagoula, MS, 61 p.
Lyczkowski-Shultz, J.
1987. Fisheries independent data on abundance and distribution of Spanish and king mackerel larvae in the northcentral Gulf of Mexico (August-November, 1983-1986). Technical Repor: submitted to the National Marine Fisheries Service, St. Petersburg, FL. 19 p. +5 tables, 5 figures.
Lyczkowski-Shultz, J., J.P. Steen Jr., and B.H. Comyns.
1988a. Early life history of red drum (Sciaenops ocellatus) in the northeentral Gulf of Mexico. Technical Report submitted to the Mississippi-Alabama Sea Grant Consortium, Ocean Springs, MS. 148 p. +24 tables, 50 figures.
Lyczkowski-Shultz, J., B.H. Comyns, and R.I. Shulman.
1988b. Red drum spawning in east Louisiana, Mississippi, and Alabama waters, 1984-1986. Technical report submitted to the National Marine Fisheries Service, Southeast Regional Office, St. Petersburg, FL. 19 p. +7 figs., 6 tabs.
McGowan, M.F., and W.J. Richards.
1986. Distribution and abundance of bluefin tuna (Thunnus thyn$n u s$ ) larvae in the Gulf of Mexico in 1982 and 1983 with estimates of the biomass and population size of the spawning stock for $\cdot 977$, 1978, and 1981-1982. Collective Volume of Scientific Papers, Int. Comm. Cons. Atlantic Tunas, Vol. 24(1986):182-186.
Moser, H.G., W.J. Richards, D.M. Cohen, M.P. Fahay, A.W.
Kendall Jr., and S.L. Richardson. (eds.)
1984. Ontogeny and systematics of fishes - Ahlstrom symposium. Am. Soc. Ichthyol. Herpetol., Spec. Pub. 1, 759 p.
Nichols, S.
1988. An estimate of the size of the red drum spawning stock using
mark/recapture. Tech. Rep.. National Marine Fisheries Service, Southeast Fisheries Center, Mississippi Laboratories, Pascagoula, MS, 24 p.
Overstreet, R.M.
1983. Aspects of the biology of the red drum, Sciaenops ocellatus, in Mississippi. Gulf Res. Rep., Suppl. 1:45-68.
Parker, K.
1980. A direct method for estimating northern anchovy, Engraulis mordax, spawning biomass. Fish. Bull., U.S. 78:541-544.
Peebles, E.B., and S.G. Tolley.
1988. Distribution, growth and mortality of larval spotted seatrout, Cynoscion nebulosus: a comparison between two adjacent estuarine areas of southwest Florida. Bull. Mar. Sci. 42(3):397-410.
Peters, K.M., and R.H. McMichael Jr.
1987. Early life history of the red drum, Sciaenops ocellatus, (Pisces: Sciaenidae), in Tampa Bay, Florida. Estuaries 10:92-107.
Picquelle, S.J., and R.P. Hewitt.
1983. The northern anchovy spawning biomass for the 1982-83 California fishing season. CalCOFI Rep., Vol. XXIV:16-28.
Poss, S.G.
1988. Identification of red drum fishery stock and establishment of a multivariate model for growth and body condition. Technical report submitted to the National Marine Fisheries Service, Southeast Regional Office, St. Petersburg, FL.
Ramsey, P.R., and J.M. Wakeman.
1987. Population structure of Sciaenops ocellatus and Cynoscion nebulosus (Pisces: Sciaenidae): biochemical variation, genetic subdivision and dispersal. Copeia 1987(3):682-695.
Ricker, W.E.
1975. Computation and interpretation of biological statistics of fish populations. Fish. Res. Board Can. Bull. 191, 382 p.
Schroeder, W.W., O.K. Huh, L.J. Rouse Jr., and W.J. Wiseman Jr. 1985. Satellite observations of the circulation east of the Mississippi Delta: cold-air outbreak conditions. Remote Sensing of Environ. 18:49-58.
Sette, O.E., and E.H. Ahlstrom.
1948. Estimations of abundance of the eggs of the Pacific pilchard (Sardinops caerulea) off southern California during 1940 and 1941. J. Mar. Res. 7:511-542.

Smith, P.E.
1973. The mortality and dispersal of sardine eggs and larvae. Rapp. P.-v. Réun. Cons. Int. Explor. Mer 164:282-292.
Smith, P.E., and S.L. Richardson.
1977. Standard techniques for pelagic fish egg and larva surveys. FAO Fish. Tech. Pap. 175, 100 p.
Smith, P.E., and R.P. Hewitt.
1985. Sea survey design and analysis for an egg production method of northern anchovy biomass assessment. In An egg production method for estimating spawning biomass of pelagic fish: application to the northern anchovy, Engraulis mordax (R. Laster, ed.), p. 17-26. NOAA Tech. Rep. NMFS 36.

Wakeman, J.M., and P.R. Ramsey.
1988. Population structure and genetic variation in red drum. Contrib. Mar. Sci., Suppl. to Vol. 30:49-56.
Wilson, C.A., D.W. Beckman, D.L. Nieland and A.L. Stanley. 1988. Age, growth, and the reproductive biology of schooling red drum from the northern Gulf of Mexico. Technical Report LSU-CFI-88-18 submitted to MARFIN and the Louisiana Dept. of Wildlife and Fisheries, $40 \mathrm{p} .+10$ figures.

# Larval Distribution and Abundance of the Scombridae in Campeche Sound, with Emphasis on the Frigate Tunas (Auxis spp.) 

AURORA RAMÍREZ-ESTÉVEZ<br>Centro Regional de Investigación Pesquera<br>Apartado Postal 580<br>Cancun, Q. Roo, México, C.P. 77500<br>MARGARITA ORNELAS-ROA<br>Centro de Investigaciones y Estudios Avanzados del Instituto Politécnico Nacional<br>Apartado Postal 73<br>Mérida, Yucatán, México, C.P. 97310


#### Abstract

Ichthyoplankton samples were obtained using double oblique tows of a paired $61-\mathrm{cm}$ bongo net containing filter meshes of 0.505 and 0.333 mm during the COSMA 16-72 cruise undertaken 8-14 August, 1972 aboard the RV Virgilio Uribe. Sample depths varied between 7 and 167 m . Temperature and salinity at a depth of 5 m averaged $27.7^{\circ} \mathrm{C}$ and 36.9 ppt , respectively, at the sampling stations. Eight scombrid species or species complexes were represented in the following percentages per $10 \mathrm{~m}^{2}$ of sea surface: frigate tunas (Auxis spp.) $77 \%$; little tunny (Euthynnus alletteratus) $7 \%$; bigeye tuna (Thunnus obesus) $2 \%$; blackfin tuna (T. atlanticus) $1 \%$; king mackerel (Scomberomorus cavalla) 1\%; Spanish mackerel (S. maculatus) $1 \%$; yellowfin tuna ( $T$. albacares) $<1 \%$; and skipjack tuna (Katsuwonus pelamis) $<1 \%$. Thunnus spp. identifiable only to genus comprised $11 \%$ of the larvae. The frigate tunas were the most abundant group, with a total of 4473 larvae per $10 \mathrm{~m}^{2}$ of sea surface. Based on larval abundance of the smallest size class represented, the reproductive biomass of the frigate tuna complex was estimated to be 163868 metric tons. The daily mortality coefficient for the frigate tuna larvae was estimated to be 0.346 . The results showed that the frigate tunas are a potential fishery resource in the area.


## Introduction

Much interest has been shown in ichthyoplankton research for the detection and evaluation of potentially important fishery resources. Studies of eggs and larvae provide a means of establishing breeding times and areas of the adults. They also provide, within reliable limits, estimates of adult biomass and natural mortality during the early life phases.

The tunas are targets of one of the most economically important fisheries in the world. Large schools of bluefin, yellowfin, blackfin, and skipjack tunas have been reported in the Gulf of Mexico (Bullis 1955; Carranza 1956; Montolio and Juárez 1976; Richards et al. 1981). However, from 1964 to 1975 tuna harvests in the Mexican Exclusive

Economic Zone of the Gulf of Mexico totaled only $50 \%$ of the average annual catch by the Mexican tuna fleet in the Pacific Ocean (Secretaría de Pesca 1987).

This study provides information on the distribution and abundance of the scombrid species in Campeche Sound, and establishes, based on an estimate of the reproductive biomass, that the frigate tunas are a resource with potential for commercial exploitation in the region.

## Materials and Methods

Ichthyoplankton samples were obtained during the COSMA 16-72 cruise on the RV Virgilio Uribe on 8-14 August, 1972. The cruise was part of the international


Figure 1
Map of the Campeche Sound showing the 80 sampling stations used during the COSMA 16-72 cruise. Depth contours are in fathoms.
agreement on Cooperative Research in the Caribbean and Adjacent Regions (CICAR). A total of 80 stations were located between lat. $18^{\circ} 40^{\prime} 00^{\prime \prime}-20^{\circ} 53^{\prime} 12^{\prime \prime} \mathrm{N}$ and long. $90^{\circ} 50^{\prime} 06^{\prime \prime}-92^{\circ} 58^{\prime} 00^{\prime \prime} W$ (Fig. 1).

Samples were collected using oblique tows according to methods described by Kramer et al. (1972) and Smith and Richardson (1977). A bongo net with mouth diameter of 0.61 cm and filter meshes of 0.505 mm and 0.333 mm was towed to a maximum depth of 200 m , or to the depth dictated by the maximum depth of the station. The net was towed at an average velocity of $1.25 \mathrm{~m} \cdot \mathrm{~s}^{-1}$ ( 2.5 knots ), and at deployment and recovery rates of $50 \mathrm{~m} \cdot \mathrm{~min}^{-1}$ and $20 \mathrm{~m} \cdot \mathrm{~min}^{-1}$, respectively. Wire angles were determined for each 10 m of wire recovered throughout the tow. Calibrated flowmeters were used in the mouth of each net.

The plankton samples were fixed in a $10 \%$ formalinseawater solution buffered with saturated sodium borate.

The scombrid larvae were separated from the total sample and identified to the lowest taxon possible based on pigmentation patterns, primarily on the upper and lower parts of the caudal peduncle, the fore- and mid-brain, and the extreme upper and lower area of the snout. The height and number of spines on the first dorsal fin, and the number of myomeres and vertebrae were taken into account (Jones 1959; Matsumoto 1959, 1962; Ueyanagi 1963; Juárez 1972, 1976; Matsumoto et al. 1972; Nishikawa and Rimmer 1987). The clearing and staining technique of Hollister (1934) was used when necessary, especially for separation of bigeye tuna, Thunnus obesus, blackfin tuna, T. atlanticus, and yellowfin tuna, T. albacares. The frigate tunas (Auxis thazard and $A$. rochei) are difficult to distinguish as larvae (Uchida 1981), so identification was made only to the genus level. Standard length (SL) was measured to the nearest 0.1 mm with a stereoscopic microscope equipped with an ocular micrometer.

The absolute abundance of larvae at each station was standarized relative to $10 \mathrm{~m}^{2}$ of sea surface according to Kramer et al.'s (1972) formula:

$$
\begin{equation*}
N_{j}=\frac{C_{j} Z_{j}}{V_{j}} 10 \tag{1}
\end{equation*}
$$

where $N_{j}$ is the number of larvae per $10 \mathrm{~m}^{2}$ of sea surface at station $j, C_{j}$ is the catch of larvae at station $j, Z_{j}$ is the maximum tow depth in meters at station $j$, and $V_{j}$ is the volume of water filtered in cubic meters at station $j$. Chisquare analysis was employed to test for differences in the standarized catches of each species in daytime versus nighttime hauls.

## Larval Abundance

The standardized abundance estimates based on the stations positive for frigate tunas were plotted as contours over the sampling area. The area of each contour was measured with a planimeter and divided among the positive stations to obtain the area represented by each station (Sette and Ahlstrom 1948). The total number of larvae $\left(P_{j}\right)$ in the area represented by station $j$ was calculated by length class as

$$
\begin{equation*}
P_{j}=\frac{C_{j} Z_{j}}{V_{j}} A_{j} \tag{2}
\end{equation*}
$$

where $A_{j}$ is the area in square meters represented by station $j$, and all other parameters are as in Equation 1.

The total number of larvae $\left(P_{i}\right)$ by length class over the entire area represented by the positive stations was calculated as

$$
\begin{equation*}
P_{i}=\sum_{j=1}^{J} P_{j} \tag{3}
\end{equation*}
$$

where $J$ is the number of stations positive for frigate tunas during the cruise (Richards et al. 1981).

An estimate of the total larval production $\left(P_{a}\right)$ of frigate tuna of the smallest length class, $2.25-2.75 \mathrm{~mm}$ (the $2.5-\mathrm{mm}$ class), during the spawning season was obtained by

$$
\begin{equation*}
P_{a}=\frac{P_{i} D}{d} \tag{4}
\end{equation*}
$$

where $D$ is the number of days in the spawning season, and $d$ is the residence time in days of the larvae in the $2.5-\mathrm{mm}$ length class (Richards et al. 1981). The duration of the reproductive period was assumed to be 180 days (April-September) based on larval occurrence data in the
western Atlantic, Gulf of Mexico, and adjacent regions reported by Klawe and Shimada (1959), Tibbo and Beckett (1972), Juárez (1976), and Ramírez and Ornelas (in press). Residence time, $d$, was calculated using a modification of a regression, $L=2.66 e^{0.06 T}$ where $L$ is length in millimeters and $T$ is age in days, fitted to size at age data from the rearing studies of little tunny (Euthynnus alletteratus) larvae by Houde and Richards (1969). Since the smallest frigate tuna larvae collected in our study were about 2.00 mm , we assumed that the hatch length and the equation's intercept equal 2.00 and that the exponent is the same as for little tunny. The modified growth equation converted to predict age from length was

$$
\begin{equation*}
\mathrm{T}=\frac{\ln \mathrm{L}-\ln 2.0}{0.06} \tag{5}
\end{equation*}
$$

The $P_{a}$ estimate was adjusted to age $0\left(P_{a d j}\right)$ using our estimate of the daily mortality rate (below) by substituting $P_{a}$ for $P$ in Equation 7 and solving for $P_{o}$.

## Reproductive Biomass

The reproductive biomass was determined based on the relative fecundity, sex ratio, and number of larvae produced during the spawning season (Saville 1964; Ahlstrom 1968). The following equation relates these variables:

$$
\begin{equation*}
B_{a}=\frac{P_{a d j}}{F_{r} K}, \tag{6}
\end{equation*}
$$

where $B_{a}$ is the biomass of reproductive adults in grams, $P_{a d j}$ is the $P_{a}$ estimate adjusted to age $0, F_{r}$ is the average relative fecundity (eggs produced per gram of adult female), and $K$ is the proportion of adults that are females. The average relative fecundity was estimated based on Simmons and McDade's (1973) equation for female Auxis sp. of 36.0 to 45.5 cm fork length, $F=-127.262+0.542 F L$, where $F$ is the number of eggs $(\times 1000)$ in the most advanced mode, and $F L$ is the fork length in millimeters. Body weights of adult frigate tunas were obtained using the length-weight relationship of Ishida (1971) for Auxis rochei captured in Japanese waters, $W=6.05 \times 10^{-3} \mathrm{FL}^{3.3}$, where $W$ is the body weight in grams and $F L$ is the fork length in centimeters. There are no similar data for frigate tuna in the Atlantic. Thus, female frigate tuna averaging $40.2 \mathrm{~cm} F L(W=1190 \mathrm{~g})$ in the Campeche Sound were assumed to have a relative fecundity of $76 \mathrm{eggs} \cdot \mathrm{g}^{-1}$. Sex ratios have not been determined for the frigate tunas in the western Atlantic. Therefore, a 1:1 male to female relationship ( $K=0.5$ ) was assumed based on studies in other areas, although significant deviations from this value have been reported (Uchida 1981).

## Larval Mortality Rate

The daily mortality rate ( $M$ ) for frigate tuna larvae was calculated using an exponential function

$$
\begin{equation*}
P=P_{0} e^{-M T} \tag{7}
\end{equation*}
$$

fitted to the number of larvae caught $(P)$ by length class summed over the positive stations versus the age $(T)$ corresponding to the midpoint of each length class estimated using Equation 5. $P_{o}$ is the absolute abundance at age 0.

## Results

## Larval Distribution and Abundance

The 80 stations sampled during the COSMA 16-72 cruise were located on the continental shelf (Fig. 1). Station locations, depths of tows, water volumes filtered, and environmental data are presented in Table 1. Capture data for the Scombridae are presented in Table 2. The capture percentages indicate that the most abundant categories were the frigate tunas, Thunnus spp. not identifiable to

Table 1
Station locations, sampling data, and environmental data for the COSMA 16-72 cruise. Only stations positive for Scombridae are included.

| Station | Position |  | Date | Hour | Volume of water strained ( $\mathrm{m}^{3}$ ) | Depth of tow (m) | Physical and chemical date |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Dep |  |  |  | Tem | Salinity |
|  | Lat. N. | Long. W. |  |  |  |  | (m) | $\left({ }^{\circ} \mathrm{C}\right)$ | (ppt) |
| 3 | $18^{\circ} 55^{\prime} 58^{\prime \prime}$ | $92^{\circ} 13^{\prime} 24^{\prime \prime}$ |  | 09/08/72 | 14:00 | 152.7 | 20.0 | 5.0 | 28.08 | 37.12 |
|  |  |  |  |  |  |  | 10.0 | 27.72 | 37.09 |
| 4 | $18^{\circ} 51^{\prime} 36^{\prime \prime}$ | $92^{\circ} 22^{\prime} 30^{\prime \prime}$ | 09/08/72 | 11:15 | 190.4 | 18.0 | 5.0 | 27.84 | 36.99 |
|  |  |  |  |  |  |  | 10.0 | 27.62 | 36.01 |
| 8 | $18^{\circ} 52^{\prime} 48^{\prime \prime}$ | $92^{\circ} 54^{\prime} 48^{\prime \prime}$ | 08/08/72 | 13:55 | 207.5 | 44.0 | 5.0 | 29.20 | 34.70 |
|  |  |  |  |  |  |  | 10.0 | 27.52 | 37.06 |
|  |  |  |  |  |  |  | 20.0 | 24.77 | 36.82 |
|  |  |  |  |  |  |  | $40.0$ | $23.40$ | $36.84$ |
| 9 | $18^{\circ} 57^{\prime} 00^{\prime \prime}$ | $92^{\circ} 45^{\prime} 48^{\prime \prime}$ | 08/08/72 | 20:23 | 170.7 | 50.4 | 5.0 | 27.62 | 36.67 |
|  |  |  |  |  |  |  | 10.0 | 27.50 | 37.03 |
|  |  |  |  |  |  |  | 20.0 | 27.60 | 37.10 |
|  |  |  |  |  |  |  | 40.0 | 27.78 | 36.86 |
| 10 | $18^{\circ} 58^{\prime} 12^{\prime \prime}$ | $92^{\circ} 36^{\prime} 00^{\prime \prime}$ | 09/08/72 | 00:40 | 136.3 | 25.2 | 5.0 | 27.72 | 37.05 |
|  |  |  |  |  |  |  | 10.0 | 27.61 | 37.07 |
|  |  |  |  |  |  |  | 20.0 | 27.60 | 37.05 |
| 12 | $19^{\circ} 05^{\prime} 00^{\prime \prime}$ | $92^{\circ} 17^{\prime} 00^{\prime \prime}$ | 09/08/72 | 15:20 | 149.9 | 19.8 | 5.0 | 28.03 | 37.05 |
|  |  |  |  |  |  |  | $10.0$ | $28.81$ | $37.05$ |
| 13 | $19^{\circ} 05^{\prime} 00^{\prime \prime}$ | $92^{\circ} 08^{\prime} 00^{\prime \prime}$ | 10/08/72 | 15:18 | 122.0 | 22.0 | 5.0 | 27.92 | 36.00 |
|  |  |  |  |  |  |  | 10.0 | 27.92 | 36.92 |
| 14 | $19^{\circ} 15^{\prime} 18^{\prime \prime}$ | $92^{\circ} 12^{\prime} 12^{\prime \prime}$ | 10/08/72 | 03:59 | 133.3 | 21.6 | 5.0 | 27.92 | 36.92 |
|  |  |  |  |  |  |  | 10.0 | 27.92 | 36.92 |
| 15 | $19^{\circ} 29^{\prime} 12^{\prime \prime}$ | $92^{\circ} 07^{\prime} 30^{\prime \prime}$ | 10/08/72 | 15:07 | 141.5 | 65.0 | 5.0 | - | 37.16 |
|  |  |  |  |  |  |  | 10.0 | 28.50 | 37.16 |
|  |  |  |  |  |  |  | 20.0 | 28.95 | 36.87 |
|  |  |  |  |  |  |  | 40.0 | 28.01 | 37.13 |
| 16 | $19^{\circ} 20^{\prime} 12^{\prime \prime}$ | $92^{\circ} 20^{\prime} 12^{\prime \prime}$ | 10/08/72 | 13:50 | 253.9 | 36.9 | 5.0 | 28.70 | 37.02 |
|  |  |  |  |  |  |  | 10.0 | 28.50 | 37.06 |
|  |  |  |  |  |  |  | $20.0$ | $28.30$ | $37.05$ |
| 17 | $19^{\circ} 13^{\prime} 00^{\prime \prime}$ | $91^{\circ} 59^{\prime} 18^{\prime \prime}$ | 10/08/72 | 12:30 | 201.6 | 25.0 | 5.0 | 28.32 | 36.08 |
|  |  |  |  |  |  |  | 10.0 | 28.33 | 36.98 |
|  |  |  |  |  |  |  | 20.0 | 28.25 | 37.05 |
| 18 | $19^{\circ} 03^{\prime} 12^{\prime \prime}$ | $91^{\circ} 54^{\prime} 00^{\prime \prime}$ | 10/08/72 | 11:03 | 216.0 | 21.0 | 5.0 | 28.20 | 37.08 |
|  |  |  |  |  |  |  | 10.0 | 27.85 | 37.08 |
| 24 | $19^{\circ} 14^{\prime} 42^{\prime \prime}$ | $91^{\circ} 48^{\prime} 30^{\prime \prime}$ | 11/08/72 | 02:43 | 159.2 | 22.0 | 5.0 | $28.33$ | $37.19$ |
|  |  |  |  |  |  |  | $10.0$ | $28.33$ | $36.98$ |
|  |  |  |  |  |  |  | 20.0 | 28.23 | 36.97 |
| 25 | $19^{\circ} 20^{\prime} 42^{\prime \prime}$ | $91^{\circ} 51^{\prime} 54^{\prime \prime}$ | 11/08/72 | 01:43 | 122.5 | 23.6 | 5.0 | 28.31 | 37.05 |
|  |  |  |  |  |  |  | 10.0 | 28.21 | 37.02 |
|  |  |  |  |  |  |  | 20.0 | 28.22 | 36.94 |


| Station | Table 1 (continued) |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Position |  | Date | Hour | Volume of water strained ( $\mathrm{m}^{3}$ ) | Depth of tow (m) | Physical and chemical date |  |  |
|  |  |  | Depth (m) |  |  |  | Temp. $\left({ }^{\circ} \mathrm{C}\right)$ | Salinity (ppt) |
|  | Lat. N. | Long. W. |  |  |  |  |  |  |
| 27 | $19^{\circ} 40^{\prime} 30^{\prime \prime}$ | $92^{\circ} 02^{\prime} 06^{\prime \prime}$ | 11/08/72 | 22:45 | 127.6 | 63.0 | 5.0 | 28.53 | 37.01 |
|  |  |  |  |  |  |  | 10.0 | 28.54 | 37.06 |
|  |  |  |  |  |  |  | 20.0 | 28.53 | 36.97 |
|  |  |  |  |  |  |  | 40.0 | 28.23 | 36.87 |
| 29 | $20^{\circ} 10^{\prime} 30^{\prime \prime}$ | $92^{\circ} 05^{\prime} 30^{\prime \prime}$ | 12/08/72 | 06:58 | 144.8 | 70.0 | 6.0 | 28.42 | 37.12 |
|  |  |  |  |  |  |  | 10.0 | 28.54 | 37.13 |
|  |  |  |  |  |  |  | 14.0 | 28.42 | 37.13 |
|  |  |  |  |  |  |  | 20.0 | 28.42 | 37.13 |
|  |  |  |  |  |  |  | 40.0 | 28.11 | 37.10 |
| 30 | $20^{\circ} 02^{\prime} 00^{\prime \prime}$ | $92^{\circ} 02^{\prime} 06^{\prime \prime}$ | 12/08/72 | 05:45 | 138.2 | 62.0 | 6.0 | 28.42 | 37.14 |
|  |  |  |  |  |  |  | 10.0 | 28.52 | 37.14 |
|  |  |  |  |  |  |  | 14.0 | 28.42 | 37.14 |
|  |  |  |  |  |  |  | 20.0 | 28.42 | 37.12 |
|  |  |  |  |  |  |  | 40.0 | 28.21 | 36.95 |
| 31 | $20^{\circ} 53^{\prime} 12^{\prime \prime}$ | $91^{\circ} 58^{\prime} 06^{\prime \prime}$ | 12/08/72 | 02:20 | 142.1 | 50.0 | 6.0 | 28.45 | 37.08 |
|  |  |  |  |  |  |  | 10.0 | 28.55 | 37.09 |
|  |  |  |  |  |  |  | 14.0 | 28.55 | 37.10 |
|  |  |  |  |  |  |  | 20.0 | 28.42 | 37.11 |
|  |  |  |  |  |  |  | 40.0 | 25.31 | 36.93 |
| 32 | $19^{\circ} 44^{\prime} 00^{\prime \prime}$ | $91^{\circ} 53^{\prime} 30^{\prime \prime}$ | 11/08/72 | 23:27 | 154.2 | 54.2 | 6.0 | 28.75 | $36.92$ |
|  |  |  |  |  |  |  | $10.0$ | $28.42$ | $37.10$ |
|  |  |  |  |  |  |  | 14.0 | 28.02 | 37.03 |
|  |  |  |  |  |  |  | 20.0 | 28.42 | 37.01 |
|  |  |  |  |  |  |  | 40.0 | 28.03 | 36.99 |
| 33 | $19^{\circ} 37^{\prime} 00^{\prime \prime}$ | $91^{\circ} 50^{\prime} 12^{\prime \prime}$ | 11/08/72 | 20:43 | 154.5 | 45.0 | $6.0$ | $28.43$ | $36.93$ |
|  |  |  |  |  |  |  | $10.0$ | $28.43$ | $36.02$ |
|  |  |  |  |  |  |  | 14.0 | 28.43 | $36.99$ |
|  |  |  |  |  |  |  | 20.0 | 28.43 | 36.99 |
|  |  |  |  |  |  |  | 40.0 | 28.87 | 36.88 |
| 35 | $19^{\circ} 18^{\prime} 48^{\prime \prime}$ | $91^{\circ} 41^{\prime} 00^{\prime \prime}$ | 11/08/72 | 15:29 | 158.7 | 21.6 | $6.0$ | $28.50$ | $36.97$ |
|  |  |  |  |  |  |  | $10.0$ | $28.50$ | $36.97$ |
|  |  |  |  |  |  |  | 14.0 | 28.30 | 36.97 |
|  |  |  |  |  |  |  | 20.0 | 28.00 | 36.96 |
| 36 | $19^{\circ} 10^{\prime} 06^{\prime \prime}$ | $91^{\circ} 36^{\prime} 48^{\prime \prime}$ | 11/08/72 | 13:07 | 150.6 | 18.0 | 5.0 | 28.51 | 36.98 |
|  |  |  |  |  |  |  | 10.0 | 28.51 | 36.98 |
|  |  |  |  |  |  |  | 11.0 | 28.30 | 36.97 |
|  |  |  |  |  |  |  | 16.5 | 28.21 | 36.95 |
| 41 | $19^{\circ} 16^{\prime} 30^{\prime \prime}$ | $91^{\circ} 29^{\prime} 00^{\prime \prime}$ | 11/08/72 | 14:18 | 187.5 | 16.2 | $5.0$ | 27.32 | $36.96$ |
|  |  |  |  |  |  |  | $10.0$ | $28.50$ | $36.96$ |
| 42 | $19^{\circ} 25^{\prime} 12^{\prime \prime}$ | $91^{\circ} 33^{\prime} 06^{\prime \prime}$ | 11/08/72 | 16:35 | 154.2 | 22.0 | 5.0 | $28.20$ | $36.95$ |
|  |  |  |  |  |  |  | 10.0 | 28.60 | $36.92$ |
|  |  |  |  |  |  |  | 20.0 | 28.31 | 36.90 |
| 43 | $19^{\circ} 33^{\prime} 36^{\prime \prime}$ | $91^{\circ} 37^{\prime} 00^{\prime \prime}$ | 09/08/72 | 19:10 | 229.7 | 30.6 | 5.0 | 28.40 | 36.97 |
|  |  |  |  |  |  |  | 10.0 | 28.45 | 36.97 |
|  |  |  |  |  |  |  | 20.0 | 28.25 | 37.03 |
| 45 | $19^{\circ} 50{ }^{\prime} 48^{\prime \prime}$ | $91^{\circ} 45^{\prime} 24^{\prime \prime}$ | 12/08/72 | 01:15 | 186.6 | 39.0 | 5.0 | 28.43 | 36.99 |
|  |  |  |  |  |  |  | 10.0 | 28.33 | 37.02 |
|  |  |  |  |  |  |  | 20.0 | 28.23 | 37.00 |
| 46 | $19^{\circ} 59^{\prime} 48^{\prime \prime}$ | $91^{\circ} 49^{\prime} 48^{\prime \prime}$ | 12/08/72 | 04:05 | 121.5 | 39.0 | $5.0$ | 28.01 | $36.08$ |
|  |  |  |  |  |  |  | 10.0 | 28.02 | 36.97 |
|  |  |  |  |  |  |  | 20.0 | 28.02 | 36.99 |
| 48 | $20^{\circ} 08^{\prime} 30^{\prime \prime}$ | $91^{\circ} 43^{\prime} 00^{\prime \prime}$ | 12/08/72 | 21:44 | 91.0 | 32.4 | 5.0 | 28.02 | 36.88 |
|  |  |  |  |  |  |  | 10.0 | 28.02 | 36.82 |
|  |  |  |  |  |  |  | 20.0 | 27.62 | 36.83 |
| 51 | $19^{\circ} 41^{\prime} 12^{\prime \prime}$ | $91^{\circ} 31^{\prime} 12^{\prime \prime}$ | 13/08/72 | 01:04 | 124.7 | 25.0 | 5.0 | 28.33 | 36.94 |
|  |  |  |  |  |  |  | 10.0 | 28.33 | - |
|  |  |  |  |  |  |  | 20.0 | 27.93 | 36.99 |


| Station | Table 1 (continued) |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Position |  | Date | Hour | Volume of water strained ( $\mathrm{m}^{3}$ ) | Depth of tow (m) | Physical and chemical date |  |  |
|  |  |  |  |  |  |  |  |  |
|  | Lat. N. | Long. W. |  |  |  |  | (m) | $\left({ }^{\circ} \mathrm{C}\right)$ | (ppt) |
| 52 | $19^{\circ} 37^{\prime} 30^{\prime \prime}$ | $91^{\circ} 26^{\prime} 30^{\prime \prime}$ |  | 13/08/72 | 02:20 | 111.9 | 12.0 | 5.0 | 28.24 | 36.95 |
|  |  |  |  |  |  |  | 10.0 | 28.22 | 36.94 |
| 53 | $19^{\circ} 25^{\prime} 12^{\prime \prime}$ | $91^{\circ} 22^{\prime} 18^{\prime \prime}$ | 13/08/72 | 03:15 | 65.9 | 15.0 | 5.0 | 28.23 | 36.94 |
| 55 | $19^{\circ} 06^{\prime} 48^{\prime \prime}$ | $91^{\circ} 16^{\prime} 12^{\prime \prime}$ | 13/08/72 | 05:15 | 78.8 | 10.8 | 5.0 | 28.23 | 37.33 |
| 57 | $19^{\circ} 22^{\prime} 12^{\prime \prime}$ | $91^{\circ} 11^{\prime} 06^{\prime \prime}$ | 14/08/72 | 12:30 | 148.0 | 14.0 | 5.0 | 28.71 | 36.83 |
|  |  |  |  |  |  |  | 10.0 | 28.81 | 36.86 |
| 58 | $19^{\circ} 30^{\prime} 42^{\prime \prime}$ | $91^{\circ} 14^{\prime} 12^{\prime \prime}$ | 14/08/72 | 10:59 | 148.0 | 14.0 | 5.0 | - | 36.99 |
|  |  |  |  |  |  |  | 10.0 | - | 36.93 |
| 59 | $19^{\circ} 37^{\prime} 18^{\prime \prime}$ | $91^{\circ} 18^{\prime} 54^{\prime \prime}$ | 14/08/72 | 10:09 | 183.5 | 25.0 | 5.0 | 28.21 | 36.87 |
|  |  |  |  |  |  |  | 10.0 | 28.21 | 36.87 |
|  |  |  |  |  |  |  | 20.0 | 25.85 | 36.90 |
| 60 | $19^{\circ} 47^{\prime} 30^{\prime \prime}$ | $91^{\circ} 22^{\prime} 30^{\prime \prime}$ | 14/08/72 | 06:24 | 123.2 | 23.4 | 5.0 | 28.13 | 36.91 |
|  |  |  |  |  |  |  | 10.0 | 28.02 | 36.95 |
|  |  |  |  |  |  |  | 20.0 | 27.31 | - |
| 61 | $19^{\circ} 55^{\prime} 30^{\prime \prime}$ | $91^{\circ} 26^{\prime} 00^{\prime \prime}$ | 14/08/72 | 04:50 | 164.1 | 30.0 | 5.0 | 28.02 | 36.90 |
|  |  |  |  |  |  |  | 10.0 | 28.02 | 36.95 |
|  |  |  |  |  |  |  | 20.0 | 26.51 | 36.90 |
| 72 | $19^{\circ} 45^{\prime} 00^{\prime \prime}$ | $91^{\circ} 59^{\prime} 18^{\prime \prime}$ | 1.3/08/72 | 12:09 | 192.4 | 14.8 | 5.0 | 28.61 | 37.00 |
|  |  |  |  |  |  |  | 10.0 | 27.91 | 36.98 |
| 80 | $18^{\circ} 59^{\prime} 30^{\prime \prime}$ | $92^{\circ} 58^{\prime} 00^{\prime \prime}$ | 08/08/72 | 15:39 | 248.1 | 112.0 | 5.0 | $27.74$ | $37.03$ |
|  |  |  |  |  |  |  | $10.0$ | $28.03$ | $36.01$ |
|  |  |  |  |  |  |  | 20.0 | 27.43 | 36.97 |
|  |  |  |  |  |  |  | 40.0 | 23.70 | 36.86 |
|  |  |  |  |  |  |  | 80.0 | 21.92 | 36.89 |
| 81 | $19^{\circ} 14^{\prime} 36^{\prime \prime}$ | $92^{\circ} 53^{\prime} 06^{\prime \prime}$ | 08/08/72 | 17:50 | 261.5 | 138.0 | 5.0 | $28.00$ | $36.75$ |
|  |  |  |  |  |  |  | $10.0$ | $27.40$ | $36.72$ |
|  |  |  |  |  |  |  | 20.0 | 27.51 | 36.75 |
|  |  |  |  |  |  |  | 40.0 | 27.51 | 37.04 |
|  |  |  |  |  |  |  | 80.0 | 22.29 | 36.87 |
| 82 | $19^{\circ} 04^{\prime} 48^{\prime \prime}$ | $92^{\circ} 50^{\prime} 12^{\prime \prime}$ | 08/08/72 | 19:10 | 243.8 | 91.0 | 5.0 | 28.42 | 33.57 |
|  |  |  |  |  |  |  | 10.0 | 27.31 | 36.95 |
|  |  |  |  |  |  |  | 20.0 | $27.91$ | $37.07$ |
|  |  |  |  |  |  |  | $40.0$ | $23.27$ | $36.87$ |
|  |  |  |  |  |  |  | $80.0$ | $22.38$ | $36.88$ |
| 83 | $19^{\circ} 07^{\prime} 42^{\prime \prime}$ | $92^{\circ} 41^{\prime} 12^{\prime \prime}$ | 11/08/72 | 01:45 | 133.0 | 72.0 | $5.0$ | $28.03$ | $37.07$ |
|  |  |  |  |  |  |  | $10.0$ | $27.91$ | $37.08$ |
|  |  |  |  |  |  |  | 20.0 | 27.81 | 37.07 |
|  |  |  |  |  |  |  | 40.0 | 23.85 | 36.86 |
| 84 | $19^{\circ} 16^{\prime} 30^{\prime \prime}$ | $92^{\circ} 45^{\prime} 00^{\prime \prime}$ | 11/08/72 | 03:06 | 277.7 | 135.0 | 5.0 | 27.51 | 36.99 |
|  |  |  |  |  |  |  | 10.0 | 27.05 | 36.93 |
|  |  |  |  |  |  |  | 20.0 | 27.51 | 36.96 |
|  |  |  |  |  |  |  | 40.0 | 27.50 | 37.02 |
|  |  |  |  |  |  |  | 80.0 | 21.99 | 36.98 |
| 85 | $19^{\circ} 26^{\prime} 42$ | $92^{\circ} 39^{\prime} 12$ | 11/08/72 | 06:47 | 298.5 | 167.0 | 5.0 | 28.43 | 37.09 |
|  |  |  |  |  |  |  | 10.0 |  | $37.09$ |
|  |  |  |  |  |  |  | 20.0 | 28.22 | 37.15 |
|  |  |  |  |  |  |  | 40.0 | 22.58 | 36.86 |
|  |  |  |  |  |  |  | 80.0 | 18.64 | 36.96 |
| 86 | $19^{\circ} 17^{\prime} 00^{\prime \prime}$ | $92^{\circ} 35^{\prime} 00^{\prime \prime}$ | 11/08/72 | 06:47 | 174.7 | 99.0 | 5.0 | 27.90 | 37.13 |
|  |  |  |  |  |  |  | 10.0 | 25.84 | 36.97 |
|  |  |  |  |  |  |  | 20.0 | 26.85 | 37.13 |
|  |  |  |  |  |  |  | $40.0$ | $27.02$ | $37.11$ |
|  |  |  |  |  |  |  | $80.0$ | 28.01 | 36.07 |
| 87 | $19^{\circ} 08^{\prime} 42^{\prime \prime}$ | $92^{\circ} 33^{\prime} 30^{\prime \prime}$ | 11/08/72 | 08:40 | 258.3 | 54.0 | 5.0 | - | 36.99 |
|  |  |  |  |  |  |  | 10.0 | 28.22 | 37.00 |
|  |  |  |  |  |  |  | 20.0 | 28.20 | 37.02 |
|  |  |  |  |  |  |  | 40.0 | 23.10 | 37.06 |


| Station | Table 1 (continued) |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Position |  | Date | Hour | Volume of water strained ( $\mathrm{m}^{3}$ ) | Depth of tow (m) | Physical and chemical date |  |  |
|  |  |  | Depth |  |  |  | Temp. | Salinity |
|  | Lat. N. | Long. W. |  |  |  |  | $(\mathrm{m})$ | $\left({ }^{\circ} \mathrm{C}\right)$ | (ppt) |
| 90 | $19^{\circ} 21^{\prime} 30^{\prime \prime}$ | $92^{\circ} 26^{\prime} 00^{\prime \prime}$ |  | 11/08/72 | 18:04 | 167.1 | 84.0 | 5.0 | 28.50 | 37.03 |
|  |  |  |  |  |  |  | 10.0 | - | 37.05 |
|  |  |  |  |  |  |  | 20.0 | 28.40 | 37.05 |
|  |  |  |  |  |  |  | 40.0 | 28.00 | 37.09 |
|  |  |  |  |  |  |  | 80.0 | 21.91 | - |
| 92 | $19^{\circ} 42^{\prime} 42^{\prime \prime}$ | $92^{\circ} 24^{\prime} 42^{\prime \prime}$ | 10/08/72 | 24:18 | 244.8 | 140.0 | 5.0 | 28.32 | 36.98 |
|  |  |  |  |  |  |  | 10.0 | 27.00 | 37.11 |
|  |  |  |  |  |  |  | 20.0 | 28.32 | 37.14 |
|  |  |  |  |  |  |  | 40.0 | 28.22 | 37.21 |
|  |  |  |  |  |  |  | 80.0 | 22.60 | 37.15 |
| 93 | $19^{\circ} 32^{\prime} 00^{\prime \prime}$ | $92^{\circ} 19^{\prime} 48^{\prime \prime}$ | 09/08/72 | 19:43 | 180.1 | 86.4 | 5.0 | 28.51 | 37.12 |
|  |  |  |  |  |  |  | 10.0 | 30.44 | 37.12 |
|  |  |  |  |  |  |  | 20.0 | 28.41 | 37.15 |
|  |  |  |  |  |  |  | 40.0 | 28.31 | 37.16 |
|  |  |  |  |  |  |  | 80.0 | 21.90 | 36.86 |
| 94 | $19^{\circ} 23^{\prime} 24^{\prime \prime}$ | $92^{\circ} 16^{\prime} 18^{\prime \prime}$ | 10/08/72 | 02:41 | 127.3 | 39.0 | 5.0 | 28.31 | $37.04$ |
|  |  |  |  |  |  |  | 10.0 | 28.21 | 36.86 |
|  |  |  |  |  |  |  | 20.0 | 28.22 | 37.05 |
| 95 | $19^{\circ} 37^{\prime} 00^{\prime \prime}$ | $92^{\circ} 13^{\prime} 00^{\prime \prime}$ | 10/08/72 | 16:53 | 181.5 | 108.0 | 5.0 | 28.50 | 37.14 |
|  |  |  |  |  |  |  | 10.0 | - | 37.13 |
|  |  |  |  |  |  |  | 20.0 | 28.40 | 37.15 |
|  |  |  |  |  |  |  | 40.0 | 28.21 | 37.15 |
|  |  |  |  |  |  |  | 80.0 | 22.60 | 37.15 |
| 96 | $19^{\circ} 51^{\prime} 06^{\prime \prime}$ | $92^{\circ} 29^{\prime} 36^{\prime \prime}$ | 10/08/72 | 18:53 | 173.4 | 90.0 | 5.0 | - | - |
|  |  |  |  |  |  |  | 10.0 | - | - |
|  |  |  |  |  |  |  | 20.0 | 33.29 | 36.95 |
|  |  |  |  |  |  |  | 40.0 | - | - |
|  |  |  |  |  |  |  | 80.0 | 28.2 | 36.97 |

Table 2
Relative abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) of the Scombridae captured during the COSMA 16-72 cruise. Positive stations are stations at which scombrid larvae were collected.

| Species | Number of positive stations | Number per $10 \mathrm{~m}^{2}$ |  |  | Percentage of total |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Total | Daytime | Nighttime |  |
| Frigate tuna | 37 | 4473 | 370 | 4103 | 7 |
| Thunnus spp. | 36 | 651 | 180 | 471 | 11 |
| Little tunny | 21 | 417 | 113 | 304 | 7 |
| Bigeye tuna | 6 | 120 | 51 | 69 | 2 |
| Blackfin tuna | 2 | 58 | - | 58 | 1 |
| King mackerel | 8 | 35 | 16 | 19 | 1 |
| Spanish mackerel | 2 | 31 | 27 | 4 | 1 |
| Yellowfin tuna | 2 | 24 | 24 | - | 1 |
| Skipjack tuna | 1 | 14 | 14 | - | 1 |

species, the little tunny, the bigeye tuna, and the blackfin tuna, respectively.

A total of 4473 frigate tuna larvae were captured per $10 \mathrm{~m}^{2}$ of sea surface, with a mean of 86 over 37 positive sampling stations. Significantly more frigate tuna larvae
were caught during the nighttime than in the daytime ( $P<0.05$ ) (Table 2). The frigate tunas captured ranged from 2.00 to 7.75 mm ; those smaller than 2.25 mm were under represented in the samples, and were eliminated from further analysis. Frigate tunas in the 2.76-3.25 mm

Table 3
Absolute abundance by standard length (mm) of frigate tuna larvae captured during the COSMA 16-72 cruise. (The total for each positive station $=C_{j}[$ Equation 1] and the total for each length class $=P$ [Equation 7].)

| Station | Standard length range (midpoint) |  |  |  |  |  |  |  |  |  |  | Total $\left(C_{j}\right)$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $\begin{gathered} 2.25-2.75 \\ (2.50) \end{gathered}$ | $\begin{gathered} 2.76-3.25 \\ (3.00) \end{gathered}$ | $\begin{gathered} 3.26-3.75 \\ (3.50) \end{gathered}$ | $\begin{gathered} 3.76-4.25 \\ (4.00) \end{gathered}$ | $\begin{gathered} 4.26-4.75 \\ (4.50) \end{gathered}$ | $\begin{gathered} 4.76-5.25 \\ (5.00) \end{gathered}$ | $\begin{gathered} 5.26-5.75 \\ (5.50) \end{gathered}$ | $\begin{gathered} 5.76-6.25 \\ (6.00) \end{gathered}$ | $\begin{gathered} 6.26-6.75 \\ (6.50) \end{gathered}$ | $\begin{gathered} 6.76-7.20 \\ (7.0) \end{gathered}$ | $\begin{gathered} 7.25-7.75 \\ (7.50) \end{gathered}$ |  |
| 4 | 3 | - | - | - | - | - | - | - | - | - | - | 3 |
| 9 | - | - | - | - | - | 1 | - | - | - | - | - | 1 |
| 10 | 6 | 1 | - | - | - | - | - | - | - | - | - | 7 |
| 15 | - | - | - | - | - | - | - | 2 | - | - | - | 2 |
| 16 | - | 17 | 1 | - | - | - | - | - | 1 | - | - | 19 |
| 17 | - | - | - | - | - | - | - | 1 | - | - | - | 1 |
| 18 | - | 1 | - | - | - | - | - | - | - | - | - | 1 |
| 24 | - | - | - | - | - | 1 | - | - | - | - | - | 1 |
| 25 | 5 | 1 | - | - | - | - | - | - | - | - | - | 6 |
| 27 | 5 | 11 | 3 | - | - | - | - | - | - | - | - | 19 |
| 29 | - | - | - | 1 | - | - | - | - | - | - | - | 1 |
| 30 | - | - | - | - | - | - | - | 1 | - | - | - | 1 |
| 31 | 21 | 14 | 5 | 6 | 2 | 3 | 1 | - | - | - | - | 52 |
| 32 | - | - | 4 | 10 | 3 | 2 | - | - | - | - | - | 19 |
| 33 | - | - | - | 2 | 1 | 1 | - | - | - | - | - | 4 |
| 36 | - | 5 | - | - | - | - | - | - | - | - | - | 5 |
| 42 | - | - | - | 1 | 1 | - | - | - | 1 | - | - | 3 |
| 43 | 18 | 26 | 1 | 1 | - | - | - | - | - | - | - | 46 |
| 45 | - | - | - | 1 | 1 | 1 | - | - | - | - | - | 3 |
| 46 | - | - | 1 | - | - | - | - | - | - | - | - | 1 |
| 48 | 1 | 1 | - | - | - | - | - | - | - | - | - | 2 |
| 51 | - | - | - | - | - | 2 | - | 1 | - | - | - | 3 |
| 53 | 4 | - | 1 | - | 1 | 1 | 1 | - | 1 | - | - | 9 |
| 60 | 20 | 6 | 1 | - | - | - | - | - | - | - | - | 27 |
| 61 | 11 | 7 | 1 | - | - | - | - | - | - | - | - | 19 |
| 81 | - | - | - | - | - | 2 | - | - | - | - | - | 2 |
| 82 | - | - | - | - | - | 2 | - | - | - | - | - | 2 |
| 83 | 2 | 6 | 3 | - | - | 4 | 2 | - | - | - | - | 17 |
| 84 | 10 | 13 | 3 | - | - | - | - | - | - | - | - | 26 |
| 85 | - | 5 | 6 | 2.5 | 2 | - | - | - | - | - | 1 | 39 |
| 86 | - | - | - | - | - | - | - | 1 | - | - | - | 1 |
| 87 | 4 | 5 | 1 | 1 | - | - | - | - | - | - | - | 11 |
| 90 | 6 | 6 | 1 | - | - | - | - | - | - | - | - | 13 |
| 92 | - | - | - | 1 | - | - | - | 1 | - | - | - | 2 |
| 93 | - | - | - | 6 | 3 | 7 | 4 | 1 | - | - | - | 21 |
| 94 | 391 | 616 | 28 | 2 | - | - | - | - | - | - | - | 137 |
| 95 | - | - | - | - | - | 1 | - | - | - | - | - | 1 |
| Total | 507 | 741 | 60 | 57 | 14 | 28 | 8 | 8 | 3 | 0 | 1 | 1427 |

length class were the most abundant (Table 3). Sampling Stations 94, 85, and 31 had the greatest standardized densities (Fig. 2), with 3173, 218, and 183, respectively, per $10 \mathrm{~m}^{2}$ of sea surface (Table 4). These stations were located between 70 and 90 km ( 40 and 50 nautical miles [nmi]) from Puerto Frontera and Punta Frontera.

The little tunny was the second most abundant species (Table 2). Its larval distribution pattern indicated that the adults spawned in areas greater than $54 \mathrm{~km}(30 \mathrm{nmi})$ from the coast (Fig. 3). In contrast, some frigate tuna larvae were found somewhat closer to the coastline. Maximum densities were obtained at Stations 93 and 32, with 81 and 88
larvae per $10 \mathrm{~m}^{2}$ of sea surface, respectively. These stations were located 95 and 117 km from Punta Frontera and Ciudad del Carmen. Catches of little tunny larvae were significantly higher $(P<0.05)$ at night than in the daytime (Table 2). Sizes ranged from $3.0-12.5 \mathrm{~mm}$, and the $3.76-4.25 \mathrm{~mm}$ length class was the modal class.

The third most abundant species was the bigeye tuna (Table 2) at Stations 30 and 31 located 144-189 km (80-105 nmi) from Atasta and Campeche (Fig. 4). Standardized densities at these stations averaged 25 and 50 larvae, respectively, per $10 \mathrm{~m}^{2}$ of sea surface. Larval sizes ranged from $3.0-7.0 \mathrm{~mm}$, with most individuals being in the


Figure 2
Distribution and standardized abundance of frigate tuna larvae during the COSMA 16-72 cruise. Depth contours are in fathoms.
$3.76-4.25 \mathrm{~mm}$ class. No significant difference was observed between diurnal and nocturnal capture rates.

Blackfin tuna larvae were captured at only two stations, 92 and 27 (Fig. 4). These stations were located about 115 and 106 km ( 62 and 57 nmi ) from Punta Frontera and Atasta. Numbers captured did not exceed 30 larvae per $10 \mathrm{~m}^{2}$ of sea surface. The larval sizes ranged from $6.0-11.5 \mathrm{~mm}$, with specimens in the $5.76-6.25 \mathrm{~mm}$ class predominating. Capture of this species took place only in nocturnal tows (Table 2).

A total of 35 king mackerel, Scomberomorus cavalla, larvae were caught, with a maximum of 10 larvae per $10 \mathrm{~m}^{2}$ of sea surface at Station 81 located $72 \mathrm{~km}(40 \mathrm{nmi})$ from Puerto Frontera (Fig. 4). Larvae were captured at stations located as close as $36 \mathrm{~km}(20 \mathrm{nmi})$ from the coastline. King mackerel larvae ranged from $2.5-11.0 \mathrm{~mm}$, with the 4.76 5.25 mm length class predominating. No significant difference was found between diurnal and nocturnal catches.

Spanish mackerel, S. maculatus, larvae were found at Stations 82 and 80 located $48-50 \mathrm{~km}$ (27-28 nmi) from Puerto Frontera (Fig. 5). A total of 31 larvae were sampled. Catches were significantly higher in the daytime than at night ( $P<0.05$ ). The maximum density was 27 larvae per $10 \mathrm{~m}^{2}$ of sea surface at Station 80. Sizes ranged from $8.0-15.5 \mathrm{~mm}$, with no length class predominating. Judging by the size of the specimens captured, spawning took place a considerable time before sampling.

Yellowfin tuna larvae measuring $3.0-10.0 \mathrm{~mm}$ were captured at Stations 95 and 30 located at 104 and 144 km (58 and 80 nmi ) from Punta Frontera and Atasta (Fig. 5). Yellowfin larvae were caught only in daytime tows (Table 2).

The skipjack tuna, Katsuwonus pelamis, was the least abundant species in the samples (Table 2). The only positive station was Station 15 located $86 \mathrm{~km}(40 \mathrm{nmi})$ from Atasta (Fig. 5). The larvae captured ranged from 6.0-7.0

Table 4
Relative abundance (number per $10 \mathrm{~m}^{2}$ of sea surface [ $N_{j}$, Equation 1]) by standard length ( mm ) of frigate tuna larvae captured during the COSMA 16-72 cruise. The standard haul factor (SHF) was obtained by the method of Kramer et al. (1972).

Standard length range (midpoint)

| Sta- <br> tion | S.H.F. | $\begin{aligned} & 2.25-2.75 \\ & (2.50) \end{aligned}$ | $\begin{gathered} 2.76-3.25 \\ (3.00) \end{gathered}$ | $\begin{gathered} 3.26-3.75 \\ (3.50) \end{gathered}$ | $\begin{gathered} 3.76-4.25 \\ (4.00) \end{gathered}$ | $\begin{gathered} 4.26-4.75 \\ (4.50) \end{gathered}$ | $\begin{gathered} 4.76-5.25 \\ (5.00) \end{gathered}$ | $\begin{gathered} 5.26-5.75 \\ (5.50) \end{gathered}$ | $\begin{gathered} 5.76-6.25 \\ (6.00) \end{gathered}$ | $\begin{gathered} 6.26-6.75 \\ (6.50) \end{gathered}$ | $\begin{gathered} 6.76-7.20 \\ (7.0) \end{gathered}$ | $\begin{gathered} 7.25-7.75 \\ (7.50) \end{gathered}$ | Total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 4 | 0.94 | 2.82 | - | - | - | - | - | - | - | - | - | - | 2.82 |
| 9 | 2.95 | - | - | - | - | - | 2.95 | - | - | - | - | - | 2.95 |
| 10 | 1.84 | 11.04 | 1.84 | - | - | - | - | - | - | - | - | - | 12.88 |
| 15 | 4.59 | - | - | - | - | - | - | - | 9.18 | - | - | - | 9.18 |
| 16 | 1.41 | - | 23.93 | 1.41 | - | - | - | - | - | 1.41 | - | - | 26.79 |
| 17 | 0.81 | - | - | - | - | - | - | - | 0.81 | - | - | - | 0.81 |
| 18 | 0.97 | - | 0.97 | - | - | - | - | - | - | - | - | - | 0.97 |
| 24 | 1.38 | - | - | - | - | - | 1.38 | - | - | - | - | - | 1.38 |
| 25 | 1.92 | 9.60 | 1.92 | - | - | - | - | - | - | - | - | - | 12.52 |
| 27 | 4.94 | 24.70 | 54.34 | 14.82 | - | - | - | - | - | - | - | - | 93.86 |
| 29 | 4.83 | - | - | - | 4.83 | - | - | - | - | - | - | - | 4.83 |
| 30 | 4.48 | - | - | - | - | - | - | - | 4.48 | - | - | - | 4.48 |
| 31 | 3.51 | 73.71 | 49.14 | 17.55 | 21.06 | 7.02 | 10.53 | 3.51 | - | - | - | - | 182.52 |
| 32 | 3.50 | - | - | 14.00 | 35.00 | 10.50 | 7.00 | - | - | - | - | -- | 66.50 |
| 33 | 2.91 | - | - | - | 5.82 | 2.91 | 2.91 | - | - | - | - | - | 11.64 |
| 36 | 1.19 | - | 5.95 | - | - | - | - | - | - | - | - | - | 5.95 |
| 42 | 1.43 | - | - | - | 1.43 | 1.43 | - | - | - | 1.43 | - | - | 4.29 |
| 43 | 1.33 | 23.94 | 34.58 | 1.33 | 1.33 | - | - | - | - | - | - | - | 61.18 |
| 45 | 1.87 | - | - | - | 1.87 | 1.87 | 1.87 | - | - | - | - | - | 5.61 |
| 46 |  | - | - | 3.21 | - | - | - | - | - | - | - | -- | 3.21 |
| 48 | 3.56 | 3.56 | 3.56 | - | - | - | - | - | - | - | - | - | 7.12 |
| 51 | 2.00 | - | - | - | - | - | 4.00 | - | 2.00 | - | - | - | 6.00 |
| 53 | 2.27 | 9.08 | - | 2.27 | - | 2.27 | 2.27 | 2.27 | - | 2.27 | - | - | 20.43 |
| 60 | 1.90 | 38.00 | 11.40 | 1.90 | - | - | - | - | - | - | - | - | 51.30 |
| 61 | 1.83 | 20.13 | 12.81 | 1.83 | - | - | - | - | - | - | - | - | 34.77 |
| 81 | 5.27 | - | - | - | - | - | 10.54 | - | - | - | - | - | 10.54 |
| 82 | 3.73 | - | - | - | - | - | 7.46 | - | - | - | - | - | 7.46 |
| 83 | 5.41 | 10.82 | 32.46 | 16.23 | - | - | 21.64 | 10.82 | - | - | - | - | 91.97 |
| 84 | 4.86 | 48.60 | 63.18 | 14.58 | - | - | - | - | - | - | - | - | 126.36 |
| 85 | 5.59 | - | 27.95 | 33.54 | 139.75 | 11.18 | - | - | - | - | - | 5.59 | 218.01 |
| 86 | 5.66 | - | - | - | - | - | - | - | 5.66 | - | - | - | 5.66 |
| 87 | 2.10 | 8.40 | 10.50 | 2.10 | 2.10 | - | - | - | - | - | - | - | 23.10 |
| 90 | 5.03 | 30.18 | 30.18 | 5.03 | - | - | - | - | - | - | - | - | 65.39 |
| 92 | 5.72 | - | - | - | 5.72 | - | - | - | 5.72 | - | - | - | 11.44 |
| 93 | 4.80 | - | - | - | 28.80 | 14.40 | 33.60 | 19.20 | 4.80 | - | - | - | 100.80 |
| 94 | 3.06 | 1196.46 | 1884.96 | 85.68 | 6.12 | - | - | - | - | - | - | - | 3173.22 |
| 95 |  | - | - | - | - | - | 5.95 | - | - | - | - | - | 5.95 |
| Total |  | 1511.04 | 2249.71 | 215.48 | 253.83 | 51.58 | 112.10 | 35.80 | 32.65 | 5.11 | 0 | 5.59 | 4472.89 |

mm . Skipjack larvae were caught only in daytime tows (Table 2).

The abundance $\left(\mathrm{P}_{i}\right)$ of frigate tuna larvae calculated by 0.5 mm length classes using equations 2 and 3 varied between $0.166 \times 10^{9}$ and $48.154 \times 10^{9}$ larvae in the area covered by the positive stations (Table 5). Ages of the larvae corresponding to the midpoints of each length class were estimated to be 3.7-22.0 days.

Larval production ( $P_{a}$, Equation 4) during the spawning season for the $2.5-\mathrm{mm}$ length class was $1.731 \times 10^{12}$ larvae (Table 5). Residence time was 3.3 d for larvae spawning that size class. $P_{a d j}$ (Equation 6) was $6.227 \times 10^{12}$ larvae.

## Reproductive Biomass

The biomass of reproductive frigate tuna adults was calculated using the $P_{a d j}$ estimate (above), $F_{r}$ of 76 eggs $\cdot \mathrm{g}^{-1}$, and $K$ of 0.5 in Equation 6. The reproductive biomass was estimated to be 163868 mt based on abundance of the $2.5-\mathrm{mm}$ length class.

## Larval Mortality Rate

Absolute abundance of the frigate tuna larvae decreased exponentially with age (Fig. 6). P values for each length


Figure 3
Distribution and standardized abundance of little tunny larvae during the COSMA 16-72 cruise. Depth contours are in fathoms.
class (Table 3) and the ages corresponding to the midpoints of the length classes (Table 5) fitted to Equation 7 yielded the equation given in Figure 6. The daily mortality coefficent, $M$, was 0.346 .

## Discussion and Conclusions

Identification of larval scombrids is not unambiguous (Richards and Potthoff 1973; Graves et al. 1988). The pigmentation pattern present on the frigate tunas in this study suggests that they are Auxis thazard. Nevertheless, owing to the difficulty in distinguishing the frigate tunas as larvae (Uchida 1981), our results are presented for the Auxis spp. complex. There is a conflict in the literature concerning the diagnostic pigmentation of larval Thunnus atlanticus (Juárez 1972; Richards and Potthoff 1973). However, we are confident of the species identifications in this study
because the Thunnus spp. were cleared and stained, and the number of vertebrae were counted.

The distribution and abundance of larval scombrids in the study area suggest that Campeche Sound is an important tuna spawning area, especially for the frigate tunas and the little tunny. It is noteworthy that, although low in abundance, the presence of larvae of the larger tunas, such as yellowfin and bigeye, indicates that these species also spawn in the area. Although it has been reported that the frigate tunas spawn in almost the entire Gulf of Mexico (Idyll and de Sylva 1963), the frigate tunas and the little tunny are found mainly in waters adjacent to land masses (Williams 1963; Juárez 1974). Larval distribution and abundance of the Scombridae in the Gulf of Mexico reported by Juárez $(1974,1976)$ and Ramírez and Ornelas (in press) have shown that spawning activity is lower in oceanic than coastal waters. We believe that this is related to the temperature and productivity of the area, both


Figure 4
Distribution and standardized abundance of bigeye tuna, blackfin tuna, and king mackerel larvae during the COSMA 16-72 cruise. Depth contours are in fathoms.
important factors in determining breeding areas. It is important to relate future ichthyoplankton studies to information on current dynamics, physical-chemical conditions, hydrological conditions, and biological productivity, and to obtain information on the role of each species in the trophicdynamics of the ecosystem.

The extent of the frigate tunas' spawning area in the Gulf of Mexico is not known, and the population boundaries would be difficult to determine. Although the COSMA 16-72 cruise was not long enough to give an adequate representation of larval production over the spawning season, we consider these data to be important for designing a future survey that would yield an adequate representation of the production dynamics over population sectors.

Our estimate of 163868 t of frigate tuna reproductive biomass suggests that they are a potential fishery resource in the region. In other countries, such as India and Ceylon
in the Indian Ocean, this species is considered to be an important fishery resource, and it is utilized in the same form as other small tunas (Williams 1963; Juárez 1976). Our large spawning population estimate is supported by the fact that the Campeche Sound is a highly productive area where the frigate tunas suffer no significant exploitation. On the other hand, our estimates may be low for the following reasons. The portion of the Campeche Sound that was sampled is probably at the fringes of the populations' spawning area in the Gulf of Mexico. In using larval abundance data for computing spawning biomass and in not accounting for mortality between recently spawned eggs and the larval stage, we have computed only a fraction of the initial egg production which is directly related to spawning stock. Egg mortality of pelagic fishes is typically high, although the gestation period is likely short at these temperatures (Table 1). This information will be improved when more is known of the duration of spawning, fecun-


Figure 5
Distribution and standardized abundance of Spanish mackerel, yellowfin tuna, and skipjack tuna larvae during the COSMA 16-72 cruise. Depth contours are in fathoms.

## Table 5

Abundance by standard length (mm) of frigate tuna larvae for the total area represented by positive stations ( $P_{i}$, Equation 3), total larval production ( $P_{a}$, Equation 4) throughout the spawning season, and the reproductive biomass estimate ( $B_{a}$, Equation 6).

| Length Class <br> $(\mathrm{SL} \mathrm{mm})$ | Range <br> Midpoint | $P_{i}$ <br> $\left(\right.$ no. $\left.\times \quad 10^{9}\right)$ | Age <br> $(\mathrm{d})$ | $P_{a}$ <br> $\left(\right.$ no. $\left.\times 10^{12}\right)$ | $B_{a}$ <br> $(\mathrm{t})$ |
| :--- | :---: | :---: | :---: | :---: | :---: |
| $2.25-2.75$ | 2.5 | 31.741 | 3.7 | - | - |
| $2.76-3.25$ | 3.0 | 48.154 | 6.8 | - | - |
| $3.26-3.75$ | 3.5 | 9.028 | 9.3 | - | - |
| $2.76-4.25$ | 4.0 | 7.478 | 11.6 | - | - |
| $4.26-4.75$ | 4.5 | 1.407 | 13.5 | - | - |
| $4.76-5.25$ | 5.0 | 3.548 | 15.3 | - | - |
| $5.26-5.75$ | 5.5 | 0.980 | 16.9 | - | - |
| $5.76-6.25$ | 6.0 | 1.025 | 1.83 | - | - |
| $6.26-6.75$ | 6.5 | 0.166 | 19.6 | - | - |
| $7.26-7.75$ | 7.5 |  |  |  | Total |
|  |  |  | 1.73 | - |  |
|  |  |  |  |  | - |



Figure 6
Absolute abundance of frigate tuna larvae ( $P$, Table 3 ) versus age corresponding to the midpoints of each size class (Table 5).
dity, sex ratios, growth rates, and mortality during the egg and larval stages.

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## Citations

Ahlstrom, E.H.
1968. An evaluation of the fishery resources available to California fishermen. In The future of the fishing industry of the United States (De Witt Gilbert, ed.), p. 65-80. Univ. Wash. Publ. Fish., New Ser, Vol. 4.
Bullis, H.R. Jr.
1955. Preliminary report on exploratory longline fishing for tuna
in the Gulf of Mexico and the Caribbean Sea. Part I: Exploratory fishing by the Oregon. Commer. Fish. Rev. 17(10):1-15.
Carranza, F.J.
1956. La pesca del atún y sus posibilidades en el Golfo de México. Inst. Mex. Rec. Renov., A.C. 11:1-33.
Graves, J.E., M.A. Simovich, and K.M. Schaefer.
1988. Electrophoretic identification of early juvenile yellowfin tuna, Thunnus albacares. Fish. Bull., U.S. 86:835-838.
Hollister, G.
1934. Clearing and dyeing fish for bone study. Zoologica. 12(10): 89-101.
Houde, E.D., and W.J. Richards.
1969. Rearing larval tunas in the laboratory. Commer. Fish. Rev. 31(12):32-34.
Idyll, C.P., and D.P. de Sylva.
1963. Synopsis of biological data on the frigate mackerel Auxis thazard (Lacépéde) 1802 (Western Atlantic). FAO. Fish. Rep. 6:778-781. Ishida, Y.
1971. Investigation of frigate mackerel (Auxis rochei), 1968 season.

Rep. Kochi Pref. Fish. Exp. Sta. 66,67:119-140. (In Japanese; unedited English Transl. in files of SW Fish. Ctr., Natl. Mar. Fish. Serv., NOAA, Honolulu, HI 96812.)
Jones, S.
1959. Notes on eggs, larvae and juveniles of fishes from Indian waters. III: Katsuwonus pelamis (Linnaeus) and IV: Neothunnus macropterus (Temminck and Schlegel). Indian J. Fish. 6:360-373. Juárez, M.
1972. Las formas larvarias de Thunnus atlanticus. Cuba. Rev. Mar y pesca 78:26-29.
1974. Dónde desova el atún? Mar y Pesca 106:44-47.
1976. Distribución de las formas larvarias de algúnas especies de
la familia Scombridae en aguas del Golfo de México. INP/CIP. Cuba. Rev. Invest. 2(1):33-65.
Klawe, W.L., and B.M. Shimada.
1959. Young scombroid fishes from the Gulf of Mexico. Bull. Mar. Sci. Gulf Carib. 9:100-115.
Kramer, D., M.J. Kalin, E.G. Stevens, J.R. Trailkill, and J.R. Zweifel.
1972. Collecting and processing data on fish eggs and larvae in the California Current Region. U.S. Dep. Commer., NOAA Tech. Rep. NMFS SSRF 370:1-38.
Matsumoto, W.M.
1959. Description of Euthynnus and Auxis larvae from the Pacific and Atlantic oceans and adjacent seas. DANA Rep. Carlsberg Found., No. 50.
1962. Identification of larvae of four species of tunas from the IndoPacific region. DANA Rep. Carlsberg Found., No. 55.
Matsumoto, W.M., E.H. Ahlstrom, S. Jones, W.L. Klawe, W.J.
Richards, and S. Ueyanagi.
1972. On the clarification of larval tuna identification particularly in the genus Thunnus. Fish. Bull., U.S. 70:1-12.
Montolio, M., and M. Juárez.
1976. El desove de Thunnus thynnus thynnus en el Golfo de Méxio - Estimado preliminar de la magnitud de la población en desove a partir de la abundancia de larvas. Int. Comm. Cons. Atl. Tunas, Collect. Vol. Sci. Pap. 6(2):337-344.
Nishikawa, Y., and D.W. Rimmer.
1987. Identification of larval tunas, billfishes and other Scombroid fishes (Suborder Scombroidei): an illustrated guide. Commonw. Sci. Ind. Res. Org., Australia. Mar. Lab. Rep. 186.
Ramírez, A.E., and M. Ornelas.
(In press). Distribución y abundancia de larvas de la familia Scombridae en el Golfo de México y Mar Caribe. Inst. Nal. Pesca. Ciencia Pesquera.
Richards, W.J., and T. Potthoff.
1973. Analysis of the taxonomic characters of young scombrid fishes, genus Thunnus. In The early life history of fish (J.H.S. Blaxter (ed.), p. 623-648. Springer-Verlag, NY, Heidelberg, Berlin.
Richards, W.J., T. Potthoff, and E.D. Houde.
1981. Abundance of bluefin tuna larvae and estimates of spawn-
ing stock sizes in the Gulf of Mexico in 1977 and 1978. Int. Comm. Cons. Atl. Tunas, Collect. Vol. Sci. Pap. 15(2):272-277.
Saville, A.
1964. Estimation of the abundance of a fish stock from egg and larval surveys. Rapp. P.-v. Reun. Cons. Perm. Int. Explor. Mer 155:164-170.
Secretaría de Pesca.
1987. Consulta popular para laplaneación democrática de la pesca en materia de pesquerias. Departamento de Administración de Pesquerías. Sría. Pesca, México, 50 p .
Sette, O.E., and E.H. Ahlstrom.
1948. Estimations of abundance of the eggs of the pacific pilchard Sardinops caerulea off Southern California during 1940 and 1941. J. Mar. Res. 7:511-542.

Simmons, D.C., and L. McDade.
1973. Contribution on the spawning of Auxis sp. (Pisces, Scombridae) in the Atlantic Ocean. Fish. Bull., U.S. 71:321-324.
Smith, P.E., and S.L. Richardson.
1977. Standard techniques for pelagic fish egg and larva surveys. FAO. Fish. Tech. Pap. 175:1-100.
Tibbo, S.N., and J.S. Beckett.
1972. Canadian research activities on tunas and tuna-like fishes in the Atlantic Ocean. 1970-1971. In Int. Comm. Cons. Atlantic Tunas. Rep. for biennial period, 1970-1971, Part III:148-151.

## Ueyanagi, S.

1963. Methods of identification for the young stages of tunas and spearfishes. I: Methods of identifying larval and postlarval tunas. Material for the Tuna Fisheries Research Council. Nankai Reg. Fish. Res. Lab. (In Japanese; English Transl. by W. G. Van Campen 1964, 10 p., SW Fish. Ctr., Natl. Mar. Fish. Serv., NOAA, Honolulu, HI 96812.)
Uchida, R.N.
1964. Synopsis of biological data on frigate tuna, Auxis thazard, and bullet tuna, A. rochei. U.S. Dep. Commer., NOAA Tech. Rep. NMFS Circ., 63 p. (FAO Fish. Synop. 124.)
Williams, F.
1965. Synopsis of biological data on frigate mackerel Auxis thazard (Lacepède) 1802 (Indian Ocean). FAO Fish. Biol. Synop. 47: 157-166.

# Abundance and Distribution of Fish Larvae in the Channel Area of the Patos Lagoon Estuary, Brazil 

JOSÉ H. MUELBERT* and GRACIELA WEISS<br>Departamento de Oceanografia<br>Universidade do Rio Grande<br>C.P. 474<br>96200 Rio Grande RS, Brazil


#### Abstract

Distribution and abundance of fish larvae in the channel area of the Patos Lagoon, Brazil are reported. The Patos Lagoon is a shallow coastal lagoon with a very narrow connection with the Atlantic Ocean. Its southernmost region, characterized by a very dynamic estuary, was the site of 32 ichthyoplankton surveys conducted at 5 stations from April 1981 to March 1983. Samples were taken at 3 depth levels (surface, mid-water and bottom) and results indicated that larval density and distribution were associated with temperature. Larvae occurred year round and at all stations, but were most abundant and diverse during spring and summer with mean densities of $61 / 100 \mathrm{~m}^{3}$ and $189 / 100 \mathrm{~m}^{3}$, respectively. However, this distribution was sometimes altered by meteorological conditions prevailing in the estuary. Twenty families of fishes were represented, including 27 taxa, 19 of which were identified to species. The three most abundant taxa overall were Micropogonias furnieri, Brevoortia pectinata, and Lycengraulis sp., whose larvae accounted for 22.9, 22.6, and $20 \%$ of the total, respectively. Parapimelodus valenciennis; Blennidae; Trichiurus lepturus; Gobiesox strumosus; Paralonchurus brasiliensis; Macrodon ancylodon; Paralichthys sp.; Atherinidae; Gobiosoma parri; and Merticirrrus spp. had relative abundances between 1 and $5 \%$.


## Introduction

Estuaries and coastal waters play an important role in the life cycles of various marine organisms. They provide an abundance of food and protection from predators for larval and juvenile fishes which use these areas as nursery grounds (Pearcy and Myers 1974; Chao and Musick 1977; Misitano 1977; Weinstein 1979; Weiss 1981; Castello 1986). Many marine fish species inhabit regions close to estuaries and rivers. Their eggs and larvae are transported into these areas where they develop and grow. In the North Atlantic, $70 \%$ of commercially important fish species spend part of their life cycle in or near estuaries (McHugh 1966, 1967; Clark 1967). In Mexico, Yañez-Arancibia (1978) reported that $80 \%$ of the coastal ichthyofauna was related to coastal lagoons or areas influenced by them. In addition to these marine species, there are some groups that have their entire life cycles confined to estuarine areas (Weiss 1981; Chao et al. 1982a).

[^2]According to Flores-Coto et al. (1983), the association between ichthyofauna and estuarine environments cannot be completely understood without ichthyoplankton studies which provide information about the early life histories of these organisms. Despite their importance, only a few surveys have been conducted in Brazil to study the relationship between estuarine environments and the development of fish eggs and larvae (Phonlor 1975; Castello 1976, 1977, 1978; Weiss and Krug 1977; Sinque 1980; Weiss 1981; Muelbert 1986).

The Patos Lagoon, located in the southernmost state of Brazil, is the largest coastal lagoon of South America, covering an area of $10360 \mathrm{~km}^{2}$. Its southern region (900 $\mathrm{km}^{2}$ ) is characterized by a very dynamic estuary (Fig. 1). Its dynamic characteristics are determined by its topography and prevailing meteorological conditions, since the tidal range within the estuary is limited to a low diurnal tidal amplitude (mean of 0.47 m ). Most of the estuarine area is very shallow (mean depth of 2 m ), and a channel in its center (mean depth of 15 m ) leads to a very narrow inlet ( 750 m ) connecting the estuary to the Atlantic Ocean. Wind pattern and precipitation in the highlands drained by the lagoon determine the salinity regime of the estuary


Figure 1
Study area and location of the sampling stations. Dotted area represents estuarine region of the Patos Lagoon.
(Castello and Möller 1978). Southerly winds are associated with salt water intrusion whereas northerly winds are related to oligohaline conditions (Costa et al. 1988).

Chao et al. (1982b) cited 110 estuarine and marine fish species inhabiting the estuary and the adjacent coastal waters of the Patos Lagoon. Brevoortia pectinata, Micropogonias furnieri, Cynoscion striatus, and Macrodon ancylodon are commercially important species whose larvae use the Patos Lagoon estuary as a nursery ground (Weiss 1981).

In this paper, we present results on abundance and distribution of fish larvae in the channel of the Patos Lagoon estuary. The study was restricted to the channel area of the estuary to account for the influence of salt water intrusion and freshwater run-off in the assemblage of fish larvae.

## Materials and Methods

Fish eggs and larvae were collected during 32 ichthyoplankton surveys at five sampling stations in the Patos Lagoon estuary from April 1981 to March 1983 (Fig. 1; Table 1). The sampling stations were distributed along the channel area of the estuary from its connection with the ocean to its most interior limit. Stations 1, 2, and 3 were sampled at surface, mid-water and bottom depths, whereas only surface and bottom were sampled at stations 4 and 5 , which were shallower. All five stations were sampled on the same day.

A $61-\mathrm{cm}$ conical net without a closing system was used to take the samples. Owing to the high concentration of

Table 1
Absolute and relative density of fish larvae, temperature, and salinity (Mean $\pm \mathrm{SE}$ ) for the entire study.

| Survey | Date | Season | Temperature $\left({ }^{\circ} \mathrm{C}\right)$ | Salinity (ppt) | Larval density |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | ( $\mathrm{N} / 100 \mathrm{~m}^{3}$ ) | (\%) |
| 1 | 06/04/81 | Autumn | $22.5 \pm 0.3$ | $18.9 \pm 2.0$ | $44.3 \pm 9.4$ | 1.72 |
| 2 | 01/06/81 | Autumn | $18.5 \pm 0.03$ | $8.1 \pm 2.0$ | $6.4 \pm 1.0$ | 0.25 |
| 3 | 19/06/81 | Autumn | $15.1 \pm 0.4$ | $25.2 \pm 0.8$ | $24.0 \pm 10.0$ | 0.93 |
| 4 | 16/07/81 | Winter | $15.3 \pm 0.3$ | $18.9 \pm 3.0$ | $34.0 \pm 9.0$ | 1.32 |
| 5 | 31/07/81 | Winter | $16.3 \pm 0.3$ | $2.2 \pm 0.4$ | $40.3 \pm 8.9$ | 1.57 |
| 6 | 20/08/81 | Winter | $18.4 \pm 0.4$ | $8.4 \pm 2.0$ | $18.7 \pm 4.5$ | 0.73 |
| 7 | 28/08/81 | Winter | $19.6 \pm 0.5$ | $16.3 \pm 2.9$ | $11.4 \pm 2.2$ | 0.44 |
| 8 | 08/09/81 | Winter | $19.6 \pm 0.4$ | $1.2 \pm 0.5$ | $34.0 \pm 10.0$ | 1.32 |
| 9 | 22/09/81 | Winter | $18.7 \pm 1.2$ | $3.2 \pm 1.8$ | $9.3 \pm 1.7$ | 0.36 |
| 10 | 16/10/81 | Spring | $21.7 \pm 1.0$ | $8.0 \pm 3.0$ | $8.0 \pm 2.2$ | 0.31 |
| 11 | 11/11/81 | Spring | $24.3 \pm 1.5$ | $12.6 \pm 3.1$ | $91.8 \pm 39.5$ | 3.57 |
| 12 | 25/11/81 | Spring | $24.0 \pm 0.6$ | $6.9 \pm 2.0$ | $44.5 \pm 7.4$ | 1.73 |
| 13 | 04/12/81 | Spring | $21.2 \pm 0.8$ | $19.5 \pm 2.0$ | $30.4 \pm 9.6$ | 1.18 |
| 14 | 23/12/81 | Summer | $22.6 \pm 0.3$ | $23.7 \pm 2.8$ | $202.3 \pm 39.1$ | 7.87 |
| 15 | 11/01/82 | Summer | $21.4 \pm 0.4$ | $27.2 \pm 3.0$ | $145.9 \pm 19.4$ | 5.68 |
| 16 | 22/01/82 | Summer | $23.3 \pm 1.2$ | $22.1 \pm 4.0$ | $474.4 \pm 66.6$ | 18.46 |
| 17 | 08/02/82 | Summer | $23.9 \pm 1.1$ | $31.5 \pm 1.5$ | $79.0 \pm 23.8$ | 3.08 |
| 18 | 02/04/82 | Autumn | $23.8 \pm 0.2$ | $32.2 \pm 1.0$ | $84.8 \pm 16.4$ | 3.30 |
| 19 | 13/05/82 | Autumn | $20.1 \pm 0.3$ | $27.5 \pm 2.0$ | $5.6 \pm 1.4$ | 0.22 |
| 20 | 01/06/82 | Autumn | $15.8 \pm 0.4$ | $26.4 \pm 2.0$ | $2.0 \pm 0.8$ | 0.08 |
| 21 | 25/06/82 | Winter | $13.7 \pm 0.6$ | $9.6 \pm 0.5$ | $1.1 \pm 0.4$ | 0.04 |
| 22 | 09/08/82 | Winter | $14.5 \pm 0.4$ | $1.5 \pm 0.5$ | $156.0 \pm 41.9$ | 6.07 |
| 23 | 30/08/82 | Winter | $16.5 \pm 0.7$ | $3.7 \pm 1.8$ | $118.8 \pm 24.7$ | 4.62 |
| 24 | 06/09/82 | Winter | $16.0 \pm 0.0$ | $12.2 \pm 3.0$ | $44.0 \pm 17.9$ | 1.71 |
| 25 | 20/09/82 | Winter | $16.2 \pm 0.4$ | $13.2 \pm 2.8$ | $19.0 \pm 17.5$ | 0.74 |
| 26 | 13/10/82 | Spring | $14.8 \pm 0.3$ | $21.9 \pm 2.8$ | $29.7 \pm 10.8$ | 1.16 |
| 27 | 17/11/82 | Spring | $19.1 \pm 0.3$ | $0.0 \pm 0.1$ | $72.2 \pm 14.0$ | 2.81 |
| 28 | 13/12/82 | Spring | $24.3 \pm 0.3$ | $2.0 \pm 1.8$ | $140.4 \pm 22.9$ | 5.47 |
| 29 | 23/12/82 | Summer | $24.6 \pm 0.8$ | $0.3 \pm 0.7$ | $69.5 \pm 10.9$ | 2.70 |
| 30 | 07/01/83 | Summer | $22.8 \pm 0.6$ | $1.7 \pm 1.5$ | $176.1 \pm 43.1$ | 6.85 |
| 31 | 21/01/83 | Summer | $23.9 \pm 0.8$ | $24.9 \pm 3.0$ | $169.4 \pm 35.9$ | 6.59 |
| 32 | 10/02/83 | Summer | $23.4 \pm 2.1$ | $9.7 \pm 3.6$ | $181.5 \pm 51.5$ | 7.06 |

suspended material, a $500-\mu \mathrm{m}$ mesh size was used to minimize clogging of the net. Tows, lasting three minutes each, were taken during daylight hours at a speed of $1 \mathrm{~m} / \mathrm{s}$ against the water current. This procedure resulted in an overestimation of the bottom and mid-water samples of approximately 10 and $5 \%$, respectively. A self-constructed digital flowmeter (identical to General Oceanics Model 2030) was placed in the mouth of the net to estimate volume filtered. Salinity values were obtained using an American Optical refractometer and expressed as ppt (parts per thousand). Water temperature, recorded in degrees Centigrade, was obtained by thermometer. Water samples were collected with a bucket from the surface and with a modified Niskin bottle from mid-water and bottom. All sampled material was preserved with $5 \%$ buffered formalin and processed in the laboratory. Fish eggs and larvae were sorted and counted using a binocular dissecting microscope and their densities were standardized to $100 \mathrm{~m}^{3}$ of water filtered.

Larvae and egg identification was done following Weiss (1981), and determined to the lowest taxonomic level possible. Standard length was measured for the most important species, Brevoortia pectinata, Lycengraulis sp., and Micropogonias furnieri, with an ocular micrometer. Seasonal designations were based on the solar calendar for the Southern Hemisphere. Only the results regarding fish larvae will be addressed in this paper.

A three-way analysis of variance (ANOVA) (Kim and Kohout 1975) was used to assess the effect of season, sampling station, depth strata, and the interactions of these factors on the distribution of temperature, salinity, and larval density. This analysis was combined with the Multiple Comparisons Test of Scheffé ( $P<0.05$ ) to determine the significant differences observed in the spatial and temporal distribution of the dependent variables. Normality of the variables was obtained with the use of the following mathematical transformation of the raw data: salinity and


Figure 2
Distribution of salinity (ppt) at each station for the 32 surveys. (-) station 1; (- -) station 2; ( $\cdots \cdots$ ) station 3; (- . - . -) station 4; and (- $-\cdots-$ ) station 5. A = Autumn; $\mathrm{W}=$ Winter; $\mathrm{Sp}=$ Spring; $\mathrm{S}=$ Summer.
fish larval density were transformed with the expression $y=\ln (x \pm 1)$. Normality was confirmed with the use of Student's $t$ test $(t<0.025)$ for the values of asymmetry and kurtosis of the transformed variable, and homogeneity of variance was tested using Bartlett-Box $F$ statistics $(P<0.05)$ (Sokal and Rohlf 1981).

## Results and Discussion

## Salinity and Temperature

During the study period the mean ( $\overline{\mathrm{x}}$ ) salinity (for all sample stations and depths combined) ranged from 0.0 to almost 35.0 ppt and exhibited a seasonal trend (Fig. 2; Table 1). An increase in salinity values was observed from spring to autumn, the period of maximum salinity. After autumn, the values began to decrease and reached minimal levels in winter (Table 1). However, this pattern was not constant, as shown by sudden changes in salinity at surveys 4 and 5,7 , and 8 , and 30 and 31 (Fig. 2; Table 1).

Salinity decreased significantly from station 1 to station 5 (Fig. 3; Tables 2 and 3). Mean values changed from $19.2 \pm 1.26$ to $5.9 \pm 1.06 \mathrm{ppt}$. Stations 2 and 3 were intermediate or transitional between stations 1,4 , and 5 (Table 3).

Salinity was significantly different with respect to depth strata (Table 2). The mean salinity increased from $10.6 \pm 0.94$ to $15.8 \pm 1.07 \mathrm{ppt}$ from surface to bottom, and surface values were significantly different from mid-water and bottom salinities.


Figure 3
Distribution of mean salinity (ppt) over all surveys and depth levels along the 5 sampling stations. The vertical lines indicate $\pm 1$ standard error.

Three vertical distribution patterns were observed for salinity: vertically homogeneous, with high or low salinity values in the entire water column; slightly stratified; and highly stratified (Fig. 2). Any one station, independent of its position in the estuary, could present any one of these patterns. Station 3, for example, was vertically homogeneous with low salinity during survey 32 (Summer/83) and with high salinity in survey 17 (Summer/82); it was slightly stratified during survey 13 (Spring/81); and it was highly stratified in survey 6 (Winter/81; Fig. 2).

Changes from one pattern to another could occur very rapidly, depending on the dominant meteorological con-

| Table 2 <br> Results of the three-way analysis of variance on temperature, salinity, and larval density. $\mathrm{df}=$ degrees of freedom; ${ }^{*}=$ variances were significantly different $(P<0.05)$; $^{* *}=$ variances were highly significantly different ( $P<0.01$ ) . |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Source of variation | df | Dependent variables <br> (Mean squares) |  |  |
|  |  | Temperature | Salinity | Larval density |
| (Temperature | 296 | 7.50 | 1.36 | 1.89 |
| Within $\{$ Salinity | 340 |  |  |  |
| Larval density | 347 |  |  |  |
| Season | 3 | 259.75* | 15.07* | 43.49* |
| Station | 4 | 5.23 | 7.68** | 2.51 |
| Depth | 2 | 6.42 | 9.40 ** | 1.11 |
| Season by station | 12 | 1.49 | 0.41 | 3.16 |
| Season by depth | 6 | 4.24 | 0.31 | 0.81 |
| Station by depth | 6 | 0.66 | 0.41 | 0.80 |
| Season by station by depth | 18 | 0.54 | 0.24 | 0.52 |

Table 3
Distance of the sampling stations from the ocean; temperature, salinity, and larval density at each station (Mean $\pm \mathrm{SE}$ ). ns = means are not significantly different between stations; $\mathrm{a}=$ mean salinities between stations 1,2 , and 3 are not significantly different; $\mathrm{b}=$ salinities between stations 2,3 , and 4 are not significantly different; $\mathrm{c}=$ mean salinities between stations 4 and 5 are not significantly different. (Multiple comparison test of Scheffé [ $P<0.05$ ].)

|  | Distance from <br> ocean <br> $(\mathrm{km})$ | Temperature <br> $\left({ }^{\circ} \mathrm{C}\right)$ | Salinity <br> $(\mathrm{ppt})$ | Larval density <br> $\left(\mathrm{N} / 100 \mathrm{~m}^{3}\right)$ |
| :---: | :---: | :---: | :---: | :---: |
| Station | 3.7 | $19.6 \pm 0.37(\mathrm{~ns})$ | $19.2 \pm 1.26(\mathrm{a})$ | $94.4 \pm 12.92(\mathrm{~ns})$ |
| 1 | 17.6 | $20.1 \pm 0.38(\mathrm{~ns})$ | $16.6 \pm 1.40(\mathrm{ab})$ | $68.5 \pm 10.90(\mathrm{~ns})$ |
| 2 | 30.5 | $20.4 \pm 0.42(\mathrm{~ns})$ | $13.7 \pm 1.40(\mathrm{ab})$ | $97.4 \pm 18.00(\mathrm{~ns})$ |
| 3 | 44.4 | $20.2 \pm 0.52(\mathrm{~ns})$ | $10.2 \pm 1.40(\mathrm{bc})$ | $66.7 \pm 13.26(\mathrm{~ns})$ |
| 4 | 62.0 | $20.8 \pm 0.53(\mathrm{~ns})$ | $5.9 \pm 1.06(\mathrm{c})$ | $83.4 \pm 14.86(\mathrm{~ns})$ |
| 5 |  |  |  |  |

ditions. The large Patos Lagoon basin is drained exclusively through the narrow access channel. During winter and spring, high precipitation levels (Castello and Möller 1978) combined with moderate northerly winds (Costa et al. 1988) result in intense freshwater discharge. During these seasons, only sporadic strong southerly winds may intrude seawater into the channel area (Fig. 2). An increase in the estuarine salinity results from a decrease in freshwater runoff during summer and fall (Castello and Möller 1978) combined with either weak or moderate southerly winds (Costa et al. 1988), or with an increased tidal action on the estuary which is dampened during winter and spring by high freshwater run-off (Abreu 1987). However, the pattern of the salinity distribution depends on the interaction of all these forces. During survey 4 and 5, for example, the velocity of the southerly wind changed from approximately 36 to $4.5 \mathrm{~km} / \mathrm{h}$ (Costa et al. 1988). This caused the mean
salinity of the channel to decrease from $18.9 \pm 3.0$ to $2.2 \pm 0.4$ ppt (Table 1; Fig. 2).

Temperature also followed a seasonal pattern (Fig. 4). Mean values ranged from $24.3 \pm 1.5^{\circ} \mathrm{C}$ during spring to $13.7 \pm 0.6^{\circ} \mathrm{C}$ during winter (Table 1; Fig. 4). The horizontal distribution of temperature was relatively constant along the stations sampled (Fig. 5) and no significant differences were found among the station means (Table 2; Table 3). Temperature remained relatively constant over depth and no significant differences were found (Table 2). The fact that the estuary is shallow and well mixed prevents the development of thermal stratification.

## Larval Abundance and Seasonal Distribution

Twenty families of fishes were represented in samples that included 27 taxa, 19 of which were identified to species


Figure 4
Distribution of mean temperature $\left({ }^{\circ} \mathrm{C}\right)$ over all sample stations and depth levels for the 32 surveys. The vertical lines indicate $\pm 1$ standard error. $\mathrm{A}=$ Autumn; W = Winter; $\mathrm{Sp}=$ Spring; $\mathrm{S}=$ Summer.
(Table 4). About $8.8 \%$ (primarily damaged clupeiform larvae) were unidentifiable. Micropogonias furnieri, Brevoortia pectinata, and Lycengraulis sp. were the most abundant species. Together they represented $65.4 \%$ of the total larval fish density, and their length distribution was between 1.8 and 12.0 mm SL, 2.9 and 21.0 mm SL , and 1.3 and 34.0 mm SL, respectively. The second group of taxa was less abundant and accounted for $23.4 \%$ of the total. These included Parapimelodus valenciennis; Blenniidae; Trichiurus lepturus; Gobiesox strumosus ${ }^{1}$; Paralonchurus brasiliensis; Macrodon ancylodon; Paralichthys sp.; Atherinidae; Gobiosoma parri; and Menticirrhus spp. A third group was composed of taxa whose relative abundance was less than $1 \%$ each and included Achirus garmani; Syngnathus folletti; Gobionellus spp.; Peprilus paru; Cynoscion striatus; Umbrina canosai; Hyporhamphus kronei; Parona signata; Symphurus jenynsi; Anchoa marinii; Prionotus punctatus; Synagrops sp.; Mugil spp.; and Porichthys porosissimus.

In general, the highest mean density of larvae was observed during the summer (Fig. 6; Table 1), during which time every species identified in the study, with the exception of Prionotus punctatus and Porichthys porosissimus, was present (Fig. 7). This was particularly evident during the summers of 1981 and 1982. Autumn showed the lowest mean larval density (Table 1). During this season Brevoortia pectinata, Lycengraulis sp., and Atherinidae were the dominant taxa, while Anchoa marinii, Parapimelodus valenciennis, Hyporhamphus kronei, Syngnathus folletti, Mugil spp., Micro-

[^3]

Figure 5
Distribution of mean temperature $\left({ }^{\circ} \mathrm{C}\right)$ over all surveys and depth levels along the 5 sampling stations. The vertical lines indicate $\pm 1$ standard error.
pogonias furnieri, Paralonchurus brasiliensis, and Gobiosoma parri were present, but not in large numbers (Fig. 7). During the period studied, winter and spring had a relatively higher density than autumn (Table 1). However, during winter, fewer species were present in the estuary (Fig. 7). An increase in the number of species and density was observed in spring, and the maximum density was reached in summer (Figs. 6 and 7).

The increase in larval fish density and composition, which occurred during spring and summer months, was associated with an increase in temperature (Figs. 4 and 7). Owing to its influence on spawning, temperature, rather than salinity, may influence the cycle of larval density (Flores-Coto et al. 1983; Houde and Alpern Lovdal 1984).

Two different patterns of seasonal distribution occurred in the Patos Lagoon estuary: one group of larvae occurred

## Table 4

Absolute and relative mean density of fish larvae collected per cruise in the Patos Lagoon estuary. All stations and depth levels were combined.

| Family | Species | Larval density |  |
| :---: | :---: | :---: | :---: |
|  |  | [(N/100 m ${ }^{3}$ )/cruise] | (\%) |
| CLUPEIDAE | Brevoortia pectinata | 225.9 | 22.56 |
| ENGRAULIDAE | Anchoa marinii | 0.2 | 00.2 |
|  | Lycengraulis sp. | 200.2 | 20.00 |
| PIMELODIDAE | Parapimelodus valencienis | 43.5 | 4.34 |
| BATRACHOIDIDAE | Porichthys porosissimus | 0.02 | 0.002 |
| EXOCOETIDAE | Hyporhampus kronei | 0.9 | 0.09 |
| ATHERINIDAE | Atherinidae spp. | 13.9 | 1.39 |
| SYNGNATHIDAE | Syngnathus folletti | 4.8 | 0.48 |
| TRIGLIDAE | Prionotus punctatus | 0.1 | 0.01 |
| PERCICHTHYIDAE | Synagrops sp. | 0.1 | 0.01 |
| CARANGIDAE | Parona signata | 0.6 | 0.06 |
| SCIAENIDAE | Cynoscion striatus | 2.3 | 0.23 |
|  | Macrodon ancylodon | 18.4 | 1.84 |
|  | Menticirrhus spp. | 11.0 | 1.10 |
|  | Micropogonias furnieri | 229.2 | 22.89 |
|  | Paralonchurus brasiliensis | 21.4 | 2.14 |
|  | Umbrina canosai | 1.7 | 0.17 |
| MUGILIDAE | Mugil spp. | 0.1 | 0.01 |
| BLENNIIDAE | Blenniidae spp. | 36.1 | 3.61 |
| GOBIIDAE | Gobiosoma parri | 11.2 | 1.12 |
|  | Gobionellus spp. | 3.3 | 0.33 |
| TRICHIURIDAE | Trichiurus lepturus | 34.3 | 3.43 |
| STROMATEIDAE | Peprilus paru | 2.6 | 0.26 |
| GOBIESOCIDAE | Gobiesox strumosus | 29.0 | 2.90 |
| BOTHIDAE | Paralichthys sp. | 14.9 | 1.49 |
| SOLEIDAE | Achirus garmani | 6.8 | 0.68 |
| CYNOGLOSSIDAE | Symphurus jenynsi | 0.2 | 0.02 |
| Others |  | 88.4 | 8.83 |



Figure 6
Distribution of larval density $\left(N / 100 \mathrm{~m}^{3}\right)$ at each station for the 32 surveys. (-) station 1 ; (---) station $2 ;(\cdots \cdots)$ station 3 ; (- . - . -) station 4; and (- . - . -) station 5. $\mathrm{A}=$ Autumn; $\mathrm{W}=$ Winter; $\mathrm{Sp}=$ Spring; $\mathrm{S}=$ Summer.
$\qquad$


Figure 7
Distribution of total larval density $\left(N / 100 \mathrm{~m}^{3}\right)$ expressed as $\ln (x+1)$ for each identified species along the 32 surveys. $\mathrm{A}=$ Autumn; $\mathrm{W}=$ Winter; $\mathrm{Sp}=$ Spring; $\mathrm{S}=$ Summer.
continuously throughout the year, and another exhibited a discontinuous seasonal cycle (Fig. 7). Brevoortia pectinata, Lycengraulis sp., and Atherinidae represented the first group and their presence in the estuary suggested that these species reproduce during the entire year. The second group predominated during early spring and late summer and included Micropogonias furnieri, Macrodon ancylodon, Paralonchurus brasiliensis, Blenniidae, Gobionellus spp., Peprilus paru, Trichiurus lepturus, Paralichthys sp., and Achirus garmani.

## Horizontal Distribution

Most of the species sampled showed a decrease in their density towards the interior of the estuary, whereas others, such as Parapimelodus valenciennis, increased. Brevoortia pectinata, Lycengraulis sp., Atherinidae, Micropogonias furnieri, and Achirus garmani were abundant throughout the entire estuary. Hyporhamphus kronei was most abundant in the central region of the estuary (stn. 3; Fig. 8).

The individual patterns of horizontal distribution and abundance reflected the origins of the species and the degree to which the estuary is used by them. Species with an oceanic origin, such as Anchoa marinii, Porichthys porosissimus, Prionotus punctatus, Synagrops sp., Parona signata, Cynoscion striatus, Umbrina canosai, Mugil spp., Trichiurus lepturus, Peprilus paru, and Symphurus jenynsi, occurred in the estuary during periods of strong salt water intrusion. This increase in salinity was the result of an increase in the frequency of southerly winds (Costa et al. 1988) which favor their transport into the estuary. They were generally restricted to regions influenced by the ocean (stns. 1 to 3 ), occurring only in low densities in the interior of the estuary (stns. 4 and 5; Fig. 8).

Species whose eggs and larvae originated in the adjacent coastal region were distributed along the entire estuary with a small decrease in abundance in the most interior waters. These species use the estuary as a nursery area for their larvae and juveniles, and include Brevoortia pectinata, Lycengraulis sp., Macrodon ancylodon, Menticirrhus spp., Micropogonias furnieri, and Paralonchurus brasiliensis (Fig. 8).

Atherinidae, Syngnathus folletti, Blenniidae, Gobionellus spp., and Achirus garmani are taxa that reside in the Patos Lagoon estuary and show a uniform density distribution throughout the entire channel (Fig. 8). However, the presence of more than one species in each of the above families could mask their individual distributional patterns.

Parapimelodus valenciennis is a freshwater species and showed a decrease in its density towards the ocean (Fig. 8). It is an uncommon species in the estuary and its presence is associated with periods of strong freshwater run-off. These larvae apparently cannot withstand increases in salinity, and they do not use the estuary as a nursery area.

## Vertical Distribution

While most of the species showed a homogeneous vertical distribution, a few were selectively distributed with depth (Fig. 9). Parapimelodus valenciennis, Umbrina canosai, Gobionellus spp., Trichiurus lepturus, and Achirus garmani were most abundant at the bottom, whereas Brevoortia pectinata, Lycengraulis sp., Parona signata, Gobiesox strumosus, and Symphurus jenynsi showed highest densities at the surface. Anchoa marinii was found at the surface and bottom, Porichthys porosissimus was present only at the bottom, and Mugil spp. occurred at surface and mid-water levels.

The vertical distribution of fish larvae in estuaries is generally related to tidal flow (Graham 1972; Weinstein et al. 1980; Fortier and Leggett 1982). In the Patos Lagoon estuary, astronomic tides have a very small amplitude and their flow can be amplified or reduced by the action of either wind (Costa et al. 1988) or precipitation in the drainage basin (Castello and Möller 1978). Consequently, the vertical distribution of fish larvae probably does not reflect a well-defined circulation pattern.

Different groups of larvae presented different patterns of vertical distribution (Fig. 9). Weiss (1981) found this selective distribution to be related to the salinity structure of the water column. According to her, Brevoortia pectinata, Lycengraulis sp., Parapimelodus valenciennis, and Atherinidae are abundant in less saline surface waters, while Micropogonias furnieris and Trichiurus lepturus are frequently found near the more saline bottom waters. In this study, however, Parapimelodus valenciennis and Atherinidae did not exhibit this pattern. Parapimelodus valenciennis was found to be more abundant at the bottom than near the surface (Fig. 9). This may have been the result of a large number of larvae being captured during an anomalous period of freshwater runoff in the spring of 1982 (Fig. 2). At this time, this species dominated the entire estuary (Fig. 7). The abundance of Atherinidae was high at the surface and at the bottom with a decrease in mid-water levels (Fig. 9). Three atherinid species occur in the Patos Lagoon estuary (Chao et al. 1982b) and their individual distributions could be associated with different depths.

## Conclusion

Micropogonias furnieri, Brevoortia pectinata, and Lycengraulis sp. are the most abundant fish larvae in the channel area of the Patos Lagoon estuary. Fish larvae are present throughout the year in the estuary, and abundance and diversity increases during the months of spring and summer, with an increase in temperature. The larvae are distributed along the entire estuary and in the entire water column. This general picture of seasonal, horizontal, and vertical distribution can be altered by the meteorological conditions prevailing in the estuary. Further studies should be con-
$\qquad$


Figure 8
Distribution of total larval density ( $N / 100 \mathrm{~m}^{3}$ ) expressed as $\ln (x+1)$ for each identified species at the 5 sampling stations.
ducted to understand specific distribution patterns better. These studies should take into account the biology and
ecology of individual species and the dynamic character of the Patos Lagoon estuary.


Figure 9
Distribution of total larval density ( $N / 100 \mathrm{~m}^{3}$ ) expressed as $\ln (x+1)$ for each identified species at the 3 depth levels. $\mathrm{S}=$ surface waters; $\mathrm{M}=$ mid-water; $B=$ bottom-waters.

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## Citations

Abreu, P.C.O.V.
1987. Variações temporais de biomassa fitoplanctônica (clorofila) e relações com fatores abióticos no canal de acesso ao estuário da lagoa dos Patos (RS-Brasil). M.S. thesis, Universidade do Rio Grande, Rio Grande, RS, Brasil, 107 p.
Castello, J.P.
1976. Projeto Lagoa. Relatórios. Rio Grande. FURG, BOA, ser. rel. 1-4.
1977. Projeto Lagoa. Relatórios. Rio Grande. FURG, BOA, ser. rel. 5-6.
1978. Projeto Lagoa. Relatórios. Rio Grande. FURG, BOA, ser. rel. 8-9.
1986. Distribuición, crecimiento e maduración sexual de la corvina juvenil (Micropogonias furnien) en el estuario de la "Lagoa dos Patos', Brasil. Physis (B. Aires) 44(106):21-36.
Castello, J.P., and O.O. Möller Jr.
1978. On the relationship between rainfall and shrimp production in the estuary of the Patos Lagoon (Rio Grande do Sul, Brazil).

Atlântica (Rio Grande) 3:67-74.
Chao, L.N., and J.A. Musick.
1977. Life history, feeding habits and functional morphology of juvenile sciaenid fishes in the York River estuary, Virginia. Fish. Bull., U.S. 75(4):657-702.
Chao, L.N., L.E. Pereira, J.P. Vieira, M.A. Benvenuti, and L.P.R. Cunha.

1982a. Bio-ecology of fishes in the estuary and the adjacent coastal region of the Lagoa dos Patos, Brazil. Atlântica (Rio Grande) (Resumos) 5(2):27.
1982b. Relação preliminar dos peixes estuarinos e marinkos da Lagoa dos Patos e região costeira adjacente, Rio Grande do Sul, Brasil. Atlântica (Rio Grande) 5(1):67-75.
Clark, J.
1967. Fish and man. Conflict in the Atlantic estuaries. Am. Littoral Soc. Spec. Publ. 5, 78 p.
Costa, C.S.B., U. Seeliger, and P.G. Kinas.
1988. The effect of wind velocity and direction on the salinity regime in the Lower Patos Lagoon estuary. Cienc. Cult. (São Paulo) 40(9):909-912.
Flores-Coto, C., F. Barba-Torres, and J. Sanchez-Robles.
1983. Seasonal diversity, abundance, and distribution of ict thyoplankton in Tamiahua Lagoon, Western Gulf of Mexico. Trans. Am. Fish. Soc. 112:247-256.
Fortier, L., and W.C. Leggett.
1982. Fickian transport and the dispersal of fish larvae in estuaries. Can. J. Fish. Aquat. Sci. 39:1150-1163.
Graham, J.J.
1972. Retention of larval herring within the Sheepscot Estuary of Maine. Fish. Bull., U.S. 70(2):299-305.
Houde, E.D., and J. Alpern Lovdal.
1984. Seasonality of occurrence, foods and food preferences of ichthyoplankton in Biscayne Bay, Florida. Est. Coast. Shelf Sci. 18:403-419.
Kim, J., and F.J. Kohout.
1975. Analysis of Variance and Covariance: Subprograms Anova and Oneway. In Statistical package for the social sciences (N.H. Nie, C.H. Hull, J.G. Jenkins, K. Steinbrenner, and D.H. Brent, eds.), p. 398-433. McGraw-Hill, NY.
McHugh, J.L.
1966. Management of estuarine fisheries. Am. Fish. Soc. Spec. Publ. 3:133-154.
1967. Estuarine nekton. In Estuaries (G.H. Lauff, ed.), p. 581620. Am. Assoc. Adv. Sci. Publ. 83.

Misitano, D.A.
1977. Species composition and relative abundance of the Columbia River Estuary, 1973. Fish. Bull., U.S. 75(1):218-222.
Muelbert, J.H.
1986. Estrutura e dinâmica do ictioplâncton da área de canal, no estuário da Lagoa dos Patos, RS, no período outono/81 à verão/83. M.A. thesis, Universidade do Rio Grande, Rio Grande, RS, Brasil, 88 p.
Pearcy, W.G., and S.S. Myers.
1974. Larval fishes of Yaquina Bay, Oregon: A nursery ground for marine fishes? Fish. Bull., U.S. 72:201-213.
Phonlor, G.
1975. Ictioplâncton da região de Tramandaí e adjacências. GEDIP-Projeto Tramandaí Documentos Ocasionais, 2: 1-45.
Sinque, C.
1980. Larvas de Sciaenidae (Teleostei) identificadas na região estuarino-lagunar de Cananéia. Bolm. Zool. (Univ. S. Paulo) 5:39-77.
Sokal, R.R., and F.J. Rohlf.
1981. Biometry: the principles and practice of statistics in biological research. W.H. Freeman and Co., San Francisco, (2nd edition), 776 p.
Weinstein, M.P.
1979. Shallow marsh habitats as primary nurseries for fishes and shellfishes, Cape Fear River, North Carolina. Mar. Biol. (Berl.) 58:227-243.
Weinstein, M.P., S.L. Weiss, R.G. Hodson, and L.R. Gerry.
1980. Retention of three taxa of postlarval fishes in an intensively flushed tidal estuary, Cape Fear River, North Carolina. Fish. Bull., U.S. 78:419-436.
Weiss, G.
1981. Ictioplancton del Estuario de Lagoa dos Patos, Brasil. Ph.D. thesis, Faculatad de Ciencias Naturales y Museo, Universidad Nacional de La Plata, 164 p.
Weiss, G., and L.C. Krug.
1977. Características do desenvolvimento e metarmorfose de Lycengraulis olidus (Engraulidae) e Brevoortia pectinata (Clupeidae) no estuário da Lagoa dos Patos, Brasil. Atlântica (Rio Grande) 2(1):83-117.
Yañez-Arancibia, A.
1978. Taxonomía, ecología y estructura de las comunidades de peces en las lagunas costeras con bocas efímeras del Pacífico de México. Instituto de Ciencias del Mar y Limnología, Univ. Nal. Auton. de México, Publ. Esp. 2, México, Distrito Federal.

# Larval Distribution and Abundance of Myctophidae, Gonostomatidae, and Sternoptychidae from the Southern Gulf of Mexico 

CÉSAR FLORES-COTO and URIEL ORDOÑEZ-LÓPEZ<br>Instituto de Ciencias del Mar y Limnología<br>Universidad Nacional Autónoma de México<br>Apdo. Postal 70-305<br>México 04510 D.F.


#### Abstract

Larval distribution and abundance of Myctophidae, Gonostomatidae, and Sternoptychidae from the southern Gulf of Mexico were studied. The material was collected during four cruises from 1983 to 1984. The family Myctophidae with 2618 larvae, was represented by 12 genera and at least 19 species; Gonostomatidae, with 1101 larvae, by 10 genera and at least 13 species; and Sternoptychidae, with 146 larvae, by two genera. Diaphus spp. ( 47.5 individuals $/ \mathrm{m}^{2}$ sea surface [ $=\mathrm{L}]$ ), Cyclothone spp. (27.6/L), Benthosema suborbitale ( $29.1 / \mathrm{L}$ ) and Maurolicus muelleri (23.3/L) were the most abundant taxa. The highest frequency of occurrence and greatest abundance of larvae were recorded over the slope and at oceanic stations, which likely corresponded to the principal spawning areas. Only the most abundant taxa occurred frequently over the outer shelf, and occasionally in shallow areas ( $\langle 50 \mathrm{~m}$ ). Only 10 out of the 28 species showed evidence of seasonality, presenting their highest larval abundance only during some of the cruises; Hygophum hygomii, Centrobranchus nigroocellatus, Vinciguerria poweriae and Pollichthys mauli were most abundant in winter, Myctophum asperum in spring, Bonapartia pedaliota in summer, Hygophum reinhardtii and Maurolicus muelleri in spring and summer, and Notoscopelus resplendens and Myctophum selenops in the winter and spring.


## Introduction

The ichthyoplanktonic composition of mesopelagic fishes is not well known on a worldwide scale (Hopkins and Lancraft 1984; Roe and Badcock 1984), the southern Gulf of Mexico being among the least studied regions. An investigation of larvae of the families Gonostomatidae, Sternoptychidae, and Myctophidae in the southern Gulf of Mexico was undertaken in the present study as part of the research program of the Laboratorio de Zooplancton, Instituto de Ciencias del Mar y Limnología, Universidad Nacional Autónoma de México. These families were chosen because of their high relative abundance in the area.
Larval stages of the species in these families are only partially known, owing in part to their great specific diversity, as about 60 genera and more than 330 species are represented (Ahlstrom et al. 1984; Moser et al. 1984).
For the southern Gulf of Mexico and Caribbean Sea, 16 genera and 43 species of Myctophidae, 15 genera and 19 species of Gonostomatidae, and 3 genera and 8 species of Sternoptychidae have been recorded (Rass 1971; Naf-
paktitis 1975; Shiganova 1977; Houde et al. 1979; Fahay 1983; Richards 1984; Romero and del Castillo 1984; Gartner et al. 1987). Studies directed at the larval stages in these families in the southern Gulf and Caribbean Sea have been scarce. The most extensive has been that by Houde et al. (1979) who recorded 16 genera and 25 species from the northeast part of the Gulf. There have been no specific investigations into our study area, except for one which Flores-Coto et al. 1988 composed a species list.

The main objectives of the our study were to contribute to the knowledge of the composition, distribution, and abundance of the species comprised by these families and to define their spawning seasons and areas.

## Study Area

The study area included the southwestern portion of the Gulf of Mexico, limited to the north by lat. $21^{\circ} \mathrm{N}$ and comprising the continental shelves of Veracruz, Tabasco, and Campeche, and adjacent oceanic zones. The slope is delineated as the 182 m isobath (Fig. 1).


Figure 1
Study area and location of stations during the cruises. Subsequent figures are for sampling stations of specific cruises.

## Material and Methods

Zooplankton samples were collected aboard the oceanographic ship Justo Sierra, during four cruises: PROGMEX I (31 March-8 April 1983, '‘spring’'); IMECO (15-25 February 1984, '‘winter''); PROGMEX II (25 April4 May 1984, '‘spring'’); and PROGMEX III (7-17 August 1984, 'summer'’).

A sampling pattern was established mainly over the continental shelf, covering a larger portion of the oceanic zone during the IMECO cruise (Fig. 1). Zooplankton sampling consisted of a double-oblique plankton tow following a circular course using a $60-\mathrm{cm}$ bongo net with 333and $505-$ micron mesh nets. The water volume filtered was
calculated using flowmeters placed in each net. The depth and tow time varied from 18 to 200 min and 5 to 23 min , respectively, according to bathymetry. Ship speed was held between 1.5 and 2 kn . Each sample was fixed in $4 \%$ formalin buffered with sodium borate.

Fish larvae were sorted from each sample and myctophids, gonostomatids, and sternoptychids separated for study. Larval density was standardized as number of larvae per $10 \mathrm{~m}^{2}$ of sea surface ('L'’) (Houde et al. 1979; Richards 1984),

$$
L_{j}=\frac{C_{j} Z_{j}}{V_{j}} \times 10
$$

Table 1
Abundance of Myctophidae larvae. Number of positive stations (S), number of larvae (N), and mean larval density (L) (per $10 \mathrm{~m}^{2}$ of sea surface) are given for each cruise.

|  | PROGMEX-I <br> 49 stations |  |  | IMECO <br> 29 stations |  |  | PROGMEX-II <br> 40 stations |  |  | PROGMEX-III <br> 55 stations |  |  | Mean |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Taxa | S | N | L | S | N | L | S | N | L | S | N | L | No. | \% |
| Diaphus spp. | 29 | 326 | 52.59 | 18 | 142 | 41.13 | 24 | 233 | 57.70 | 34 | 222 | 38.43 | 47.46 | 18.35 |
| Benthosema suborbitale | 18 | 86 | 29.51 | 16 | 105 | 36.23 | 20 | 74 | 20.83 | 26 | 119 | 29.80 | 29.09 | 11.25 |
| Notolychnus valdiviae | 13 | 36 | 15.69 | 10 | 31 | 17.50 | 12 | 37 | 19.99 | 22 | 95 | 26.28 | 19.86 | 7.68 |
| Lampanyctus spp. | 10 | 25 | 11.86 | 14 | 85 | 35.64 | 7 | 11 | 9.22 | 13 | 19 | 10.27 | 16.75 | 6.48 |
| Hygophum taaningi | 19 | 45 | 11.81 | 7 | 11 | 8.21 | 11 | 22 | 10.47 | 12 | 20 | 10.61 | 10.27 | 3.97 |
| Diogenichthys atlanticus | 10 | 14 | 8.90 | 7 | 10 | 8.05 | 14 | 31 | 12.31 | 22 | 40 | 10.80 | 10.01 | 3.87 |
| Hygophum macrochir | 3 | 7 | 12.00 | 8 | 14 | 10.03 | 4 | 4 | 5.58 | 1 | 2 | 12.04 | 9.91 | 3.83 |
| Myctophum nitidulum-affine | 17 | 34 | 10.53 | 11 | 17 | 8.42 | 14 | 25 | 11.78 | 18 | 22 | 8.35 | 9.77 | 3.78 |
| Hygophum reinhardtii | 14 | 25 | 8.76 | 8 | 9 | 6.69 | 10 | 14 | 8.19 | 22 | 39 | 12.01 | 8.91 | 3.45 |
| Myctophum asperum | 2 | 3 | 8.44 | 3 | 3 | 5.88 | 6 | 10 | 13.70 | 3 | 3 | 5.83 | 8.46 | 3.27 |
| Myctophum obtusirostre | 3 | 6 | 6.85 | 3 | 6 | 12.50 | 3 | 3 | 3.10 | 5 | 7 | 9.47 | 7.98 | 3.09 |
| Centrobranchus nigroocellatus | 3 | 3 | 5.21 | 3 | 7 | 12.82 | 3 | 4 | 8.14 | 1 | 1 | 4.47 | 7.66 | 2.96 |
| Hygophum hygomii | 1 | 1 | 2.69 | 9 | 24 | 15.79 | 3 | 4 | 4.96 | 1 | 1 | 6.27 | 7.43 | 2.87 |
| Notoscopelus resplendens | 7 | 11 | 9.18 | 11 | 22 | 10.73 | 3 | 3 | 7.15 | 0 | 0 | 0.00 | 6.76 | 2.62 |
| Myctophum selenops | 9 | 15 | 9.17 | 4 | 5 | 6.62 | 8 | 14 | 9.96 | 0 | 0 | 0.00 | 6.44 | 2.49 |
| Myctophum spp. | 9 | 16 | 10.33 | 0 | 0 | 0.00 | 10 | 12 | 6.43 | 8 | 11 | 7.30 | 6.01 | 2.33 |
| Lepidophanes spp. | 5 | 13 | 9.23 | 0 | 0 | 0.00 | 5 | 6 | 9.02 | 2 | 2 | 4.49 | 5.68 | 2.20 |
| Hygophum sp. | 1 | 2 | 7.94 | 0 | 0 | 0.00 | 0 | 0 | 0.00 | 1 | 2 | 6.42 | 3.59 | 1.39 |
| Lobianchia gemellarii | 0 | 0 | 0.00 | 1 | 1 | 7.31 | 0 | 0 | 0.00 | 0 | 0 | 0.00 | 1.83 | 0.71 |
| Ceratoscopelus warmingii | 0 | 0 | 0.00 | 1 | 1 | 5.39 | 0 | 0 | 0.00 | 0 | 0 | 0.00 | 1.35 | 0.52 |
| Hygophum benoiti | 0 | 0 | 0.00 | 0 | 0 | 0.00 | 2 | 2 | 3.16 | 0 | 0 | 0.00 | 0.79 | 0.31 |
| Myctoph-indeter. | 10 | 102 | 44.51 | 17 | 189 | 61.10 | 10 | 29 | 15.65 | 15 | 23 | 9.07 | 32.58 | 12.60 |
| Total |  | 770 | 275.18 |  | 682 | 310.02 |  | 538 | 237.32 |  | 628 | 211.90 | 258.60 | 100 |

where $\quad L_{j}=$ number of larvae at station j under $10 \mathrm{~m}^{2}$ of sea surface
$C_{j}=$ the catch of larvae at station j
$Z_{j}=$ the depth (in meters) at station j
$V_{j}=$ the volume filtered by the net (in cubic meters) at station j

As two samples were available for each station, resulting from the 333- and 505-micron nets, the highest value of larval density was always taken for each taxon, independently of the mesh. This treatment was based on the assumption that some taxa may be more efficiently sampled by either the 505 - or 333 -micron mesh.

## Results

A total of 3863 larvae were identified. Of those, 2618 corresponded to myctophids with 12 genera and at least 19 species; 1101 gonostomatids with 10 genera and at least 13 species, and 146 sternotychids, represented by 2 genera and at least 2 species (Tables 1 and 2).
Species identifications in the genera Diaphus, Lampanyctus, Lepidophanes, Cyclothone, Sternoptyx, Argyropelecus, and in
some specimens of Myctophum and Hygophum were difficult and will, therefore, be treated at the generic level. The larvae of Myctophum nitidulum and Myctophum affine could not be distinguished from each other and will be treated as the complex " $M$. nitidulum-affine." Species are considered by family and are listed in decreasing order of generic abundance.

## Family Myctophidae

Diaphus spp. (47.5/L), Benthosema suborbitale (29.1/L), Notolychnus valdiviae ( 19.0 L ), and Lampanyctus spp. (161.7/L) were the most abundant taxa, accounting for about $43.7 \%$ of all myctophids collected (Table 1). Larvae of these taxa occurred during all four oceanic cruises, where they were widely distributed, and caught at virtually all oceanic and slope stations (Figs. 2, 3, 4, and 5). Lampanyctus spp. occurred at a somewhat lower frequency than the other taxa, and its larvae were scarce over the continental shelf, generally occurring at stations greater than 100 m in depth (Fig. 5). Hygophum was the next most abundant taxon, accounting for about $15.8 \%$ of the family, although no one species was especially common (maximum 10.3 L). Hygophum species had a wide distribution at the slope and
$\qquad$

## Table 2

Abundance of Gonostomatidae and Sternoptychidae larvae. Number of positive stations (S), number of larvae (N), mean larval density (L) (per $10 \mathrm{~m}^{2}$ of sea surface) are given for each cruise.

|  | $\begin{aligned} & \text { PROGMEX-I } \\ & \text { (spring) } \\ & 49 \text { stations } \end{aligned}$ |  |  | IMECO <br> (winter) 29 stations |  |  | PROGMEX-II <br> (spring) 40 stations |  |  | PROGMEX-III <br> (summer) <br> 55 stations |  |  | Mean |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Taxa | S | N | L | S | N | L | S | N | L | S | N | L | No. | \% |
| Cyclothone spp. | 22 | 102 | 22.32 | 17 | 100 | 32.87 | 23 | 150 | 37.06 | 28 | 87 | 18.20 | 27.62 | 19.65 |
| Maurolicus muelleri | 9 | 46 | 36.36 | 4 | 8 | 9.38 | 14 | 47 | 17.98 | 25 | 121 | 29.61 | 23.33 | 16.60 |
| Vinciguerria poweriae | 10 | 17 | 9.06 | 4 | 27 | 36.41 | 10 | 24 | 13.80 | 14 | 24 | 10.30 | 17.39 | 12.38 |
| Pollichthys mauli | 8 | 16 | 10.66 | 7 | 28 | 17.95 | 4 | 4 | 6.84 | 0 | 0 | 0.00 | 8.86 | 6.30 |
| Gonostoma atlanticum | 6 | 11 | 8.99 | 6 | 8 | 6.82 | 8 | 10 | 7.28 | 15 | 20 | 7.84 | 7.73 | 5.50 |
| Vinciguerria attenuata | 4 | 4 | 6.39 | 9 | 11 | 6.78 | 8 | 14 | 10.53 | 11 | 13 | 6.78 | 7.62 | 5.42 |
| Vinciguerria nimbaria | 7 | 9 | 5.58 | 6 | 8 | 7.71 | 7 | 11 | 8.68 | 6 | 6 | 7.68 | 7.41 | 5.28 |
| Gonostoma elongatum | 1 | 1 | 11.23 | 8 | 11 | 8.05 | 0 | 0 | 0.00 | 12 | 14 | 7.51 | 6.70 | 4.77 |
| Bonapartia pedaliota | 0 | 0 | 0.00 | 1 | 1 | 4.65 | 1 | 1 | 6.30 | 4 | 7 | 12.34 | 5.82 | 4.14 |
| Ichthyococcus ovatus | 0 | 0 | 0.00 | 0 | 0 | 0.00 | 1 | 1 | 5.34 | 2 | 2 | 6.67 | 3.00 | 2.14 |
| Margrethia obtusirostra | 0 | 0 | 0.00 | 1 | 1 | 6.02 | 0 | 0 | 0.00 | 3 | 3 | 5.53 | 2.89 | 2.05 |
| Valenciennellus tripunctulatus | 0 | 0 | 0.00 | 2 | 2 | 4.86 | 1 | 1 | 6.39 | 0 | 0 | 0.00 | 2.81 | 2.00 |
| Diplophos taenia | 1 | 1 | 2.58 | 0 | 0 | 0.00 | 0 | 0 | 0.00 | 0 | 0 | 0.00 | 0.65 | 0.46 |
| Gonost-indeter. | 10 | 34 | 22.58 | 12 | 32 | 16.75 | 9 | 27 | 19.21 | 12 | 36 | 16.33 | 18.72 | 13.32 |
| Total-Gonostomatidae |  | 241 | 135.76 |  | 257 | 158.25 |  | 290 | 139.41 |  | 333 | 128.77 | 140.55 | 100 |
| Sternoptyx sp. | 3 | 5 | 13.40 | 14 | 55 | 21.36 | 11 | 25 | 15.36 | 19 | 42 | 14.31 | 16.11 | 72.80 |
| Argyropelecus spp. | 2 | 2 | 7.37 | 0 | 0 | 0.00 | 6 | 7 | 7.68 | 6 | 10 | 9.03 | 6.02 | 27.20 |
| Total-Sternoptychidae |  | 7 | 20.77 |  | 55 | 21.36 |  | 32 | 23.04 |  | 52 | 23.34 | 22.13 | 100 |



Figure 2
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Diaphus spp., southern Gulf of Mexico. Smallest dots represent stations with no larvae.



Figure 3 (top)
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Benthosema suborbitale, southern Gulf of Mexico. Smallest dots represent stations with no larvae.

Figure 4 (bottom)
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Notolychnus valdiviae, southern Gulf of Mexico. Smallest dots represent stations with no larvae.


Figure 5
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Lampanyctus spp., southern Gulf of Mexico. Smallest dots represent stations with no larvae.
oceanic zones, but were only occasionally found over the continental shelf. Hygophum taaningi and $H$. reinhardtii had relatively high frequencies of occurrence.

The genus Myctophum represented $14.9 \%$ of the family, although, as for Hygophum, the individual species were not very abundant (maximum 9.8 L ). All species had a low frequency of occurrence except for $M$. nitidulum-affine. The larvae of Myctophum had a wide distribution and were principally caught at oceanic and slope stations; over the continental shelf they were scarce and larvae of $M$. asperum were not recorded in this area.

Diogenichthys atlanticus represented 3.9 \% of the farnily (7.7/L), and was sixth in frequency of occurrence. All larvae were captured at slope and oceanic stations, except for three stations over the continental shelf.

The remaining myctophid taxa, Centrobranchus nigroocellatus, Notoscopelus resplendens, Lobianchia gemellarii, and Ceratoscopelus warmingii, were scarce. These species were recorded at oceanic stations and over the continental slope. Six myctophid species appeared to show seasonal peaks of abundance (Table 1). They included Hygophum hygomii and Centrobranchus nigrocellatus (winter peak); Hygophum reinhardtii (spring/summer peak); Myctophum asperum (spring peak); and Notoscopelus resplendens and Myctophum selenops (winterspring peak).

## Family Gonostomatidae

Of the gonostomatids, Cyclothone spp. (27.6/L), Maurolicus muelleri (23.3/L), Vinciguerria poweriae (17.4/L), Vinciguerria attenuata (7.6/L), Vinciguerria nimbaria (7.4/L), Pollichthys mauli (8.9/L), Gonostoma atlanticum (7.7/L), and Gonostoma elongatum ( $6.7 / \mathrm{L}$ ) were most frequent and had the highest mean densities.

Cyclothone spp. were widely distributed in the oceanic zone and over the continental slope, where they were caught at almost all stations sampled; over the shelf they were present less frequently but were found at some stations less than 50 m depth (Fig. 6).

Maurolicus muelleri and the three Vinciguerria species were distributed widely along the slope and oceanic zones; they were occasionally recorded over the shelf (Figs. 7 and 8).

Larvae of $G$. atlanticum and $G$. elongatum were not captured over the continental shelf, but they were broadly distributed in the slope and oceanic areas, although $G$. elongatum was only found at one station during the spring cruises. Pollichthys mauli larvae were mostly restricted to stations over the continental slope and the nearby oceanic areas, being found only once over the shelf.

The remaining gonostomatids, Bonapartia pedaliota, Margrethia obtusirostra, Valenciennellus tripunctulatus, Ichthyococcus



Figure 6 (top)
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Cyclothone spp., southern Gulf of Mexico. Smallest dots represent stations with no larvae.

Figure 7 (bottom)
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Maurolicus muelleri, southern Gulf of Mexico. Smallest dots represent stations with no larvae.


Figure 8
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Vinciguerria poweriae ( $\bullet$ ), Vinciguerria attenuata ( $\boldsymbol{\square}$ ) and Vinciguerria nimbaria ( $\mathbf{\Delta}$ ), southern Gulf of Mexico. Smallest dots represent stations with no larvae.
ovatus, and Diplophos taenia, were scarce or rare. With the exception of $D$. taenia, which was collected over the shelf, these species were present only in samples from oceanic and slope stations. Seasonal peaks of abundance were apparent for 4 gonostomatid species (Table 2). They included Vinciguerria poweriae and Pollichthys mauli (winter peak), Maurolicus muelleri (spring/summer peak), and Bonabartia pedaliota (summer peak).

## Family Sternoptychidae

Sternoptyx sp. was the most abundant and frequent taxon of this family, with a mean density of 16.0 L (Fig. 9). Larvae of Argyropelecus spp. were scarce and were not caught during the February cruise.

The higher abundance and frequency of occurrence of both taxa were recorded in the oceanic area; they were scarce over the shelf. Highest densities of Sternoptyx sp. were recorded in winter; however, this taxon was common year-round.

## Discussion and Conclusions

The relative abundance and composition of taxa found in
this study closely corresponded to data collected from the northeast Gulf of Mexico (Houde et al. 1979) and the Caribbean Sea (Richards 1984).

In the present study, the family Myctophidae had the highest number of larvae and taxa. Diaphus spp., B. suborbitale, $N$. valdiviae, and Lampanyctus spp. accounted for more than $43 \%$ of the myctophids, whereas Cyclothone spp., $M$. muelleri, and $V$. poweriae were most abundant among the gonostomatids, accounting for about $40 \%$ of this family, and Sternoptyx sp. made up approximately $70 \%$ of the sternoptychids.

The relative abundances of these taxa closely resembled the records of Houde et al. (1979) for the northeast Gulf, where Diaphus spp. were the most abundant taxa; the relative abundances of the remaining taxa differed only slightly. Species differing in relative abundance between the southern and northeastern Gulf of Mexico included Maurolicus muelleri (second in abundance among gonostomatids in the present study but the most abundant gonostomatid in the northeast Gulf) and Gonostoma atlanticum (common in the our study but rare in the northeast Gulf).

Differences between the southern Gulf and the Caribbean Sea (Richards 1984) were large in terms of relative


Figure 9
Abundance (number of larvae per $10 \mathrm{~m}^{2}$ of sea surface) and distribution of Sternoptyx sp., southern Gulf of Mexico.
abundances of taxa; however, species compositions were very similar. In contrast to our findings, Richards (1984) reported C. warmingii, P. mauli, and G. elongatum to be abundant; whereas $B$. suborbitale, D. atlanticus, M. muelleri and $V$. poweriae were found to be rare. These differences may have been related to sampling periodicity, however, since the Caribbean area was sampled only during winter and summer.

The distribution patterns of all species collected in this study can be considered to be primarily oceanic. However, many species, particularly the most common ones, had at least some larvae in the neritic zone, principally at the outer zone of the wide shelf of Tabasco-Campeche or over the narrow shelf of Veracruz, where mixing of neritic and oceanic water takes place. Some of these species were occasionally captured in areas less than 50 m depth. The distribution of scarce or rare species, such as M. asperum, L. gemellarii, C. warmingii, B. pedaliota, V. tripunctulatus, $M$. obtusirostra, I. ovatus, and Argyropelecus spp., was restricted to the oceanic area.

The wide distribution of species in the oceanic area suggests that spawning usually takes place in deep zones and that migration of larvae to neritic waters occurs by means of currents. Thus similar distributions should be expected over geographically distant shelves such as those of Florida
and Tabasco-Campeche. However, the narrowness of the Florida shelf allows most of the species in these families to occur at depths less than 100 m .

Only 10 out of the 28 recorded species showed evidence of seasonality, presenting their highest abundances only during some of the cruises. Hygophum hygomii, Centrobranchus nigroocellatus, Vinciquerria poweriae and Pollichthys mauli were most abundant in winter, Myctophum asperum in spring, Bonapartia pedaliota in summer, and Hygophum reinhardtii and Maurolicus muelleri in the period of spring and summer; whereas the greatest abundance of Notoscopelus resplendens and Myctophum selenops larvae were recorded in the winter and spring.

The seasonal patterns of these species do not correspond with the data for these northeast Gulf (Houde et al. 1979) or for the Caribbean (Richards 1984), except for M. muelleri which was principally recorded in summer in the northeast Gulf, and during spring and summer in this study.

Most of the species which exhibited seasonal abundance peaks in the northeast Gulf and Caribbean were most abundant in the summer. In our study area, B. pedaliota was the only species which had its highest occurrence in this period, but this species was rare in the other study areas.

Identification and comparison of seasonal trends in this and other studies may have been limited by the low
numbers of cruises made (four in this study and two in the Caribbean). However, differing environmental conditions between the study areas should also be considered as a potential factor for influencing the spawning periods of these species.

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## Citations

Ahlstrom, E.H., W.J. Richards, and S.H. Weitzman.
1984. Families Gonostomatidae, Sternoptychidae and associated stomiiform Groups: Development and Relationships. In Ontogeny and Systematics of Fishes (H.G. Moser, W.J. Richards, D.M. Cohen, M.P. Fahay, A.W. Kendall Jr., and S.L. Richardson, eds.), p. 184-198. Am. Soc. Ichthyol. Herpetol. Spec. Pub. No. 1.
Fahay, M.P.
1983. Guide to the early stages of marine fishes occurring in the western North Atlantic Ocean, Cape Hatteras to the southern Scotian shelf. J. Northwest At. Fish. Sci. 4:1-432.
Flores-Coto, C., L. Sanvicente-Añorve, R. Pineda-López, and
M. Rodríguez-Vanlier.
1988. Composición, distribución y abundancia ictioplanctónica del Sur del Golfo de México. Univ. Cienc. Univ. Juárez Autón. de Tabasco 5 (9):65-84.
Gartner, J.V., T.L. Hopkins, R.C. Baird, and D.M. Milliken.
1987. The lanternfishes (Pisces: Myctophidae) of the eastern Gulf
of Mexico. Fish Bull., U.S. 85:81-98.
Hopkins, 'T.L., and T.M. Lancraft.
1984. The composition and stock of mesopelagic micronecton at $27^{\circ} \mathrm{N}, 86^{\circ} \mathrm{W}$ in the eastern Gulf of Mexico. Contrib. Mar. Sci. 27:143-158.
Houde, E.D., J.C. Leak, C.E. Dowd, S.A. Berkely, and W.J. Richards. 1979. Ichthyoplankton abundance and diversity in the eastern Gulf of Mexico. Report to Bureau of Land Management, Contract AA550-CT7-28, 546 p.
Moser, H.G., E.H. Ahlstrom, and J.R. Paxton.
1984. Myctophidae: Development. In Ontogeny and Systematics of Fishes (H.G. Moser, W.J. Richards, D.M. Cohen, M P. Fahay, A.W. Kendall Jr., and S.L. Richardson, eds.), p. 218-239. Am. Soc. Ichthyol. Herpetol. Spec. Pub. No. 1.
Nafpaktitis, B.G.
1975. Review of the lanternfish genus Notoscopelus (family Myctophidae) in the North Atlantic and Mediterranean. Bull. Mar. Sci. 25:75-87.
Rass, T.S.
1971. Deep sea fish in the Caribbean Sea and the Gulf of Mexico (The American Mediterranean Region). In Coloquio sobre investigaciones y recursos del Mar Caribe y regiones adyacentes (UNESCO eds.), p. 509-525; 18-26 Nov. 1968, Willemstand, Curacao, Antillas Holandesas.
Richards, W.J.
1984. Kinds and abundances of the larvae in the Caribbean Sea and adjacent areas. U.S. Dep. Commer., NOAA Tech. Rep., NMFS SSRF-776:1-54.
Roe, H.S.J., and J. Badcock.
1984. The diel migrations and distributions within a mesopelagic community in the northeast Atlantic. 5. Vertical migrations and feeding of fish. Prog. Oceanog. 13:389-424.
Romero, M., and J. del Castillo.
1984. Distribución y abundancia de larvas y juveniles de peces mictófidos en el Mar Caribe. Acad Cienc. Cuba. 26:1-16.
Shiganova, T.A.
1977. Larvae and juveniles of the lanternfish (Myctophidae, Pisces) of the Atlantic Ocean. Okeanol. Akad. Nauk SSSR. 109:42-112.

# Larval Migration and Mortality Rates of Bay Anchovy in the Patuxent River 

JULES J. LOOS<br>Water Quality Dept., Potomac Electric Power Company<br>1900 Pennsylvania Ave.<br>N.W. Washington, DC 20068

ELGIN S. PERRY
Dept. of Animal Sciences
University of Maryland
College Park, MD 20742


#### Abstract

The distribution, migration patterns, and mortality of bay anchovy (Anchoa mitchill) were studied in the Patuxent River subestuary of Chesapeake Bay from 1 June to 17 August 1987. The sizespecific distribution of larvae indicated hydrodynamic transport from downriver spawning areas to upriver nursery areas. This distribution was evidenced by many small larvae and few large larvae downriver opposed to relatively more large larvae and few small larvae upriver. An alternative hypothesis, that this pattern was caused solely by differential mortality between upriver and downriver portions of the estuary, was eliminated. Mortality rates were computed with a decreasing rate function of length from the Pareto family using abundance data that are censored for a range of growth rates found in the literature ( $3 \mathrm{~mm} / \mathrm{wk}$ and $4 \mathrm{~mm} / \mathrm{wk}$ ). Mortality rate estimates varied from 0.161 to 0.162 per mm of growth at 10 mm total length (TL) and from 0.064 to 0.079 per mm of growth at 20 mm TL . The mortality rate function derived here is compared to a range of constant mortality rates estimated for larval bay anchovies in Biscayne Bay, Florida. Mortality rates for the Pareto model were greater than the constant estimates for larvae less than two days old and became less than the constant mortality rates for larvae greater than five days old. As a result the net mortalities estimated by the Pareto model for the Patuxent River were much less by the time larvae were eight days old.


## Introduction

This study is a component of an entrainment impact assessment for the Chalk Point Electric Generating Station on the Patuxent River, a tributary of the Chesapeake Bay Estuary. The assessment determines the geographic boundaries of the larval bay anchovy, Anchoa mitchilli, population in the Patuxent River and estimates recruitment. Geographic boundaries of the population are determined by analysis of the migration patterns of the larvae. Recruitment is estimated through standing crop estimates and a mortality function. An independent assessment has been done for Maryland's Power Plant Research Program (Polgar et al. 1988) and is the subject of ongoing studies. Here, we present the findings relative to the migration of
larval bay anchovy in the Patuxent River and an improved methodology for estimating larval mortality rates.

The bay anchovy is a suitable subject for the study of fisheries dynamics because it is abundant over a wide geographical area and it is accessible to sampling. Eggs are free floating; small larvae are initially planktonic; and larger larvae, juveniles and adults, are pelagic. Bay anchovy range from the Gulf of Maine to the Yucatan Peninsula (Hildebrand 1963), is one of the dominant species in the Chesapeake Bay area (Hildebrand and Schroeder 1928), and is an important forage species (Hollis 1952; Merriner 1975; Chao and Musick 1977). Bay anchovy populations appear to be regulated largely by events occurring during the first few months of life (Vouglitois et al. 1987)


Figure 1
Map of the Patuxent River estuary with segment boundaries used to refer to sampling locations in this study. Segment boundaries are nautical river miles of Cronin and Pritchard (1975). The following abbreviations are used in the inset: BAL $=$ Baltimore, MD; D.C. = Washington, District of Columbia; CHES BAY $=$ Chesapeake Bay; PAX R. = Patuxent River.

Bay anchovy spawning in the Chesapeake Bay extends from the Atlantic Ocean (Olney 1983) nearly to freshwater at the upper end of tributary rivers (Dovel 1971). The extent of spawning upriver is thought to be controlled by salinity. The bay anchovy can spawn in waters with salinity as low as 1 part per thousand (ppt), but they typically do not spawn in waters less than 6 ppt (Dovel 1971). Maximum spawning occurs in waters of 13 to 15 ppt . This study was designed to sample the region of the Patuxent that begins above and extends below the area of maximum spawning activity.

Bay anchovy spawning in the upper Chesapeake Bay and Patuxent River occurs from April to September with a peak in July (Dovel 1971). Spawning occurs at night (Ferraro 1980a, 1980b) and the eggs require approximately 20 to 24 hours to hatch (J. Cowan, Chesapeake Biological Laboratory, Solomons, MD. pers. commun. Nov. 1989; Kuntz 1914) and produce larvae approximately 2 to 3 mm total length (TL) (Wang and Kernehan 1979).

The Patuxent River is located in Maryland, USA. Its confluence with Chesapeake Bay is on the West side in the middle portion of the bay above that of the Potomac River

|  | Table 1 <br> Methods employed in three sampling programs. |  |  |
| :---: | :---: | :---: | :---: |
|  |  | Sampling Program |  |
|  | Regional study | Diel study | Egg study |
| Patuxent River segments | 4 to 35 | 35 and 10 to 14 | 4 to 35 |
| Chesapeake Bay | One transect | Not sampled | Not sampled |
| Sampling dates (1987) | 1 June to 19 August ${ }^{\text {a }}$ | 15 June to 17 August | 29 June to 19 August |
| Frequency | Weekly | Biweekly | Weekly |
| Diel periods included | Night | Day and nıght | Day |
| Sampling gear | 0.25-m² Tucker Trawl | $1.0-\mathrm{m}^{2}$ Tucker Trawl | 0.25-m Tucker Trawl |
| Type of tow | Discrete depth | Discrete depth | Stepped oblique |
| Sampling design | Random stratified | Random stratitified | Fixed station |

(Fig. 1). The Patuxent River Estuary is 61 km long and has a volume of 695 million $\mathrm{m}^{3}$ and a surface area of $113 \mathrm{~km}^{2}$ (Edinger et al. 1989).

Freshwater inflow to the Patuxent River Estuary during the summer spawning season averages $5 \mathrm{~m}^{3} / \mathrm{sec}$. Most of the estuary is mesohaline (Academy of Natural Sciences, Philadelphia (ANSP) 1983). During this period, the lower deep portion of the estuary (maximum depth, 29 m ) is partially mixed with a gradual longitudinal salinity gradient (Edinger et al. 1989). Near river segment 14 (Fig. 1) the estuary becomes shallow (maximum depth, 4 m ), has a steep longitudinal salinity gradient. Depth increases in segment 11 (maximum depth, 6 m ), and the water column remains partially mixed.

This paper gives results of studies of various aspects of bay anchovy recruitment. First, a description of sampling methods is presented, followed by a description of analytical methods for assessing temporal and spatial distribution. A mortality model based on the assumption that mortality rate is an inverse function of larval length is introduced. This is followed by results and discussion to assess the merits of the decreasing mortality rate model relative to a constant mortality rate model and to evaluate the hypothesis that bay anchovy larvae originate in the lower section of the Patuxent River estuary and migrate to the middle section of the estuary.

## Sampling Methods

Bay anchovies in the Patuxent River estuary reportedly spawn near the river mouth and move to nursery areas upriver (Dovel 1971). Vertical and longitudinal distributions of eggs and larvae during this study verified this migration phenomenon. Distribution of eggs and larvae were observed in three sampling programs (Table 1). The geographic location of samples was determined with refer-
ence to 35 river segments (Fig. 1). There was no sampling above segment 4. Earlier studies (Setzler et al. 1979) indicated few bay anchovy larvae above segment 4.

The regional study specified that collections be made over a 12 -week period from 1 June to 19 August 1987. The 35 segments were grouped into 9 regions, each with up to 4 depth layers (Fig. 1, Table 2), resulting in a total of 32 sampling cells. From 29 June to 19 August collections were made at four depths at randomly selected locations along a transect across the Chesapeake Bay at the mouth of the Patuxent River. During each week two nighttime collections were made from each of the Patuxent river cells and at each depth along the Chesapeake Bay transect. These collections were made with a $0.25-\mathrm{m}^{2}$ Tucker trawl with $253-\mu$ mesh. A General Oceanics model 2030R flow meter was mounted in the mouth of the net to quantify volume filtered. Tows were of 2-min duration and the mean sample volume was $25 \mathrm{~m}^{3}$. Salinity was measured at the surface and bottom in each segment.

The diel study specified day and night sampling and employed a larger net. A $1.0-\mathrm{m}^{2}$ Tucker trawl with $253-\mu$ mesh was used to sample each of three depth layers in segment 35 at the mouth of the river and in segments 10 to 14 upriver. Three depth layers were sampled at each location and 6 to 9 samples were collected in each depth layer. Sampling was done every two weeks at alternate locations during the period 15 June to 17 August. These tows were also of 2-min duration and the mean sample volume in both areas was about $100 \mathrm{~m}^{3}$.

Densities of eggs and newly hatched larvae were monitored by sampling weekly during the day for the period 29 June to 3 August. Samples were collected from five to six fixed locations within each of three river sections: lower river (segments 26 to 35 ), middle river (segments 14 to 25), and upper river (segments 4 to 13). Oblique tows of variable duration were taken and the mean sample volume was $65 \mathrm{~m}^{3}$. Sampling was done between 0900 and 1500

Table 2
Hypsographic data for sampling regions in the Patuxent River Estuary (From Cronin and Pritchard 1975 and Edinger et al. 1989).

| Region <br> (segments included) $)^{a}$ | Length <br> $(\mathrm{m} \times 10)$ | Surface area <br> $\left(\mathrm{m}^{2} \times 10\right)$ | Volume <br> $\left(\mathrm{m}^{3} \times 10\right)$ | Mean depth <br> $(\mathrm{m})$ | Sampling depth <br> layers |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 to 3 | 5.56 | 1.95 | 4.8 | 2.46 | $1,2,3$ |
| to 7 | 7.41 | 1.67 | 8.3 | 4.97 | $1,2,3$ |
| 8 to 11 | 7.41 | 5.50 | 13.6 | 2.47 | $1,2,3$ |
| 12 to 13 | 3.70 | 5.59 | 15.1 | 2.70 | $1,2,3$ |
| 14 to 17 | 7.41 | 12.74 | 42.4 | 3.33 | $1,2,3,4$ |
| 18 to 21 | 7.41 | 19.04 | 96.6 | 5.07 | $1,2,3,4$ |
| 22 to 25 | 7.41 | 20.82 | 122.2 | 5.87 | $1,2,3,4$ |
| 26 to 29 | 7.41 | 15.43 | 136.7 | 8.87 | $1,2,3,4$ |
| 3 to 33 | 7.41 | 15.28 | 155.2 | 10.16 | $1,2,3,4$ |
| 34 to 35 | 3.70 | 14.74 | 99.6 | 6.76 | $1,2,3,4$ |

${ }^{a}$ Each segment is 1.85 km (1 nautical mile) in length. There was no sampling above segment 4. Earlier studies (Setzler et al. 1979) indicated few bay anchovy larvae above segment 4.
${ }^{b}$ Layer 1 represents depth from 0 to 1 m ; layer 2, 1 to 3 m ; layer 3, 3 to 8 m ; Layer $4,8 \mathrm{~m}$ to bottom. Layers 3 and 4 were pooled in analysis.
hours. Sampling during the day allowed some mixing to occur between the spawning event the night before and sampling, and reduced the patchiness of the egg distribution resulting in less variance.

Larvae 4- to $23-\mathrm{mm}$ TL were considered to be sampled effectively. Larvae less than $4-\mathrm{mm}$ TL were subject to loss owing to extrusion (Houde and Lovdal 1984) through the sampling net and to poor sorting efficiency during sample processing. Thus only larvae greater than 3-mm TL were used in mortality computations. Larvae greater than 7-mm TL exhibited gear avoidance and appropriate corrections were implemented before making standing crop estimates.

## Analytical Methods

## Distribution and Migration

Analysis of variance (ANOVA) followed by a Tukey multiple range test was used to evaluate longitudinal and vertical differences in densities of eggs and larvae grouped in $4-\mathrm{mm}$ length categories. Densities were transformed by the logarithmic expression

$$
\begin{equation*}
z=\log _{\mathrm{e}}\left(\frac{n}{V}+1\right) \tag{1}
\end{equation*}
$$

where $z=$ response variable analyzed
$n=$ count of larvae in the sample
$V=$ volume sampled by the gear.
In this transformation, densities were standardized to the mean sample volume, thus retaining the approximate
lognormal distribution of the counts. A graphical check (Tukey 1962; Draper and Smith 1981) found that normality assumptions and homogeneous variance assumptions were met when data were standardized by this transformation.

Size distributions of larvae collected at the mouth of the river (segment 35) during the diel study were compared graphically with those collected in the upper portion of the river (segments 10 to 14 ).

Standing crop estimates were based on data from $0.25-\mathrm{m}^{2}$ Tucker trawl collections. Adjustments in densities obtained with the $0.25-\mathrm{m}^{2}$ Tucker trawl were made to reflect size specific sampling efficiency. This correction was derived by comparison of densities of larvae collected with the $0.25-\mathrm{m}^{2}$ trawl with those obtained with a $1.0 \mathrm{~m}^{2}$ trawl. Larvae less than 7 mm TL apparently did not avoid the smaller net. This correction factor, estimated by least squares for larvae greater than 7 mm , was based on the following expression:

For $L>7$ :
$\log _{e}\left(D_{1.0}\right)-\log _{e}\left(D_{0.25}\right)=-0.2656+0.0418 L ;$
where
$\log _{\mathrm{e}}\left(D_{0.25}\right)=$ logarithm transform as shown above for larval density in $0.25 \mathrm{~m}^{2}$ Tucker trawl (density units-number per $100 \mathrm{~m}^{3}$ );
$\log _{e}\left(D_{0.1}\right)=$ logarithm transform as shown above for larval density in $1.0 \mathrm{~m}^{2}$ Tucker trawl; (density units-number per $100 \mathrm{~m}^{3}$ );
$L=$ total length (mm).

## Table 3

Selected growth rates of the bay anchovy reported in the literature.

| Growth rate ( $\mathrm{mm} / \mathrm{d}$ ) | Type of study | State | Source |
| :---: | :---: | :---: | :---: |
| 0.43 to 0.55 | laboratory | FL | Saksena and Houde 1972 |
| 0.48 | field | NC | Fives et al. 1986 |
| 0.43 to 0.56 | field | FL | Leak and Houde 1987 |
| 0.39 to 0.62 | mesocosm | MD | Cowan and Houde (1989) ${ }^{\text {a }}$ |
| 0.58 to 0.62 | field | NY | Castro and Cowen (in press) |

The correction factor was obtained by exponentiating the difference:

$$
\begin{array}{ll}
\frac{D_{1.0}}{D_{0.25}}=\mathrm{e}^{(-0.2656+0.0418 L)}, & \text { for } L>7 \\
\frac{D_{1.0}}{D_{0.25}}=1, & \text { for } L \leqslant 7 \tag{3}
\end{array}
$$

Standing crops were estimated by multiplying the adjusted density of larvae in each cell by the volume of each of the 32 sampling cells. Cell volumes were estimated from the geometry of the estuary (Cronin and Pritchard 1975). The total standing crop is reported by river section.

## Mortality

The larval mortality model assumed that the mortality rate was inversely proportional to length scaled by a power transformation, i.e.,

$$
\begin{equation*}
\frac{d N_{L}}{d L}=\frac{-\alpha N_{L}}{L^{\beta}} ; \tag{4}
\end{equation*}
$$

where $L=$ length;
$N_{L}=$ the number of larvae of length $L$;
$\alpha, \beta=$ model parameters, $\alpha>0, \beta \geqslant 0$.
This differential equation integrates to the following survival model:

$$
\begin{equation*}
\log _{\mathrm{e}}\left(N_{L}\right)=\frac{-\alpha}{1-\beta} L^{(1-\beta)}+c \tag{5}
\end{equation*}
$$

where $c=$ a constant of integration.

This model is a member of the Pareto family of survival models (Miller 1981; Arnold 1985). For nonlinear least
squares estimation new parameters were assigned to yield the following equation:

$$
\begin{equation*}
\log _{e}\left(N_{L}\right)=\alpha^{\prime} L^{\beta^{\prime}}+c \tag{6}
\end{equation*}
$$

where $\quad \alpha^{\prime}=-\alpha /(1-\beta)$;

$$
\beta^{\prime}=(1-\beta)
$$

The mortality rate per individual was obtained by dividing both sides of the initial differential equation by $N_{L}$ and by changing the sign to express the fractional decrease in $N_{L}$ as positive. Thus the mortality rate per individual is $\alpha L^{-\beta}$.

The assumptions required to apply this model are 1) there can be no immigration or emigration of larvae from the system; 2) length must be a valid proxy for age; and 3 ) sampling is sufficiently frequent to sample all cohorts within the spawning season. If the mortality model is applied to individual cohorts within the spawning season, an additional assumption that survival should remain constant over the study period is required. If all larvae spawned in the season are considered as a cohort (year class), this assumption is not required.

The regional study data were used for the mortality analysis because it included the whole estuary and more adequately represented all sizes of larvae. Length was emphasized in analyses rather than age to facilitate the processing of the large number of samples. A range of growth rates for bay anchovy from several sources was used in discussion to enhance comparisons with published rates.

Abundances for $1-\mathrm{mm}$ length intervals were summed over sample weeks. Abundance estimates used in these analyses were adjusted for sampling efficiency as appropriate. Data for bay anchovies were censored to include complete weekly cohorts only. Estimates of length versus age based on otolith analysis for bay anchovy indicated growth of about 3 to 4 mm per week (Table 3 ). Thus the model was fitted to the data using two censoring schemes: one assuming $3-\mathrm{mm}$ growth per week, and one assuming $4-\mathrm{mm}$ growth per week. The mortality model was estimated

| Table 4 <br> Average weekly salinity (ppt) in the Patuxent River. NS $=$ No salinity measurement. |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | Segme |  |  |  |  |
| Week | 4-7 | 8-11 | 12-13 | 14-17 | 18-21 | 22-25 | 26-29 | 30-33 | 34-35 |
| 6/01/87 | 3.0 | 5.0 | NS | 10.0 | 10.0 | 11.5 | 13.0 | 12.5 | 13.5 |
| 6/08/87 | 0.8 | 1.0 | 7.3 | 11.3 | 11.0 | 12.0 | 13.0 | 13.0 | 13.5 |
| 6/15/87 | 2.5 | 6.5 | 10.0 | 11.5 | 12.5 | 12.5 | 12.8 | 13.0 | 13.3 |
| 6/22/87 | 3.8 | 3.8 | 9.3 | 11.0 | 11.8 | 12.8 | 13.0 | 13.5 | 14.0 |
| 6/29/87 | 2.0 | 4.5 | 8.8 | 10.5 | 11.3 | 12.5 | 12.5 | 13.0 | 12.3 |
| 7/06/87 | 2.0 | 5.0 | 9.8 | 11.5 | 12.0 | 12.5 | 13.0 | 13.0 | 13.8 |
| 7/13/87 | 4.0 | 5.5 | 10.0 | -1.3 | 12.3 | 12.8 | 12.8 | 13.0 | 13.0 |
| 7/20/87 | 2.3 | 4.5 | 9.0 | 10.3 | 11.5 | 11.5 | 12.3 | 12.3 | 11.8 |
| 7/27/87 | 5.3 | 6.0 | 10.3 | -1.8 | 12.8 | 13.0 | 13.0 | 14.3 | 14.5 |
| 8/03/87 | 6.0 | 6.5 | 10.8 | 2.0 | 12.8 | 12.8 | 12.8 | 12.8 | 13.0 |
| 8/10/87 | 7.5 | 8.0 | 11.0 | 11.8 | 12.3 | 12.8 | 12.5 | 13.0 | 13.0 |
| 8/17/87 | 5.0 | 6.8 | 10.8 | 12.0 | 12.8 | 12.8 | 11.5 | 12.5 | 13.0 |

using the nonlinear regression procedure of SAS Institute (1985).

## Results

## Distribution and Migration

Vertically averaged salinity at the mouth of the river ranged from 11.8 to 14.5 . ppt. Salinity upriver was more variable; in segments 4 to 7 salinity ranged from 0.8 ppt in June to 7.5 ppt in August (Table 4).
Anchovy spawning occurred further up river as the season progressed. Egg densities in 1987 showed an interaction between river longitude and date ( $P=0.001$ ). From 29 June to 13 July, greatest densities were in the lower and middle portions of the estuary. During the week of 20 July, densities became more uniform in the river. By 27 July and 3 August, the trend had reversed and the greatest densities were upriver (Table 5). A similar trend was observed for newly hatched larvae. In spite of this shift in densities, standing crops of eggs and newly hatched larvae upriver remained low because of the small river volume in the upper portion of the estuary (Table 6).
The larval densities from the diel data supported the hypothesis that larvae migrate up the estuary as they grow. The size distribution of larvae collected at the mouth of the estuary was compared to those collected in the upper portion of the estuary. Densities of 2- to 3-mm TL larvae were greater ( $P<0.05$ ) in the lower segment (Table 7), whereas densities of each of the $4-\mathrm{mm}$ size classes between 8 to 25 mm were greater upriver. These differences were the same for day and night collections. There were no significant differences in densities of larvae 4 to 7 mm .
Viewing the distribution by $3-\mathrm{mm}$ size classes, there was a single large mode for the smallest sized larvae at the

Table 5
Mean densities (Mean log (density +1 ) per $100 \mathrm{~m}^{3}$ ) of bay anchovy eggs and newly hatched larvae in the Patuxent River.

| Week | Upper | Section |  |
| :---: | :---: | :---: | :---: |
|  |  | Middle | Lower |
| Eggs |  |  |  |
| 6/29/87 | 2.7 | 6.9 | 6.8 |
| 7/06/87 | 2.6 | 6.7 | 6.9 |
| 7/13/87 | 1.7 | 6.5 | 7.2 |
| 7/20/87 | 3.0 | 5.8 | 6.6 |
| 7/27/87 | 4.3 | 4.3 | 6.2 |
| 8/03/87 | 5.5 | 3.1 | 4.3 |
| Larvae (2 to 3 mm TL ) |  |  |  |
| 6/29/87 | 2.1 | 1.5 | 3.8 |
| 7/06/87 | 1.1 | 2.8 | 5.7 |
| 7/13/87 | 2.6 | 4.0 | 4.8 |
| 7/20/87 | 3.3 | 3.8 | 5.1 |
| 7/27/87 | 2.6 | 0.6 | 3.6 |
| 8/03/87 | 3.5 | 0.0 | 0.4 |

mouth of the estuary (Fig. 2); whereas upriver there were two modal lengths, one of small larvae associated with the late season spawning and another at $20-\mathrm{mm}$ TL.

This trend was more pronounced in the standing crop estimates based on data from the regional study. A plot of mean standing crops for bay anchovy larvae versus length for the three sections of the Patuxent River (Fig. 3 ) showed that the standing crop of small larvae found in the lower section of the estuary was several orders of magnitude greater than the number found in the middle and upper sections. This high standing crop of small larvae in the lower section resulted from high densities com-

Table 6
Estimated standing crops (millions) of bay anchovy eggs and newly hatched larvae in the Patuxent River.

|  |  | Section |  |
| :--- | :---: | :---: | :---: |
| Week | Upper | Middle | Lower |
| Eggs |  |  |  |
| 6/29/87 | 371 | 3263 | 5470 |
| $7 / 06 / 87$ | 15 | 2742 | 7384 |
| $7 / 13 / 87$ | 60 | 4800 | 6371 |
| $7 / 20 / 87$ | 89 | 2889 | 8378 |
| $7 / 27 / 87$ | 620 | 589 | 2323 |
| $8 / 03 / 87$ | 328 | 124 | 469 |
|  |  |  |  |
| Larvae (2 to 3 mm TL) | 7 | 29 |  |
| $6 / 29 / 87$ | 4 | 241 | 619 |
| $7 / 06 / 87$ | 8 | 533 | 2076 |
| $7 / 13 / 87$ | 39 | 220 | 753 |
| $7 / 20 / 87$ | 16 | 5 | 2530 |
| $7 / 27 / 87$ | 14 | 0 | 313 |
| $8 / 03 / 87$ |  |  | 5 |

## Table 7

Mean diel densities (Mean log (density +1 ) per $100 \mathrm{~m}^{3}$ ) of bay anchovy larvae in the Patuxent River at upriver stations versus downriver stations averaged over depth layers and over the sampling season.

| Total length <br> $(\mathrm{mm})$ | Day |  |  | Night |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | Upriver | Downriver |  | Upriver | Downriver |
|  | 2.06 | 3.76 |  | 1.99 | 2.60 |
| $4-7$ | 2.68 | 3.07 |  | 2.74 | 2.72 |
| $8-11$ | 2.75 | 1.67 |  | 2.56 | 1.84 |
| $12-15$ | 2.00 | 0.38 |  | 2.18 | 0.60 |
| $16-19$ | 1.37 | 0.04 |  | 2.05 | 0.24 |
| $20-23$ | 1.36 | 0.02 |  | 2.19 | 0.36 |

bined with the large volume of that portion of the estuary (Tables 1 and 8 ). Numbers of large larvae, on the other hand, were greatest in the middle section of the estuary.
An evaluation of the vertical distribution of larvae indicated that at the mouth of the river larvae of lengths less than 12-mm TL remain in deep water during day and night ( $P<0.05$ ) (Table 9). Larvae in each length class greater than $11-\mathrm{mm}$ TL showed no significant vertical patterns downriver. In contrast, in segments 10 to 14 , during the day larvae in each length class greater than 8 mm were near the surface $(P<0.05)$ layer while at night larvae in each length class greater than 11 mm were near the bottom layer ( $P<0.05$ ). The smaller larvae did not exhibit vertical patterns in the upper river.


Figure 2
Comparison of length-density distributions of bay anchovy larvae at the lower (L) and upper (U) part of the river.


Figure 3
Mean standing crop estimates for bay anchovy larvae for three portions of the Patuxent River estuary in 1987.

## Mortality

The parameter estimates for a range of growth rates are:

|  | $3 \mathrm{~mm} /$ week | $4 \mathrm{~mm} /$ week |
| :---: | :---: | :---: |
| $\alpha$ | 1.7631 | $\alpha=3.4505$ |
| $\beta$ | 1.0375 | $\beta=1.3320$ |
|  | $=-37.5911$ | $c=0.7849$. |

Table 8
Average densities per $100 \mathrm{~m}^{3}$ of bay anchovy larvae ( $>3 \mathrm{~mm} \mathrm{TL}$ ) in the Patuxent River and the adjacent portion of Chesapeake Bay. NS $=$ Not sampled.

|  | Segment ${ }^{\text {a }}$ |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Week | 4-7 | 8-11 | 12-13 | 14-17 | 18-21 | 22-25 | 26-29 | 30-33 | 34-35 | 35 | Bay |
| Larvae (4 to 9 mm TL ) |  |  |  |  |  |  |  |  |  |  |  |
| 6/01/87 | 0 | 10 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | NS | NS |
| 6/08/87 | 0 | 10 | 10 | 0 | 0 | 0 | 0 | 0 | 0 | NS | NS |
| 6/15/87 | 0 | 10 | 0 | 0 | 10 | 0 | 10 | 0 | 20 | 10 | NS |
| 6/22/87 | 20 | 0 | 0 | 0 | 10 | 30 | 20 | 30 | 40 | NS | NS |
| 6/29/87 | 40 | 30 | 110 | 20 | 30 | 20 | 30 | 130 | 300 | 210 | 720 |
| 7/06/87 | 40 | 40 | 40 | 20 | 10 | 50 | 80 | 250 | 590 | NS | 250 |
| 7/13/87 | 50 | 30 | 20 | 50 | 60 | 200 | 100 | 170 | 520 | 570 | 1450 |
| 7/20/87 | 50 | 220 | 70 | 150 | 90 | 130 | 150 | 270 | 480 | NS | 520 |
| 7/27/87 | 520 | 360 | 70 | 40 | 10 | 10 | 20 | 50 | 170 | 120 | 90 |
| 8/03/87 | 120 | 280 | 20 | 10 | 0 | 0 | 30 | 30 | 40 | NS | 30 |
| 8/10/87 | 90 | 50 | 10 | 0 | 0 | 0 | 10 | 10 | 0 | 0 | 0 |
| 8/17/87 | 170 | 330 | 110 | 10 | 0 | 0 | 0 | 0 | 20 | NS | 0 |
| Larvae ( $>9 \mathrm{~mm} \mathrm{TL}$ ) |  |  |  |  |  |  |  |  |  |  |  |
| 6/01/87 | 0 | 10 | 0 | 0 | 10 | 0 | 0 | 0 | 0 | NS | NS |
| 6/08/87 | 20 | 10 | 20 | 0 | 0 | 0 | 0 | 0 | 0 | NS | NS |
| 6/15/87 | 0 | 10 | 0 | 10 | 0 | 0 | 0 | 0 | 0 | 0 | NS |
| 6/22/87 | 20 | 10 | 20 | 20 | 0 | 20 | 0 | 0 | 10 | NS | NS |
| 6/29/87 | 10 | 30 | 70 | 150 | 20 | 10 | 0 | 10 | 0 | 0 | 150 |
| 7/06/87 | 50 | 130 | 280 | 170 | 0 | 20 | 20 | 20 | 40 | NS | 20 |
| 7/13/87 | 60 | 140 | 80 | 960 | 70 | 80 | 10 | 30 | 0 | 10 | 60 |
| 7/20/87 | 50 | 60 | 190 | 420 | 60 | 40 | 80 | 50 | 40 | NS | 50 |
| 7/27/87 | 130 | 130 | 220 | 220 | 110 | 70 | 20 | 30 | 50 | 40 | 20 |
| 8/03/87 | 100 | 160 | 160 | 480 | 270 | 100 | 110 | 20 | 60 | NS | 40 |
| 8/10/87 | 90 | 220 | 110 | 150 | 120 | 30 | 20 | 10 | 20 | 50 | 0 |
| 8/17/87 | 170 | 240 | 150 | 280 | 80 | 40 | 60 | 50 | 0 | NS | 0 |

${ }^{a}$ Estimates for segment 35 are from the diel study (averaged over day and night periods). Estimates for other segments are from the regional study and have been adjusted for sampling efficiency.

Table 9
Vertical diel distributions (Mean log [density +1] per $100 \mathrm{~m}^{3}$ ) of bay anchovy larvae in two portions of the Patuxent River averaged over the sampling season.

| Length (mm) | Diel period | Depth zone (m) |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Upriver portion of Patuxent River |  |  | Mouth of Patuxent River |  |  |
|  |  | 0-1 | 1-2 | $>2$ | 0-1 | 1-3 | $>3$ |
| 2-3 | Day | 1.66 | 2.21 | 2.40 | 2.88 | 3.36 | 4.61 |
|  | Night | 1.84 | 2.03 | 2.14 | 1.97 | 2.31 | 3.23 |
| 4-7 | Day | 2.68 | 2.85 | 2.44 | 2.57 | 2.62 | 3.71 |
|  | Night | 2.58 | 2.68 | 2.03 | 2.20 | 2.29 | 3.36 |
| 8-11 | Day | 3.13 | 2.73 | 2.26 | 1.23 | 1.46 | 2.10 |
|  | Night | 2.37 | 2.70 | 2.61 | 1.20 | 1.62 | 2.44 |
| 12-15 | Day | 2.51 | 1.97 | 1.34 | 0.32 | 0.31 | 0.47 |
|  | Night | 1.65 | 2.40 | 2.60 | 0.26 | 0.61 | 0.83 |
| 16-19 | Day | 2.46 | 1.00 | 0.35 | 0.00 | 0.07 | 0.04 |
|  | Night | 0.96 | 2.53 | 2.85 | 0.17 | 0.51 | 0.10 |
| 20-23 | Day | 2.21 | 1.23 | 0.38 | 0.06 | 0.02 | 0.00 |
|  | Night | 1.19 | 2.52 | 3.08 | 0.23 | 0.37 | 0.44 |



Figure 4
Mortality curve with respect to length for bay anchovy using 1987 data censored according to $3 \mathrm{~mm} /$ week growth and the observed abundances from which the curve was estimated.

The fit of the curve assuming 4 mm growth per week was better than one assuming 3 mm growth per week (Figs. 4 and 5).

Estimated mortality rates per individual $\left(\alpha L^{-\beta}\right)$ based on these curves ranged from 0.161 to 0.162 per mm of growth at 10 mm TL and from 0.064 to 0.079 per mm of growth at 20 mm TL. These rates correspond to $15 \%$ per mm at 10 mm TL and $6.2 \%$ and $7.6 \%$ per mm at 20 mm TL.

## Discussion

## Sampling Considerations

Information from the random stratified sampling design implemented in this study was used to address the hypothesis that bay anchovy eggs are spawned downriver and the young larvae move upriver to a nursery area. To analyze changes in abundance, vertical, longitudinal, and temporal stratification was required. Randomly selecting sampling locations within strata satisfies the underlying assumptions of hypothesis testing through ANOVA.

The importance of sampling a broad spectrum of strata can not be overemphasized. For example, the geographic area considered by Polgar et al. (1988) excluded the lower section of the Patuxent River estuary. Their estimate of the mortality rate for postlarval bay anchovies in the Patuxent River in 1978 ( 0.016 per tidal cycle or 0.032 per day) was considerably less than our range of estimates. Our data ('lower estuary'' in Figure 3) show that excluding data from this section of the estuary leads to an underestimate of mortality.


Figure 5
Mortality curve with respect to length for bay anchovy using 1987 data censored according to $4 \mathrm{~mm} /$ week growth and the observed abundances from which the curve was estimated.

Furthermore, when making population assessments, it is important to consider the volumes of the river segments. Early studies in the Patuxent River estuary (Dovel 1971) considered only density data. Note that the density of large larvae upriver seems relatively high, but the volumes of upriver segments are relatively small. If one considered larval density only, larval mortality would be underestimated and the extent of upriver migration would be overestimated.

## Distribution and Migration

The longitudinal distribution of the eggs indicates that spawning occurs primarily in the lower section of the river. Collections in the Patuxent River in 1978 by Setzler et al. (1979) did not extend below segment 20, but the distribution pattern for eggs is consistent with the data for 1987 in that the highest densities were found downriver. The late season spawning observed upriver in 1987 may have been partially associated with an increase in salinity. Salinity in the upper river segments increased approximately 2 ppt during the period 20 July to 3 August (Table 4).

From the pattern of many small larvae and few large larvae downriver opposed to relatively more large larvae and few small larvae upriver (Fig. 2) we infer either that the larvae hatch downstream and migrate upriver or that the larvae suffer much greater mortality downriver than upriver. However, the differential mortality argument is not sufficient to explain the data. For any cohort abundance is a monotonically decreasing function of length or age. Thus the sum or average of cohort abundances also has this property. For the upriver segment, the number
of larvae with lengths 17 to 23 mm TL averaged over the spawning season (i.e. averaged over cohorts) actually exceeded the number of larvae with lengths 8 to 14 mm . Since it is impossible to explain this phenomenon solely by differential mortality, one must conclude that the increase in density of 17 to 23 mm larvae upriver resulted wholly or partially from migration of larvae from downriver.

This same issue is examined in terms of avoidance corrected abundance (Fig. 3) with data from three sections of the river. It is clear that there were more larvae in the middle section (segments 14 to 25) in the 15 to 21 mm length category than can be explained without invoking migration from the lower section (segments 26 to 35). Small larvae originate in the lower third of the estuary and move to the middle section of the estuary as they grow. Few larvae enter the upper section (segments 4 to 13) of the river.
While the data indicate that larval migration occurred, the mechanism for the migration is not clear. Many studies have found that larvae use water currents to move to nursery areas (Norcross and Shaw 1984; Weinstein 1988). Current velocity estimates for tidally averaged water flow in the Patuxent River indicate upriver flow in deeper water (ANSP 1983; Edinger et al. 1989). Thus the vertical position of larvae will determine the direction of transport based on the net direction of flow at that vertical position. Since small larvae downriver remain in lower depths, they could be transported to their nursery areas upriver with baroclinic circulation.

## Mortality

We use a model based on the simple but appealing assumption that larval mortality rate decreases with increasing length. Swimming speed and thus ability to avoid predators is a direct function of length rather than age (Fuiman and Webb 1988) The mortality rate is believed to be greatest for eggs and very small larvae because they are unable to avoid predation. Larger larvae should be better able to obtain food and avoid predators and their survival should be greater. Other assumptions of this application need to be examined.

Use of length as a surrogate for age is justified because of the close correspondence between length and age (Fives et al. 1986; Leak and Houde 1987). The logic is that if survival $=g$ (time) and length $=f($ time $)$ then survival $=$ $g\left(f^{-1}\right.$ (length)). For convenience the plot of abundance versus length is referred to as a "survival curve" or a "mortality curve."

Immigration/emigration of bay anchovy larvae $\geqslant 10 \mathrm{~mm}$ TL would not be expected to be a major source of error in mortality estimates because densities at the mouth of the river are relatively low compared to densities in the middle portion of the study area (Table 8). Our data suggest that juvenile bay anchovies ( $\geqslant 23 \mathrm{~mm} \mathrm{TL}$ ) move down river, but these were not included in our mortality analysis.

In contrast, one would expect that small larvae would be transported into the river because of high densities in deeper layers at the mouth of the river. This would cause an underestimation of mortality for these small larvae. An evaluation of the emigration of small larvae from Chesapeake Bay requires further study.

Based on published results, we have assumed a growth rate of 3 to 4 mm per week (Table 3). With cohorts defined by using length classes approximately equivalent to one weeks growth, weekly sampling is sufficient to represent all cohorts of the spawning season.

This model was motivated by finding that the commonly used constant mortality rate model (Houde 1987; Leak and Houde 1987) was inadequate for these data. Residual plots for our data indicated that a constant rate model underestimated mortality for small larvae and overestimated mortality for large larvae. The Weibull mortality model can be used for modeling increasing or decreasing mortality rate curves (Pinder et al. 1978). The Weibull model has been employed for modeling a decreasing mortality rate in a larval fish population (Saila and Lough 1981). However, use of the Weibull model seems to be founded on empirical curve fitting rather than biological assumptions. We prefer a model founded on the commonly observed phenomenon that mortality decreases with increasing length (Ware 1975; Hackney 1977; Saila and Lough 1981).

As noted, we may have underestimated mortality of larvae $<10 \mathrm{~mm}$ TL because the immigration of these small larvae into the river from the Chesapeake Bay was not taken into account. Thus the difference in mortality rates between large and small larvae may have been even larger than we estimated. If this is the case, there is even greater need for a decreasing mortality model.

The Pareto model is also applicable when ages of larvae are known. Lo (1985) used an age-based Pareto model for eggs and larvae of northern anchovy. The length-based Pareto model presented here can also be used. One need only substitute the growth equation for length in the denominator of the first differential equation and perform the calculus to obtain a model based on age which uses the same assumptions. If the growth equation is assumed linear, a simple substitution of the growth equation in the final model yields

$$
\begin{equation*}
\log _{\mathrm{e}}\left(N_{t}\right)=\frac{-\alpha}{1-\beta}(h+g t)^{(1-\beta)}+c . \tag{7}
\end{equation*}
$$

The expression $(h+g t)$ is a linear model for length as a function of age, where $t=$ age
$h=$ hatching size
$g=$ growth rate.
This equation indicates that the number of surviving larvae at age $t$ increases as hatching size and growth rate increase. The importance of hatching size and growth rate


Figure 6
Constant mortality rates published by Leak and Houde (1987) compared to decreasing mortality rates published here. The two solid curves show the range of values reported by Leak and Houde. The two dashed lines show the range of values that result from using 3 and $4 \mathrm{~mm} /$ week growth rates in this study.


Figure 7
Survival curves with respect to age for bay anchovy larvae estimated here compared to those computed from estimates by Leak and Houde (1987). The two solid curves show the range of values reported by Leak and Houde. The two dashed lines show the range of values that result from using 3 and $4 \mathrm{~mm} /$ week growth in this study.
in the evaluation of mortality has been emphasized by Ware (1975) and Houde (1987), respectively.

The mortality rate derived here is compared to a constant mortality rate estimated for larval bay anchovies in

Biscayne Bay by Leak and Houde (1987). The Pareto function is transformed to a function of age by compositing the mortality with respect to length function with a length with respect to age function. Two growth rates for the Patuxent River, $3 \mathrm{~mm} /$ week and $4 \mathrm{~mm} /$ week, are used for comparison. Mortality rates for the Pareto model are greater than the constant estimates for larvae less than two days old and become less than the constant mortality rates for larvae greater than five days old (Fig. 6). As a result the net mortalities estimated by the Pareto model for the Patuxent River are much less by the time larvae are eight days old (Fig. 7).

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## Citations

Academy of Natural Sciences, Phladelphia (ANSP).
1983. Hydrographic characteristics of the Patuxent estuary. In Chalk Point Station 316 Demonstration - Technical Reports, Vol. 1, p. 69-124. Acad. Nat. Sci., Philadelphia, PA.
Arnold, B.C.
1985. Pareto distribution. In Encyclopedia of statistical sciences, Vol. 6 (S. Kotz and N.L. Johnson, eds.), p. 569-574. John Wiley \& Sons, NY.
Chao, L.N., and J.A. Musick.
1977. Life history, feeding habits, and functional morphology of juvenile sciaenid fishes in the York River estuary, Virginia. Fish. Bull., U.S. 75(4):657-701.
Cowan, J.H. Jr., and E.D. Houde.
1989. Growth, growth variability, and survival of bay anchovy (Anchoa mitchilli) larvae in mesocosm enclosures. Int. Cons. Explor. Sea C.M. 1989/L: 3 Ref H, 11 p.
Castro, L.R., and R.K. Cowen.
In press. Growth rates of bay anchovy (Anchoa mitchilli) in Great South Bay under recurrent brown tide conditions, summers 1987 and 1988. In Novel phytoplankton blooms: causes and impacts of recurrent brown tides and other unusual blooms (E.M. Cosper, E.J. Carpenter, and M.V. Bricelj, eds.) Lecture Notes on Coastal and Estuarine Studies, Springer-Verlag, Berlin.
Cronin, W.B., and D.W. Pritchard.
1975. Additional statistics on the dimensions of the Chesapeake Bay and its tributaries: cross-section widths and segment volumes per meter depth. Chesapeake Bay Institute, Ref. 75-3. Johns Hopkins Univ., Baltimore, MD.
Dovel, W.L.
1971. Fish eggs and larvae of the upper Chesapeake Bay. Natural Resources Institute Univ. Maryland Contrib. No. 460, 71 p.
Draper, N.R., and H. Smith Jr.
1981. Applied regression analysis, 2nd ed. John Wiley \& Sons, NY, 709 p.

## Edinger, J.E., E.M. Buchak, and N.C.L. Huang.

1989. Chalk Point Steam Electric Station Patuxent estuary hydrodynamic and transport model verification and real time intake entrainment rates for 1985 sampling. J.E. Edinger and Assoc., Wayne, PA, 22 p.
Ferraro, S.P.
1980a. Pelagic fish eggs and larvae of the Peconic Bays, New York: 1972-1974. Ph.D. thesis, State Univ. of New York at Stony Brook.
1980b. Daily time of spawning of 12 fishes in the Peconic Bays, New York. Fish. Bull., U.S. 78(2):455-464.
Fives, J.M., S.M. Warlen, and D.E. Hoss.
1990. Aging and growth of larval bay anchovy, Anchoa mitchilli, from the Newport River estuary, North Carolina. Estuaries 9(4B): 362-367.
Fuiman, L.A., and P.W. Webb.
1991. Ontogeny or routine swimming activity and performance in zebra danios (Teleostei: Cyprinidae). Anim. Behav. 36:250-261.
Hackney, P.A.
1992. Methods for calculating survival rate, biomass production, growth rate and assessing entrainment of lacustrine ichthyoplankton. In Proceedings of the first symposium on freshwater larval fish (L.L. Olmsted, ed.), p. 212-249. Hosted by Duke Power Co., Charlotte, NC.
Hildebrand, S.F.
1993. Family Engraulidae. In Fishes of the western North Atlantic, Part 3 (H.B. Bigelow, ed.), p. 152-249. Sears Found. Mar. Res. No. 1.
Hildebrand, S.F., and W.C. Schroeder.
1994. Fishes of the Chesapeake Bay. Fish. Bull., U.S. 53:1-366.

Hollis, E.H.
1952. Variation in the feeding habits of the striped bass, Roccus saxatilis (Walbaum). Bull. Bingham Oceanogr. Collect., Yale Univ. 14:(1):111-131.
Houde, E.D.
1987. Fish early life dynamics and recruitment variability. Am. Fish. Soc. Symp. 2:17-29.
Houde, E.D., and J.A. Lovdal.
1984. Seasonality of occurrence, foods and food preferences of ichthyoplankton in Biscayne Bay, Florida. Estuarine, Coastal and Shelf Sci. 18:403-419.
Kuntz, A.
1914. The embryology and larval development of Bairdiella chrysura and Anchoa mitchilli. Fish. Bull., U.S. 35:87-134.
Leak, J.C., and E.D. Houde.
1987. Cohort growth and survival of bay anchovy, Anchoa mitchilli, in Biscayne Bay, Florida. Mar. Ecol. Prog. Ser. 37(2-3):109-122.
Lo, N.C.
1985. Modeling life-stage-specific instantaneous mortality rates, an application to nothern anchovy, Engraulis mordax, eggs and larvae. Fish. Bull., U.S. 84(2):395-416.

Merriner, J.V.
1975. Food habits of the weakfish, Cynoscion regalis, in North Carolina waters. Chesapeake Sci. 16(1):74-76.
Miller, R.G.
1981. Survival analysis. John Wiley \& Sons, N.Y, 238 p.

Norcross, B.L., and R.F. Shaw.
1984. Oceanic and estuarine transport of fish eggs and larvae: a review. Trans. Am. Fish. Soc. 113:153-167.
Olney, J.E.
1983. Eggs and larvae of the bay anchovy, Anchoa mitchilli, and the weakfish, Cynoscion regalis, in lower Chesapeake Bay with notes on associated ichthyoplankton. Estuaries 6(1):20-35.
Pinder, J.E., J.G. Wiener, M.H. Smith.
1978. The Weibull distribution: a new way of summarizing survivorship data. Ecology 59:175-179.
Polgar, T.T., M.A. Turner, and J.K. Summers.
1988. Effect of power plant entrainment on the population dynamics of the bay anchovy (Anchoa mitchilli). Ecol. Modelling 41(3/4): 201-218.
Saila, S.B., and R.G. Lough.
1981. Mortality and growth estimation from size data - an application to some Atlantic herring larvae. Rapp. P.-v. Reun. Cons. Int. Explor. Mer 178:7-14.
Saksena, V.P., and E.H. Houde.
1972. Effect of food level on the growth and survival of laboratory reared larvae of bay anchovy (Anchoa mitchilli Valenciennes) and scaled sardine (Harengula pensacolae Goode \& Bean). J. Exp. Mar. Biol. Ecol. 8:249-258.
SAS Institute.
1985. SAS user's guide: statistics. Version 5 edition. SAS Institute, Cary, NC, 956 p.
Setzler, E.M., K.V. Wood, D. Shelton, G. Drewry, and J.A. Mihursky. 1979. Chalk Point Steam Electric Studies. Patuxent Estuary Studies. Ichthyoplankton Population Studies. 1978 Data Report. Univ. Maryland, Chesapeake Biol. Lab. Ref. No. UMCEES 79-20-CBL, 111 p .
Tukey, J.W.
1962. The future of data analysis. Annals of mathematical statistics. Ann. Math. Stat. 33:1-67.
Wang, J.C.S., and R.J. Kernehan.
1979. Fishes of the Delaware estuaries. Ecological Analysts, Inc., Towson, MD, 410 p.
Ware, D.M.
1975. Relationship between egg size, growth and natural mortality of larval fish. J. Fish. Res. Board Can. 32:2503-2512.
Weinstein, M.P., ed.
1988. Fish and shellfish transport through inlets. Am. Fish. Soc. Symp. 3, 165 p.
Vouglitois, J.J., K.W. Able, R.J. Kurtz, and K.A. Tighe.
1987. Life history and population dynamics of the bay anchovy in New Jersey. Trans. Am. Fish. Soc. 116(2):141-153.

# Hypothetical Northern Spawning Limit and Larval Transport of Spot* 

BRENDA L. NORCROSS** and DEBORAH A. BODOLUS<br>College of William and Mary<br>School of Marine Science<br>Virginia Institute of Marine Science<br>Gloucester Point, VA 23062


#### Abstract

The exact northern limit of the spawning grounds of spot (Leiostomus xanthurus) has not been determined. Previous reports of spot spawning during the winter/spring in the Middle Atlantic Bight (MAB) are refuted based on the presence of low bottom-water temperatures at that time. Analyses of historic bottom isotherms in the MAB during winter/spring show that the most northerly occurrence of required $17^{\circ} \mathrm{C}$ bottom temperatures from December to May is on the outer continental shelf off North Carolina near the Gulf Stream. It is therefore suggested that spot recruiting to Chesapeake Bay are spawned near Cape Hatteras at the shelf break in winter. A mechanism must then exist to transport larvae approximately 250 km northward in the MAB before recruiting to Chesapeake Bay one to three months later. Episodes of southerly winds, interspersed between seasonal spring northerly winds, were identified from 1978 through 1988. These southerly winds could cause current reversals over the inner shelf areas of the MAB and may transport waves of new spot recruits northward from the southern spawning site. These winds may have been related to waves of juvenile spot recruiting to Virginia estuaries in 1987 and 1988. The duration and frequency of southerly wind events during the months when larvae are being transported could have a major impact on year-class success.


## Introduction

Spot (Leiostomus xanthurus) is a common sciaenid along the middle and southern Atlantic and Gulf coasts of the United States. They appear to have a protracted spawning season which probably begins in September and ends in April or May with reported times varying with location (Hildebrand and Cable 1930; Dawson 1958; Lewis and Judy 1983). It has been suggested that spawning occurs off Chesapeake Bay from late fall through early spring (Welsh and Breder 1923; Hildebrand and Schroeder 1928; Lippson and Moran 1974). The season extends from October through March off the coasts of North and South Carolina (Dawson 1958) with peak spawning occurring in December and January (Hildebrand and Cable 1930; Dawson 1958; Berrien et al. 1978; Lewis and Judy 1983; Warlen and Chester 1985).

In fall, spot are generally distributed in the Middle Atlantic Bight (MAB) nearshore from Long Island, NY

[^4]to Cape Lookout, NC (NMFS/NEFC 1983-87). In the winter, from Cape Hatteras to central Florida, spot may move offshore to occupy the shelf edge (Manooch and Raver 1984). Although spot have been recorded from depths of 165 m (Squire 1958) and 204 m (Springer and Bullis 1956), they are usually found in depths less than 100 m (Pearson 1932; Dawson 1958). Spot tagged in Chesapeake Bay (Pacheco 1962) and Delaware Bay (Pearson 1932) were returned by trawlers operating south of Cape Hatteras, NC in winter. This suggests that spot from these northern areas may have a common coastal feeding or spawning ground during winter (Chao and Musick 1977). There are so few spot captured in the MAB in the spring that the species is not reported by groundfish cruises (NMFS/NEFC 1984-88) undertaken in March, lending further support for a common winter ground offshore or south of Cape Hatteras.

Since spawning has not been observed, many conclusions about spawning time and place are based on collections of larvae and juveniles (Hildebrand and Cable 1930; Lewis and Wilkens 1971; Berrien et al. 1978; Lewis and Judy 1983). Spawning probably occurs on the mid to outer continental shelf (Warlen and Chester 1985) over deeper

Table 1
Literature records of initial spot recruitment dates and sizes in Chesapeake Bay.

| Spot reference | Place | Date | Size ( mm ) |
| :---: | :---: | :---: | :---: |
| Welsh and Breder 1923 | Chesapeake Bay | Jan.-April | 19-37 |
| Hildebrand and Schroeder 1928 | Chesapeake Bay | 1st wk March | 15-19 |
| Pacheco 1957 | York River | April | 15-22 |
| Pacheco 1962 | York River | April/May | 22-85 |
| Richards and Castagna 1970 | eastern shore | May-Sept. | 21-99 |
| Chao and Musick 1977 | York River | Early April | 14-45 |
| Schauss 1977 | Lynnhaven Bay | April | - |
| Orth and Heck 1980 | Chesapeake Bay | March | - |
| Olney 1983 | Chesapeake Bay | Feb./March | 11-20 |
| Weinstein and Brooks 1983 | Chesapeake Bay | April | 20 |
| Heck and Thoman 1984 | York River | April | - |
| McCambridge and Alden 1984 | James River | April | 16-22 |
| Smith et al. 1984 | York River | April | - |
| Cowan and Birdsong 1985 | Wachapreague | April/May | 15-33 |
| O'Neill and Weinstein 1987 | York River | April | - |
| Olney and Boehlert 1988 | Chesapeake Bay | April/May | - |
| Seigfried 1989 | York River | April | 14-19 |

bottoms (Dawson 1958). Off North Carolina, the size and age of spot larvae increased from offshore toward the coastline (Lewis and Judy 1983; Warlen and Chester 1985). Primarily, time of spawning has been inferred from time of juvenile recruitment to the estuaries. Juveniles recruit to Georgia estuaries from February through April (Rogers et al. 1984) and to South Carolina from January through June (Beckman and Dean 1984; McGovern 1986), peaking in February (McGovern 1986). Recruitment to North Carolina may occur October through March, but appears to be concentrated from January to March (Tagatz and Dudly 1961; Williams and Deubler 1968; Lewis and Mann 1971; Turner and Johnson 1973; Lewis and Judy 1983). Spot are approximately two months of age when entering estuaries in North Carolina (Warlen and Chester 1985) and South Carolina (Beckman and Dean 1984). Recruitment to Chesapeake Bay usually occurs in April and May (Table 1) but has been reported as early as January (Welsh and Breder 1923), February (Olney 1983), or March (Hildebrand and Schroeder 1928; Orth and Heck 1980). Variation in time of juvenile recruitment indicates a change in time or place of spawning or in mechanisms responsible for transport of larvae to inshore nurseries.

These observations indicate that spot spawn over the mid to outer shelf, inshore of the Gulf Stream, from at least North Carolina southward. However, since spot recruit to Chesapeake Bay as post-larvae or juveniles in the spring, it has been concluded that spawning also takes place as far north as the Chesapeake area of the MAB (Welsh and Breder 1923; Hildebrand and Schroeder 1928; Johnson 1978; Lippson and Moran 1984). Mechanisms responsible for transport of larvae from the spawning ground to the nursery have not been investigated. It is the intention of
this paper to analyze historic and recent data to formulate hypotheses which exclude the MAB (Fig. 1) as a spawning location and propose theoretical transport pathways from a southerly spawning site to Chesapeake Bay. Additionally, we will suggest direction for future investigations to test these hypotheses.

## Methods

Bottom isotherms were obtained from the National Marine Fisheries Service (NMFS)/Northeast Fisheries Center (NEFC) bottom-trawl survey cruise publications (Edwards et al. 1962; Davis 1979; Nickerson and Wright 1980; NMFS/NEFC 1983-1988) or as unpublished plots (L. Lierheimer, NMFS/NEFC, Woods Hole, MA 02543, December 1987). Bottom temperatures ( ${ }^{\circ} \mathrm{C}$ ) from cruises 8701 (February-March 1987), 8801 (February 1988), and 8901 (February-March 1989) were collected aboard the NOAA ship Ferrel by the authors. Because laboratory studies of spot show spawning to take place between $17.5^{\circ} \mathrm{C}$ and $25.0^{\circ} \mathrm{C}$ (Hettler and Powell 1981), the $17^{\circ} \mathrm{C}$ isotherm was selected as an index of spawning temperature. Bottom water temperatures were examined between $39^{\circ} \mathrm{N}$ and $33.5^{\circ} \mathrm{N}$ latitude (Fig. 2A) from November through May, 1965-1989. Data were aggregated seasonally based on timing of most collections and applicability to spot life history: November-December, January-February and March-May (Table 2).

Wind data from Norfolk International Airport, Virginia has previously been validated for application in the MAB (Norcross and Austin 1988). Daily resultant wind speed and direction for 1978-1988 (National Weather Service,


Figure 1
The area of study on the east coast of the United States including the Middle Atlantic Bight and Chesapeake Bay. Note: southern extent of bottom temperatures collected on NMFS cruises (Table 2): Chincoteague Island ( $38^{\circ} \mathrm{N}$ ), Chesapeake Bay mouth $\left(37^{\circ} \mathrm{N}\right)$, Oregon Inlet $\left(36^{\circ} \mathrm{N}\right)$, and Cape Hatteras $\left(35^{\circ} \mathrm{N}\right)$. Sites of collections of juvenile spot on the eastern shore of Virginia, (Wachapreague Inlet, Sand Shoal Inlet, and Fisherman's Island), and within Chesapeake Bay, (Occohannock Creek, York River, James River, and Lynnhaven Bay). Norfolk International Airport, site of recording of wind data.

1978-88) were processed by smoothing with seven-day running averages which corresponded to the physical time range applicable for transport in the South Atlantic Bight (Yoder 1983). The daily resultant winds were analyzed for the time period from November through June, the time of spawning and recruitment of spot to Chesapeake Bay. The total number of southerly wind events (i.e., the number of wind vectors having a northward component) and the number of southerly events lasting four consecutive days were compared for different years and months.

Ichthyoplankton samples were collected near the shore to 200 m offshore in the southern MAB as far north as New

Jersey and as far south as Cape Lookout, NC. Larvae were collected by stepped oblique tows using Gulf-V samplers aboard the RV Dolphin in 1965 and 1966 (Berrien et al. 1978). Later collections of spot larvae (1977-1986), available from NMFS/NEFC cruises (M. Fahay, NMFS/ NEFC, Highlands, NJ 07732, pers. commun. Nov. 1988), used double oblique bongo-net tows following standard MARMAP guidelines (Jossi and Marak 1983). These sampling techniques do not yield information on the vertical distribution of larvae. The horizontal distribution of larval spot was plotted for November-December, January-February, March, April, and May by combining data from all cruises. The mean length of larvae was calculated for each station. Station locations from each cruise did not overlap since those from the RV Dolphin were located on transects perpendicular to the coastline and those from the NMFS/NEFC cruises were selected by stratified random sampling techniques.

As part of a larger study, estuarine areas within Chesapeake Bay and locations in lagoons on the seaside of Virginia's eastern shore were sampled twice monthly from September 1986 through August 1987 and monthly from September 1987 through August 1988. Sample locations consisted of Wachapreague Channel and Inlet (five stations) Sand Shoal Channel and Inlet (five stations) and Fisherman's Island (one station) on the seaside of the Virginia eastern shore; Occohannock Creek (four stations) on the bayside of the eastern shore; and the mouth of the York River (eight stations) on the west side of Chesapeake Bay (Fig. 1). Stations were located in channels of creeks and rivers and directly behind the barrier islands, and in shallow areas within the marsh complex. Sites were similar with regard to depth, proximity to houses and creek size. Weather permitting, two samples were taken at all 23 stations on each of 36 trips. A $4.9-\mathrm{m}$ semi-balloon trawl with a codend mesh of 6.4 mm and a $3.2-\mathrm{mm}$ codend liner was used at 18 of the stations. At the other five stations, one at each site, a $6.1-\mathrm{m}$ bag seine with a $3.2-\mathrm{mm}$ mesh was used (Norcross and Hata, in press). All spot captured were preserved in either $10 \%$ formalin or $95 \%$ ethanol and returned to the laboratory where they were measured to the nearest 1 mm increment of total length (TL).

## Results and Discussion

Examination of 48 sets of bottom isotherms- 10 for November-December, 14 for January-February, and 24 for March-May (Table 2)—produced plots of the $17^{\circ} \mathrm{C}$ isotherm by season (Fig. 2A). Water as warm as $17^{\circ} \mathrm{C}$ is present in the MAB during November-December, but the location exhibits interannual variation in position. Rarely is there any water as warm as $17^{\circ} \mathrm{C}$ present in winter. Of the 14 sets of data examined for January-February, there was a very restricted amount of water warmer than $17^{\circ} \mathrm{C}$


Figure 2
The continental shelf area to 200 m (dotted line) in the Middle Atlantic Bight and around Cape Hatteras. (A) Seasonal distribution of the $17^{\circ} \mathrm{C}$ isotherm. Numbers, located on the side of the isotherm greater than $17^{\circ} \mathrm{C}$, indicate the year in which the isotherm was in that position.
Source is data in Table 2. (B) Bottom water temperatures on 1 May generated from average temperatures of 10 years of NMFS data (D. Mountain and T. Holzwarth, NMFS/NEFC, Woods Hole, MA 02543, unpubl. data 1988).

Table 2
Bottom-water temperature data, 1965-1989 (Davis 1979; Nickerson and Wright 1980; NMFS/NEFC 1983-1988; Norcross and Bodolus, unpub. data.) All cruises started at least as far north as New Jersey; southern extent of cruise track listed here.

| Year | Dates | Cruise I.D. \# | Southern extent | Highest temp. |
| :---: | :---: | :---: | :---: | :---: |
| November-December |  |  |  |  |
| 1965 | 3-23 Dec | 6504 | Cape Lookout | $22^{\circ} \mathrm{C}$ |
| 1966 | 9 Nov-4 Dec | 6614 | Cape Lookout | $22^{\circ} \mathrm{C}$ |
| 1971 | 8-19 Nov | 7103 | Cape Lookout | $24^{\circ} \mathrm{C}$ |
| 1978 | Dec | 7807 | Chesapeake Bay | $16^{\circ} \mathrm{C}$ |
| 1980 | 19 Nov-21 Dec | 8012 | Oregon Inlet | $14^{\circ} \mathrm{C}$ |
| 1982 | 15 Nov-22 Dec | 8209 | Chincoteague | $14^{\circ} \mathrm{C}$ |
| 1983 | 14 Nov-21 Dec | 8309 | Cape Henry | $18^{\circ} \mathrm{C}$ |
| 1984 | 29 Nov-7 Dec | 8409 | Cape Hatteras | $24^{\circ} \mathrm{C}$ |
| 1985 | 5 Nov-12 Dec | 8510 | Cape Hatteras | $24^{\circ} \mathrm{C}$ |
| 1986 | 3 Nov-12 Dec | 8610 | Cape Hatteras | $24^{\circ} \mathrm{C}$ |
| January-February |  |  |  |  |
| 1959 | 23 Jan-3 Feb | 126 | Cape Hatteras | $15^{\circ} \mathrm{C}$ |
| 1966 | 26 Jan-9 Feb | 6601 | Cape Lookout | $17^{\circ} \mathrm{C}$ |
| 1978 | Jan-Feb | 7801 | Oregon Inlet | $13^{\circ} \mathrm{C}$ |
| 1978 | 16 Feb-17 Mar | 7802 | Cape Hatteras | $12^{\circ} \mathrm{C}$ |
| 1979 | Jan | 7901 | Chesapeake Bay | $10^{\circ} \mathrm{C}$ |
| 1979 | $23 \mathrm{Feb}-15 \mathrm{Mar}$ | 7903 | Cape Hatteras | $12^{\circ} \mathrm{C}$ |
| 1980 | $3 \mathrm{Jan}-10 \mathrm{Feb}$ | 8001 | Oregon Inlet | $12^{\circ} \mathrm{C}$ |
| 1981 | $17 \mathrm{Feb}-24 \mathrm{Mar}$ | 8101 | Chesapeake Bay | $11^{\circ} \mathrm{C}$ |
| 1984 | 9 Jan-10 Feb | 8401 | Chesapeake Bay | $12^{\circ} \mathrm{C}$ |
| 1985 | 7 Jan-8 Feb | 8501 | Chesapeake Bay | $12^{\circ} \mathrm{C}$ |
| 1986 | 7 Jan-12 Feb | 8601 | Cape Hatteras | $17^{\circ} \mathrm{C}$ |
| 1987 | 5 Jan-13 Feb | 8701 | Cape Hatteras | $16^{\circ} \mathrm{C}$ |
| 1987 | 23 Feb-6 Mar | 8701 | Cape Hatteras | $21^{\circ} \mathrm{C}$ |
| 1988 | 8 Feb-26 Feb | 8801 | Cape Hatteras | $18^{\circ} \mathrm{C}$ |
| 1989 | $13 \mathrm{Feb}-2 \mathrm{Mar}$ | 8901 | Cape Hatteras | $21^{\circ} \mathrm{C}$ |
| March-May |  |  |  |  |
| 1966 | 6-21 Apr | 6603 | Cape Lookout | $21^{\circ} \mathrm{C}$ |
| 1968 | 4 Mar-16 May | 6803 | Cape Hatteras | $10^{\circ} \mathrm{C}$ |
| 1969 | $5 \mathrm{Mar}-10$ Apr | 6902 | Cape Hatteras | $10^{\circ} \mathrm{C}$ |
| 1970 | $12 \mathrm{Mar-29}$ Apr | 7003 | Oregon Inlet | $14^{\circ} \mathrm{C}$ |
| 1971 | 9 Mar-5 May | 7101 | Cape Hatteras | $18^{\circ} \mathrm{C}$ |
| 1972 | 8 Mar-24 Apr | 7202 | Oregon Inlet | $13^{\circ} \mathrm{C}$ |
| 1972 | 20-31 Mar | 7208 | Cape Lookout | $20^{\circ} \mathrm{C}$ |
| 1973 | 16 Mar-15 May | 7303 | Oregon Inlet | $11^{\circ} \mathrm{C}$ |
| 1974 | $12 \mathrm{Mar}-4 \mathrm{May}$ | 7404 | Oregon Inlet | $14^{\circ} \mathrm{C}$ |
| 1975 | 14 Mar-12 May | 7503 | Oregon Inlet | $10^{\circ} \mathrm{C}$ |
| 1976 | 4 Mar-5 May | 7605 | Oregon Inlet | $12^{\circ} \mathrm{C}$ |
| 1977 | 19 Mar-27 Apr | 7703 | Oregon Inlet | $11^{\circ} \mathrm{C}$ |
| 1978 | 20 Mar-25 May | 7804 | Cape Hatteras | $14^{\circ} \mathrm{C}$ |
| 1978 | Apr-May | 7804 | Cape Hatteras | $17^{\circ} \mathrm{C}$ |
| 1979 | 20 Mar-12 May | 7904 | Cape Lookout | $18^{\circ} \mathrm{C}$ |
| 1979 | 6-19 May | 7904 | Cape Hatteras | $20^{\circ} \mathrm{C}$ |
| 1980 | 29 Feb-4 Apr | 8002 | Cape Hatteras | $18^{\circ} \mathrm{C}$ |
| 1980 | 16 Mar-2 May | 8003 | Cape Lookout | $18^{\circ} \mathrm{C}$ |
| 1981 | 17 Mar-14 May | 8100 | Cape Lookout | $17^{\circ} \mathrm{C}$ |
| 1981 | 18 Mar-9 Apr | 8103 | Cape Hatteras | $11^{\circ} \mathrm{C}$ |
| 1981 | 20 May-18 Jun | 8103 | Chincoteague | $12^{\circ} \mathrm{C}$ |
| 1985 | 1 Apr-2 May | 8303 | Cape Hatteras | $14^{\circ} \mathrm{C}$ |
| 1986 | 6 May-7 Jun | 8603 | Cape Hatteras | $19^{\circ} \mathrm{C}$ |
| 1987 | 5 May-8 Jun | 8704 | Oregon Inlet | $11^{\circ} \mathrm{C}$ |

available in only five of the 12 years. While $17^{\circ} \mathrm{C}$ water was available in March-May, it was restricted to near Cape Hatteras (Fig. 2, A and B). There is a convergence zone near Cape Hatteras which causes a dramatic increase in bottom water temperature (Bumpus 1973). This zone forms in the autumn, persists throughout the winter, and breaks down in the spring (Bumpus et al. 1973). Remnants of this convergence zone can be seen in the March-May bottom temperatures (Fig. 2A), with water warmer than $17^{\circ} \mathrm{C}$ present in almost half of the years. Ten years of data taken by NMFS/NEFC cruises show this convergence zone at Cape Hatteras on 1 May (Fig. 2B).

Based on these bottom temperature distributions, we conclude that spot do not spawn in the MAB in winter, rather that they spawn near or south of Cape Hatteras where the water temperature is $17^{\circ} \mathrm{C}$ or higher. Spot could theoretically spawn in the MAB in November and early December when water warmer than $17^{\circ} \mathrm{C}$ is still present. However, since spot do not recruit to Chesapeake Bay until March and April (Table 1), it seems unlikely they they would spawn in the fall as that would require a mechanism to keep the larvae offshore in the MAB for four to six months.

If spot are spawning at the shelf break near Cape Hatteras in winter and spring, as indicated by bottom water temperatures, then a mechanism must exist to transport the larvae northward on the shelf from Cape Hatteras towards Chesapeake Bay. The mean longshore flow in the MAB, however, is southerly (Bumpus 1973). This mean flow decreases from a maximum near the shelf break toward the coast where the shallow inner shelf is prone to wind-driven forces (Beardsley and Boicourt 1981).

Eggs and larvae of spot are pelagic (Powell and Gordy 1980), though their depth distribution is unknown, and thus subject to transport by winds and currents. Year-class strength seems to vary yearly and to be determined by time of recruitment to the estuary (Joseph 1972). These nonperiodic year-to-year fluctuations are thought to be caused by environmental differences that occur on the spawning ground (Joseph 1972). Extended periods of offshoredirected winds during the pelagic phase may cause drastic reductions in a year class (Manooch and Raver 1984).

Vectors of seven-day running averaged winds from Norfolk International Airport for 1978-1988 show strong northerly winds from October through April with episodes of southerly winds from January onward. May and June show a seasonal shift to predominantly southerly winds (Fig. 3). These episodes of seasonal wind reversal could cause current reversals in the MAB (Bumpus 1973) and serve as a transport mechanism for spot recruiting to Chesapeake Bay. The number of southerly wind events and the number of southerly wind events lasting four consecutive days (Fig. 4) were analyzed for differences between months and years. There was no significant difference


Figure 3
Wind vectors from Norfolk International Airport, Virginia, (1978-1988) were processed by smoothing with 7 -day running averages. Time scale is 1 July through 30 June of the following year. Up is north. Orientation is in the direction towards which the wind is blowing, i.e., away from the center line. Note southerly events occurring from December to March. Arrows indicate appearance of cohorts of juvenile spot in the Chesapeake Bay estuaries in 1987 and 1988 (Figures 6 and 7).


Figure 4
Total number of southerly wind events and number of consecutive 4-day southerly wind events taken from vector plots in Figure 3.
between years in number of southerly winds of either type occurring between November and June. However, the total number of southerly events ( $F=10.3$; $\mathrm{df}=7,63 ; P<0.01$ ) and the number of four-day southerly events $(F=9.6$;
$\mathrm{df}=7,63 ; P<0.01$ ) occurring per month were significantly different.
As expected, the total number of southerly events per year is constant. It is the timing of these southerly events relative to periods when larvae are pelagic that is most important. Spot larvae begin entering Chesapeake Bay primarily during March and April. If these fish are approximately two months of age as found in North and South Carolina, then the time frame for wind analysis needs to be narrowed to those months where wind events are most likely to affect the transport process. April and May, the time of shift to a summer wind pattern in the MAB, consistently provide a sufficient number of southerly events to transport larvae spawned late in the season northward toward Chesapeake Bay.

Year-class strength is hypothesized to be dependent on the number of southerly events occurring prior to the wind shift. Average year classes would rely totally on the windinduced transport during April/May, very poor year classes would result when there was a reduced number of southerly events during April/May, and dominant year classes would then be dependent on the number of southerly events occurring from January through March. Analysis of the fourday southerly wind events occurring during this three month period detected a significant difference $(F=2.55$; $\mathrm{df}=9,18 ; P<0.05$ ) between years. Results from the total number of southerly wind events was not conclusive ( $F=$ $1.92 ; \mathrm{df}=9,18 ; P=0.114$ ), but still reflected a difference between years.
Analysis of monthly or seasonally averaged winds indicates that winter is not favorable for onshore transport south of Cape Hatteras (Yoder 1983). These time scales conceal short periods of wind variability which are important for larval fish transport. Transport across the southeastern U.S. shelf in winter can be explained by 'event' scale wind reversals which are most numerous in the outer regions (Yoder 1983). The most favorable shelf current regime for shoreward transport is hypothesized to be in the mid-layer during winter (Miller et al. 1984; Miller et al. 1985). A three-layer winter offshore/onshore transport regime, combined with vertical migration of larvae, may keep the fall/winter spawned larvae offshore and delay entrance to North Carolina estuaries during winter (Miller et al. 1984). Perhaps there is a similar mechanism which bypasses or only partially uses this physical regime and results in transport around Cape Hatteras and northward to Chesapeake Bay. This would be consistent with 57-82 day old fish (Siegfried 1989) entering a Chesapeake Bay estuary. Alternatively, since the water column north of Cape Hatteras is unstratified at this time of year (Beardsley et al. 1976), reducing the effect of Ekman transport on the upper water column, the transport mechanism may be
less complex. For 90 days, spot larvae are buffeted both north and south by winds and currents but the additive product of pulses of southerly winds cause a resultant northward movement.

Information on distribution of spot larvae during winter and early spring in the MAB is limited (Berrien et al. 1978; Morse et al. 1987; M. Fahay, NMFS/NEFC, Highlands, NJ 07732, pers. commun. Nov. 1988). Distribution of spot larvae (Fig. 5) in November-December was concentrated south of Cape Hatteras and spread northward to Oregon Inlet. Larvae collected during February were primarily south of Oregon Inlet with highest densities found near Cape Hatteras. By March, the larvae were widely distributed from south of Cape Lookout to the area off False Cape, VA, south of the Chesapeake Bay mouth. Distribution continued to progress northward during April when larvae were found north of the mouth of Chesapeake Bay. During May, the regional distribution was further condensed with almost all larvae found north of False Cape.

The extent of distribution varied from year to year, but several patterns can be seen. Distribution of larvae spread northward to Chesapeake Bay over time and most larvae were captured inshore where waters are prone to windinduced current reversals. Mean size of larvae increased from offshore to inshore and with distance northward, suggesting a spawning site near or south of Cape Hatteras on the outer shelf. The range of larval sizes found each month further emphasized the protracted spawning period.

Distribution of larvae, coupled with information on size and age, has been used in the South Atlantic Bight to infer time and area of spawning (Lewis and Judy 1983; Warlen and Chester 1985). From Cape Hatteras southward, the highest larval abundance in winter (NovemberMarch) is on the outer shelf in depths of 40-200 m (Lewis and Judy 1983; Yoder 1983; Warlen and Chester 1985). Concurrently, surface temperatures there are warmer and more favorable for spawning and larval survival (Yoder 1983).

Pelagic larvae are unable to regulate their environment, but they may moderate the effects through an appropriate behavioral response to reliable environmental signals (Leggett 1985). Microscale migrations of larvae of several species have been shown to influence their association with beneficial current regimes, resulting in transport into estuarine nursery grounds (Leggett 1985; Miller et al. 1985). Therefore, it is possible that spring reversals of wind patterns north of Cape Hatteras could act as an environmental signal to which spot larvae respond, resulting in northward transport to Chesapeake Bay. However, early larval survival is highly variable when dependent upon such large-scale meteorological events (Leggett et al. 1984). Such
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Figure 5
Aggregate seasonal distribution and abundance of spot larvae from the RV Dolphin cruises (1965-1966) (Berrien et al. 1978) and NMFS/NEFC surveys (1977-1986) (M. Fahay, NMFS/NEFC, November 1988). Numbers indicate mean size of larvae (mm).
variable recruitment of spot is common in Chesapeake Bay and Delaware Bay (Thomas 1978).

Data on juvenile spot recruitment in Virginia are not available from the same time period as the larval collec-
tions, but they are available from recent samples collected from 1986 to 1988. In 1987, spot first appeared during the second April collection period, 20-23 April, recruitment peaking in May and continuing through June (Fig. 6). The


Figure 6
1987 bimonthly sampling. Length/frequency histograms of newly recruited spot aggregated over five sites in Chesapeake Bay and on the eastern shore of Virginia (from top to bottom: Wachapreague, Sand Shoal, Fisherman's Island, Occohannock Creek, and York River).
trend of peak recruitment in May and June was also seen in 1988 collections, but spot appeared in Chesapeake Bay during March (Fig. 7), a month earlier than in 1987. The March/April appearance of recruits between 13 and 41 mm TL is in agreement with most other reports of spot entrance to Chesapeake Bay (Table 1).

Evidence of a protracted spawning season is seen with the presence of new recruits ( $10-25 \mathrm{~mm}$ TL) in all collections (Figs. 6 and 7). Episodes of southerly winds in 1987 and 1988 occurred prior to the initial appearance of spot within Chesapeake Bay and prior to the appearance of each successive cohort (Fig. 3). The earlier arrival of spot in March 1988 could be related to the number (12) of southerly wind events in February as opposed to 1987 when there were no southerly events in February (Fig. 4) and spot did not appear until late April.

Based on the growth rates of Warlen and Chester (1985) and Siegfried (1989), spot entering Chesapeake Bay in 1987 and 1988 were spawned between January and April. At that time of year, the only bottom water warmer than $17^{\circ} \mathrm{C}$ was just off Cape Hatteras (Fig. 2), 250 km south of Chesapeake Bay.

It is not impossible for fish larvae to be transported in excess of 250 km to reach nursery grounds. Larvae of the American eel (Anguilla rostrata) and bluefish (Pomatomus saltatrix) are also transported northward along the Atlantic coast before entering estuaries. However, unlike spot, it is thought that transport of these species is dependent upon the Gulf Stream (Kendall and Walford 1979; Kleckner and McCleave 1982). The apparent dependence of pelagic spawners on "normal"' oceanographic drift conditions suggests that anomalies in these patterns could result in variation in recruitment success (Norcross and Shaw 1984). The suggested 'normal' condition affecting spot recruitment to Chesapeake Bay is episodes of southerly reversals in spring wind patterns. The timing of these reversals is hypothesized to cause observed anomalous variations in spot recruitment abundance.

The hypothesis that wind events are responsible for a portion of the interannual variability in spot recruitment suggests the need for investigation of the offshore larval distribution coupled with the physical dynamics of the system. Circulation patterns over the MAB continental shelf and around Cape Hatteras are complex, and transport processes may also be related to the amount of estuarine discharge, tidal influence, and stratification of the water column. Current meter data for the MAB south of Chesa-

peake Bay are needed to verify inferences of water movement in conjunction with southerly wind events. Computer simulations of the shelf currents would aid in identifying physical factors that have the greatest effect on recruitment variation. This effect would be further modified by the vertical distribution of spot larvae, though this distribution is currently unknown. Discrete depth collections are there-

Figure 7
1988 monthly sampling. Length/frequency histograms of newly recruited spot aggregated over five sites in Chesapeake Bay and on the eastern shore of Virginia (from top to bottom: Wachapreague, Sand Shoal, Fisherman's Island, Occohannock Creek, and York River).
fore needed to determine depth distribution of spot larvae and associated changes which may occur as larvae develop. Field studies are needed to verify spot spawning sites and to collect concomitant physical data. The possibility exists that fish recruiting to areas north of Cape Hatteras are spawned at a later date than those recruiting to more southern estuaries. Aging of larvae, based on daily otolith increments, would yield information on spawning time, duration of transport, and growth rates during the larval phase. Sampling of new recruits on a time scale finer than the monthly and bimonthly data presented here is necessary to adequately test the hypothesized relationship with southerly wind events.

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## Citations

Beardsley, R.C., and W.C. Boicourt.
1981. On estuarine and continental shelf circulation in the Middle Atlantic Bight. In Evolution of physical oceanography (B.A. Warren and C. Wunsch, eds.), p. 198-233. MIT Press, Cambridge, MA.
Beardsley, R.C., W.C. Boicourt, and D.V. Hansen
1976. Physical oceanography of the Middle Atlantic Bight. In Middle Atlantic continental shelf and the New York Bight (M.G. Gross, ed.), p. 20-34. Am. Soc. Limnol. Oceanogr., Spec. Symp. No. 2.
Beckman, D.W., and J.M. Dean.
1984. The age and growth of young-of-the-year spot, Leiostomus xanthurus, Lacepede, in South Carolina. Estuaries 7:487-496.
Berrien, P.L.; M.P. Fahay, A.W. Kendall Jr., and W.G. Smith. 1978. Ichthyoplankton from the RV Dolphin Survey of Continen-
tal Shelf waters between Martha's Vineyard, Massachusetts and Cape Lookout, North Carolina, 1965-66. NOAA Tech. Rep. NMFS SSRF 15, 152 p.
Bumpus, D.F.
1973. A description of the circulation on the continental shelf of the East coast of the United States. Prog. Oceanogr. 6:111-157.
Bumpus, D.F., R.E. Lynde, and D.M. Shaw.
1973. Physical oceanography. Chapter 1. In Coastal and offshore environmental inventory, Cape Hatteras to Nantucket shoals. Univ. Rhode Island Mar. Publ. Ser. No. 2, 67 p.
Chao, L.N., and J.A. Musick.
1977. Life history, feeding habits, and functional morphology of juvenile sciaenid fishes in the York River estuary, Virginia. Fish. Bull., U.S. 75(4):675-702.
Cowan, J.H. Jr., and R.S. Birdsong.
1985. Seasonal occurrence of larval and juvenile fishes in a Virginia Atlantic coast estuary with emphasis on drums (Family Sciaenidae). Estuaries 8:48-59.
Davis, C.W.
1979. Bottom-water temperature trends in the Middle Atlantic Bight during spring and autumn, 1964-76. NOAA Tech. Rep. NMFS SSRF 739, 13 p.
Dawson, C.E
1958. A study of the biology and life history of the spot, Leiostomus xanthurus Lacepede, with special reference to South Carolina. Contrib. Bears Bluff Lab. 28:1-48.
Edwards, R.L., R. Livingstone Jr., and P.E. Hamer,
1962. Winter water temperatures and an annotated list of fishes - Nantucket Shoals to Cape Hatteras, Albatross III Cruise No. 126. U.S. Fish Wildl. Serv. Spec. Sci. Rep. Fish. 397, 31 p.
Heck, K.L. Jr., and T.A. Thoman.
1984. The nursery role of seagrass meadows in the upper and lower reaches of the Chesapeake Bay. Estuaries 7:70-92.
Hettler, W.F., and A.B. Powell.
1981. Egg and larval fish production at the NMFS Beaufort laboratory, Beaufort, NC, USA. Rapp. P.-v. Reun. Cons. Int. Explor. Mer 178:501-503.
Hildebrand, S.F., and L.E. Cable.
1930. Development and life history of fourteen teleostean fishes at Beaufort, North Carolina. Bull. U.S. Bur. Fish. 46:383-488.
Hildebrand, S.F., and W.C. Schroeder.
1928. The fishes of Chesapeake Bay. Bull. U.S. Bur. Fish. 43(1), 388 p.
Johnson, G.D.
1978. Leiostomus xanthurus Lacepede, Spot. In Development of fishes of the Mid-Atlantic Bight: An atlas of egg, larval and juvenile stages. Vol IV: Carangidae through Ephippidae, p. 203-208. U.S. Fish Wildl. Serv., Biol. Serv. Prog., FWS/OBS-78/12.

Joseph, E.B.
1972. The status of the sciaenid stocks of the middle Atlantic coast. Chesapeake Sci. 14:87-100.
Jossi, J.W., and R.R. Marak.
1983. MARMAP Plankton survey Manual. NOAA Tech. Memo. NMFS-F/NEC-21, 258 p.
Kendall, A.W. Jr., and L.A. Walford.
1979. Sources and distribution of bluefish, Penatomus saltatrix, larvae and juveniles of the east coast of the United States. Fish. Bull., U.S. 77:213-238.
Kleckner, R.C., and J.D. McCleave.
1982. Entry of mitrating American eel leptocephali into the Gulf Stream system. Helg. Wiss. Meeres. 35:329-339.
Leggett, W.C.
1985. The role of migrations in the life history evolution of fish. In Migration: mechanisms and adaptive significance (M. Rankin, ed.), p. 277-295. Contr. Mar. Sci., Vol. 27 (Suppl.).

Leggett, W.C., K.T. Frank, and J.E. Carscadden. 1984. Meteorological and hydrographic regulation of year class strength in capelin (Mallotus villosus). Can. J. Fish. Aquat. Sci. 41:1193-1201.
Lewis, R.M., and M.H. Judy.
1983. The occurrence of spot, Leiostomus xanthurus, and Atlantic croaker, Micropogonias undulatus, larvae in Onslow Bay and Newport River estuary, North Carolina. Fish. Bull., U.S. 81(2):405-412.
Lewis, R.M., and W.C. Mann.
1971. Occurrence and abundance of larval Atlantic menhaden, Brevoortia tyrannus, at two North Carolina inlets and a list of associated species. In Proceedings of a workshop on egg, larval and juvenile stages of fish in Atlantic coast estuaries (A.L. Pacheco, ed.), p. 142-147. U.S. Natl. Mar. Fish. Serv., Mid. Atlantic Coastal Fish. Ctr., Highlands, NJ. Publ. No. 1, 338 p.
Lewis, R.M., and E.P.H. Wilkens.
1971. Abundance of Atlantic menhaden larvae and associated species during a diel collection at Beaufort, North Carolina. Chesapeake Sci. 12:185-187.
Lippson, A.J., and R.L. Moran.
1974. Manual for identification of early developmental stages of fishes of the Potomac River estuary. Env. Tech. Cent./Martin Marietta, Baltimore, MD, 282 p.
Manooch, C.S. III, and D. Raver Jr.
1984. Fishes of the southeastern United States. North Carolina State Museum of Natural History, Raleigh, NC, 362 p.
McCambridge, J.T. Jr., and R.W. Alden III.
1984. Growth of juvenile spot, Leiostomus xanthurus Lacepede, in the nursery region of the James River, Virginia. Estuaries 7: 478-486.
McGovern, J.C.
1986. Seasonal recruitment of larval and juvenile fishes into impounded and non-impounded marshes. M.S. thesis, College of Charleston, Charleston, SC.
Miller, J.M., J.P. Reed, and L.J. Pietrafessa.
1984. Patterns, mechanisms and approaches to the study of migrations of estuarine dependent fish larvae and juveniles. In Mechanisms of migration in fishes (J.D. McCleave, G.P. Arnold, J.J. Dodson, and W.H. Neill, eds.), p. 209-225. Plenum Press, NY.
Miller, J.M., L.B. Crowder, and M.L. Moser.
1985. Migration and utilization of estuarine nurseries by juvenile fishes: an evolutionary perspective. In Migration: mechanisms and adaptive significance (M. Rankin, ed.), p. 338-352. Contr. Mar. Sci., Vol. 27 (Suppl.).
Morse, W.W., M.P. Fahay, and W.G. Smith.
1987. MARMAP surveys of the continental shelf from Cape Hatteras, North Carolina, to Cape Sable, Nova Scotia (1977-1984). Atlas No. 2. Annual Distribution patterns of fish larvae. NOAA Tech Mem. NMFS-F/NEC-47, 215 p.
National Marine Fisheries Service/Northeast Fisheries Center.
1983. Fisherman's Report, Fall bottom trawl survey, Cruise 8308. September 12-October 14, RV Albatross $I V$. NMFS/NEFC, Woods Hole, MA.
1984a. Fisherman's Report, Bottom trawl survey, Cape Lookout - Western Georges Bank. February 29-March 29, RV Albatross IV. NMFS/NEFC, Woods Hole, MA.

1984b. Fisherman's Report, Bottom trawl survey, Cape Hatteras - Western Scotian Shelf. September 10-November 9, RV Albatross IV. NMFS/NEFC, Woods Hole, MA.
1985. Fisherman's Report, Bottom trawl survey, Cape Lookout - Western Scotian Shelf. February 25-April 13, RV Albatross IV. NMFS/NEFC, Woods Hole, MA.
1986a. Fisherman's Report, Bottom trawl survey, Cape Lookout - Western Scotian Shelf. March 3-April 27, RV Albatross IV. NMFS/NEFC, Woods Hole, MA.

1986b. Fisherman's Report, Bottom trawl survey, Cape Lookout - Western Scotian Shelf. September 14-November 7, RV Albatross IV \& RV Delaware II. NMFS/NEFC, Woods Hole, MA.
1987a. Fisherman's Report, Bottom trawl survey, Cape Hatteras - Western Scotian Shelf. March 24-April 28, RV Albatross IV \& RV Delaware II. NMFS/NEFC, Woods Hole, MA.
1987b. Fisherman's Report, Bottom trawl survey, Cape Hatteras - Western Scotian Shelf. September 10-November 6, RV Albatross IV. NMFS/NEFC, Woods Hole, MA.
1988. Fisherman's Report, Bottom trawl survey, Cape Hatteras - Western Scotian Shelf. March 4-April 21, RV Albatross IV. NMFS/NEFC, Woods Hole, MA.
National Weather Service.
1978-1988. Local climatological data monthly summaries. Norfolk International Airport. Norfolk, VA.
Nickerson, S.R., and W.R. Wright.
1980. Spring and fall bottom temperatures on the continental shelf Cape Hatteras to Cape Sable, 1972 to 1979, with surface temperature and salinity for 1972 and 1979. USDOC/NOAA/ NMFS/NEFC, Woods Hole, Mass. Lab. Ref. No. 80-01.
Norcross, B.L., and H.M. Austin.
1988. Middle Atlantic Bight meridional wind component effect on bottom water temperatures and spawning distribution of Atlantic croaker. Cont. Shelf Res. 8(1):69-88.
Norcross, B.L., and D. Hata.
In press. Seasonal composition of finfish in waters behind the Virginia barrier islands. VA. J. Sci.
Norcross, B.L., and R.F. Shaw.
1984. Oceanic and estuarine transport of fish eggs and larvae: A review. Trans. Am. Fish. Soc. 113:153-165.
Olney, J.E.
1983. Eggs and early larvae of the bay anchovy, Anchoa mitchilli, and the weakfish, Cynoscion regalis, in lower Chesapeake Bay with notes on associated ichthyoplankton. Estuaries 6:20-35.
Olney, J.E., and G.W. Boehlert.
1988. Nearshore ichthyoplankton associated with seagrass beds in the lower Chesapeake Bay. Mar. Ecol. Prog. Ser. 45:33-43.
O'Neill, S., and M.P. Weinstein.
1987. Feeding habitats of spot, Leiostomus xanthurus, in polyhaline versus meso-oligohaline tidal creeks and shoals. Fish. Bull., U.S. 85:785-796.
Orth, R.J., and K.L. Heck Jr.
1980. Structural components of eelgrass (Zostera marina) meadows in the lower Chesapeake Bay - fishes. Estuaries 3:278-288.
Pacheco, A.L.
1957. The length and age composition of spot, Leiostomus xanthurus, in the pound net fishery of lower Chesapeake Bay. M.A. thesis, College of William and Mary, Williamsburg, VA, 34 p.
1962. Age and growth of spot in lower Chesapeake Bay, with notes on distribution and abundance of juveniles in the York River system. Chesapeake Sci. 3(1):18-28.
Pearson, J.C.
1932. Winter trawl fishery off the Virginia and North Carolina coasts. U.S. Bur. Fish., Invest. Rep. No. 10, 31 p.
Powell, A.B., and H.R. Gordy.
1980. Egg and larval development of the spot, Leiostomus xanthurus (Sciaenidae). Fish. Bull., U.S. 78:701-714.

Richards, C.E., and M. Castagna.
1970. Marine Fishes of Virginia's eastern shore (inlet and marsh, seaside waters). Chesapeake Sci. 11:235-248.
Rogers, S.G., T.E. Targett, and S.B. VanSant.
1984. Fish-nursery use in Georgia salt-marsh estuaries: the influence of springtime freshwater conditions. Trans. Am. Fish. Soc. 113:595-606.
Schauss, R.P Jr.
1977. Seasonal occurrence of some larval and juvenile fishes in Lynnhaven Bay, Virginia. Am. Midl. Nat. 98:275-282.
Siegtried, R.
1989. Documentation of daily rings in otoliths of young-of-the-year spot, Leiostomus xanthurus, and Atlantic croaker, Micropogonias undulatus. M.A. thesis, College of William and Mary, Gloucester Point, VA. Unpubl. manuscr.
Smith S.M., J.G. Hoff, S.P. O'Neill, and M.P. Weinstein.
1984. Community and trophic organization of nekton utilizing shallow marsh habitats, York River, VA. Fish. Bull., U.S. 82: 455-467.
Springer, S., and H.R. Bullis Jr.
1956. Collections by the Oregon in the Gulf of Mexico. U.S. Fish. Wildl. Serv., Spec. Sci. Rep. Fish. 196:1-134.
Squire J.L. Jr.
1958. Cruise report - M/V Delaware - Cruise 58-1. U.S. Fish Wildl. Serv. Bur. Comm. Fish, 9 p.
Tagatz, M.E., and D.L. Dudley.
1961. Seasonal occurrence of marine fishes in four shore habitats near Beaufort, N.C., 1957-1960. U.S. Fish Wildl. Serv., Spec. Sci. Rep. Fish. 390, 19 p.
Thomas, D.S.
1978. The early life history and ecology of six species of drum (Sciaenidae) in the lower Delaware River, a brackish tidal estuary. Ichthyol. Assoc. Delaware Prog. Rep. 3; Part III:77-81. Middletown, DE.
Turner, W.R., and G.N. Johnson.
1973. Distribution and relative abundance of fishes in Newport River, North Carolina. NOAA Tech. Rep. NMFS SSRF 666.
Warlen, S.M., and A.J. Chester.
1985. Age, growth, and distribution of larval spot, Leiostomus xanthurus, off North Carolina. Fish. Bull., U.S. 83:587-599.
Weinstein, M.P., and H.A. Brooks.
1983. Comparative ecology of nekton residing in a tidal creek and adjacent seagrass meadow: community composition and structure. Mar. Ecol. Prog. Ser. 12:15-27.
Welsh, W.W., and C.M. Breder.
1923. Contributions to the life histories of Sciaenidae of the eastern United States Coast. Bull. U.S. Bur. Fish. 39:141-201.
Williams, A.B., and E.E. Deubler Jr.
1968. Studies on macroplanktonic crustaceans and ichthyoplankton of the Pamlico Sound complex. N.C. Dept. Cons. Dev., Div. Comm. Sports Fish., Raleigh, NC, Spec. Sci. Rep. 13, 103 p.
Yoder, J.A.
1983. Statistical analysis of the distribution of fish eggs and larvae on the southeastern U.S. continental shelf with comments on oceanographic processes that may affect larval survival. Estuarine Coastal Shelf Sci. 17:637-650.

# Ichthyoplankton Assemblages Sampled by Night Lighting in Nearshore Habitats of Southwestern Puerto Rico 

GEORGE D. DENNIS,* DENIS GOULET, and JAY R. ROOKER<br>Department of Marine Sciences<br>University of Puerto Rico<br>Mayagüez, Puerto Rico 00709-5000


#### Abstract

Larval fishes were sampled in four nearshore habitats: coral reef, seagrass bed, mangrove lagoon, and mangrove prop roots in southwestern Puerto Rico. A lift net with attached night light was employed to determine seasonal abundance and species composition of the nearshore ichthyoplankton assemblage. Coral reef and seagrass bed habitats usually possessed the greatest abundance and species richness of larval fishes. Few early stages of larval fishes were collected in mangrove habitats suggesting that they were not nursery areas. The abundance of larval fishes in the open water area of the lagoon was not significantly different from the prop root habitat. Although all habitats were within close proximity (ca. 2 km ), there were different patterns in abundance between the coral reef/seagrass bed and mangrove habitats. Based on low abundance of larval fishes and few species captured, the mangroves cannot be considered an important spawning or nursery area for larval fishes in southwestern Puerto Rico.


## Introduction

Larval fishes in nearshore tropical environments have not been thoroughly studied because of the difficulty in using standard ichthyoplankton sampling gear in these areas. A major problem encountered when sampling tropical nearshore waters with an active gear, such as a towed net, is navigation among shallow reef areas. Sampling at night amplifies navigational problems, but may be particularly important in nearshore areas where larval fishes may aggregate near bottom or visually avoid towed nets during the day (Powles 1977; Thayer et al. 1983). Thus, most tropical ichthyoplankton studies have been concentrated in oceanic waters where large vessels can operate (Ahlstrom 1971, 1972; Powles 1975; Leis and Miller 1976; Richards 1984). Few studies have examined ichthyoplankton around mangroves (Wyatt 1982; Flores-Coto et al. 1983; Collins and Finucane 1984; Little et al. 1988; Powell et al. 1989) even though these areas are considered major fish nurseries (Heald and Odum 1970).

This study addresses the hypothesis that mangrove areas are spawning or larval nursery areas for fishes in southwestern Puerto Rico. We employed a lift net with attached

[^5]night light to reduce the problem of sampling with towed nets in coral reef and mangrove habitats.

## Materials and Methods

## Sampling Area

The study area was located on the southwestern coast of Puerto Rico, the most easterly large island in the Greater Antilles (Fig. 1). The coast is fringed by relatively undisturbed red mangrove forests (Rhizophora mangle) and nearshore waters are dotted with red mangrove cays. Many well-developed coral reefs are also found in the area. This portion of Puerto Rico has a dry climate with a total rainfall of 695 mm for 1988. There are no rivers and little freshwater runoff; hence water quality is good. Larval fishes were sampled in four nearshore habitats: mangrove prop root, mangrove lagoon, seagrass bed, and coral reef.
The mangrove prop root habitat included the prop root system adjacent mangrove produced muddy bottom areas (Dennis, in press). Soft mud bottom abutted the mangroves and average water depth for prop-root stations averaged 1.2 m . Four sampling stations were selected here.

A small lagoon surrounded by red mangroves served as the mangrove lagoon habitat and was the primary mangrove habitat sampling site (Fig. 1). The lagoon station


Figure 1
Map of sampling area in southwestern Puerto Rico. M is the primary mangrove sampling site (4 prop-root stations and one lagoon station). BA and BB are additional mangrove lagoons. SG is the seagrass bed station and CR is the coral reef station.
was over soft mud and had a depth of 2 m . One lagoon station was established near the center of the east entrance to the lagoon. Two additional mangrove lagoons to the west of the primary lagoon were also sampled (Fig. 1). They also had a soft mud bottom and an average depth of 1 m .

A seagrass bed station (Fig. 1) was selected off Isla Cueva on a shallow platform primarily covered with Thalassia testudinum and some small gorgonians and coral patches. Water depth at this site was 1.5 m .

A coral reef station was located on a fringing reef off the leeward end of a mangrove cay (Fig. 1). The reef was dominated by Acropora palmata and Millepora spp. The sample site was directly over coral in a water depth of 1.7 m . Adjacent to the reef the bottom was covered with seagrass.

## Sampling Methods

We used a lift-net with a night light as a sampling device, composed of a floating platform housing a circular 12-volt sealed-beam automobile headlight bulb (Fig. 2). A standard $50-\mathrm{cm}$ diameter by $165-\mathrm{cm}$ long conical plankton net of 500 -micron mesh was attached to the platform by four guide ropes. Power for the light was supplied by a 12 -volt marine battery aboard a $6-\mathrm{m}$ fiber-glass support boat.

Sampling consisted of lowering the plankton net to a depth of one meter or less depending on bottom depth, then turning on the light for 10 minutes. The net was then rapidly hoisted to the surface by hand while the light was still on and it sampled fish in the water column between the net and light source. Sampling characteristics of this device are discussed in detail (D. Goulet et al. 1988).

Three replicates were taken at each station during each sampling period. Coral reef and seagrass bed stations were sampled sequentially with a 10 minute no-light period between replicates; mangrove stations were alternately sampled. The order of station sampling was randomly selected and all samples taken within one day of the new moon from 2000 to 0100 hours.

Monthly samples were collected at the four prop-root mangrove stations in August, September, and October 1987; at one of these stations from September 1987 to February 1989; and at the mangrove lagoon station from October 1987 to February 1989. Two additional mangrove lagoons (Fig. 1) were sampled in February 1989. No samples were taken at the mangrove stations in November 1988. Seagrass bed and coral reef stations were sampled from March 1988 to February 1989. Because of inclement weather and gear failure, samples were missed in August 1988 at the seagrass bed station and in August and September 1988 at the coral reef station. No samples were


Figure 2
Lift-net sampling device with night light.
taken in January 1989. Water temperature, salinity, and dissolved oxygen were measured at 0.3 m below the surface at each station.

Samples were initially preserved in $10 \%$ formalin and transferred to $5 \%$ formalin within one week. After sorting, identification, and measurement, the fishes were stored in $70 \%$ ethanol. All fishes were identified to the lowest taxon possible. A taxon was defined as a distinct life-history stage of a species. Life-history stages included: preflexion larvae (before notochord flexion), postflexion larvae (after notochord flexion and including flexion larvae), and juveniles (fish with a full complement of adult fin ray counts) (after Leis and Rennis 1983).

We considered mangroves to be spawning areas for fishes if early preflexion larvae were abundant in that habitat. Also, if mangroves are larval nursery areas, then later stage (postflexion) larvae should be abundant there.

## Analysis

Two-way ANOVA analyses were carried out on log transformed $(x+1)$ abundance data for total number of taxa, total number of larvae, total number of non-dwarf herring (Jenkinsia spp.) larvae, and the four most abundant taxa, dwarf herring, sardine (Harengula spp.), anchovies (Anchoa sp.), and bonefish (Albula vulpes) to test for differences in abundance among habitats and months. Only the eight months when all habitats were sampled were used in this ANOVA. A two-way ANOVA among the four primary mangrove prop-root stations and months was used to test
for significant differences in number of larvae among locations within a mangrove lagoon. Samples were taken during three consecutive months (August-October 1987) for this analysis. Two additional mangrove lagoons (Fig. 1) were compared to Lagoon M during February 1989 with a two-way ANOVA by habitat (lagoon and prop roots) and location (three mangrove lagoons) to test for differences among lagoons. Tukey's HSD test was used to determine which levels of a factor were significant (Sokal and Rohlf 1981).

Additional comparisons of larval abundance were made among habitats (summed over all sampling periods) with a chi-square test assuming larval abundance was proportional to sampling effort.

Similarity among habitats was measured by the percent similarity formula, $P S=1-0.5 \Sigma\left|p_{x, i} p_{y, i}\right|$, where $p_{x, i}=$ proportion of taxa $i$ in habitat $x$ and $p_{y, i}=$ proportion of taxa $i$ in habitat $y$ based on abundance over all sampling periods (Kohn and Riggs 1982). Unweighted pair-group arithmetic average (UPGMA) clustering was used to create the similarity dendrogram (Sneath and Sokal 1973).

## Results

Habitats had significantly different patterns of larval fish abundance (Figs. 3 and 4). In all cases there was a significant interaction between habitats and months which indicated that habitats had different trends in larval fish abundance over time ( $F$-test, habitat by month interaction term, $P<0.01$ ). The mangrove lagoon and prop-root stations differed from coral reef and seagrass bed stations in their patterns of abundance. Larval fish were most abundant at seagrass and coral reef stations from February to April while larvae were most abundant in the mangroves in July (Fig. 3). There was a greater number of taxa at seagrass bed and coral reef stations from December to April, but there was little difference among the four habitats from June to October.

Seasonal patterns of abundance differed among habitats for the most abundant taxa (Fig. 4). From March to April, bonefish larvae were abundant at the coral reef station, but over the whole sampling period they were more commonly collected at the mangrove prop-root station.

Dwarf herring were the most abundant larvae taken, and were primarily collected from seagrass bed and coral reef stations (Fig. 4). Patterns in abundance of dwarf herring larvae (preflexion and postflexion) were similar over time at the coral reef station with a major abundance peak in February 1989 and smaller peaks in May 1988 and October 1988 (Fig. 4). The seagrass bed station differed from the coral reef station in lacking an abundance peak in May 1988 and having a peak in March 1988 (Fig. 4).

The sardine was most commonly collected at mangrove and seagrass bed stations, but few were taken at the coral


Figure 3
Monthly trends in arithmetic mean number of larvae excluding Jenkinsia spp., number of taxa, and number of larvae by habitat.
reef station. The anchovy was primarily collected from mangrove and seagrass bed stations with a major abundance peak in July at mangrove stations and another smaller peak between September and October at seagrass bed and mangrove lagoon stations.

The coral reef station had more preflexion and postflexion larvae than other stations (Chi-square test, $P<0.01$, in both cases). Bonefish (Albula vulpes), sea bream (Archosargus rhomboidalis), preflexion and postflexion dwarf herring (Jenkinsia spp.), preflexion and postflexion silversides (Atherinidae), Clinidae species 1 and 2, and unidentified preflexion larvae, were also significantly more abundant at this station (Table 1) (Chi-square test, $P<0.01$ in all
cases). The coral reef station was most similar to the seagrass bed station and very dissimilar to mangrove stations (Fig. 5).

The seagrass bed station had significantly more taxa, preflexion larvae, postflexion larvae, and juveniles than mangrove stations (Table 1) (Chi-square test, $P<0.01$ ). Four taxa were most common here: Gobiidae species 2, dwarf herring juveniles, and sardine postflexion larvae and juveniles. Also, dwarf herring postflexion larvae were significantly more abundant at this station than at mangrove stations (Chi-square test, $P<0.01$ ). The seagrass bed station had an intermediate assemblage of larval fishes relative to coral reef and mangroves, but was much more


Figure 4
Monthly trends in arithmetic mean number of Albula vulpes, Jenkinsia spp., Harengula spp., and Anchoa sp. by habitat.
similar to the coral reef station than the mangrove stations (Fig. 5).

Mangrove stations, in general, had fewer taxa and number of larvae than coral reef and seagrass bed stations (Table 1). The mangrove lagoon had more preflexion larvae and postflexion sardine larvae than the prop-root station (Chi-square test, $P<0.01$ in both cases). Only two taxa were more abundant at the prop-root station, bonefish and gerreid larvae, and the latter was significantly more abundant at the prop-root station than at any other station (Chi-square test, $P<0.01$ ). Mangrove stations were
very similar to each other and next most similar to the seagrass bed station (Fig. 5).

Examination of larval fish abundance within lagoon M at four prop-root stations over a three month period indicated similar trends in larval abundance among stations ( $F$-test, station by month interaction, $P=0.273$ ). Months were significantly different in larval fish abundance ( $F$-test, $P=0.012$ ), but there was only a marginally significant difference among stations ( $F$-test, $P=0.041$ ).

Comparison of larval fish abundance among three mangrove lagoons indicated no interaction between habitats


Figure 5
Similarity among habitats based on taxon abundance over all sampling periods. Dendrogram formed by UPGMA method on percent similarity (PS).
and locations ( $F$-test, $P=0.553$ ) and no significant difference between habitats ( $F$-test, $P=0.629$ ). There was a significant difference in larval fish abundance among lagoons with the primary sampling lagoon (M) having significantly more larvae than the other two ( $F$-test, $P=0.012$, Tukey's HSD test, Sokal and Rohlf 1981).

Salinity was high ( $34-37 \mathrm{ppt}$ ) at all stations, even during the rainy season. Water temperature during the study ranged from a low of $26.0^{\circ} \mathrm{C}$ in January-February to a high of $30.5^{\circ} \mathrm{C}$ in July, but differed less than $1^{\circ} \mathrm{C}$ among stations during any sampling period. Dissolved oxygen usually ranged from 5 to 7 ppm , but on one occasion measured 2.5 ppm near bottom at a mangrove prop-root station. There was little variation in environmental parameters among stations within any sampling period.

## Discussion

Almost any active method of collecting ichthyoplankton in waters with obstructions will result in sampling difficulties. One solution to this problem is the use of a passive aggregating device, such as light. Its application has been primarily relegated to a qualitative, ancillary role in the past. Using light as a quantitative method of sampling can been criticized on two main points: volume sampled is unknown and species selectivity bias.

Volume sampled is dependent on water clarity and current speed. Theoretically, more turbid water should result in fewer larvae attracted owing to a smaller area of light influence. Greater current speed should (up to the point where larvae can no longer maintain their position) result in more larvae passing within the sphere of light influence and in potentially being retained in the area for collection.

Current velocity in the nearshore environment is primarily a result of daytime wind-driven circulation in locations, such as the Caribbean, where there is a limited tidal range (ca. 0.5 m ). Nighttime, usually a period of low wind, further reduces the influence of current speed on volume sampled. In this study we attempted to control these factors by sampling in areas of similar high water clarity and keeping the duration of sampling short ( 10 minutes).
Light is selective both for taxonomic composition and size. Though taxon selectivity is not well documented, it is known that different stages of some fish species react differently to light (Bulkowski and Meade 1983). Still, there is a tremendous range of taxa collected by light methods (Doherty 1987) and this same bias is known to occur in towed gears (Thayer et al. 1983). In this study forty-five taxa represented by 7342 larvae were taken.

Size selectivity may be species specific and biased toward either smaller or larger size groups in active gears depending on gear type. Larger larvae are usually less well sampled because they avoid the net (Thayer et al. 1983; Gregory and Powles 1988). Methods using light usually catch more later-stage larvae (presettlement) and juveniles than towed-net gears (this study; Doherty 1987) making them potentially complementary methods for sampling larval fishes.

Although the four habitats sampled were only about 2 km apart, they exhibited different patterns in larval fish abundance. Abundance peaked from February to April at coral reef and seagrass bed stations, but peaked in July and August at mangrove stations when densities at coral reef and seagrass bed stations were lowest. The different patterns in these two nearby areas seemed to contradict the idea of passive dispersal of preflexion larvae within the nearshore environment. Normal wind-driven circulation from the southeast should push water (and preflexion larvae) from nearshore reefs and seagrass beds into mangrove areas. This circulation alone would increase the abundance of preflexion larvae in the mangroves. But preflexion dwarf herring larvae were never collected in mangroves, though juveniles and adults commonly occur there. Even between the coral reef and seagrass bed stations there was little coherence in peaks of abundance for preflexion dwarf herring larvae. This species, instead, makes use of epibenthic and benthic areas and therefore may not be subjected to passive transport by currents (Powles 1977; pers. obs.).

The mangrove prop-root habitat had a low density of larval fish, as did the 'open water' lagoon station. Yet larval gerreids, bonefish, and sardine made use of the mangrove habitat. The generally low density of larval fishes in mangrove habitats did not support the hypothesis that these areas are nurseries for larval fishes at least in southwestern Puerto Rico. Several studies support our findings. In the Florida Everglades, there were fewer taxa and larvae taken by towed net in mangrove estuaries than in nearshore areas (Collins and Finucane 1984). Highest diver-

Table 1
Number of fishes by taxa collected by night lighting in four nearshore habitats off southwestern Puerto Rico. Mangrove habitats include only the primary mangrove sampling site (see Fig. 1). Life-history stages are PR: preflexion, PO: postflexion, and J: juvenile.

| Taxa | Stage | Habitat |  |  |  | Total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Coral Reef | Seagrass Bed | Mangrove Lagoon | Mangrove <br> Prop Root |  |
| ALBULIDAE Albula vulpes | PO | 90 | 51 | 36 | 101 | 278 |
| APOGONIDAE | PO | - | 1 | - | - | 1 |
| ATHERINIDAE | PR | 393 | 35 | 15 | 2 | 445 |
|  | PO | 29 | 3 | - | 1 | 33 |
| CARANGIDAE | PO | 1 | - | - | - | 1 |
| Oligoplites saurus | J | - | - | 3 | 8 | 11 |
| Trachinotus sp. | PO | - | - | - | 1 | 1 |
| CLINIDAE $\begin{aligned} & \text { Species } 1 \\ & \\ & \text { Species } 2\end{aligned}$ | PO | 22 | 4 | - | - | 26 |
|  | PO | 58 | 14 | - | - | 72 |
| CLUPEIDAE Harengula spp. | PO | 27 | 90 | 91 | 65 | 273 |
|  | J | 24 | 129 | 106 | 112 | 371 |
| Jenkinsia spp. | PR | 2216 | 115 | - | - | 2331 |
|  | PO | 1106 | 1043 | 8 | 3 | 2160 |
|  | J | 189 | 324 | 3 | 5 | 521 |
| Opisthonema oglinum | PO | - | - | 6 | - | 6 |
|  |  | - | - | 2 | - | 2 |
| DACTYLOSCOPIDAE | PO | 1 | - | - | - | 1 |
| ELOPIDAE Elops saurus | PO | 1 | - | - | - | 1 |
| ENGRAULIDAE Anchoa sp. | PR | , | 2 | 1 | - | 6 |
|  | PO | 2 | 47 | 67 | 54 | 170 |
|  | J | 2 | 7 | 4 | 10 | 23 |
| GERREIDAE | PO | 3 | 5 | 3 | 25 | 36 |
| GOBEISOCIDAE $\begin{aligned} & \text { Species 1 } \\ & \\ & \\ & \text { Species 2 }\end{aligned}$ | PO | 6 | - | - | - | 6 |
|  | PO | 13 | 2 | - | - | 15 |
|  | PO | 4 | 12 | - | - | 16 |
| GOBIIDAE Species 1 <br> Species 2 | PR | 2 | 10 | - | 1 | 13 |
|  | PO | 10 | 71 | 1 | 3 | 85 |
| Species 3 | PO | 1 | 3 | - | - | 4 |
| Species 4 | PO | - | 1 | 1 | 2 | 4 |
| Species 5 | PO | - | 1 | - | - | 1 |
| LUTJANIDAE | PO | 3 | - | - | 1 | 4 |
| OPHIDIIDAE | PO | 2 | 1 | - | - | 3 |
| HAEMULIDAE | PO | 8 | 4 | - | - | 12 |
| MUGILIDAE Mugil sp. | PO | 1 | 1 | 6 | 2 | 10 |
| POMACENTRIDAE | PO | - | - | 1 | - | 1 |
| SCARIDAE | PO | 2 | 2 | - | - | 4 |
| SCORPAENIDAE | PO | 1 | - | - | - | 1 |
| $\begin{array}{ll}\text { SERRANIDAE } & \text { Epinephelus itajara } \\ & \text { Hypoplectrus sp. }\end{array}$ | PO | - | - | 3 | - | 3 |
|  | PO | - | 1 | - | - | 1 |
| SPARIDAE Archosargus rhomboidalis | PO | 101 | 24 | 11 | 21 | 157 |
| SPHYRAENIDAE Sphyraena barracuda | J | - | - | 2 | 2 | 4 |
| SYNGNATHIDAE | PO | 7 | 1 | - | - | 8 |
| SYNODONTIDAE Synodus sp. | PO | 2 | - | - | - | 2 |
| Undetermined larvae | PR | 1000 | 16 | 8 | 1 | 1025 |
|  | PO | 3 | 1 | - | - | 4 |
| Total No. of Taxa |  | 34 | 31 | 21 | 20 | 45 |
| Total No. of Larvae |  | 5118 | 1561 | 258 | 405 | 7342 |
| Preflexion |  | 3614 | 178 | 24 | 4 | 3820 |
| Postflexion |  | 1504 | 1383 | 234 | 279 | 3400 |
| Juveniles |  | 215 | 460 | 120 | 137 | 932 |
| No. of Samples |  | 27 | 30 | 45 | 51 | 127 |

sity and abundance of larval fishes was also found at nonestuarine stations by Powell et al. (1989) in Florida Bay, where spawning (based on preflexion larval abundance) occurred in intermediate to high salinities. Flores Coto et al. (1983) found most larvae in Tamiahua Lagoon (western Gulf of Mexico) to have originated there and few larvae entered the lagoon from nearshore waters. Within Tamiahua Lagoon the greatest abundance of larvae was in the center of the lagoon away from shoreline habitats. The number of fish eggs decreased with increasing estuarine conditions in mangrove areas of Kenya (Little et al. 1988) and India (Krishnamurthy and Jeyaseelan 1981). There was also a gradient from high to low abundance of larval fish from the mouth to upper reaches of a mangrove creek in Kenya (Little et al. 1988).

Patterns of larval fish abundance in tropical nearshore island habitats may be different from tropical or temperate estuaries where migration into the estuaries from nearshore waters is more typical (Weinstein 1979; Weinstein et al. 1980; Shaw et al. 1988). The predominance of reef-associated species and high salinity conditions in tropical nearshore island habitats may account for a limited coupling with the shelf fish assemblage.

Haemulid and lutjanid larvae were noticeably absent from our collection, even though they comprised about $55 \%$ of juvenile fishes in mangroves of this area (Dennis, in press). It is possible that these taxa were not attracted to light at the stage they entered the prop-root habitat or settled in other nearby habitats and migrated to the prop roots. In Florida Bay, few snapper larvae were collected in shallow water areas, but preflexion larvae were found at the shelf edge near coral reefs (Powell et al. 1989).

It is also possible that many taxa may recruit to nearshore habitats in short duration periods (ca. 1-3 days) which are easily missed by monthly sampling (Doherty 1987). We observed this phenomenon in September 1988 when three jewfish (Epinephelus itajara) larvae were collected at the mangrove lagoon station. The following night we sampled at the marine station dock on Isla Magueyes (a mangrove-fringed island about 3.5 km east of lagoon M) (Fig. 1) and collected three additional jewfish larvae. The third night no jewfish larvae were collected. Though the complete duration of the jewfish recruitment event is not known, probably only fortuitous sample timing leads to capture of this and possibly other species.

Smith et al. (1987) described the nearshore assemblage of fish larvae as differing from that found offshore by being composed of morphologically unspecialized forms that may spend their complete larval phase nearshore. The nearshore larval fish assemblage collected off southwestern Puerto Rico fits this description as it was composed of common shallow-water fish families with few specialized larval forms (e.g., bonefish leptocephali, jewfish larvae). Without synoptic sampling across the shelf we are unable to estimate what proportion of the larval fish assemblage
might have originated and remained in the nearshore environment, but several taxa, such as bonefish, sardine, jewfish, and lutjanids were collected only at late development stages, an incident which suggests recruitment from outside the nearshore environment. Cross-shelf sampling will be needed before the source of some taxa can be determined.

The lack of many larval fish taxa in mangroves might be attributed to the numerous piscivore predators that reside in mangrove prop roots (Dennis, in press) and to periodically poor environmental conditions. Sluggish water movement and high biological oxygen demand can result in occasionally low oxygen conditions in the mangroves. One incident of depressed dissolved oxygen level ( 2.5 ppm ) was measured near bottom at night in the mangrove prop roots. The effect of low oxygen on ichthyoplankton should be ascertained before there is further judgment on the quality of mangrove areas as nursery grounds for larval fishes.

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## Citations

## Ahlstrom, E.H.

1971. Kinds and abundance of fish larvae in the eastern tropical Pacific based on collections made on EASTROPAC I. Fish. Bull., U.S. 69:3-77.
1972. Kinds and abundance of fish larvae in the eastern tropical Pacific on the second multi-vessel EASTROPAC survey, and observations on the annual cycle of larval abundance. Fish. Bull., U.S. 70:1153-1242.

Bulkowski, L., and J.W. Meade.
1983. Changes in phototaxis during early development of walleye. Trans. Am. Fish. Soc. 112:445-447.
Collins, I.A., and J.H. Finucane.
1984. Ichthyoplankton survey of the estuarine and inshore waters of the Florida Everglades, May 1971 to February 1972. NOAA Tech. Rep. NMFS SSRF 6, 75 p.
Dennis, G.D.
In press. Island mangrove habitats as spawning and nursery areas for commercially important fishes in the Caribbean. Proc. Gulf Carib. Fish. Inst. 41.
Doherty, P.J.
1987. Light-traps: selective but useful devices for quantifying the
distributions and abundances of larval fishes. Bull. Mar. Sci. 41:423-431.
Flores-Coto, C., F. Barba-Torres, and J. Sánchez-Robles.
1983. Seasonal diversity, abundance, and distribution of ichthyoplankton in Tamiahua Lagoon, western Gulf of Mexico. Trans. Am. Fish. Soc. 112:247-256.
Goulet, D., G.D. Dennis, and J.R. Rooker.
1988. The use of a night-light, lift-net device to sample larval fishes in tropical nearshore waters. Dep. Mar. Sci., Univ. Puerto Rico, Mayagüez, Puerto Rico. Unpubl. manuscr., 26 p.
Gregory, R.S., and P.M. Powles.
1988. Relative selectivities of Miller high-speed samplers and light traps for collecting ichthyoplankton. Can. J. Fish. Aquat. Sci. 45:993-998.
Heald, E.J., and W.E. Odum.
1970. The contribution of mangrove swamps to Florida fisheries. Proc. Gulf Carib. Fish. Inst. 22:130-135.
Kohn, A.J., and A.C. Riggs.
1982. Sample size dependence in measures of proportional similarity. Mar. Ecol. Prog. Ser. 9:147-151.
Krishnamurthy, K., and M.J. Jeyaseelan.
1981. The early life history of fishes from the Pichavarum mangrove system of India.Rapp. P.-v. Reun. Cons Int. Expl. Mer 187: 416-423.
Leis, J.M., and J.M. Miller.
1976. Offshore distribution patterns of Hawaiian fish larvae. Mar. Biol. (Berl.) 36:359-367.
Leis, J.M., and D.S. Rennis.
1983. The larvae of Indo-pacific coral reef fishes. Univ. Hawaii, Honolulu, 269 p.
Little, M.C., P.J. Reay, and S.J. Grove.
1988. Distribution gradients of ichthyoplankton in a eastern African mangrove creek. Estuarine Coastal Shelf Sci. 26:669-677.
Powell, A.B., D.E. Hoss, W.F. Hettler, D.S. Peters, and S. Wagner. 1989. Abundance and distribution of ichthyoplankton in Florida Bay and adjacent waters. Bull. Mar. Sci. 44:35-48.
Powles, H.
1975. Abundance, seasonality, distribution, and aspects of the ecology of some larval fishes off Barbados. Ph.D. Diss., McGill

Univ., 227 p.
1977. Description of larval Jenkinsia lamprotaenia (Clupeidae, Dussumieriinae) and their distribution off Barbados, West Indies. Bull. Mar. Sci. 27:788-801.
Richards, W.J.
1984. Kinds and abundances of fish larvae in the Caribbean Sea and adjacent areas. NOAA Tech. Rep. NMFS SSRF 776, 54 p.
Shaw, R.F., B.D. Rogers, J.H. Cowan, and W.H. Herke.
1988. Ocean-estuary coupling of ichthyoplankton and nekton in the northern Gulf of Mexico. Am. Fish. Soc. Symp. 3:77-89.
Smith, C.L., J.C. Tyler, and L. Stillman.
1987. Inshore ichthyoplankton: a distinctive assemblage? Bull. Mar. Sci. 41:432-440.
Sneath, P.H.A., and R.R. Sokal.
1973. Numerical taxonomy. W.H. Freeman \& Co., San Francisco, CA, 53 p.
Sokal, R.R., and F.J. Rohlf.
1981. Biometry. 2nd Ed. W.H. Freeman \& Co., San Francisco, CA, 859 p.
Thayer, G.W., D.R. Colby, M.A. Kjelson, and M.P. Weinstein.
1983. Estimates of larval-fish abundance: diurnal variation and influences of sampling gear and towing speed. Trans. Am. Fish. Soc. 112:272-279.
Weinstein, M.P.
1979. Shallow marsh habitats as primary nurseries for fishes and shellfishes, Cape Fear River, North Carolina. Fish. Bull., U.S. 77:339-357.
Weinstein, M.P., S.L. Weiss, R.G. Hodson, and L.R. Gerry.
1980. Retention of three taxa of postlarval fishes in an intensively flushed estuary, Cape Fear River, North Carolina. Fish. Bull., U.S. 78:419-436.

Wyatt, J.R.
1982. Survey of ichthyoplankton Part I. Section 2. Results of oceanographic and ichthyoplankton sampling. The distribution, abundance, and development of young Jamaican reef fishes. Scientific Report of the UWI-ODA Fisheries Ecology Research Project 1974-1979. Res. Rep. No. 6, Dep. Zool. Univ. West Indies, 122 p.

# Microhabitat and Diet Segregation among Coexisting Young-of-Year Sunfishes (Centrarchidae) 

JAMES B. LAYZER* and MICHAEL D. CLADY**<br>U.S. Fish and Wildlife Service<br>Oklahoma Cooperative Fish and Wildlife Research Unit<br>Oklahoma State University<br>Stillwater, Oklahoma 74078


#### Abstract

Young-of-year (yoy) sunfishes (Lepomis spp.) often remain in vegetated habitats in order to avoid predators and thereby limit opportunities to partition resources. Minnow traps were used to collect yoy of five species of sunfishes to determine their diet and use of microhabitats (type of vegetation, water depth, and substrate type) in Lake Rush, Oklahoma. The relative importance of each of these niche dimensions in separating species varied between years (1980 and 1981) in response to changes in the abundance and composition of aquatic macrophytes. In 1980 the type of vegetation was important in separating nine of ten species pairs, but only three pairs had low overlap along this dimension in 1981. Use of water depth effectively separated warmouth (L. gulosus) from redear sunfish (L. microlophus), longear sunfish (L. megalotis), and green sunfish (L. cyanellus) in 1980; but depth distributions of redear sunfish, longear sunfish, and warmouth were more similar in 1981. Longear sunfish was the only species that used mainly coarse substrates in both years. Although estimates of species segregation by diet may be conservative, dietary differences among five species pairs in 1980, and four pairs in 1981, were substantial. Niche complementarity was evident in both years. Bluegills ( $L$. macrochirus) and redear sunfish had nearly total overlap for substrate type but overlapped little for use of vegetation in 1980 and diet in 1981.


## Introduction

Resource partitioning among coexisting fish species in aquatic communities has been widely studied (Ross 1986). Such differential use of resources may be due to many factors. Among sunfishes (Lepomis spp.), adults and subadults clearly partition resources (Werner et al. 1977; Laughlin and Werner 1980), but there is little evidence that young-of-year (yoy) of these species do so. In fact, Laughlin and Werner (1980) and Mittelbach (1984) found that young of all species of sunfish were confined to vegetation. They suggested that vegetation provided a refuge from predators, and that confinement to vegetation could increase the intensity of interspecific interactions. Because it is always profitable for species to avoid competition when possible (Pianka 1976), overt competition is rarely observed in

[^6]established communities; instead, postcompetitive relationships, whereby coexisting species divide resources, are more evident than instances of apparent competition. The common coexistence of sunfishes in North American waters suggests that these species, including their young, may have resolved conflicts in resource exploitation.

Five species of yoy sunfishes often were collected from Lake Rush, southwestern Oklahoma, in the same seine haul over a seemingly homogeneous area: bluegill ( $L$. macrochirus); redear sunfish (L. microlophus); longear sunfish (L. megalotis); warmouth (L. gulosus); and green sunfish ( $L$. cyanellus). A cursory examination of the stomach contents from several individuals of each species suggested that they ate similar foods. Growth of centrarchids during the first year of life in Lake Rush is slow; mean total lengths of yoy sunfishes collected in late October varied from 34 mm for bluegills to 43 mm for longear sunfish (Layzer 1982). Since these sizes are below average for Oklahoma (Mense 1976), it could be inferred that some resource (presumably food) was limiting fish production. If so, resources might be partitioned to allow continued coexistence. We here attempt to resolve the apparent contra-
diction between our preliminary observations of similar patterns of resource use by sympatric yoy sunfishes and the expected partitioning of resources under an assumed regime of a limited resource.

## Study Area

Lake Rush, located in the Wichita Mountains National Wildlife Refuge in southwest Oklahoma, is a 20.9-ha impoundment on Blue Beaver Creek. Maximum depth is 10 m and average depth is 4.8 m . Typically, flow from Blue Beaver Creek and other tributaries stop by early to midJuly. Lake Rush becomes thermally stratified by mid-June, with near anoxic conditions in the hypolimnion (depth 3 $\mathrm{m})$. Secchi disc depth averaged $1.45 \pm 0.05 \mathrm{~m}$ during the summers of 1979 and 1980. The elongated basin of the impoundment lies along an east-west axis; littoral areas are primarily along the northern and southern shorelines. The southern shoreline has a steeply sloping bottom; a firm, heterogeneous substratum consisting of a mixture of particles ranging from coarse sand to large boulders, and a few small isolated patches of water milfoil (Myriophyllum sp.). The northern side has a gently sloping bottom with a sandy-silt substratum overlain with fine organic detritus and dense stands of aquatic macrophytes.

The distribution and composition of the submergent vegetation changed markedly during the study as a result of an application of 2,4-D herbicide crystals by refuge personnel in April 1981. In 1980 strikingly apparent zones of the plant community were each dominated by a single species. The emergent zone was 5 to 10 m wide and extended from the shoreline out to a depth of about 75 cm . This area was densely populated by spikerush (Eleocharis sp.). Unlike lakes with stable water levels, the emergent zone was bordered by a submergent zone 3 to 5 m wide out to a depth of 110 cm , dominated by muskgrass (Chara sp.). This zone was apparently created by low lake levels which occurred from late summer to early spring during most years. Continuous with this zone and out to a depth of 1.8 to 2.5 m was a 30 to 60 m wide band of water milfoil interrupted by isolated patches of pondweed (Potamogeton spp.); American lotus (Nelumbo lutea); and coontail (Ceratophyllum demersum). The outer band of vegetation, which was more irregular, was composed entirely of coontail to a depth of 2.5 to 2.8 m . These water depths and associated plant distributions existed only during portions of 1980 when the lake was full. Since the source of water is primarily surface runoff, the lake level varies seasonally. In 1979 and 1980, levels were maximum in the spring and decreased throughout the summer, gradually exposing the emergent zone and most of the muskgrass by early fall.

The herbicide $2,4-\mathrm{D}$, which was applied only over the submergent zone, eliminated virtually all of the water milfoil in 1981 but had little effect on coontail. In June

1981, dense patches of coontail were noted in the same locations where it grew in 1980, including areas previously surrounded by water milfoil. During the summer of 1981, the zone which was occupied by water milfoil in 1980 was gradually colonized by muskgrass in the shallower areas and by coontail in deeper areas until nearly all of the suitable habitat (fine substrate) was occupied. In addition to categorizing each zone by the dominant species of macrophyte, zones of coontail were separated into dense ( $>8 \mathrm{~kg} / \mathrm{m}^{2}$; wet weight) and sparse stands ( $<3 \mathrm{~kg} / \mathrm{m}^{2}$ ).

Lake Rush contained a diverse fish assemblage of 17 species dominated by centrarchids (Layzer 1982). Largemouth bass (Micropterus salmoides) was the primary piscivorous fish and made up 5 to $23 \%$ of all fish collected by electrofishing from 1979 to 1981 (Layzer and Clady 1981). The majority of largemouth bass were 175 to 275 mm (total length) and dietary analysis ( $n=114$ ) revealed that sunfishes made up $91 \%$ of the total fishes consumed (Layzer et al. 1983).

## Materials and Methods

## Fish Collections

Sunfishes were collected during eight periods from June to October, 1980, and during seven periods from June to September, 1981. Each sampling period lasted from 2 to 5 days and periods were separated by 7 to 18 days. Fish were caught in unbaited cylindrical minnow traps ( $6.4-\mathrm{mm}$ wire mesh) which had a funnel opening ( 2.5 cm ) at both ends. Initially, traps were set on the bottom for approximately 24 hours; but beginning in mid-August 1980, and continuing for the remainder of the study, traps were set for an average of 1.62 hours (S.D. $\pm 0.53$ ).

When each trap was set, water depth was measured to the nearest 10 cm with a metal measuring rod. In shallow areas, substrates were classified visually as fine (principally organic) or coarse (mixture of particle sizes ranging from sand to boulder), and vegetation was pulled up by hand and identified to genus using keys of Fassett (1969) and Muenscher (1967). In deeper areas, substrate was determined by sounds transmitted by the metal measuring rod. Typically, substantial amounts of water milfoil or coontail (the only plants occurring in deeper water) were retrieved with the traps. In 1981 quantitative plant samples ( $0.2 \mathrm{~m}^{2}$ ) were collected at all depths (Layzer 1982). These vegetation samples were dominated by a single species; in fact, $92 \%$ of the samples contained only one species. Samples taken in areas classified as having no vegetation did not contain any macrophytes. Distances between adjacent traps varied considerably but were at least 15 m . Traps were set only in habitats that appeared to be homogeneous over an area $10 \mathrm{~m}^{2}$.

Fish collected from each trap were preserved separately in $10 \%$ formalin; later they were measured to the nearest

Table 1
Mean number ( $\pm 1$ S.E.) of young-of-year sunfishes collected per trap for each sampling period in 1980 (sampling period: E $=$ early; $\mathrm{M}=$ middle; $\mathrm{L}=$ late $)$.
$\left.\begin{array}{lcccccc} \\ & & & & & & \text { Mean catch per trap }\end{array}\right]$
millimeter (total length), and identified using descriptions and keys in Cross (1967); Miller and Robison (1973); Pflieger (1975); as well as species-specific characteristics described in Layzer (1982).

## Resource Use

Except for six yoy green sunfish captured by electrofishing, all sunfishes used to analyze diets were captured in minnow traps set for 2.2 hours or less. Stomach contents were examined under a dissecting microscope and identified to the lowest possible taxon. Widths of all prey items in a stomach (up to a maximum of 20 of one type) were measured with an ocular micrometer to the nearest 0.1 mm .

For analysis of depth utilization, all data for each year were combined into five depth categories (cm): $\leqslant 50$, $51-100,101-150,151-200$, and $>200$. To compensate for unequal sampling effort, data were standardized by calculating a catch per unit effort (CPUE) for each of the depth categories for each species as follows: CPUE $_{i k}=$ number of species $k$ collected in the $i$ th resource state/ number of traps set in the $i$ th resource state. Chi-square tests for differences in probability distributions (Conover 1971) were used to test for differences in the depth and vegetational distributions among species. When more than $20 \%$ of the expected values in a contingency table were $<5$, adjacent categories were combined. For each species, chi-square goodness of fit tests (Conover 1971) were used to test the null hypothesis that the observed catch distributions were the same as the distributions of trapping effort among resource states. The proportion of use $\left(p_{i k}\right)$ of a given resource state was found by

$$
p_{i k}=\mathrm{CPUE}_{i k} / \sum_{i}^{n} \mathrm{CPUE}_{i k},
$$

where $n$ is the number of states in a resource set. The same procedure was used for determining the proportional use of the substrate and vegetation dimensions. The $p_{i k}$ 's obtained in this manner were then used to calculate niche breadth and overlap for each of the habitat dimensions. Niche breadth ( $B$ ) was calculated by the formula of Levins (1968) as modified by Pianka (1973):

$$
B=\left(1 / \sum_{i}^{n} p i k^{2}\right) / n
$$

A proportional overlap value $(O)$ was calculated following Schoener (1968) to determine the similarity in resource usage between pairs of species:

$$
O=1-0.5 \sum_{i}^{n}\left|p_{i j}-p_{i k}\right|
$$

## Results

## Relative Abundance

In 1980, 1857 yoy sunfishes were caught in minnow traps (Table 1). Bluegills and redear sunfish were codominant, together making up $91 \%$ of the catch. These species also dominated seine collections in 1979 and 1980, but bluegills were 30 to $50 \%$ more abundant than redear sunfish (Layzer and Clady 1981). Differences in relative abundance between gears apparently were due to size selectivity. Seines captured more yoy bluegills $<30 \mathrm{~mm}$ TL, while larger yoy bluegills were more abundant in traps (Layzer 1982). Minnow traps also selected for slightly larger redear sunfish; however, by late August 1980, only $24 \%$ of the

Table 2
Mean number ( $\pm 1$ S.E.) of young-of-year sunfishes collected per trap for each sampling period in 1981 (sampling period: E
$=$ early; $\mathrm{M}=$ middle; $\mathrm{L}=$ late $)$.

| Sampling period | Number of traps | Mean catch per trap |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Bluegill | Redear sunfish | Longear sunfish | Warmouth | Green sunfish |
| June (M) | 123 | $0.04 \pm 0.03$ |  |  |  |  |
| July (E) | 96 | $0.10 \pm 0.04$ |  |  |  |  |
| July (M) | 142 | $0.43 \pm 0.07$ | $0.09 \pm 0.03$ | $0.02 \pm 0.01$ | $0.06 \pm 0.02$ |  |
| August (E) | 142 | $1.54 \pm 0.17$ | $0.44 \pm 0.09$ | $0.15 \pm 0.04$ | $0.15 \pm 0.04$ | $0.02 \pm 0.01$ |
| August (M) | 96 | $1.97 \pm 0.29$ | $0.86 \pm 0.18$ | $0.22 \pm 0.08$ | $0.21 \pm 0.05$ |  |
| September (E) | 94 | $2.81 \pm 0.35$ | $1.28 \pm 0.22$ | $0.10 \pm 0.05$ | $0.15 \pm 0.05$ | $0.02 \pm 0.02$ |
| September (L) | 96 | $3.41 \pm 0.45$ | $1.07 \pm 0.23$ | $0.08 \pm 0.04$ | $0.36 \pm 0.11$ |  |
| Total fish collected |  | 1074 | 381 | 62 | 100 | 5 |



Figure 1
Depth distributions (adjusted for trapping effort) of five species of sunfish collected in minnow traps. Sample sizes are in parentheses.
redear sunfish collected by seining were $<30 \mathrm{~mm}$, while $74 \%$ of the bluegills in seine hauls were $<30 \mathrm{~mm}$. No evidence of selection against the largest yoy sunfishes was observed. In fact, minnow traps captured yearling sunfish of all species up to about 80 mm TL. If size were the only
component of gear selectivity, then CPUE of minnow traps probably was an underestimation of the relative abundance of bluegills $<30 \mathrm{~mm}$.

In 1981, 1622 yoy sunfishes were captured in minnow traps. Bluegills and redear sunfish again dominated, but
bluegills were nearly three times as abundant as redear sunfish (Table 2). Differences in relative abundance of species between years were primarily due to the significant decrease ( $P<0.001$ ) in numbers of redear sunfish in 1981. Abundance of longear sunfish also declined in 1981 ( $P<0.01$ ), but the GPUE of bluegills and warmouths did not change ( $P>0.10$ ).

Young-of-year sunfishes were first captured in minnow traps in late July 1980 (Table 1). Except for what may have been a sampling artifact in September, CPUE of redear sunfish reached a peak in early August and declined steadily thereafter. In contrast to the trend for redear sunfish, abundance of bluegills steadily increased from early in the season until early October.

Bluegills were captured earlier in 1981 than in 1980, but otherwise their numbers increased as in 1980 (Table 2). In 1981 redear sunfish first appeared in trap catches in midJuly; numbers then increased progressively, peaking in early September. The differences between years suggest that redear sunfish fry hatched later in 1981 than in 1980. Although adult redear sunfish were observed on nests in early April of both years, nests were deserted for an unknown period of time in 1981 when water temperatures dropped from $21^{\circ} \mathrm{C}$ to $18^{\circ} \mathrm{C}$ between 9 and 10 April. During this same time, refuge personnel applied the herbicide 2,4-D to Lake Rush. Although male bluegills also deserted nests when the temperature dropped, bluegills reproduced normally during this same period (Layzer 1982). No discernible pattern of abundance was apparent for longear sunfish, warmouth, and green sunfish during either year. Fluctuations in CPUE for these species were relatively small and may have resulted from the combined effects of recruitment, mortality, and sampling error.

## Habitat Utilization

The depth distributions of yoy sunfishes differed among species and between years (Fig. 1). In 1980, all species except warmouth were caught proportionately more often in traps set at depths of 50 cm or less. Distribution of bluegills was bimodal with high abundances in the shallowest habitats and at depths between 151 and 200 cm . The concentration of warmouths in deep water generally complemented the distributions of other species. In 1981, all species made greater use of deeper habitats. Although the distribution of redear sunfish was inversely related to depth each year, the proportion collected at depths of 50 cm or less was only 0.32 in 1981 compared to 0.75 in 1980. Longear sunfish used decidedly different depths each year; in 1980 their distribution was unimodal and inversely related to depth, whereas in 1981 their distribution was bimodal and fish were most abundant at depths of 50 cm or less and between 101 and 150 cm . Although few green sunfish were collected each year, they apparently did not use depths greater than 150 cm in 1981. In 1980

Table 3
Proportional use (adjusted for trapping effort) of substrate types by young-of-year sunfishes in Lake Rush in 1980 and 1981.

|  |  | Substrate |  |
| :--- | :--- | :--- | :--- |
|  |  | Fine | Coarse |
| Sluegill | Year | 0.72 | 0.28 |
|  | 1980 | 0.76 | 0.24 |
| Redear sunfish | 1981 | 0.75 | 0.25 |
|  | 1980 | 0.71 | 0.29 |
| Longear sunfish | 1981 | 0.03 | 0.97 |
|  | 1980 | 0.17 | 0.83 |
| Warmouth | 1981 | 0.79 | 0.21 |
|  | 1980 | 0.83 | 0.17 |
| Green sunfish | 1981 | 1.00 | 0.00 |
|  | 1980 | 0.83 | 0.17 |

green sunfish were collected only from the shallowest habitats.

Longear sunfish were clearly segregated from other species on the basis of substrate (Table 3). Each year longear sunfish used primarily habitats with a coarse substrate; other species were most abundant in areas having a fine organic substrate.

Vegetation was also important in the habitat segregation of yoy sunfishes in Lake Rush. In 1980 most bluegills were collected in habitats containing coontail and water milfoil, but many bluegills were also collected from areas where muskgrass and quillwort (Isoetes sp.) grew (Table 4). With the disappearance of water milfoil in 1981, bluegills greatly increased their use of coontail stands. Each year the greatest densities of redear sunfish occurred in habitats where spike rush and muskgrass grew. Microhabitats lacking vegetation were used extensively by longear sunfish each year; in 1981, however, proportionately more longear sunfish were collected by traps set in coontail stands. Although longear sunfish used sparse stands of coontail twice as much as dense stands, the difference was not significant ( $\chi^{2}=2.25,1 \mathrm{df}, P>0.10$ ).

In 1980 proportionately more warmouths were collected from habitats containing coontail and American lotus. In 1981 significantly more warmouths were collected from dense stands (wet biomass $>8 \mathrm{~kg} / \mathrm{m}^{2}$ ) of coontail compared to sparse stands (wet biomass $\left.<3 \mathrm{~kg} / \mathrm{m}^{2}\right)\left(\chi^{2}=\right.$ $13.10,1 \mathrm{df}, P<0.001$ ). Nearly all green sunfish were collected from habitats containing muskgrass or spikerush.

## Feeding Periodicities

All fishes used in dietary analyses were collected between 0600 and 2100 hours. The mean number of prey per

Table 4
Proportional use (adjusted for trapping effort) of vegetation types by young-of-year sunfishes in Lake Rush in 1980 and 1981.

Figure 2
Mean number of prey items in sunfish stomachs (solid lines), and proportion (adjusted for trapping effort) of each species of sunfish collected in minnow traps over time of day (broken line).
stomach was used as a measure of foraging intensity. Data were separated into five 3-h-long periods based on the time of day traps were set. Because few green sunfish were examined and most were captured during the first and last
time periods, this species was excluded from this analysis. The remaining species consumed high numbers of prey in mid-morning and late afternoon or evening (Fig. 2). The fewest prey were found in fish of all species caught in the

Table 5
Variation in diets ( $\%$ numbers) among three individual bluegills and a comparison of diets for 10 individual bluegills collected on the same date and diets of all bluegills collected in 1981. The three individual bluegills were collected by the same minnow trap fished between 1850 and 2005 hours at a depth of 110 cm over a fine substrate in a dense stand of coontail on 1 September 1981.

|  | Total length (mm) | $N$ | Taxa |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Daphnia |  | Ostracoda | Chironomidae | Copepoda | Chydoridae | Other |
| Individual A | 28 | 1 | 0.17 |  | 0.02 | 0.47 | 0.12 | 0.20 | 0.02 |
| Individual B | 31 | 1 | 0.23 |  | 0.03 | 0.12 | 0.48 | 0.17 | 0.01 |
| Individual C | 35 | 1 | 0.12 |  | 0.14 | 0.38 | 0.16 | 0.18 | 0.01 |
| All bluegills on September 1, 1981 |  | 9 | 0.14 |  | 0.08 | 0.26 | 0.22 | 0.26 | 0.05 |
| All bluegills collected in 1981 |  | 62 | 0.16 |  | 0.15 | 0.23 | 0.17 | 0.25 | 0.04 |

Table 6
Percentage abundance of each prey item in diets of young-of-year sunfishes in Lake Rush by year.

| Species | Year | $N$ |  | Taxa |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Daphnia sp. | Ostracoda | Chironomidae | Copepoda | Chydoridae | Other |
| Bluegill | 1980 | 75 | 14.4 | 19.2 | 20.1 | 6.8 | 33.5 | 6.0 |
|  | 1981 | 62 | 16.0 | 15.2 | 22.6 | 16.6 | 25.5 | 4.1 |
| Redear sunfish | 1980 | 66 | 0.7 | 55.9 | 12.9 | 4.6 | 24.1 | 1.8 |
|  | 1981 | 54 | 1.1 | 84.2 | 7.7 | 3.2 | 1.7 | 2.1 |
| Longear sunfish | 1980 | 43 | 0.0 | 62.2 | 12.4 | 2.6 | 22.0 | 0.8 |
|  | 1981 | 43 | 1.1 | 55.3 | 17.4 | 8.1 | 15.3 | 2.8 |
| Warmouth | 1980 | 45 | 18.9 | 13.7 | 58.4 | 3.2 | 1.6 | 4.2 |
|  | 1981 | 48 | 50.1 | 4.7 | 11.2 | 23.6 | 1.7 | 8.7 |
| Green sunfish | 1980 | 4 | 0.0 | 0.0 | 40.0 | 0.0 | 0.0 | 60.0 |
|  | 1981 | 8 | 0.0 | 57.5 | 5.5 | 4.8 | 28.3 | 3.9 |

middle of the day. Only bluegills fed mostly near dusk; other species foraged earlier in the evening.

Movement of fishes (as measured by trap catches) generally paralleled trends in feeding intensity, though the greatest catches of bluegills occurred one time period before the peak in stomach contents (Fig. 2). Maximum numbers of prey in warmouth occurred in the morning; in contrast, catches of warmouth in traps were greatest late in the day.

## Prey Utilization

A total of 24 types of prey were identified from stomach contents. Most prey items were identified to family (e.g., Chironomidae and Chydoridae) or subclass (e.g., Ostracoda, Copepoda). This absence of taxonomic detail in description of stomach contents yielded conservative estimates of diet separation among sunfishes. Consequently, species with high diet overlap may have greater differences
in diets than we estimated and, in like manner, dietary differences may be more pronounced.

No seasonal trends in diets were observed. However, the relative and absolute importance of various prey items did fluctuate over time for a given species and between species for the same sampling period. The variable diets of individual fish collected together (Table 5), suggested that dietary differences between sampling periods could have resulted from chance. Since diets varied among individuals within a sample as much as between samples (Layzer 1982), all food habits data were combined by year. Five prey types generally accounted for over $90 \%$ of all food items (Table 6). They were the same prey types that we observed in stomachs of fishes collected by seining but we did not count them. All species of sunfishes ate similar prey, though in different proportions. Ostracods were the most abundant item in the diets of both redear sunfish and longear sunfish. Other sunfishes also ate many ostracods. Diets of warmouths and green sunfish were the most

| Niche breadth of yoy sunfishes in Lake Rush for various resource dimensions. |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Resource dimension |  |  |  |
| Species | Year | Substrate | Vegetation | Depth | Diet |
| Bluegill | 1980 | 0.83 | 0.56 | 0.71 | 0.76 |
|  | 1981 | 0.78 | 0.49 | 0.94 | 0.86 |
| Redear sunfish | 1980 | 0.81 | 0.38 | 0.29 | 0.44 |
|  | 1981 | 0.86 | 0.27 | 0.80 | 0.23 |
| Longear sunfish | 1980 | 0.53 | 0.16 | 0.41 | 0.37 |
|  | 1981 | 0.71 | 0.35 | 0.62 | 0.49 |
| Warmouth | 1980 | 0.76 | 0.31 | 0.60 | 0.42 |
|  | 1981 | 0.71 | 0.49 | 0.68 | 0.51 |
| Green sunfish | 1980 | 0.50 | 0.26 | 0.17 | 0.32 |
|  | 1981 | 0.71 | 0.30 | 0.59 | 0.40 |

variable between years. For instance, daphnids made up only $19 \%$ of the diet of warmouths in 1980 but accounted for $50 \%$ in 1981. The five major food items were consumed more evenly by bluegills each year; no single item made up more than $34 \%$ of their diets.
All species ate prey of similar size, and differences between species in their utilization functions were simply a reflection of the type of prey consumed rather than a result of selecting different sizes of the same prey type. For example, the average width of food items in redear sunfish stomachs was 0.35 mm because ostracods, which made up 56 to $84 \%$ of the diet of redear sunfish each year, averaged 0.39 mm in width (Layzer 1982).

## Niche Breadth

Bluegills generally showed the broadest resource utilization although niche breadth changed between years (Table 7). Change in depth use was most pronounced for redear sunfish; their niche breadth for depth increased from 0.29 to 0.80 . The increase in niche breadth of longear sunfish along the vegetation axis reflected substantially greater use of vegetated habitats in 1981 than in 1980.

The proportional use of the various prey types by redear sunfish changed the most of any species between years, but this change was in contrast to the changes noted in their relative use of habitats. Redear sunfish seemingly foraged over a greater range of habitats in 1981 than in 1980, but became more selective in their feeding.

## Niche Overlap

Longear sunfish showed little overlap (all values $\leqslant 0.31$ ) in 1980, and only slightly higher overlap in 1981 with other species along the substrate axis (Table 8). In contrast, overlap values for use of substrate between all other pairs
of species were high (all $\geqslant 0.72$ ) in both years. Overlap in depth utilization was moderately high ( 0.55 to 0.75 ) for six species pairs in 1980 and increased in 1981. There was strong complementarity between the use of vegetation and depth in 1980; five of the ten species pairs overlapped greatly along one habitat axis but little along the other. For example, redear sunfish and longear sunfish used similar depths ( $O=0.74$ ), but overlapped little ( $O=0.13$ ) in their use of vegetation. Spatial segregation was less obvious in 1981.

Longear sunfish and redear sunfish had nearly identical diets in 1980; though their diets diverged somewhat in 1981, overlap remained high. Most other species showed changes in diet overlap between years. For example, overlap between redear sunfish and bluegills decreased from 0.63 in 1980, to 0.31 in 1981. Most species appeared to segregate along the vegetation axis in 1980 when nine species pairs had overlaps $<0.42$. In contrast, only three species pairs had low overlap for use of vegetation in 1981.

## Discussion

Partitioning of resources among yoy sunfishes was evident in Lake Rush even though most species were confined to the vegetation. The importance of vegetation type, depth, and diet in segregating species changed between years, indicating a high level of behavioral plasticity for these species. In both years, depth utilization for redears and warmouths yielded essentially mirror images: redear sunfish were most abundant in the shallowest areas, and warmouths were more abundant in the deepest areas. In 1980 overlap between bluegills and redears was moderately high ( 0.61 and 0.63 ) for both depth and prey type but low for vegetation type. Moreover, diets of these species varied somewhat at different depths indicating that diets of these

Table 8
Habitat and diet overlaps among young-of-year sunfishes in Lake Rush. ( $\mathrm{BG}=$ bluegill; $\mathrm{RE}=$ redear sunfish; $\mathrm{LE}=$ longear sunfish; $W M=$ warmouth; GS $=$ green sunfish).

| Species | 1980 |  |  |  | 1981 |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Substrate | Vegetation | Depth | Diet | Substrate | Vegetation | Depth | Diet |
| BG vs. RE | 0.97 | 0.34 | 0.61 | 0.63 | 0.95 | 0.67 | 0.73 | 0.31 |
| BG vs. LE | 0.31 | 0.18 | 0.62 | 0.57 | 0.41 | 0.73 | 0.73 | 0.60 |
| BG vs. WM | 0.93 | 0.41 | 0.66 | 0.57 | 0.93 | 0.69 | 0.64 | 0.52 |
| BG vs. GS | 0.72 | 0.27 | 0.36 | 0.26 | 0.93 | 0.37 | 0.62 | 0.53 |
| RE vs. LE | 0.28 | 0.13 | 0.74 | 0.94 | 0.46 | 0.57 | 0.71 | 0.71 |
| RE vs. WM | 0.96 | 0.32 | 0.34 | 0.34 | 0.88 | 0.50 | 0.46 | 0.20 |
| RE vs. GS | 0.75 | 0.62 | 0.75 | 0.15 | 0.88 | 0.56 | 0.82 | 0.69 |
| LE vs. WM | 0.24 | 0.10 | 0.33 | 0.31 | 0.34 | 0.53 | 0.46 | 0.28 |
| LE vs. GS | 0.03 | 0.11 | 0.55 | 0.13 | 0.34 | 0.31 | 0.66 | 0.82 |
| WM vs. GS | 0.79 | 0.19 | 0.09 | 0.44 | 1.00 | 0.17 | 0.29 | 0.19 |

species diverged more when they occurred in the same habitats (Layzer 1982). In 1981 bluegills and redear sunfish were distinctly separated on the basis of diet even though their use of habitat was similar.
High overlap occurred between bluegills and warmouths for both habitat and diet. Perhaps these overlap values approximated the maximum tolerable overlap between these species (Pianka 1972) or these two species may have segregated along an additional dimension or by a finer resolution of diet or microhabitat than recognized. The second alternative is supported by the significantly greater use of dense versus sparse stands of coontail by warmouth while bluegills showed no such difference in use. Moreover, because our estimates of diet segregation are conservative, diets may have been more divergent than we recognized.
In both years, extensive use by longear sunfish of nonvegetated habitats with coarse substrate clearly separated them from other congeners. Together these two habitat dimensions effectively separated longear sunfish from the other four species of sunfishes, which lived primarily in vegetated areas with fine substrates. Use of exposed habitats in Lake Rush by yoy longear sunfish differs from their restriction to shallow vegetated areas in Michigan lakes (Laughlin and Werner 1980). Differences in the distribution of longear sunfish may reflect differences in community composition, specifically within the genus Lepomis. The changes in depth distribution of longear sunfish in 1981 indicated that the explanation for their distribution in vegetated and nonvegetated habitats is complex. Increased use of vegetated habitats by longear sunfish in 1981 was probably related both to the decrease in relative abundance of redear sunfish and to changes in the macrophyte community. The differences in depth distribution of redear sunfish and longear sunfish in 1981, and, more importantly, the greater shift towards deeper water by longear sunfish suggest they may have been avoiding areas of high redear abundance. Furthermore, longear sunfish also increased
the breadth of their diet in 1981, whereas redear sunfish restricted their diet, thus somewhat reducing niche overlap.

Young-of-year sunfish species, other than longear sunfish, in 1980 remained primarily in vegetation, presumably limiting their risk of predation by largemouth bass. Mittelbach (1988) demonstrated that confinement to the vegetation can result in exploitative competition among juvenile sunfishes. The low growth rates of sunfishes reported in Lake Rush by Layzer et al. (1983) suggest that intra- or inter-specific competition may have been occurring despite the observed partitioning of resources. Sunfish diets may be reflective of exploitative competition. The abundance of ostracods in sunfish stomachs was surprising because Vinyard (1979) found that only $30 \%$ of the ostracods ingested by bluegills were digested. Because of their apparent resistance to digestion, it is unlikely that ostracods were a preferred food item even though they were a major item in sunfish diets. Although differential digestion rates of prey may bias food-habits analysis (Gannon 1976), we do not believe that the observed high occurrence of ostracods in sunfish stomachs was due to their resistance to digestion. Head capsules of chironomids are also resistant to digestion; yet, they were uncommon in stomachs of most fishes. Further, ostracods also were abundant in the stomachs of seine caught fishes. We suggest that if yoy sunfishes were competing for food, it was on the basis of quality, and not necessarily quantity, of food items. Perhaps sunfishes are severely limited by food in Lake Rush but receive sufficient nutritional value from ostracods and other prey to meet their basic metabolic demands. Foraging on nutritionally poor prey may be a seasonal survival strategy.

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## Citations

Conover, W.J.
1971. Practical nonparametric statistics. John Wiley \& Sons Inc., New York, NY, 462 p.
Cross, F.B.
1967. Handbook of fishes of Kansas. Univ. Kansas, Museum of Natural History Misc. Publ. No. 45, 357 p.
Fassett, N.C.
1969. A manual of aquatic plants. Univ. Wisconsin Press, Madison, WI, 405 p .
Gannon, J.E.
1976. The effects of differential digestion rates of zooplankton by alewife, Alosa pseudoharengus, on determinations of selective feeding. Trans. Am. Fish. Soc. 105:89-95.
Laughlin, D.R., and E.E. Werner.
1980. Resource partitioning in two coexisting sunfish: pumpkinseed (Lepomis gibbosus) and northern longear sunfish (Lepomis megalotis peltastes). Can. J. Fish. Aquat. Sci. 37:1411-1420.
Layzer, J.B.
1982. Resource partitioning among young-of-year sunfish (Lepomis spp.), and phenotypic variation of bluegills (L. macrochirus). Ph.D. diss., Oklahoma State Univ., Stillwater, OK, 176 p.
Layzer, J.B., and M.D. Clady.
1981. Evaluation of the striped bass $\times$ white bass hybrid for controlling stunted bluegills. Proc. Ann. Conf. Southeast. Assoc. Wild. Fish Comm. 35:297-310.
Layzer, J.B., M.D. Clady, and O.E. Maughan.
1983. Factors influencing fish populations in Oklahoma lakes and
ponds. Final Report, Fed. Aid Proj. No. F-41-R, 117 p.
Levins, R.
1968. Evolution in changing environments, some theoretical explorations. Princeton Univ. Press, Princeton, NJ, 120 p.
Mense, J.B.
1976. Growth and length-weight relationships of twenty-one reservoir fishes in Oklahoma. Oklahoma Fish. Res. Lab. Bull. No. 13, 155 p .
Miller, R.J., and H.W. Robison.
1973. The fishes of Oklahoma. Oklahoma State Univ. Press, Stillwater, OK, 246 p .
Mittelbach, G.G.
1984. Predation and resource partitioning in two sunfishes (Centrarchidae). Ecology 65:499-513.
1988. Competition among refuging sunfishes and effects of fish density on littoral zone invertebrates. Ecology 69:614-623.
Muenscher, W.C.
1967. Aquatic plants of the United States. Cornell Univ. Press, Ithaca, NY, 374 p .
Pflieger, W.L.
1975. The fishes of Missouri. Missouri Dep. of Cons., Springfield, MO, 342 p .
Pianka, E.R.
1972. $r$ and $k$ selection or $b$ and $d$ selection? Am. Nat. 106: 581-588.
1973. The structure of lizard communities. Ann. Rev. Ecol. System. 4:53-74.
1976. Competition and niche theory. In Theoretical ecology: principles and applications (R.M. May, ed.), p. 114-141. W.B. Saunders, Philadelphia, PA.
Ross, S.T.
1986. Resource partitioning in fish assemblages: a review of field studies. Copeia 1986:352-388.
Schoener, T.W.
1968. The Anolis lizards of Bimini: resource partitioning in a complex fauna. Ecology 49:704-726.
Vinyard, G.
1979. An ostracod (Cypriodopsis vidua) can reduce predation from fish by resisting digestion. Am. Midl. Nat. 102:188-190.
Werner, E.E., D.J. Hall, D.R. Laughlin, D.J. Wagner, L.A. Wilsmann, and F.C. Funk.
1977. Habitat partitioning in a freshwater fish community. J. Fish Res. Board Can. 34:360-370.

# Comparative Scanning Electron Microscopy Studies of the Egg Envelopes of Six African Barbus and Three Pseudobarbus Species (Cyprinidae)* 

J.A. CAMBRAY<br>Albany Museum<br>Grahamstown 6140, South Africa


#### Abstract

The external morphology of the egg envelopes of six Barbus (B. aeneus, B. andrewi, B. anoplus, B. capensis, B. kimberleyensis, and B. viviparus) and three Pseudobarbus (P. afer, P. asper, and P. quathlambae) species (Cyprinidae) were studied using scanning electron microscopy. Conventional specimen preparation was used as well as the Cryo technique. Electron photomicrographs and a comparative description of the egg surface sculpturing of these nine species is given. The egg surface ultrastructure of all species is distinct. The honeycomb lattice surface structure of the whitefish ( $B$. andrewi), is more similar to that of $P$. afer and $P$. asper than to those of the three yellowfish species studied (B. aeneus, B. kimberleyensis, and B. capensis). The relict redfin minnow, P. quathlambae, which only occurs in the upper catchment of the Orange River in Lesotho, does not have a distinct honeycomb lattice. The results indicate that a knowledge of the ultrastructure of the egg envelopes of African cyprinids could be useful for identification of eggs collected from the wild.


## Introduction

Cambray and Teugels (1988) have noted that the early life history of the some 2000 African freshwater fishes has received very little attention. The reproductive behaviour of some of the species has been recorded (Lévêque and Daget 1984), however little is known of the ultrastructure of the egg envelopes of the majority of the 2000 species. Even in the North American Great Lakes only $34 \%$ of the fish species have had their eggs described (Werner 1976; Johnson and Werner 1986). Studies on the ultrastructure of the envelopes of eggs spawned in freshwater have been somewhat limited (Johnson and Werner 1986) compared to studies on marine fishes.

The egg envelopes of different species have been shown to be ornamented with filaments, spines, patterns of ridges, loops, blebs and pustules (Ahlstrom and Moser 1980; Laale 1980; Boehlert 1984; Matarese and Sandknop 1984). Possibly the morphogenetic activity of individual follicle cells generates these elaborate structures (Wourms 1976).

[^7]It has been shown for some taxa that these ornamentations and the ultrastructure of the egg envelopes are species specific (Lönning 1972; Ivankov and Kurdyayeva 1973). Therefore the surface structure of the egg envelope in some species can be a useful character for the identification of eggs of different species (Mikodina 1987). However, for many fish species the most difficult ontogenetic stage to identify is the egg and Markle and Frost (1985) argued that few can be identified to the species level. If eggs are found to be distinctive then they can be identified. Researchers will then be able to establish where spawning sites are for specific species, data which will be of value in ecological impact work. Characters associated with the egg envelope have been shown to be highly adapted to the environmental conditions in which the embryo develops (Ivankov and Kurdyayeva 1973; Laale 1980). Matarese and Sandknop (1984) have noted that the main goal of taxonomy with respect to fish eggs is identification and not speculation on the systematic significance of any characters present.

Since there is very little known about the early life history of African freshwater fishes the identification of the egg envelopes of some of the species may provide an important aid in the study of their life histories. In this paper one of the main goals has been to add another set of characters, that is the egg surface ultrastructure, to the present knowledge of African cyprinids. This paper
describes and compares the surface structure of the egg envelope of four large and two small Barbus species as well as three Pseudobarbus species. Scanning Electron Microscopy (SEM) was used as the analytical tool in this investigation as it provides the depth of field necessary to permit visualization of surface structures which are difficult to see using light microscopy.

## Methods

Laale (1980), Robertson (1981), Balon (1985), among others have pointed out the confusion and inconsistency in the literature concerning the terminology of fish egg coats. The chorion is the egg coat which envelopes the egg and, in protecting the developing embryo from the external environment (Johnson and Werner 1986), is the major protective coating of the ovum. In many papers the outermost egg envelope of fish eggs is called the chorion without regard to its origin (Shelton 1978). Balon (1985) cautioned that "If the existence of two coats is known, the inner coat should be called zona radiata (cortical coat) and the second or outer coat, chorion (Ivankov and Kurdyayeva 1973; Avni and Soin 1974). If not known, an indifferent term 'egg shell(s) or membrane(s)' should be used.'" In this paper, to prevent confusion, the covering of the egg is called the egg envelope or egg coat, as it is not known if there are inner and outer egg coats for the species studied. The term egg membrane is not used here, as a true membrane is a thin trilaminar cytoplasmic structure of lipids, proteins, and some carbohydrates that forms the boundary between a cell and its environment (Allaby 1985), whereas the covering of fish eggs is very thick and extracellular.

Eggs for the present study were taken from the growing special collection of early life-history stages housed at the Albany Museum. The eggs were taken from the following developmental series: smallmouth yellowfish (B. aeneus Burchell, 1822) (AMG/P 12248); whitefish (B. andrewi Barnard, 1937) (AMG/P 12249); chubbyhead barb ( $B$. anoplus Weber, 1897) (AMG/P 12255); Clanwilliam yellowfish (B. capensis Smith, 1840) (AMG/P 12250); largemouth yellowfish ( $B$. kimberleyensis Gilchrist \& Thompson, 1913) (AMG/P 12251); bowstripe barb (B. viviparus, Weber, 1897) (AMG/P 12252); Eastern Cape redfin ( $P$. afer (Peters, 1864)) (AMG/P 12253); Oreodaimon ( $P$. quathlambae (Barnard, 1938)) (AMG/P 11224); and the smallscale redfin ( $P$. asper (Boulenger, 1911)) (AMG/P 12254). All these series were from known parentage except Oreodaimon eggs which were collected from the wild. Oreodaimon eggs could not be confused with any other fish eggs in the Upper Orange River system in Lesotho (Cambray and Meyer 1988).

Only mature, fertilized ova were used and they were all obtained by stripping gravid females and fertilizing them with the milt from ripe-running males, except for Oreo-

Table 1
Egg envelope 'pore' size and 'pore' density of four Barbus species.

|  | 'Pore' size <br> $(\mu \mathrm{m})$ | 'Pore' density <br> $\left(\right.$ per $\left.5 \mu \mathrm{~m}^{2}\right)$ |
| :--- | :---: | ---: |
| Barbus anoplus | $0.29 \pm 0.05$ | $29.8 \pm 2.58$ |
| Barbus capensis | $0.20 \pm 0.02$ | $50.4 \pm 3.58$ |
| Barbus kimberleyensis | $0.18 \pm 0.02$ | $9.8 \pm 0.45$ |
| Barbus viviparus | $0.27 \pm 0.04$ | $18.2 \pm 1.48$ |

daimon where wild-collected eggs were used. In the present study only water-hardened, fertilized eggs were used to reduce variability of the observed structural features caused by these two factors. All eggs were fixed in $5 \%$ phosphate-buffered formalin. It is noted that in many egg envelope ultrastructure studies the eggs are often fixed in a solution of glutaraldehyde and post-fixed in osmium tetroxide. The majority of the eggs collected during ecological studies by either fisheries biologists or researchers collecting invertebrates are put into buffered or unbuffered formalin. It may therefore be more useful to study formalin fixed material, as the objective of the present study was to assess whether ultrastructural differences in the egg envelopes of African fish eggs can be used as an aid in species identification. Other workers have used a similar approach (Markle and Frost 1985; Olivar 1987).

The eggs were prepared using conventional techniques, involving a graded series of ethanol, transitional solvents, amyl acetate, and critical point drying (Cross 1987). The specimens were sputter-coated with gold and examined in a JEOL JSM-840 Scanning Electron Microscope operating at 10 KV .

Distortion during dehydration of biological specimens is a major problem when viewing SEM material, therefore a second method was used. The phosphate-buffered forma-lin-preserved eggs of three species were prepared using the Cryo technique. This technique is usually used to observe small delicate specimens in a frozen hydrated state. This method minimizes distortion. However, distortion can occur with ice crystal formation and the deposition of frost on the specimen surface (Cross 1987). Sub-cooled nitrogen was prepared in a vacuum chamber. Air was admitted to the vacuum chamber and the surface-dried eggs were fixed on stubs, rapidly quench-frozen in the nitrogen, and left to freeze for two minutes. The eggs were then taken to a chamber of the JEOL JSM-840 SEM unit which had been cooled by liquid nitrogen. The specimens were metal coated and viewed (Cross 1987). This method does not remove the mucous from the egg surface and was used as a comparative method to the more conventional method outlined above.

The measurement of the polygons of the honeycomb patterns of the egg envelopes was done from the photomicro-

Table 2

| The size of the honeycomb cells (polygon units), pattern units, and cell walls of $B$. andrewi, |
| :--- |
| $P$. afer, and $P$ asper. |

graphs and taken along the greatest axis, using a calibrated eyepiece micrometer in a binocular microscope. The thickness of the cell walls and the diameters of the pores were measured with the eyepiece micrometer. The number of pores per square unit were counted using a five $\mu \mathrm{m}$ square grid randomly put on the photomicrograph $(6000 \times)$ for five independent counts of all the pores within the grid.

## Results

## Ultrastructure of the Egg Envelope

Whitefish (B. andrewi)-Water-hardened eggs of whitefish are spherical and $2.09 \pm$ SD 0.07 mm in diameter. The surface of the egg envelope is characterized by a honeycomb lattice pattern (Fig. 1a). The individual pattern units are usually five-sided although there are cells with three to six sides and there are also several round and oblong cells. The actual cell walls are better outlined than those of the two Pseudobarbus species. There is also a greater cell depth than that of the Pseudobarbus species.

Chubbyhead barb (B. anoplus)-Cambray (1983) gave the size of water-hardened chubbyhead barb eggs as 1.1 mm in diameter. The egg envelopes of $B$. anoplus are highly adhesive (Cambray 1983). The surface of the egg envelope is covered with roundish 'pores' or pits. Since cross sections of these eggs have not been done it is not possible to distinguish between pores and pits, therefore in this paper the descriptive term 'pore' is used. Manner et al. (1977) used electron microscopy to determine that the pores did not penetrate the egg envelope of Pimephales promelas. The 'pores' are $0.29 \pm$ SD $0.05 \mu \mathrm{~m}$ in diameter and are sunken on the surface of the egg envelope (Fig. 1b). The 'pores' are fairly evenly distributed and there are approximately $29.8 \pm$ SD 2.58 per $5 \mu^{2}$ (Table 1 ).

Eastern Cape redfin (P. afer) -Water-hardened eggs of the Eastern Cape redfin are $1.56 \pm 0.08 \mathrm{~mm}$ in diameter with a small perivitelline space. The surface of the egg
envelope is covered with a relatively shallow honeycomb network (Fig. 1c) compared with whitefish (Fig. 1a). The dimensions of the cells and cell walls are given in Table 2. The individual pattern units are fairly uniformly 'round' surrounded by five to six other cells.

Smallscale redfin (P. asper) -Water-hardened eggs of the smallscale redfin are $1.12 \pm 0.05 \mathrm{~mm}$ in diameter with a small perivitelline space. The surface of the egg envelope is covered with a honeycomb network of cells similar to the Eastern Cape redfin. The dimensions of the cells and cell walls are given in Table 2. Each cell has either five or six sides (Fig. 1d).

Oreodaimon ( $\boldsymbol{P}$. quathlambae) -Water-hardened eggs of Oreodaimon are $1.8 \pm \mathrm{SD} 0.09 \mathrm{~mm}$ in diameter with a small perivitelline space (Cambray and Meyer 1988). Unlike the Eastern Cape redfin and the smallscale redfin, the surface of the egg envelope of Oreodaimon is not covered by a honeycomb structure. There are faint round markings on the surface of the egg envelope (Fig. 1e), which are $0.75 \pm$ SD $0.10 \mu \mathrm{~m}$ in diameter.

Clanwilliam yellowfish (B. capensis)—Water-hardened eggs of the Clanwilliam yellowfish are $2.76 \pm$ SD 0.13 mm in diameter. The egg has no surface ornamentation except for 'pores' which are $0.20 \pm$ SD $0.02 \mu \mathrm{~m}$ in diameter and fairly evenly spaced with 50.4 (46-55) 'pores' per square unit (Table 1; Fig. 1f).

Largemouth yellowfish (B. kimberleyensis) - Water-hardened eggs of the largemouth yellowfish are $3.31 \pm$ SD 0.26 in diameter. The surface of the egg envelope is relatively smooth and there are comparatively few 'pores' on the egg envelope (Fig. 1g). The 'pores' have a diameter of 0.18 $\pm$ SD $0.02 \mu \mathrm{~m}$ and are quite evenly distributed with 9.8 (9-10) 'pores' per square unit (Table 1).

Bowstripe barb (B. viviparus) -Water-hardened eggs of the bowstripe barb are 1.0 mm in diameter with a 0.2 mm


Figure 1
SEM photomicrographs of the ultrastructure of the egg envelope of the following species, eggs (a-h) prepared by conventional means, (i) prepared by Cryo technique: (a) Whitefish (B. andrewi); (b) Chubbyhead barb (B. anoplus); (c) Eastern Cape redfin (P. afer); (d) Smallscale redfin ( $P$. asper); (e) Oreodaimon ( $P$. quathlambae); ( f ) Clanwilliam yellowfish (B. capensis); (g) Largemouth yellowfish ( $B$. kimberleyensis); (h) Bowstripe barb ( $B$. viviparus); (i) Smallmouth yellowfish (B. aeneus). Scalebar $=1 \mu \mathrm{~m}$.


Figure 1-Continued
$\qquad$


Figure 1-Continued
perivitelline space (Ferguson 1987). The scattered 'pores' (Fig. 1h) have a diameter of $0.27 \pm$ SD $0.04 \mu \mathrm{~m}$ with 18.2 (16-19) 'pores' per square unit (Table 1).

Smallmouth yellowfish (B. aeneus)-Water-hardened eggs of smallmouth yellowfish are approximately 3.1-3.4 mm in diameter (Groenewald 1961). The conventional preparation technique was not successful and the Cryo technique was useful for this species. The surface of the egg was smooth and had pores or pits (Fig. 1i). The 'pores' on the smallmouth yellowfish were not compared with the size and density of the 'pores' on the eggs of other species because only the Cryo technique was successful for this species.

## Comparison of SEM Techniques

The Cryo technique was used on the eggs of a few species to compare with the conventional critical point drying (CPD) method. CPD prepared whitefish eggs show the typical honeycomb pattern (Fig. 1a), while the mucous remains on eggs prepared with the Cryo method (Fig. 2a). The same is seen for the redfin minnows (Figs. 1, c and $\mathrm{d} ; 2, \mathrm{~b}$ and c ). It has been noted that the fixation methods used in electron microscopy tend to dissolve the mucous coating more or less completely (Lönning and Hagström 1975). The mucous layer on the highly adhesive eggs of B. anoplus prepared with CPD is not present (Fig. 1b). In addition the structures are left intact using the Cryo technique and are not as 'eroded' as they were for the conventionally prepared eggs (compare Figs. 1d and 2c).

## Discussion

## Egg Envelope Microstructure

The outer egg envelope microstructure in some species is formed by follicle cells during oogenesis (Sponaugle and Wourms 1979; Stehr 1979; Stehr and Hawkes 1983) giving rise to patterns which may be seen in ovarian eggs. However, the pattern seen on ovarian eggs of certain species can be radically different from those observed on water-hardened, fertilized eggs (Lönning et al. 1984). Morphological changes may occur in the ultrastructure of the egg envelope of certain species during embryogenesis. Manner et al. (1977) found these changes were observed on the outer and inner surfaces as well as in the cross section.

There is great diversity in the structure of the egg envelopes from different species of teleosts in thickness, number of lamella, and surface structure. Most fish eggs have smooth unornamented egg envelopes; however, in some species it can be elaborately ornamented (Ahlstrom and Moser 1980). The most widespread type of egg envelope ornamentation is a noneycomb-like polygonal net-
work found on the outer surface of the egg envelope (Ahlstrom and Moser 1980). The most common network structure is hexagonal but in some species the polygons can have 4,5 , or 7 sides which are intermixed with the more usual 6 -sided pattern. In the present study this pattern was observed in whitefish (Fig. 1a), which has deep cell units compared with the shallow honeycomb structure of the redfin minnows examined (Figs. 1, c and d).

Boehlert (1984) suggested that studying the structure of the egg envelope on ovarian eggs could prove useful in the identification of marine plankton samples. Since the egg envelope ultrastructure of most African freshwater fishes is not known, SEM work of developing ovarian eggs may be useful in discerning differences in eggs collected from the wild and may aid in the identification of these samples. But care should be taken in case there are major changes after fertilization and water-hardening.

In a freshwater study, Riehl (1979) prepared identification keys based on the structural peculiarities of the egg envelopes of several freshwater fishes. The egg envelopes of the silver carp (Hypophthalmichthys molitrix), bighead (Aristichthys nobilis), the grass carp (Ctenopharyngodon idella) and the black carp (Mylopharyngodon piceus) are distinguished by their species-specific structural features (Mikodina and Makeyeva 1980). In another study on the eggs of freshwater fishes, Keevin et al. (1980) used egg envelope ornamentation to distinguish genera of killifishes. Johnson and Werner's (1986) study on the ultrastructure of the egg envelopes of five freshwater fish species was initiated so that researchers could distinguish between the eggs of key species in order to assess the environmental impact of energy-generating facilities and other perturbations on spawning and nursery grounds. Knowledge of egg envelope ultrastructure in African fishes would also be of value in environmental impact studies.

The envelopes of certain marine teleost eggs have a morphology which is species specific (Lönning and Hagström 1975). Mikodina (1987) found that the structural characteristics of the egg envelope surface can be used as taxonomic traits for the pelagic eggs of several marine fish species. Some authors contend that in many groups of fishes the structure of the egg envelope may be an indication of the systematic status of the species (Rass 1953; Ivankov and Kurdyayeva 1973). In this preliminary paper the egg envelope ultrastructural characters discussed are only considered for egg identification.

## 'Pore' Size and Density

Olivar (1987) found that in some marine fish eggs, pore size varied little between species, but pore density differed markedly. In the species Lönning (1981) examined, both pore diameter and the number of pores varied. In the present study both 'pore' diameter and density varied between species (Table 1). However, 'pore' density dif-


Figure 2
SEM photomicrographs of the ultrastructure of the egg envelope prepared using the Cryo technique for the following species: (a) Whitefish (B. andrewi); (b) Eastern Cape redfin ( $P$. afer); (c) Smallscale redfin ( $P$. asper). Scalebar $=1 \mu \mathrm{~m}$.
fered more markedly than did 'pore' diameter, a finding which is similar to the findings of Olivar (1987). The appearance of the 'pores' is distinct in the Barbus species studied here, varying in size and number per square unit. The 'pore' diameter is smallest in the largemouth yellowfish egg envelopes $(0.18 \mu \mathrm{~m})$ and the largest 'pores' were observed on the egg envelopes of $B$. anoplus eggs $(0.29 \mu \mathrm{~m})$ (Table 1). The greatest number of 'pores' per square unit were found on the whitefish eggs $\left(50.4 / 5 \mu \mathrm{~m}^{2}\right)$ and the least number on the largemouth yellowfish eggs $\left(9.8 / 5 \mu \mathrm{~m}^{2}\right)$ (Table 1).

Care must be taken with the interpretation of pore structure because morphological changes after fertilization can be accompanied by changes in the surface structure of the egg coat (Perry 1984). Different preparation techniques (CPD vs. Cryo) can cause varying distortions. In the present study, smallmouth yellowfish eggs were prepared using only the Cryo technique owing to distortions using the CPD method. Mikodina (1987) cautioned that the structural characteristics of pores on the egg envelope are considered unreliable criteria for the identification of eggs (Mikodina 1987). In a study of the egg envelope structure of Pseudopleuronectes americanus, Perry (1984) found that the average distance between the pores and pore diameter decrease after fertilization, possibly because of the final levelling and hardening of the egg envelope after fertilization. Hagström and Lönning (1968) have also demonstrated that there is a change in the pores of certain species after insemination of the egg.

After fertilization and hardening in sea water the surface of the egg envelopes of lumpsucker eggs (Cyclopterus lumpus) were no longer smooth and covered with regularly distributed pores but became characterized by ridges and almost completely covered by papillae (Lönning et al. 1984). Obviously one needs to take care in the interpretation of surface structures and indicate whether the eggs have been fertilized and water-hardened.

## Pseudobarbus Species

The redfin minnows of southern Africa are distinctive and at a specific level are not obviously similar to other southern African Barbus species; therefore the flexible-rayed redfins were removed from the genus Barbus and put into the genus Pseudobarbus (Skelton 1980, 1988). Oellermann (1988) showed that chromosome counts of the Eastern Cape redfin was 96 . He therefore placed the redfin minnows with the group of cyprinids showing tetraploidic origins. The Eastern Cape redfin and the smallscale redfin are closely related (sister species: Skelton 1988) and it has been shown in the present study that their egg envelope ultrastructure is similar and distinctive (Fig. 1, c and d). In contrast the egg envelope structure of the more distantly related redfin minnow, Oreodaimon, does not have a honeycomb pattern of ridges (Fig. 1e). The redfin minnow egg envelope
surface patterns are unlike the patterns observed on the two small Barbus species examined here. Skelton (1980) suggested that of all the Barbus species in southern Africa the chubbyhead barb; Marico barb (B. motebensis); and the redtail barb (B. gurneyi) are most similar to the flexible-rayed redfins (for example, the Eastern Cape, smallscale, and Oreodaimon redfins). The chubbyhead barb does not have the honeycomb structured egg envelope as do two of the redfins, but a more simple egg surface covered with 'pores.' In addition the chubbyhead barb is clearly diploid compared to the tetraploid redfins (Oellermann 1988).

## Ecological Notes

It has been suggested that superficial similarity of egg envelope structure on phylogenetically distant genera supports a functional role (Robertson 1981) and independent derivation (Boehlert 1984). One would expect differences in egg surface ultrastructure because of the range in habitat and breeding habits of teleosts (Kuchnow and Scott 1977). Surprisingly, in their study of the egg envelopes of 40 fish species belonging to 8 orders, Ivankov and Kurdyayeva (1973) demonstrated that it is not correct to put overriding significance on the mode of spawning and the nature of the spawning grounds on the structure of the egg envelope thereby supporting the findings of Rass (1953).

Some authors have suggested that there can be differences in the egg envelope structures in fishes of the same ecological group. For example, a thin gelatinous membrane attaches eggs to submerged vegetation in fishes of the family Cyprinidae whereas villi serve this purpose in the "lake minnow" and a long filament in the "rotan"' (Ivankov and Kurdyayeva 1973). Whereas Mikodina (1987) has shown that different fish species developing in similar ecological conditions have similar egg envelope structures.

The conservative structure of the egg envelope has been noted in fishes with different spawning ecology and different types of eggs (pelagic or benthic). Examples given by Ivankov and Kurdyayeva (1973) were several silurids which lay eggs on a different substrate for each species yet in all species the egg envelope was represented by a thick honeycomb pattern. The structure of the egg envelope may be an indication of the systematic status of some species because they found a similarity in the structure of the egg envelope of fishes of the same systematic group although the fish lay their eggs on different substrates (Rass 1953; Ivankov and Kurdyayeva 1973). Ivankov and Kurdyayeva (1973) suggested that the structure of the vitelline 'membrane' was due to its protective function whereas the structure of the outer egg envelope, the chorion, was an indication of the systematic status of the species.

Ivankov and Kurdyayeva (1973) noted that fishes of the family Cyprinidae are mainly phytophilous. The fish species examined in the present study were riverine species and in the family Cyprinidae. All the eggs examined were
demersal (substratum) eggs. The chubbyhead barb (Cambray 1983) and the bowstripe barb (Ferguson 1987) have adhesive egg envelopes. The three redfin minnows have slightly or nonadhesive egg envelopes (Cambray and Meyer 1988; pers. obs.). Species such as the redfin minnows breed in the riffle areas after rains (Cambray and Meyer 1988). The redfin minnows put their nonadhesive eggs under boulders whereas the chubbyhead and bowstripe barbs are more typically phytophilous (Cambray and Bruton 1984; Ferguson 1987).

It has often been said that the primary function of the outer egg envelope is to protect the embryo from the external environment (Manner et al. 1977; Groot and Alderdice 1985). In many species egg envelope thickness can often be related to the environment where the eggs are found, such as the thick envelope of some species of demersal eggs compared with thinner envelopes in certain pelagic eggs (Lönning 1972; Stehr and Hawkes 1983). It is now necessary to examine the depth and number of egg envelopes of African cyprinids.

## Conclusion

The main objective of this preliminary paper on the ultrastructure of the egg envelopes of several African Barbus and Pseudobarbus species was to assess whether or not Scanning Electron Microscopy would prove a useful analytical tool in identifying eggs collected from the wild. The results from this study provide encouragement for further work in that each of the nine species examined showed distinctive egg envelope ultrastructures, most notably the redfin minnows and the whitefish. For example, if egg collections of several minnow species (such as the co-occurring species, smallscale redfin and the chubbyhead barb) were made at the same time in the Groot River, Gamtoos River System of the Cape Province, the eggs could be positively identified by using Scanning Electron Microscopy. The Cryo technique is a useful method to prepare eggs for viewing in the SEM, especially if one wants to observe the mucous covering and its position on the egg coat. The Albany Museum has initiated a collection of SEM photomicrographs of the ultrastructure of the egg envelopes of African freshwater fish species to augment their early life-history collections.

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## Citations

Ahlstrom, E.H., and H.G. Moser.
1980. Characters useful in identification of pelagic marine fish eggs. Calif. Coop. Oceanic Fish. Invest. Rep. 21:121-131.
Allaby, M.
1985. The Oxford dictionary of natural history. Oxford Univ. Press, Oxford.
Avni, A.A., and S.G. Soin.
1974. Adaptive peculiarities of Nothobranchius quentheri (Pfeffer) embryogenesis in connection with life in periodically drying out tropical waters. Voprosy ichtiologii 14:846-858. (In Russian.)
Balon, E.K.
1985. Early ontogeny of Labeotropheus Ahl, 1927 (Mbuna, Cichlidae, Lake Malawi), with a discussion on advanced protective styles in fish reproduction and development. In Early life histories of fishes (E.K. Balon, ed.), p. 207-236. Junk, Dordrecht.

Boehlert, G.W.
1984. Scanning electron microscopy. In Ontogeny and Systematics of Fishes (H.G. Moser, ed.), p. 42-50. Am. Soc. Ichthyol. Herpetol. Spec. Publ. No. 1.
Cambray, J.A.
1983. Early development and larval behaviour of a minnow, Barbus anoplus (Pisces; Cyprinidae). S. Afr. J. Zool. 18:331-336.
Cambray, J.A., and M.N. Bruton.
1984. The reproductive strategy of a barb, Barbus anoplus (Pisces: Cyprinidae) colonizing a man made lake in South Africa. J. Zool. (Lond.) 204:143-168.
Cambray, J.A., and K. Meyer.
1988. Early ontogeny of an endangered, relict, cold-water cyprinid from Lesotho, Oreodaimon quathlambae (Barnard, 1938). Rev. Hydrobiol. Trop. 21(4):309-333.
Cambray, J.A., and G.G. Teugels.
1988. Selected annotated bibliography of early developmental studies of African freshwater fishes. Ann. Cape Prov. Mus. Nat. Hist. 18(2):31-56.
Cross, R.H.M.
1987. A handbook on the preparation of biological material for electron microscopy. Electron Microscopy Unit, Rhodes Univ., Grahamstown, South Africa.
Ferguson, S .
1987. Propagation of Barbus viviparus under hatchery conditions. Lammergeyer Dec. 1987:55-59.
Groenewald, A.A.v.J.
1961. A progress report on the culture of Barbus holubi, the Vaal River yellowfish, at the Provincial Fisheries Institute, Lydenburg. Res. Rep. Prov. Fish. Inst., Transvaal Prov. Admin.
Groot, E.P., and D.F. Alderdice.
1985. Fine structure of the external egg membrane of five species of Pacific salmon and steelhead trout. Can. J. Zool. 63:552-566. Hagström, B.E., and S. Lönning.
1968. Electron microscopic studies of unfertilized and fertilized eggs from marine teleosts. Sarsia 33:73-80.
Ivankov, V.N., and V.P. Kurdyayeva.
1973. Systematic differences and the ecological importance of the membranes in fish eggs. J. Ichthyol. (Eng. transl. Vopr. Ikhtiol.) 13:864-873.

Johnson, E.Z., and R.G. Werner.
1986. Scanning electron microscopy of the chorion of selected freshwater fishes. J. Fish. Biol. 29:257-265.
Keevin, T.M., G. Phillips, J.E. Thomerson, and D.C. Taphorn.
1980. Egg chorion ornamentation and taxonomy of the annual killifishes Austrofundulus, Rachovia, and Terranatos. (Abstract.) Am. Soc. Ichthyol. Herpetol., 60th Annual meeting, Fort Worth, TX.
Kuchnow, K.P., and J.R. Scott. 1977. Ultrastructure of the chorion and its micropyle apparatus in the mature Fundulus heteroclitus (Walbaum) ovum. J. Fish. Biol. 10:197-201.
Laale, H.W.
1980. The perivitelline space and egg envelopes of bony fishes: a review. Copeia 1980:210-226.
Lévêque, C., and J. Daget.
1984. Cyprinidae. In Check-list of the freshwater fishes of Africa (J. Daget, J.P. Grosse, and D.F.E. Thys van den Audenaerde, eds.), p. 217-342. ORSTOM Paris, MRAC Tervuren.
Lönning, S.
1972. Comparative electron microscopic studies of teleostean eggs with special reference to the chorion. Sarsia 49:41-48.
1981. Comparative electron microscope studies of the chorion of the fish egg. Rapp. P.-v. Reun. Cons. Int. Explor. Mer 178: 560-564.
Lönning, S., and B.E. Hagström.
1975. Scanning electron microscope studies of the surface of the fish eggs. Astarte 8:17-22.
Lönning, S., E. Kjorsvik, and J. Davenport.
1984. The hardening process of the egg chorion of the cod, Gadus morhua L., and lumpsucker, Cyclopterus lumpus L. J. Fish. Biol. 24:505-522.
Manner, H.W., M. VanCura, and C. Muehleman.
1977. The ultrastructure of the chorion of the fathead minnow, Pimephales promelas. Trans. Am. Fish. Soc. 106:110-114.
Markle, D.F., and L.-A. Frost.
1985. Comparative morphology, seasonality, and a key to planktonic fish eggs from the Nova Scotian shelf. Can. J. Zool. 63: 246-257.
Matarese, A.C., and E.M. Sandknop.
1984. Identification of fish eggs. In Ontogeny and systematics of fishes (H.G. Moser, ed.), p. 27-31. Am. Soc. Ichthyol. Herpetol. Spec. Pub. No. 1.
Mikodina, E.V.
1987. Surface structure of the egg membranes of teleostean fishes. J. Ichthyol. (Eng. transl. Vopr. Ikhtiol.) 27(3):19-26.

Mikodina, E.V., and A.P. Makeyeva.
1980. Structure and some properties of egg membranes in the freshwater pelagophilous fishes. J. Ichthyol. (Eng. transl. Vopr. Ikhtiol.) 20(2):86-94.
Oellermann, L.K.
1988. The karyology and taxonomy of the southern African yellow-
fish (Pisces: Cyprinidae). M.S. thesis, Rhodes Univ., Grahamstown, South Africa, 127 p.
Olivar, M.P.
1987. Chorion ultrastructure of some fish eggs from the South-East Atlantic. S. Afr. J. Mar. Sci. 5:659-671.
Perry, D.M.
1984. Post-fertilization changes in the chorion of winter flounder Pseudopleuronectes americanus Walbaum, eggs observed with scanning electron microscopy. J. Fish. Biol. 25:83-94.
Rass, T.S
1953. The importance of the structure of eggs and larvae to the systematics of fishes. In Ocherki po obshchim voprosam ikhtiologii (Essays on general aspects of ichthyology). Moscow, Leningrad, Acad. Sci. USSR Press.
Riehl, R.
1979. Ein erweiterber und verbesserter bestimmungsschlussel für die eier deutscher Susswasser - Teleosteer. Z. angew. Zool. 66(2): 199-216.
Robertson, D.A.
1981. Possible functions of surface structure and size in some planktonic eggs of marine fishes. N.Z. J. Mar. Freshwater Research 15:147-153.
Shelton, W.L.
1978. Fate of the follicular epithelium in Dorosoma petenense (Pisces: Clupeidae). Copeia 1978:237-244.
Skelton, P.H.
1980. Systematics and biogeography of the redfin Barbus species (Pisces: Cyprinidae) from southern Africa. Ph.D. thesis, Rhodes Univ., Grahamstown, South Africa, 416 p.
1988. A taxonomic revision of the redfin minnows (Pisces, Cyprinidae) from southern Africa. Ann. Cape Prov. Mus. Nat. Hist. 16(10):201-307.
Sponaugle, D.L., and J.P. Wourms.
1979. Secondary egg envelope pre-patterning by follicle cells during fish oogenesis. Am. Zool. 19:946.
Stehr, C.M.
1979. The development of the hexagonal surface structure of the C-O sole (Pleuronichthys coenosus). Am. Zool. 19:976.
Stehr, C.M., and J.W. Hawkes.
1983. The development of the hexagonally structured egg envelope of the C-O sole (Pleuronichthys coenosus). J. Morphol. 178: 267-284.
Werner, R.R.
1976. Current level of taxonomic information on Great Lakes fish eggs and larvae. In Great Lakes fish egg and larvae identification: proceedings of a workshop (J. Boreman, ed.). U.S. Fish and Wildl. Serv., FWS/OBS-76.23.
Wourms, J.P.
1976. Annual fish oogenesis. I: Differentiation of the mature oocyte and formation of the primary envelopes. Develop. Biol. 50:338-354.

# Effects of Fixation and Dehydration on Shrinkage and Morphology in Common Snook Yolk-Sac Larvae 

MARIE F. DeLEON, RUTH O. REESE, and WALTER J. CONLEY<br>Florida Marine Research Institute<br>Florida Department of Natural Resources<br>100 Eighth Avenue S.E.<br>St. Petersburg, FL 33701


#### Abstract

Five fixatives (Bouin's, Davidson's, phosphate buffered glutaraldehyde and paraformaldehyde, and $10 \%$ formalin/seawater) were compared to evaluate mean notochord length (MNL) shrinkage of yolk-sac larvae of the common snook (Centropomus undecimalis). Measurements were taken at initial fixation, after 90 days' storage in fixative, and after serial dehydration. Larvae fixed in Bouin's solution and Davidson's solution shrank $9.05 \%$ and $11.49 \%$ in mean notochord length. The MNL of larvae fixed in glutaraldehyde for 90 days was not significantly different from the MNL of live larvae; however, MNL decreased $12.67 \%$ following dehydration. Paraformaldehydefixed larvae had a maximum shrinkage of $5.06 \%$ at 90 days; dehydration resulted in total shrinkage of $16.74 \%$. Larvae fixed in formalin/seawater shrank $5.86 \%$ at 90 days; dehydration resulted in total shrinkage of $12.08 \%$. Observations were made of changes in the morphological appearance of larvae preserved in various fixatives.


## Introduction

The length of fishes collected in the field is one of the most common parameters used in examining the life history of a species. Length-frequency analyses have been used to determine growth (MacDonald and Pitcher 1979; Schnute and Fournier 1980) as well as ontogenetic shifts in habitat (Gilmore et al. 1983; Brandt 1986); diet (Laroche 1982; Govoni et al. 1983; Peters and McMichael 1987); and feeding chronology (Brandt 1986). Stock-assessment methods and determination of recruitment are often based on length-frequency analyses (Jones 1984; Pauly and Morgan 1987), although it is realized there are serious shortcomings with length-frequency analysis (Basson et al. 1988) because the differential treatment of collections may compromise the accurate assessment of true (live) length.
Most marine fish larvae are fixed and preserved in formalin-based solutions that may be buffered, mixed with acids and/or alcohol, or diluted with seawater. Phosphatebuffered formalin solutions are preferred to solutions buffered with borax or limestone because they minimize shrinkage; dampen pH changes; and prevent pigment loss, decalcification, and demineralization (Farris 1963; Taylor 1977; van der Veer 1982; Lavenberg et al. 1984; Markle 1984; Tucker and Chester 1984). Formalin-mixed solutions such as Bouin's and Davidson's fluids, which contain acid
components, also have been recommended because they maintain cellular integrity at the light microscope level and because they ensure minimal shrinkage of fixed tissue (Stickland 1975; Theilacker 1980; Hinton et al. 1984). Formalin solutions in seawater are used extensively because of their convenience in the field. Although distortion of preserved tissue may be minimal initially, shrinkage of tissue in fixative increases over time and increases significantly with subsequent dehydration (Parker 1963; Rosenthal et al. 1978; Schnack and Rosenthal 1978; Hay 1981, 1982; Tucker and Chester 1984).

Few studies have investigated the fixative-induced shrinkage in yolk-sac larvae. Farris (1963) observed the effects of borax-buffered 3\% formalin/seawater solution over a 6 -week period; Schnack and Rosenthal (1978) and Rosenthal et al. (1978) studied the effects of $2 \%$ and $4 \%$ formalin/seawater solutions over a 20 -day period and for a period of more than 1 year, respectively. Formalin has also been found to have a differential effect on yolk and tissue (Heming and Preston 1981).

In this study we evaluate the shrinking of newly hatched snook larvae (Centropomus undecimalis) at initial fixation, after storage in fixative, and after a graded alcohol dehydration. Five fixative solutions commonly used at field stations and laboratories are compared. Larvae used in this study came from a single spawn and had not absorbed their yolk sacs.

This homogeneous group provided a means for gathering shrinkage data from larvae in a primary developmental stage prior to first feeding. In live larvae observed at this stage, tissues are translucent with visible dendritic and condensed chromatophores. Internal organs are easily observed. Two regimes were used to evaluate shrinkage: A) larvae fixed and subsequently dehydrated within 6 days of initial fixation for storage in $70 \%$ ethanol and B) la:vae retained in fixative for 90 days.

## Materials and Methods

Four hundred laboratory-spawned common snook larvae were collected live 18-24 hours after hatching in 2-liter beakers and were individually placed by pipette onto ring slides. Care was taken to minimize damage during handling because such damage could promote shrinkage prior to live measurement and fixation (Theilacker 1980). Anesthesia, which has been shown to contribute to shrinkage in live larvae (Parker 1963; Theilacker 1980), was not used. Measurements were taken to the nearest 0.01 mm of notochord length (NL). Each larva was measured in a small drop of seawater of sufficient size to keep the larva alive; an ocular micrometer was used at $40 \times$ for measurements, and all live and subsequent measurements were made by the same person. Individual larvae were placed live in numbered vials of approximately $1-\mathrm{mL}$ fixative volume.

Fixatives used in the study were as follows:

- Bouin's fluid (Sheehan \& Hrapchak 1980)

Saturated picric acid ( 21 gm to 1 liter) in distilled water $\quad 1500 \mathrm{~mL}$ Formalin $\quad 500 \mathrm{~mL}$ Glacial acetic acid 100 mL

- Davidson's fluid (Yevich and Barszcz 1977)

Formaldehyde ( $40 \%$ solution) 200 mL Glycerin 100 mL Ethanol (95\% solution) 300 mL Glacial acetic acid 100 mL Water (distilled or seawater) 300 mL
(Davidson's is cloudy when mixed with seawater with high salinity. Distilled water was used in preparation of fixative for this study.)

- $3 \%$ glutaraldehyde in 0.1 M phosphate buffer (made with monobasic and dibasic sodium phosphate - pH 7.4)
- $10 \%$ paraformaldehyde in 0.1 M phosphate buffer (made with monobasic and dibasic sodium phosphate - pH 7.4 )
- $10 \%$ formalin in seawater (seawater salinity $-29.5 \%$, collected from larval incubation cone).

Fixatives were prepared within 24 hours of use, and pH was determined initially and after 90 days. The total sample was divided into two groups of 200 each (Groups A and B). Each group was composed of 5 subgroups of 40 individuals; each subgroup represented one of the five fixatives being studied. The vials were stored at room temperature for subsequent measurements.

## Group A

After 3 days in fixative, larvae were remeasured. Fixative solution was removed by pipette, and serial dehydration was begun in $30 \%$ ethanol, which was changed to $50 \%$ and then $70 \%$ ethanol, and 24 hours allowed between changes. Each larva was measured at the time of alcohol change and after the larva had been in $70 \%$ ethanol for 24 hours. Group A samples were then stored at room temperature.

## Group B

Group B samples were remeasured 30, 60, and 90 days after fixation. No solutions were changed.

## Statistical Testing

Initial shrinkage was tested for treatment effect using a Model I one-way analysis of variance (ANOVA; Sokal and Rohlf 1969). Because subsequent measurements were made on the same individuals, we could not assume independence of the error term. Overall treatment effect was, therefore, tested using a repeated measures model (Winer 1971). Data analyses were completed using SAS utility programs on Florida Department of Natural Resources' IBM 4341 Group II System computer. All tests for significance were performed at the $95 \%$ level $(P<0.05)$.

## Results

## Bouin's and Davidson's Fluids

Effect of Fixation and Subsequent Dehydration-Fixation in Bouin's resulted in significant larval shrinkage ( $10.04 \%$; Table 1); there was, however, little change in mean notochord length (MNL) following serial dehydration (Fig. 1). Larvae fixed in Davidson's and serially dehydrated to $70 \%$ alcohol shrank $13.48 \%$ (Table 1; Fig. 1). Larvae shrank approximately one percent following each treatment.

Effect over Time-Larvae fixed in Bouin's and Davidson's fluids shrank $9.05 \%$ and $10.21 \%$ (Table 1) by 30 days after fixation, but little change in MNL was recorded in subsequent measurements for either fixative (Fig. 2). Initial shrinkage was significant. A maximum shrinkage

Table 1
Shrinkage due to fixation and ethanol dehydration (Group A) and over time without fixative change (Group B). Mean notochord length in millimeters (percent shrinkage).

|  | Group A |  |  |  |  | Group B |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Live | 3 days | 30\% Etoh | 50\% Etoh | 70\% Etoh | Live | 30 days | 60 days | 90 days |
| Bouin's | 2.39 | $\begin{gathered} 2.15 \\ (10.04) \end{gathered}$ | $\begin{gathered} 2.14 \\ (10.46) \end{gathered}$ | $\begin{gathered} 2.13 \\ (10.88) \end{gathered}$ | $\begin{gathered} 2.13 \\ (10.88) \end{gathered}$ | 2.32 | $\begin{gathered} 2.11 \\ (9.05) \end{gathered}$ | $\begin{gathered} 2.10 \\ (9.48) \end{gathered}$ | $\begin{gathered} 2.11 \\ (9.05) \end{gathered}$ |
| Davidson's | 2.30 | $\begin{gathered} 2.10 \\ (8.70) \end{gathered}$ | $\begin{gathered} 2.05 \\ (10.87) \end{gathered}$ | $\begin{gathered} 2.02 \\ (12.17) \end{gathered}$ | $\begin{gathered} 1.99 \\ (13.48) \end{gathered}$ | 2.35 | $\begin{gathered} 2.11 \\ (10.21) \end{gathered}$ | $\begin{gathered} 2.09 \\ (11.06) \end{gathered}$ | $\begin{gathered} 2.08 \\ (11.49) \end{gathered}$ |
| Phosphate Buffered Glutaraldehyde | 2.37 | $\begin{gathered} 2.32 \\ (2.11) \end{gathered}$ | $\begin{gathered} 2.33 \\ (1.69) \end{gathered}$ | $\begin{gathered} 2.20 \\ (7.17) \end{gathered}$ | $\begin{gathered} 2.07 \\ (12.67) \end{gathered}$ | 2.35 | $\begin{gathered} 2.35 \\ (0.00) \end{gathered}$ | $\begin{gathered} 2.35 \\ (0.00) \end{gathered}$ | $\begin{gathered} 2.35 \\ (0.00) \end{gathered}$ |
| Phosphate Buffered Paraformaldehyde | 2.39 | $\begin{gathered} 2.30 \\ (3.77) \end{gathered}$ | $\begin{gathered} 2.30 \\ (3.77) \end{gathered}$ | $\begin{gathered} 2.16 \\ (9.62) \end{gathered}$ | $\begin{gathered} 1.99 \\ (16.74) \end{gathered}$ | 2.37 | $\begin{gathered} 2.27 \\ (4.22) \end{gathered}$ | $\begin{gathered} 2.26 \\ (4.64) \end{gathered}$ | $\begin{gathered} 2.25 \\ (5.06) \end{gathered}$ |
| $10 \%$ Formalin/ Seawater | 2.40 | $\begin{gathered} 2.32 \\ (3.33) \end{gathered}$ | $\begin{gathered} 2.28 \\ (5.00) \end{gathered}$ | $\begin{gathered} 2.18 \\ (9.17) \end{gathered}$ | $\begin{gathered} 2.11 \\ (12.08) \end{gathered}$ | 2.39 | $\begin{gathered} 2.29 \\ (4.18) \end{gathered}$ | $\begin{gathered} 2.29 \\ (4.18) \end{gathered}$ | $\begin{gathered} 2.25 \\ (5.86) \end{gathered}$ |

of $9.05 \%$ in Bouin's and $11.49 \%$ in Davidson's was recorded at 90 days. The initial pH of Bouin's fluid was 1.9 and had changed to 2.3 after 90 days. The initial pH of Davidson's fluid was 2.7; it had fallen to 2.5 after 90 days.

Morphological Changes-After fixation in Bouin's, formerly translucent larvae turned golden yellow and pigment markings disappeared. Viscera visible in live animals were no longer distinguishable. After these larvae were dehydrated to $70 \%$ ethanol, the yellow color lightened somewhat and internal morphology was more discernible. Tissue color of larvae fixed in Davidson's changed to light brown and pigment markings disappeared, but external morphology and viscera were more visible than in Bouin's.

## 3\% Glutaraldehyde in 0.1 M Phosphate Buffer ( pH 7.4 )

Effect of Fixation and Subsequent Dehydration-No significant difference in MNL was found between live larvae and the same animals measured 3 days after fixation (Fig. 1). Mean notochord length, however, decreased by $7.17 \%$ in $50 \%$ ethanol and by $12.67 \%$ in $70 \%$ ethanol (Table 1).

Effect over Time-Mean notocord length decreases were not observed (Fig. 2; Table 1). Measurements made at 90 days may have been compromised because the posterior end of the notochord was not clearly distinguishable because of tissue opacity. The initial fixative pH was 7.6 ; it had changed to 7.1 after 90 days.

Morphological Changes-A heavy precipitate formed during fixation, adhering to larvae and obscuring them in the fixative solution. Serially dehydrated larvae that had
been in fixative three days were rinsed in distilled water but retained a coating of precipitate. Clarity of larval external and internal morphology after 90 days in alcohol was similar to that of live specimens but had degenerated considerably when observed at 7 months. No additional body color was noted initially in larvae stored in fixative, but pigment markings disappeared. External and internal morphology were clearly distinguishable at 60 days after fixation. As previously noted, larvae stored in fixative for 90 days had become noticeably opaque, making the posterior end of the notochord difficult to distinguish. The larvae had also become brittle, and damage due to handling, including breakage and loss of finfold sections, was greater.

## $10 \%$ Paraformaldehyde in 0.1 M Phosphate Buffer ( pH 7.4 )

Effect of Fixation and Subsequent Dehydration-Little change in MNL was noted for larvae fixed in paraformaldehyde and dehydrated in $30 \%$ ethanol, but shrinkage increased approximately $6 \%$ with each subsequent alcohol change. When dehydrated in $70 \%$ ethanol, larvae had decreased in MNL by $16.74 \%$ (Fig. 1).

Effect over Time-No significant change in MNL was noted for larvae preserved in buffered paraformaldehyde for 30 days (Fig. 2), but maximum MNL shrinkage of $5.06 \%$ was observed 90 days after fixation (Table 1). The initial fixative pH was 7.4 ; it had not changed after 90 days.

Morphological Changes-Precipitate also formed in buffered paraformaldehyde, making it difficult to locate specimens in the vials. Pigment markings were indistinguishable in both treatment groups. There was no change in body color in animals stored in paraformaldehyde, and their


Figure 1
Group A: Effect of fixation and ethanol dehydration on changes in notochord length (mm). Error bars represent standard error.
fixed appearance was similar to that of live specimens. Dehydrated larvae turned a light brown color in alcohol, and viscera were no longer distinguishable.

## $10 \%$ Formalin in Seawater

Effect of Fixation and Subsequent Dehydration-Larvae fixed in formalin/seawater shrank significantly upon initial fixation. Little additional shrinkage occurred after transfer
of larvae to $30 \%$ ethanol, but shrinkage increased approximately $4 \%$ when larvae were transferred to $50 \%$ ethanol and increased another $2.91 \%$ when they were transferred to $70 \%$ ethanol (Fig. 1; Table 1).

Effect over Time-Larvae stored in $10 \%$ formalin/seawater shrank significantly ( $4.18 \%$ ) after 30 days in fixative (Fig. 2; Table 1), but little difference in MNL was observed in subsequent measurements. A total shrinkage of $5.86 \%$


Figure 2
Group B: Effect of fixation over time on changes in notochord length (mm). Error bars represent standard error.
was recorded at 90 days. The initial fixative pH was 6.6 ; it had changed to 7.1 after 90 days.

Morphological Changes-Tissue clarity was reduced in larvae stored in fixative, and internal morphology became indistinguishable in dehydrated larvae. Pigment markings disappeared in both treatments, and body color changed to light brown in alcohol. A slight precipitate formed in the fixative, but larvae were not obscured.

## Overall Treatment Effects

Glutaraldehyde was the only fixative that did not cause significant larval shrinkage over time (Table 2; Fig. 2). However, when followed by an ethanol dehydration series, significant shrinkage occurred (Fig. 1). All other treatments (Group B) and treatment combinations (Group A) caused significant shrinkage of larvae (Table 2).

Table 2
Overall treatment effect of fixation and ethanol dehydration (Group A) and fixation over time (Group B).

|  | Group A |  |  | Group B |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  | $F$ value | Significance |  | $F$ value | Significance |
| Phosphate buffered <br> paraformaldehyde | 267.38 | 0.0001 | 58.45 | 0.0001 |  |
| Phosphate buffered <br> glutaraldehyde | 138.86 | 0.0001 | 0.07 | 0.9756 |  |
| $10 \%$ Formalin/ <br> seawater | 152.10 | 0.0001 | 80.55 | 0.0001 |  |
| Davidson's solution <br> Bouin's solution | 161.98 | 0.0001 | 188.69 | 0.0001 |  |

## Discussion

Size, as indicated by notochord or standard length, is important in the evaluation of larval growth and development. The amount of shrinkage as a result of fixation and dehydration must be considered for an accurate estimate of live parameters. Thus, the choice of fixative can play a key role in obtaining reliable growth data. However, fixative and preservative choice will depend upon the goals of the particular study, and no single fixative has been found to be ideal for all types of fixation and preservation.

Bouin's and Davidson's solutions are often used for histological analyses; Bouin's, however, can render tissues brittle. Davidson's is the preferred fixative for gonad examination. The acid component of these solutions makes them unsuitable for osteological examinations. Both solutions caused significant shrinkage in the present study, whether larvae were stored in fixative or dehydrated. Theilacker (1980) reported an $8 \%$ decrease in standard length (SL) of feeding larvae fixed in Bouin's, although shrinkage of other body dimensions ranged up to $25 \%$. Leslie and Moore (1986) examined freshwater larvae fixed in Davidson's B and reported mixed results, with shrinkage ranging from unmeasurable for yolk-sac larvae to $4.3 \%$ for larger larvae. These investigators suggested that freshwater larvae may respond differently to fixation and that real differences in the extent of length changes may be related mainly to osmotic processes.

Larvae fixed in phosphate-buffered glutaraldehyde did not shrink significantly during either initial fixation or during storage in fixative for more than 90 days, but pigment markings disappeared. Oozeki and Hirano (1988) reported similar results for glutaraldehyde-fixed sea-bream larvae. In contrast, Stickland (1975) reported significant shrinkage of muscle fibers fixed in phosphate-buffered glutaraldehyde; he did, however, note the superior preservation of muscle tissue for histological studies. Taylor (1977) claimed
that glutaraldehyde is superior to formaldehyde for fixing soft tissue because glutaraldehyde is a more efficient crosslinking agent.

Larvae stored in buffered paraformaldehyde had shrunk $3.76 \%$ at 3 days after fixation and $4.97 \%$ at 90 days, at which time tissue transparency was sufficient to allow observation of internal morphology. Dehydration to $70 \%$ ethanol resulted in shrinkage of $16.72 \%$, the maximum of all fixatives evaluated, and also resulted in change in body color and loss of tissue transparency.
Formaldehyde solutions are by far the most common means of fixation and preservation for larval fishes and were recommended by some authors (Richards and Berry 1973; Ahlstrom 1976). Snook larvae fixed in $10 \%$ formalin/seawater had initial shrinkage of $4.13 \%$ after 30 days and a total shrinkage of $5.81 \%$ after 90 days. Shrinkage of clupeoid larvae fixed in formalin/seawater solutions varied according to the strength of the solution (Hay 1982); salinity (Hay 1982); length of time the larvae stayed in the net (Theilacker 1980; Hay 1981); and whether the larvae were field-caught or laboratory-reared (Blaxter 1971). Larvae of the southern flounder, Paralichthys lethostima, fixed in unbuffered and buffered $10 \%$ formalin/seawater shrank $5.2 \%$ and $7.3 \%$, respectively, when stored in fixative and periodically evaluated over 6 years (Tucker and Chester 1984). In two size classes of freshwater fish, length increased initially following fixation in freshwater formaldehyde solutions; after 65 days in isopropyl alcohol, length approximated live length (Billy 1982).

Alcohol fixation and storage is necessary when otolith examination is desired; however, pH should be carefully monitored (Radtke and Waiwood 1980). Alcohol is still used extensively as a long-term preservative for fish larvae because it is inexpensive and does not carry the health risks of other preservatives. Nonetheless, the degree of shrinkage is greater in alcohol than it is in other fixatives. In addition, tissues become discolored and pigment mark
ings disappear. Pigment retention can be extremely important when larval indentification is required or when life-history series, sensu Moser and Ahlstrom 1970, are a research goal. It should be noted, however, that other factors such as exposure to light and fluctuating temperature may have caused the observed pigment loss.

Alcohol cannot be recommended as a preservative when health risks of alternative preservatives can be greatly reduced with mechanical ventilation of storage facilities and work areas, and safe, proper handling of potentially carcinogenic solutions. Tucker and Chester (1984) recommended $4 \%$ formalin/distilled water buffered with sodium acetate for long-term preservation and storage.

Fish larvae are collected following many different protocols. Morphological parameters such as size, chromatophore patterns, and osteology can be affected by handling, fixation, and storage methods. Evaluation of the effects of fixation and storage on morphology, as evidenced by shrinkage of animals and other morphological changes, is needed to allow critical evaluation of specimens. Choosing the best fixative for use in a study can contribute significantly to obtaining reliable results. Of the five fixatives evaluated in this study, one can not be recommended over the others based on shrinkage alone. How the preserved tissue is to be examined and/or further processed for embedding, sectioning, and staining must be considered in order to choose the most suitable fixative for the research. Different fixatives for aliquots of the same sample have been used when larvae were to be studied for more than one purpose (Oozeki et al. 1989). In the present study, phosphate-buffered glutaraldehyde caused the least amount of shrinkage, whereas live appearance was best maintained when larvae were fixed and stored in phosphate-buffered paraformaldehyde.

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## Citations

Ahlstrom, E.H.
1976. Maintenance of quality in fish eggs and larvae collected during plankton hauls. In Zooplankton fixation and preservation (H.F. Steedman, ed.), p. 313-318. The UNESCO Press, Paris.

Basson, M., A.A. Rosenberg, and J.R. Beddington.
1988. The accuracy and reliability of two new methods for estimating growth parameters from length-frequency data. J. Cons. Int. Explor. Mer 44:277-285.
Billy, A.J.
1982. The effects of formalin and isopropyl alcohol on length and weight measurements of Sarotherodon mossambicus Trewavas. J. Fish. Biol. 21:107-112.
Blaxter, J.H.S.
1971. Feeding and condition of Clyde herring larvae. Rapp. P.-v. Reun. Cons. Int. Explor. Mer 160:128-136.

Brandt, S.B.
1986. Ontogenetic shifts in habitat, diet, and diel-feeding periodicity of slimy sculpin in Lake Ontario. Trans. Am. Fish. Soc. 115: 711-715.
Farris, David A.
1963. Shrinkage of sardine (Sardinops caerulea) larvae upon preservation in buffered formalin. Copeia 1963:185-186.
Gilmore, R.G., C.J. Donohue, and D.W. Cooke.
1983. Observations on the distribution and biology of east-central Florida populations of the common snook, Centropomus undecimalis (Bloch). FL Sci. 46:306-313.
Govoni, J.J., D.E. Hoss, and A.J. Chester. 1983. Comparative feeding of three species of larval fishes in the northern Gulf of Mexico: Brevoortia patronus, Leiostomus xanthurus, and Micropogonias undulatus. Mar. Ecol. Prog. Ser. 13:189-199.
Hay, D.E.
1981. Effects of capture and fixation on gut contents and body size of Pacific herring larvae. Rapp. P.-v. Reun. Cons. Int. Explor. Mer 178:395-400.
1982. Fixation shrinkage of herring larvae: effects of salinity, formalin concentration, and other factors. Can. J. Fish. Aquat. Sci. 39:1138-1143.
Heming, T.A., and R.P. Preston.
1981. Differential effect of formalin preservation on yolk and tissue of young chinook salmon (Oncorhynchus tshawytscha Walbaum). Can. J. Zool. 59:1608-1611.
Hinton, D.E., E.R. Walker, C.A. Pinkstaff, and E.M. Zuchelkowski.
1984. Fixation of teleost tissues. Nat. Cancer Inst. Monogr. 65: 291-320.
Jones, R.
1984. Assessing the effects of changes in exploitation pattern using length composition data. FAO Fish. Tech. Pap. 256, 118 p.
Laroche, J.L.
1982. Trophic patterns among larvae of five species of sculpins (Family:Cottidae) in a Maine estuary. Fish. Bull., U.S. 80: 827-840.
Lavenberg, R.G., G.E. McGowen, and R.E. Woodsum.
1984. Preservation and curation. In Ontogeny and systematics of fishes, p. 57-59. Am. Soc. Ichthyol. and Herpetol., Spec. Publ. 1.
Leslie, J.K., and J.E. Moore. 1986.
Changes in lengths of fixed and preserved young freshwater fish. Can. J. Fish. Aquat. Sci. 43:1079-1081.
MacDonald, P.M.D., and Pitcher, T.J.
1979. Age groups from size frequency data: a versatile and efficient method of analysing distribution mixtures. J. Fish. Res. Board Can. 36:987-1001.
Markle, D.F.
1984. Phosphate buffered formalin for long term preservation of formalin fixed ichthyoplankton. Copeia 1984:525-528.
Moser, H.G., and E.H. Alhstrom.
1970. Development of lanternfishes (family Myctophidae) in the California Current. Part I: Species with narrow-eyed larvae. Bull. Los Angel. Cty. Mus. Nat. Hist. Sci. 7, 145 p.

Oozeki, Y., and R. Hirano. 1988. Effects of Glutaraldehyde fixation on the body size of red sea bream (Pagrus major) larvae. Aquaculture 71:265-269.
Oozeki, Y., T. Ishii, and R. Hirano.
1989. Histological study of the effects of starvation on reared and wild-caught larval stone flounder, Kareius bicoloratus. Mar. Biol. 100:269-275.
Parker, R.R.
1963. Effects of formalin on length and weight of fishes. J. Fish. Res. Board Can. 20:1441-1455.
Pauly, D., and G.R. Morgan (eds.).
1987. Length-based methods in fisheries research. ICLARM Conf. Proc. No. 13, Manila.
Peters, K.M., and R.H. McMichael Jr.
1987. Early life history of the red drum, Sciaenops ocellatus (Pisces: Sciaenidae), in Tampa Bay, Florida. Estuaries 10:92-107.
Radtke, R.L., and K.G. Waiwood.
1980. Otolith formation and body shrinkage due to fixation in larval cod (Gadus morhua). Can. Tech. Rep. Fish. Aquat. Sci. 929:1-10.
Richards, W.J., and F.H. Berry.
1973. Preserving and preparing larval fishes for study. In Proceedings of a workshop on egg, larval and juvenile stages of fish in Atlantic coast estuaries, p. 12-19. NOAA NEFC Sandy Hook Lab. Tech. Rep. 1.
Rosenthal, H., D. Kuhlmann, and O. Fukuhara.
1978. Shrinkage of newly hatched larvae of the Red Sea bream (Chrysophrys major Temminck \& Schlegel) preserved in formalin. Arch. Fischereiwiss. 29:59-63.
Schnack, D., and H. Rosenthal.
1978. Shrinkage of Pacific herring larvae due to formalin fixation and preservation. Ber. Dtsch. Wiss. Komm. Meeresforsch. 26: 222-226.

Schnute, J., and D. Fournier.
1980. A new approach to length frequency analysis: growth structure. Can. J. Fish. Aquat. Sci. 37:1337-1351.
Sheehan, D.C., and B.B. Hrapchak.
1980. Theory and Practice of Histotechnology, 2nd ed. C.V. Mosby Co., St. Louis.
Sokal, R.R., and F.J. Rohlf.
1969. Biometry. W.H. Freeman and Company, San Francisco.

Stickland, N.C.
1975. A detailed analysis of the effects of various fixatives on animal tissue with particular reference to muscle tissue. Stain Technol. 50:255-264.
Taylor, W.R.
1977. Observations on specimen fixation. Proc. Biol. Soc. Wash. 90:753-763.
Theilacker, G.H.
1980. Changes in body measurements of larval northern anchovy, Engraulis mordax, and other fishes due to handling and preservation. Fish. Bull., U.S. 78:685-692.
Tucker, J.W. Jr., and A.J. Chester.
1984. Effects of salinity, formalin concentration and buffer on quality of preservation of southern flounder (Paralichthys lethostigma) larvae. Copeia 1984:981-988.
van der Veer, J.
1982. Simple and reliable methods for fixation, mounting and staining of small and delicate marine plankton for light microscopic identification. Mar. Biol. (Berl.) 66:9-14.
Winer, B.J.
1971. Statistical principles in experimental design. McGraw-Hill, NY.
Yevich, P.P., and C.A. Barszcz.
1977. Preparation of aquatic animals for histopathological examination. U.S. Environ. Prot. Agency, Cincinnati, OH, 22 p.

# Histological Effects from Long-Term Storage of Common Snook Yolk-Sac Larvae in Fixatives and Alcohol 

RUTH O. REESE, MARIE F. DeLEON, and WALTER J. CONLEY

Florida Marine Research Institute
Florida Department of Natural Resources
100 Eighth Avenue, S.E.
St. Petersburg, FL 33701-5095


#### Abstract

Yolk-sac larvae were fixed in five fluids (Bouin's fluid, Davidson's fluid, $3 \%$ glutaraldehyde in 0.1 M phosphate buffer, $10 \%$ paraformaldehyde in 0.1 M phosphate buffer, and $10 \%$ for$\mathrm{malin} / \mathrm{seawater}$ ) and stored for more than two years in fixative or $70 \%$ ethanol. All larvae were dehydrated in $95 \%$ ethanol, infiltrated, and embedded in glycol methacrylate resin. Sagittal sections cut at $3.5 \mu \mathrm{~m}$ were stained with hematoxylin and eosin or periodic acid/Schiff's stain and evaluated for quality of histological results and possible presence of artifacts. Aldehydes provided optimum fixation for histological evaluation, but the best fluid for long-term storage depended upon the aldehyde used.


## Introduction

Histological evaluation has been used to determine the nutritional status of field-caught fish larvae through morphometric evaluation of various structures (Ehrlich et al. 1976; Theilacker 1978), gross condition of various tissues and structures (O'Connell 1976; 1980), and histological indication of active absorption of proteins (Oozeki et al. 1989). Although various fixatives have been used, the discussion of tissue appearance has usually not included possible fixation artifacts even though observations of shrinkage (Hay 1982; Tucker and Chester 1984; DeLeon et al. 1991) and loss of pigments (Ahlstrom 1976) have demonstrated that artifactual changes occur. Further, histological investigations of larval fish tissues have, with few exceptions (Govoni 1980), relied on evaluation of tissues subjected to paraffin embedment, a process that causes substantial shrinkage (Ross 1953; Baker 1958).

In the past, the requirements of paraffin embedment have dictated the use of fixatives containing coagulative chemicals (Baker 1958). Noncoagulative aldehyde fixatives have long been recognized as best for tissues examined by electron microscopy (Glauert 1978). However, most plastics that are stable under the electron beam are water insoluble and not compatible with the majority of histological stains, which are soluble only in water. Introduction of a water-soluble plastic embedding resin, glycol methacrylate, has allowed greater use of aldehyde fixatives for light
microscopy. No previous studies have specifically compared the effects of different primary and combination fixatives on the histology of larval fishes. This paper addresses the effects of commonly used tissue fixatives and preservatives on laboratory-spawned yolk-sac larvae of the common snook (Centropomus undecimalis) embedded in glycol methacrylate.

## Materials and Methods

Four hundred common snook larvae hatched from laboratory-spawned eggs were collected live 18-24 hours after hatching and were placed individually in vials containing approximately 1 mL of fixative. Five fixatives were evaluated: Bouin's fluid (Sheehan and Hrapchak 1980); Davidson's fluid (Yevich and Barszcz 1977); 3\% glutaraldehyde in 0.1 M phosphate buffer; $10 \%$ paraformaldehyde in 0.1 M phosphate buffer; and $10 \%$ formalin in seawater (formalin/seawater). The total sample was divided into groups A and B. Each group was composed of 5 subgroups of 40 individuals, each representing one of the five fixatives. After 3 days in fixative, Group A was serially dehydrated to $70 \%$ ethanol. Group B samples remained in fixative. Both groups were then stored at room temperature on open shelves under normal laboratory lighting for 31 months.

Six larvae from each fixative treatment in groups A and B were dehydrated to $95 \%$ ethanol and embedded in glycol

Table 1
Summary of quality of fixation and preservation of various tissues and structures. $P=$ poor, $A=$ acceptable, $E=$ excellent, $(+)=$ observed, $(-)=$ not observed, FAP $=$ fixed and preserved, H\&E $=$ hematoxylin and eosin.
$\left.\begin{array}{lcccccccc}\hline & \begin{array}{c}\text { Digestive } \\ \text { tract }\end{array} & \begin{array}{c}\text { Muscle } \\ \text { tissue }\end{array} & \begin{array}{c}\text { Brain \& } \\ \text { nerve cord }\end{array} & \begin{array}{c}\text { Nuclear } \\ \text { morphology }\end{array} & \text { Pituitary } & \text { Eye } & \text { Otoliths } & \text { Finfold } \\ \text { stain }\end{array}\right]$


Figure 1
Composite sagittal section of a C. undecimalis larva 24 hours after hatching.
methacrylate (JB-4 Kit, Polysciences ${ }^{\text {TM }}$ ) in BEEM $^{\text {TM }}{ }^{*}$ capsules. Larvae were serially sectioned in the sagittal plane

[^8]at $3.5 \mu \mathrm{~m}$ with glass knives using an LKB 2218 Historange ${ }^{\mathrm{TM}}$. microtome. Sections were placed on acidcleaned slides, baked overnight at $75^{\circ} \mathrm{C}$, and stained as a group with hematoxylin and eosin (H\&E) or Quintero's modified periodic acid/Schiff's stain (QPAS), which uses


Figure 2
Midsagittal sections of C. undecimalis larvae 24 hours after hatching fixed in Bouin's and Davidson's fluids and stored therein or in $70 \%$ ethanol; stained with H\&E. (a) Bouin's - stored in $70 \%$ ethanol; (b) Bouin's - stored in fixative; (c) Davidson's - stored in $70 \%$ ethanol; (d) Davidson's - stored in fixative. $1=$ lens; $m=$ muscle; $n m=$ neuromast; $n=$ nerve cord; $n t=$ notochord; $o=$ otolith; $\mathrm{p}=$ pituitary; $\mathrm{y}=$ yolk.
a counterstain of metanil yellow (Iliana Quintero, Florida Marine Res. Inst., unpubl. manuscr.).

## Results

Each of the combined fixation and preservation treatments was graded as poor, acceptable, or excellent in terms of its effect on the histological clarity of features that are present in snook yolk-sac larvae (Table 1; Fig. 1). Even at low magnification $(\times 50)$, overall differences were evident.

The quality of tissues fixed in Bouin's, whether preserved in fixative or ethanol, was poor (Table 1). The material fixed in Bouin's fluid and preserved in ethanol (Fig. 2a) revealed little tissue contrast with $H \& E$ stain, and basophilia is absent in tissue stored in fixative (Fig. 2b). Tissues fixed and preserved in Bouin's fluid exhibited variable results; e.g., some tissues showed cellular swelling, whereas others were shrunken.

Tissues fixed in Davidson's fluid were of only slightly better quality (Table 1). Yolk was more confluent in appearance (Fig. 2, c and d) than with other fixatives. Staining lacked contrast, especially in the tissue stored in alcohol (Fig. 2c), which almost totally lacked eosinophilia.

Tissues fixed in formalin/seawater and preserved in alcohol shrank substantially, thereby distorting many features (Fig. 3a). However, formalin/seawater fixation with preservation in the fixative resulted in clearer histology than either Bouin's or Davidson's (Fig. 3b) and yielded better stain contrast. Overall, this fixative resulted in poor to acceptable tissues (Table 1).

Alcohol preservation of buffered paraformaldehyde-fixed tissue resulted in preparations that were of slightly better quality than formalin/seawater-fixed tissues (Fig. 3c; Table 1). Paraformaldehyde fixation with storage of tissue in the fixative, however, allowed clear identification of many features characteristic of yolk-sac snook larvae (Table 1; Fig. 3d). Stain contrast was sharp. The globular shape of
$\qquad$


Figure 3
Midsagittal sections of $C$. undecimalis larvae 24 hours after hatching fixed in aldehydes and stored therein or in $70 \%$ ethanol; stained with H\&E. (a) Formalin/seawater - stored in $70 \%$ ethanol; (b) Formalin/seawater - stored in fixative; (c) Paraformaldehyde in phosphate buffer - stored in $70 \%$ ethanol; (d) Paraformaldehyde in phosphate buffer - stored in fixative; (e) Glutaraldehyde in phosphate buffer - stored in $70 \%$ ethanol; (f) Glutaraldehyde in phosphate buffer - stored in fixative. $\mathrm{lv}=$ liver; $\mathrm{m}=\mathrm{muscle} ; \mathrm{mc}=$ mucous cell; $\mathrm{n}=$ nerve cord; on = optic nerve; $\mathrm{p}=$ pituitary; $\mathrm{pc}=$ pancreas; $\mathrm{r}=$ retina.
the finfold mucous cells was obvious, muscle fibers exhibited minimum shrinkage, and mitotic figures were clear.

Tissues fixed in phosphate-buffered glutaraldehyde and preserved in ethanol (Table 1; Fig. 3e) provided better
structural delineation than any other treatment. Preservation in glutaraldehyde (Fig. 3f) achieved similar fixation quality, but the tissue was brittle and difficult to handle without fracture. Stain contrast was acceptable, but not

as well defined as with other aldehyde preparations (Table 1).

Examination of tissue at higher magnification ( $\times 330$ ) revealed more detailed differences among treatments. For
example, in neural tissue, small neuroepithelial cells with dark basophilic nuclei appeared in all treatments (Fig. 4). Cells with large, round, lightly basophilic nuclei and a single nucleolus, representing differentiating neurons, were
most clearly seen in tissue fixed with paraformaldehyde or glutaraldehyde (Fig. 4, d and e). In contrast, poor fixation resulted in nerve cord shrinkage (Figs. 2c; 3, a and c), which grossly distorted the appearance of individual cells.

Fixation of the digestive tract was adequate in most fixative/ preservative combinations (Table 1). In tissues fixed in Bouin's and Davidson's fluids, clumped mitotic figures were abundant and nuclei were centrally located within swollen cells with coagulated cytoplasm (Fig. 5, a and b). Preservation of formalin/seawater-fixed tissue in alcohol produced substantial shrinkage and pycnotic nuclei (Fig. 5c). In paraformaldehyde- and glutaraldehyde-fixed tissue (Fig. 5, d and e), nuclei were in the proximal third of the cuboidal cell, and the homogeneous cytoplasm revealed no evidence of inclusions. Only a small liver primordium was present, posterior and lateral to the yolk (Fig. 3e). Liver mitotic figures were clear in the aldehyde fixatives. Two to three exocrine pancreatic acini were obvious in all larvae fixed in aldehydes (Fig. 3, b and f).
In paraformaldehyde- and glutaraldehyde-fixed larvae, the chevron-shaped segmented myomeres consisted of confluent myofibers with multiple, oval, centrally located nuclei with a single nucleolus (Fig. 6) and well-defined myosepta. Shrinkage was minimal, but the cross striations (A and I bands) were not sharp. In tissues fixed in Bouin's, Davidson's, and formalin/seawater, the muscle fibers were shrunken and separated and myosepta were not clear (Fig. 4, a through c). However, cross striations (Fig. 5c) were well defined.

Most nuclei in developing snook larvae were basophilic and processed granular chromatin, which is typical of interphase. In some nuclei, single or multiple nucleoli were visible. In Bouin's and Davidson's fixatives, all nuclei had peripherally clumped chromatin with a central body resembling a nucleolus. This chromatin clumping also interfered with the observation of mitotic stages (Fig. 4, a and $b$ ). The preservation of mitotic figures was superior in glutaraldehyde-fixed tissue (Fig. 7).
In tissues fixed in glutaraldehyde, the differentiating lens fibers of the eye were fully nucleated and minimally shrunken. The retinal layer was clearly defined and its basal area contained many mitotic figures, but no pigment was observed (Fig. 8). In aldehyde-fixed tissue, the large optic nerve with its sheathing cells (Fig. 3 a, d, and f) could be followed from the choroid fissure to the contralateral side of the brain, where it terminates in the diencephalon (Fig. 3d). In poorly fixed tissues, the differentiated areas of the eye were less obvious, the retinal structure was unclear, the optic nerve was not visible, and the lens was severely shrunken (Fig. 2a).
In the otic capsule, two of the three otolith pairs, probably sagittae and lapilli, were visible when fixed in glutaraldehyde and preserved in ethanol (Fig. 9). Otoliths were present, but not optimally fixed, in all animals fixed in
aldehydes and preserved in alcohol and in all larvae fixed in Davidson's, whether preserved in alcohol or fixative (Table 1; Fig. 2c).
Structural definition of the lobular pituitary, anterior to the tip of the notochord, was best in aldehydes (Figs. 3, b and e; and 10). In Davidson's- and Bouin's-fixed tissues, a separate pituitary structure was not well defined (Fig. 2d).
The finfold is composed of a two-layered epithelium with many mucous cells subtended by an acellular matrix. Not evident in most fixed sections, however, is a dense fibrillar material seen in larval tissue that has been fixed and preserved in glutaraldehyde and stained with QPAS (Fig. 11). Hematoxylin staining of the same tissue was unremarkable (Fig. 2f). In glutaraldehyde-fixed material, the mucous cells were spherical and their basal nuclei and granular contents were well defined (Figs. 3, e and f; 7; 8 ; and 10 ). In poorly fixed tissue, the finfold epithelium appeared as a thin filamentous structure with a few poorly defined cells (Fig. 2). Many neuromasts were present along the finfold epithelium in all preparations, but structural detail was variable (Figs. 2a and 8). Nerve tracks subtending peripheral neuromasts were seen only in glutaralde-hyde-fixed tissues (Fig. 8).

## Discussion

In general, chemical fixatives are either coagulant or noncoagulant. A specific chemical may be preferred for preservation of a particular tissue or structure or to aid in diagnostic staining. However, some chemicals may have a negative impact on tissues, causing swelling, shrinkage, or other types of cellular distortion. Therefore, primary chemical fixatives are often combined in an attempt to balance the defects of one primary fixative by adding another fixative with compensating properties. For example, two of the most commonly recommended fixative mixtures for histological examination of teleost tissues are Bouin's and Davidson's solutions. In Bouin's fluid, picric acid, a coagulant fixative that facilitates the admission of paraffin by tissues, also shrinks tissue and renders chromatin acidophilic. Acetic acid, a noncoagulant component, is used to compensate for picric acid. However, acetic acid swells protein gels and fibers and produces a precipitate with nucleoprotein. The formaldehyde in Bouin's fixes cytoplasm well, defines chromosomes poorly, and hinders the penetration of paraffin into tissues (Baker 1958).

The lack of basophilia in tissues preserved for over two years in Bouin's was not surprising, because fluids intended only for rapid fixation, such as Bouin's and other coagulative mixtures, invariably lead to excessive hardening (Steedman 1976). Retention in such fixatives is not normally recommended, but was included here as part of an overall comparison.


Figures 6-11
6. Myomere of $C$. undecimalis larva 24 hours after hatching, fixed in paraformaldehyde and stored in fixative; $\mathrm{H} \& E$ stain. dt $=$ digestive tract $; \mathrm{mb}=$ myofiber; $\mathrm{ms}=$ myosepta; $\mathrm{my}=$ myomere; ne $=$ neuro-epithelial cell; $\mathrm{nt}=$ notochord.
7. Mitotic figures in liver of $C$. undecimalis larva 24 hours after hatching, fixed in glutaraldehyde and stored in ethanol; H\&E stain. mc - mucous cell; $\mathrm{mf}=$ mitotic figure; $\mathrm{y}=$ yolk.
8. Eye and neuromast of $C$. undecimalis larva 24 hours after hatching fixed in glutaraldehyde and stored in alcohol; PAS/metanil yellow stain. $1=$ lens; $\mathrm{mc}=$ mucous cell; $\mathrm{mf}=$ mitotic figure; $\mathrm{nm}=$ neuromast; $\mathrm{nn}=$ neuromast nerve; $\mathrm{r}=$ retina.
9. Otic capsule and otolith of $C$. undecimalis larva 24 hours after hatching fixed in glutaraldehyde and stored in alcohol; H\&E stain. $\mathrm{dn}=$ differentiating neuron; $\mathrm{n}=$ nerve cord; $\mathrm{o}=$ otolith.
10. Pituitary, tip of notochord, and edge of finfold with mucous cells of $C$. undecimalis larva 24 hours after hatching fixed in glutaraldehyde and stored in alcohol; H\&E stain. $\mathrm{mc}=$ mucous cell; $\mathrm{nt}=$ notochord; or $=$ oral cavity; $\mathrm{p}=$ pituitary.
11. Muscle and finfold of $C$. undecimalis larva 24 hours after hatching fixed in glutaraldehyde and stored in fixative; PAS/metanil yellow stain. em $=$ external milieu; $\mathrm{fm}=$ finfold matrix; $\mathrm{m}=$ muscle.

Davidson's fluid consists of the noncoagulant fixatives acetic acid and formaldehyde combined with the coagulant fixative ethanol. In addition to the previously mentioned effects of acetic acid and formaldehyde, ethanol denatures protein, precipitates nucleic acids without rendering them insoluble in water (Baker 1958), and shrinks and hardens tissues. Glycerol, which is not a fixative, was added to this combination as a softening agent, enhancing fixation results (Hinton et al. 1984).

The most widely used noncoagulant fixatives are aldehydes, principally formaldehyde and glutaraldehyde. The distinction between formalin and formaldehyde is often not clearly defined. Commercial formalin consists of $37-41 \%$ formaldehyde dissolved in water containing many trace elements and 7-13\% methanol, which is added to prevent polymerization (Steedman 1976). Paraformaldehyde is a polymer that can be prepared to yield a pure, relatively stable, aqueous formaldehyde without additives. The crosslink bonds of this monoaldehyde are not as stable as those of glutaraldehyde, and soaking tissues in water can remove many of the cross-linking bonds, thus destroying some of the valuable effects of formalin fixation (Stickland 1975). Glutaraldehyde is a dialdehyde and stabilizes structures by cross-linking before cellular contents are extracted (Glauert 1978). Tissue fixation in glutaraldehyde results in better structural retention and slightly less shrinkage than in formaldehyde (Stickland 1975). Long-term storage in buffered glutaraldehyde caused tissue to be brittle and difficult to handle without fracture. This dialdehyde produces strong cross-links between protein molecules and results in a progressively growing reticulum inside the cell (Geyer 1973). This continued cross-linking may cause the brittle tissue condition noted with long-term storage in glutaraldehyde. The presence of extensive aldehyde groups is also suggested by intensified staining with QPAS, which links through aldehydes (Sheehan and Hrapchak 1980). The finfold matrix, not normally evident in fixed sections, appeared in QPAS-stained material of glutaraldehyde-fixed and stored larvae as a dense fibrillar matrix. Muscle tissue also showed clumping of stain.

In the present study, three variations of aldehyde fixation were evaluated: commercial formalin buffered with seawater, commonly used for life-history studies; paraformaldehyde buffered with phosphate, the fixative of choice for histological preparation in many laboratories; and glutaraldehyde buffered with phosphate, a fixative introduced in 1963 for electron microscopy (Sabatini et al. 1963) that has also been used successfully with a variety of teleost tissues (Watanabe 1981; Kessel et al. 1985). Only recently has glutaraldehyde been used for ichthyoplankton preservation (Oozeki and Hirano 1988). Glutaraldehyde fixation causes tissues to become hard during paraffin embedding. Differences in hardness between the paraffin embedding medium and the processed tissue causes tissue fracture during sectioning. The hard plastic embedding
media used for electron microscopy infiltrate glutaralde-hyde-fixed tissue well and allow thin sectioning without fracture.

In the past, the requirements of paraffin embedding have, to a large extent, controlled the choice of fixatives. However, Baker (1958) speculated that the introduction of new embedding media was likely to result in less reliance on coagulative fixatives. Water-insoluble plastic embedding media are used for electron microscopy. Because the spectrum of histological stains comprises mostly watersoluble pigments, the tinctorial choices for tissues embedded in these early resins were greatly reduced. Introduction of glycol methacrylate, a water-soluble resin, not only provided a harder substrate easily sectioned at $2 \mu \mathrm{~m}$ but also was expected to allow the use and/or adaptation of diagnostic stains developed for paraffin. Although few diagnostic stains available for paraffin preparations have been successfully adapted for use with glycol methacrylateembedded tissues, some stain combinations have proved suitable (Govoni 1983). For this investigation, a modified $\mathrm{H} \& \mathrm{E}$ stain and PAS technique using metanil yellow as a counterstain, which yields excellent histological detail (Quintero, in prep.), were used. In addition, infiltration and embedment in glycol methacrylate can be accomplished at relatively low temperatures rather than the $55-65^{\circ} \mathrm{C}$ required for paraffin, thus reducing the degree of temperature-induced hardening and artifact.

The best histological results were achieved with fixation in the two pure buffered aldehydes, but the best preservative depended upon the aldehyde used. Fixation in glutaraldehyde and subsequent preservation in ethanol were superior for most tissues. Nerve tracks subtending peripheral neuromasts (Fig. 8) were seen only in glutaral-dehyde-fixed tissue; perhaps greater initial shrinkage in other fixative fluids (DeLeon et al. 1991) caused a separation of these fine nerves. Fixation in phosphate-buffered paraformaldehyde with subsequent preservation in the fixative also resulted in generally excellent histology.

Hopwood (1969) stated, "No one fixative is ideal for all situations." Careful selection of fixative is critical in the development of protocols for evaluation of teleost tissues. For example, the tissue may be prepared for examination of external features, histological evaluation requiring histochemical techniques or the application of other special research methodologies. In clinical situations, multiple fixation procedures are often used to provide for contingencies. For human surgical biopsies, Dawson (1973) recommended that samples be divided into three pieces: one to be fixed for standard paraffin embedding and subsequent histological staining, a second to be quenched (rapidly frozen below $-70^{\circ} \mathrm{C}$ ), and a third to be fixed in aldehyde for later examination using electron microscopy. Fisheries work seldom allows the luxury of such a complete initial preparation. The immediate processing and embedment of collected tissues is often impossible. Therefore, not only
should the fixation method be carefully evaluated, but additional consideration should be given to the choice of storage fluid because the preservative also has a significant effect on tissue quality. Whatever fixative technique is employed, researchers must be aware that the ultimate goal of histological fixation of tissues is not simply to avoid decay but also to clarify understanding of ongoing processes by obtaining a sample of 'frozen physiology."

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## Citations

Ahlstrom, E.H.
1976. Maintenance of quality in fish eggs and larvae collected during plankton hauls. In Zooplankton fixation and preservation (H.F. Steedman, ed.), p. 313-318. UNESCO Press, Paris.
Baker, J.R.
1958. Principles of biological microtechniques. John Wiley and Sons, Inc. NY, 357 p.
Dawson, I.M
1973. Fixation: what should the pathologist do? In Fixation and histochemistry (P.J. Stoward, ed.). Chapman Hall, London.
DeLeon, M.F., R.O. Reese, and W.J. Conley.
1991. Effects of fixation and dehydration on shrinkage and morphology in common snook yolk-sac larvae. In Larval fish recruitment research in the Americas: proceedings of the thirteenth annual larval fish conference; 21-26 May 1989, Merida, Mexico (R.D. Hoyt, ed.), p. 121-128. Dep. Commer., NOAA Tech. Rep. NMFS 95.
Ehrlich, K.F., J.H.S. Blaxter, and R. Pemberton.
1976. Morphological and histological changes during the growth and starvation of herring and plaice larvae. Mar. Biol. 35:105-118.
Geyer, G.
1973. Ultrahistochemie. Gustan Fischer Verlag., Stuttgart, 473 p. Glauert, A.M.
1978. Fixation, dehydration and embedding of biological specimens. North-Holland Publishing Company, NY, 207 p.
Govoni, J.J.
1980. Morphological, histological and functional aspects of alimentary canal and associated organ development in larval Leiostomus xanthurus. Rev. Can. Biol. 39:69-80.
1984. Histology. In Ontogeny and systematics of fishes, p. 40-42. Am. Soc. Ichthyol. Herpetol., Spec. Publ. 1.
Hay, D.E.
1982. Fixation shrinkage of herring larvae: effects of salinity, formalin concentration, and other factors. Can. J. Fish. Aquat. Sci.

39:1138-1143.
Hinton, D.E., E.R. Walker, C.A. Pinkstaff, and E.M. Zuchelkowski. 1984. Fixation of teleost tissues. Natl. Cancer Inst. Monogr. 65:291-320.
Hopwood, D.
1969. Fixatives and fixation: a review. Histochem. J. 1:323-360.

Kessel, R.G., H.N. Tung, R. Roberts, and H.W. Beams.
1985. The presence and distribution of gap junctions in the oocyte follicle cell complex of the zebrafish, Brachydanio rerio. J. Submicrosc. Cytol. Pathol. 17:239-253.
O'Connell, C.P.
1976. Histological criteria for diagnosing the starving condition in early post yolk-sac larvae of the northern anchovy, Engraulis mordax Girard. J. Exp. Mar. Biol. Ecol. 25:285-312.
1980. Percentage of starving northern anchovy, Engraulis mordax, larvae in the sea as estimated by histological methods. Fish. Bull., U.S. 78:475-489.

Oozeki, Y., and R. Hirano.
1988. Effects of glutaraldehyde fixation on the body size of Red Sea bream (Pagrus major) larvae. Aquaculture 71:265-269.
Oozeki, Y., T. Ishii, and R. Hirano.
1989. Histological study of the effects of starvation on reared and wild-caught larval stone flounder, Kareius bicoloratus. Mar. Biol. 100:269-275.
Ross, K.F.A.
1953. Cell shrinkage caused by fixatives and paraffin-wax embedding in ordinary cytological preparations. Q. J. Microsc. Sci. 94:125-139.
Sabatini, D.D., K. Bensch, and R.J. Barnett.
1963. Cytochemistry and electron microscopy: The preservation of cellular ultrastructure and enzymatic activity by aldehyde fixation. J. Cell Biol. 17:19-58.
Sheehan, D.C., and B.B. Hrapchak.
1980. Theory and practice of histotechnology, 2nd ed. C.V. Mosby Co., St. Louis, Toronto, London, 481 p.
Steedman, H.F
1976. General and applied data on formaldehyde fixation and preservation of marine zooplankton. In Zooplankton fixation and preservation (H.F. Steedman, ed.), p. 103-154. UNESCO Press, Paris.
Stickland, N.C.
1975. A detailed analysis of the effects of various fixatives on animal tissue with particular reference to muscle tissue. Stain Technol. 50:255-264.
Theilacker, G H.
1978. Effects of starvation on the histological and morphological characteristics of jack mackerel Trachurus symmetricus larvae. Fish. Bull., U.S. 76:403-413.
Tucker, J.W. Jr., and A.J. Chester.
1984. Effects of salinity, formalin concentration and buffer on quality of preservation of southern flounder (Paralichthys lethostigma) larvae. Copeia 1984:981-988.
Yevich, P.P., and C.A. Barszcz.
1977. Preparation of aquatic animals for histopathological examination. U.S. Environ. Prot. Agency, Cincinnati, Ohio, 22 p.
Watanabe, Y.
1981. Ingestion of horseradish peroxidase by the intestinal cells in larvae or juveniles of some teleosts. Bull. Jpn. Soc. Sci. Fish. 47:1299-1307.


[^0]:    *Miembro de la Carrera del Investigador Científico y Tecnológico, Consejo Nacional de Investigaciones Científicas y Técnicas.

[^1]:    Universidad Prat, Facultad de Ciencias del Mar, Iquique: W. Palma Saez, J.L. Pizarro, R. Correa, V. Fernandez.

[^2]:    * Present address: Department of Oceanography, Dalhousie University, Halifax, NS, B3H 4J1, Canada.

[^3]:    'Johnson and Greenfield (1983) (Northeast Gulf Science 6(1):33-49) consider Gobiesox barbatulus rather than $G$. strumosus to be the form ocurring in Brazil. Based on the fact that there is only one species of Gobiesox in the Patos Lagoon, we decided to use G. strumosus following Chao et al. (1982b). Further investigation to determine which form is present in the southern coast of Brazil is necessary, but it is beyond the scope of this study.

[^4]:    * Virginia Institute of Marine Science Contribution No. 1572.
    ** Present address: University of Alaska Fairbanks, Institute of Marine Science, Fairbanks. AK 99775.

[^5]:    * Present address: NOAA, Office of Undersea Research SSMC1 R/OR2, 1335 East-West Hwy., Silver Spring, MD 20910.

[^6]:    * Present address: U.S. Fish and Wildlife Service, Tenncssee Cooperative Fishery Research Unit, Box 5114, Tennessee Technological University, Cookeville, Tennessee 38505
    ** Present address: U.S. Forest Service, P.O. Box 1148, Corvallis, Oregon 97339.

[^7]:    * The title of NOAA Technical Report NMFS 95 unfortunately does not reflect this single contribution from South Africa. To have included the words "South Africa" in the title of the proceedings might have misled readers into assuming multiple contributions from this country. Managing Editor.

[^8]:    * The use of commercial products does not imply endorsement by Florida Department of Natural Resources.

