## ANNUAL REPORT 2001

## Monitoring the Marine Environment of Long Island Sound at Millstone Power Station Waterford, Connecticut



## Millstone Environmental Laboratory

# Monitoring the Marine Environment of Long Island Sound at Millstone Power Station 

## 2001 Annual Report

## Prepared by:

Staff of
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April 2002


We dedicate this report to Emesto Lorda, who will be retiring later this year from the Millstone Environmental Laboratory (MEL). Ernesto has an extensive and rigorous educational background that makes him an especially valued staff member. He received an B.S. in Mechanical Engineering from the University of Zaragoza in Spain, an M.S. in Ocean Engineering from the University of Massachusetts at Amherst, and a Ph. D. in Biological Oceanography and Applied Statistics from the Graduate School of Oceanography of the University of Rhode Island. Before joining the staff at MEL in 1982, Emesto had a diverse professional career, having worked as a design and field engineer in hydraulic construction and public works, as a consulting engineer, and as a university research associate. He has many professional certifications and memberships, including being a Charter Member of the Environmental Statistics Section of the American Statistical Association. He has been solicited to review numerous articles for science journals. He has authored or co-authored more than 20 papers published in the scientific literature, wrote numerous technical reports, and presented his work at symposia and national and regional conferences. His primary responsibilities at MEL, formerly for Northeast Utilities, and currently for Dominion include biometrics and statistics and population dynamics modeling. He provides oversight in these fields and guides staff scientists in data analyses, experimental design, impact assessments, and modeling. His dissertation research at the University of Rhode Island involved the stochastic modeling of the dynamics of fish populations, which he was later able to adapt specifically to an issue of interest at Millstone Power Station. This work became the foundation for the stochastic population dynamics model, which has been the primary long-term forecasting tool for the Niantic River winter flounder population in assessing the effects of Millstone Station, the environment, and fishing on this stock of fish. The model and its approach to impact assessment has received considerable acclaim and notice through the electric utility industry. He is also an accomplished writer and editor whose work has undeniably bettered our reports, including the present one. Over the years, Emesto has interacted constructively with every staff member of MEL as well as with other departments and environmental and station management. He also provides trusted advice to the Ecological Advisory Committee of MEL in critical analyses in fisheries biology, experimental design, and statistical methodologies. In particular, he worked closely for many years with his friend and mentor, Dr. Saul Saila, Professor Emeritus of the University of Rhode Island, in examining innovative methodologies to use with fisheries data and in impact assessments of power plant operations. Dr. Saila recently noted that "Ernesto was one of the most versatile and competent of the many graduate students I had the pleasure of working with." We will sorely miss Emesto's advice and wise counsel and wish him well in any and all future endeavors he may wish to pursue.

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## Acknowledgements

This report was prepared by the staff of Dominion Nuclear Connecticut, Inc. Environmental Laboratory, located at Millstone Nuclear Power Station, PO Box 128, Waterford, CT 06385.

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Special appreciation is extended to the summer intern staff from the University of Connecticut, Connecticut College, and Three Rivers Community-Technical College for their considerable efforts in the field and laboratory: Eric Brzozowski, Dave Boettcher, Kate Driscoll, Tom Hochdorfer, Brian James, Jakub Kircun, and Eric Reyer. Penelope Howell, David Molnar, and David Simpson of the Connecticut Department of Environmental Protection, Fisheries Division; David Taylor of the University of Rhode Island, Graduate School of Oceanography; Thomas Currier of the Massachusetts Division of Marine Fisheries; Tim Lynch of the Rhode Island Department of Environmental Management, Division of Fish and Wildlife; Dr. Michael Scherer of Marine Research, Inc.; and Matt Male of Old Lyme, CT kindly supplied information on finfishes and lobster used in this report. Critical reviews of this report and oversight of the monitoring programs were completed by the following members of the Millstone Ecological Advisory Committee: Dr. John Tietjen (chair; emeritus, City University of New York), Dr. W. Huntting Howell (University of New Hampshire), Dr. Nelson Marshall (emeritus, University of Rhode Island), Dr. William Pearcy (emeritus, Oregon State University), Dr. Saul Saila (emeritus, University of Rhode Island), Dr. Robert Whitlatch (University of Connecticut), and Dr. Robert Wilce (emeritus, University of Massachusetts).

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## Executive Summary

## Lobster Studies

The American lobster (Homarus americanus) is the single most valuable fisheries species in New England. In Connecticut, the regional economic value of the lobster fishery is between $\$ 12$ and $\$ 15$ million annually. Lobsters are also one of the largest mobile benthic invertebrates in the western North Atlantic and important inhabitants of coastal ecosystems. Several aspects of Millstone Power Station (MPS) operation could potentially impact the local population of American lobster. In early summer, after hatching from eggs, lobster larvae swim to the surface to begin the 6 - to 8 -week planktonic phase of their life cycle and are susceptible to entrainment through cooling water systems. Juvenile and adult lobsters can be impinged on intake travelling screens or be exposed to the heated effluent in the discharge area. Because of the economic and ecological importance of this species, lobsters have been monitored from May through October since 1978, using wire lobster traps set at three stations around MPS. Since 1984, entrainment studies have been conducted during the hatching season, to estimate the number of lobster larvae drawn through the cooling water system. The objective of the lobster monitoring program is to determine if operation of MPS has caused changes in local lobster abundance and population characteristics beyond those expected from changes in the fishery and natural variability.
Despite intense and increasing fishing effort on the species, landings made in the commercial fishery throughout New England have increased markedly over the last three decades. Commercial landings of lobsters in Connecticut waters of Long Island Sound (LIS) increased from 0.7 million pounds in 1979 to a record 5.6 million pounds in 1998. However, landings declined $70 \%$ between 1998 and 2000 to only 1.7 million pounds. Landings declined even further in 2001 when only 1.5 million pounds were harvested. The recent decline in lobster abundance has been attributed to a significant lobster mortality event in western LIS. Beginning in the fall of 1998 and throughout the summer of 1999, an unexplained highly virulent illness emerged in lobsters harvested from western LIS. Lobstermen reported that more than half of the lobsters caught in traps were dead or died before arriving at port. In some areas of western LIS landings have declined more than $90 \%$ over the past few years. Many researchers suspect that the die-off in western

LIS was caused by multiple disease pathogens exacerbated by environmental stressors, such as increased seawater temperature, anoxia and contaminants. Concurrently with the lobster die-off in western LIS, a major outbreak of shell disease began in 1998 in eastern LIS and nearshore coastal areas of Rhode Island and Massachusetts. Naturally occurring bacteria and fungi have been implicated as causative agents of shell disease. These microorganisms consume the exoskeletons of crustaceans, resulting in a shell that is pitted and marred with necrotic lesions. The unsightly appearance of the shell greatly affects lobster marketability and although shell disease is not immediately fatal, death may ultimately occur.
In response to the die-off in western LIS and shell disease outbreak in eastern LIS, the U.S. Secretary of Commerce declared a failure of the LIS commercial lobster fishery in January, 2000. In 2001, the U.S. Congress allocated $\$ 13.9$ million in federal disaster funds for the LIS lobster industry. Half of this Federal aid has been earmarked for research investigations on causes and economic impacts of the die-off and shell disease in LIS lobsters.
The most notable changes in the population characteristics of local lobsters during 2001 were observed in the proportion of egg-bearing females and their size structure, which may be related to changes in lobster growth and the size at which females become sexually mature. The benefits of earlier maturation may be significant to lobster population dynamics.
Small size at maturity and subsequent egg production from sublegal-sized females may explain why LIS lobsters are so resilient in the face of intense exploitation. : Currently, the smaller size at sexual maturity allows females to spawn once or twice before reaching the legal size and could maintain current population levels into the future. However, the longterm sustainability of the Connecticut fishery may now be threatened by the lobster die-off in western LIS and the widespread outbreak of shell disease in our area. During 2001, we found that egg-bearing female lobsters had the highest prevalence and severity of shell disease and although the effects of the disease on eggdevelopment, hatching success and larval survival are unknown, the density of lobster larvae collected in samples of the MPS cooling water was the lowest observed in 18 years of lobster larvae entrainment studies. Alternatively, if larvae found in our area originate from the west, the lobster die-off in western

LIS could be responsible for the low number of larvae found in our study area during 2001. The events of mass mortality in western LIS and the effects on recruitment and subsequent legal lobster abundance in our area due to the recent outbreak of shell disease could be significant, if egg-bearing females suffer higher mortality.
At present, the die-off and subsequent catastrophic decline in lobster abundance observed in western LIS has not been observed in our study area in eastern LIS. Although catch-per-unit-effort of 1.587 lobster/pot during 2001 continued to decline from the record high of 2.560 observed in 1999, the 2001 value was within the range of annual catch-per-unit-effort (CPUE) values observed since 1978. In addition, our results indicated a significant increasing trend in total lobster catch over the past 24 years. Furthermore, despite the fact that nearly all the legal-size lobsters are removed by fishing each year, no significant long-term trend in legal lobster abundance was observed in our area. These findings suggest that operation of the MPS cooling water system since the early 1970s has not caused a decrease in the local lobster population. Our monitoring studies indicate that the changes observed in lobster population characteristics and fluctuations in lobster abundance are related more to a combination of natural environmental processes (seawater temperature), episodic events (diseases), and overfishing, than to operations of MPS.

## Benthic Infauna

Studies of benthic infaunal communities inhabiting marine sedimentary environments in the vicinity of MPS have documented long-term and short-term trends in community and population-level parameters. Analysis of long-term data has provided evidence of acute and chronic impacts, some related to power plant construction and operation, and others, to presumably natural disturbances. Infaunal communities exhibiting evidence of MPS impacts were observed at the stations in the immediate vicinity of MPS, including Intake (IN), Effluent (EF), and Jordan Cove (JC). Community changes at these sites were closely associated with changes in sediment composition related to power plant-induced disturbance. The infaunal community at Giants Neck (GN) reference site was unaffected by MPS and provided a baseline from which variability associated with natural environmental cycles and disturbances could be assessed.

The reference site (GN), well beyond any MPS influence has exhibited changes in sediments and community structure over the entire period. Sediment
mean grain size was highly variable but siltclay was relatively consistent, with no long-term trends in either parameter. Infaunal community composition at GN station has generally been dominated by three taxa (Tharyx spp., oligochaetes and Aricidea catherinae); however, some long-term trends in abundance of these taxa were noted. A. catherinae, common at GN in the early monitoring years, has declined in abundance, and Tharyx spp. abundance has increased over the 22 -year study period. Some species at GN have exhibited relatively high year-toyear fluctuations that have proved useful in validating area-wide shifts in species abundance and community structure in response to natural causes and unrelated to MPS operation. For example, there was a pulse in abundance of the polychaete Polydora socialis at all stations in 2001 indicating a regional phenomenon. Similar population shifts occurred for the opportunistic polychaete Mediomastus ambiseta, at GN (and at stations near MPS) during 1983-88, 1994, and 1998-99. Other widespread population shifts in past study years included large increases in abundance of the amphipods Leptocheirus pinguis and Ampelisca spp., and of the polychaetes Prionospio steenstrupi and Pygospio elegans.

Infaunal community changes associated with MPS were first observed at IN and were caused by dredging and cofferdam removal during Unit 3 construction from 1983 to 1985 . Since then, sediments (primarily siluclay content. levels) have stabilized and become more similar to sediments of pre-impact years. Evidence of some infaunal community stabilization and recovery was also apparent. In particular, numbers of individuals and species richness at IN have increased over the study period, as have abundances of organisms typically more common in early study years or at other sites, such as A. catherinae and Tharjx spp. These trends, along with concomitant decreases in abundance of Nucula annulata and other opportunistic species (e.g., the amphipods $L$ pinguis and Ampelisca spp.), indicate some degree of recovery at IN. Recovery at IN is not complete, however, as other organisms which have established post-impact community dominance, such as oligochaetes and Protodorvillea gaspeensis, maintained or increased their degree of dominance in recent years and may persist indefinitely.

Silt deposition at JC attributed to sediment scouring from MPS discharge area following Unit 3 start-up in 1986 resulted in increased sediment silt/clay content and abrupt changes in infaunal community structure. The altered community has exhibited changes in recent years, moving toward pre-impact community structure. For example, A. catherinae and Tharyx spp. abundances have approached levels observed
prior to 1986. However, continued trends toward recovery noted in previous years (e.g., reduced siluclay content during 1994 and 1995, and rebounding abundances of oligochaetes and Polycirrus eximius through 1993) turned out to be short-term community changes no longer evident in recent data. Additionally, the opportunistic molluse N. annulata has maintained a population at JC through 2001 that is still well above pre-Unit 3 levels. This biological evidence of disturbance is consistent with trends in sediment characteristics; elevated sildclay levels in sediments at JC were still evident in 2001.

The benthic infaunal site that experiences chronic MPS operational impact processes is the EF station. Current scour from the MPS discharge directly impacted both the sediments and the infaunal community 100 m from the discharge at EF , particularly after Unit 3 startup in 1986. Relatively coarse sediments with low silt/clay levels were characteristic of the 3 -unit benthic habitat at EF. While sediment characteristics remain different from those observed prior to 1986, the altered sedimentary environment at EF, and the infaunal community it supports, stabilized under the new environmental conditions created by the 3 -unit discharge. Discharge flow from Units 2 and 3 since 1998 has had little effect on this stabilization,' which has allowed for rebounds of Tharyx spp. and A. catherinae, taxa common prior to 1986. In addition, oligochaete and P. gaspeensis abundances have generally decreased from high abundances during the early 3 -unit period. Oligochaete abundances at EF during the last two years ( 2000 and 2001) were among the lowest observed during the study period, possibly related to the heavy mussel set observed in the discharge area in 2000.

## Eelgrass

Eelgrass (Zostera marina) is an ecologically important component of shallow water habitats. Eelgrass population monitoring has been part of marine environmental studies in the vicinity of MPS since 1985. Eelgrass populations at three locations were monitored during 2001: Jordan Cove (JC), White Point (WP) and Niantic River (NR). . The JC and WP sites have been consistently sampled since 1985, while the NR station has been changed five times because of declines in the overall abundance of eelgrass in the Niantic River.
Eelgrass beds at the two monitoring sites nearest MPS (JC and WP) have exhibited slight, but statistically significant, long-term declines in most population parameters. However, compared to the

Niantic River, they have generally supported healthy populations, based on parameters monitored throughout the 17 -year study. These two populations are exposed to variability in water temperature more indicative of natural solar warming and hydrodynamic conditions in Jordan Cove than the MPS discharge. These natural factors are particularly influential at the JC sampling site, which is the most shallow of the siudy sites, and is immediately adjacent to extensive shallow sand flats vulnerable to solar warming. In addition to temperature, there does not appear to be any relationship between WP and JC population fluctuations and power plant discharge flow and heat output. With Unit 1 permanently retired, the likelihood of thermal plume incursion at JC and WP in the future is further reduced.

Eelgrass shoot density and standing stock biomass estimates have been most variable during the study period in the Niantic River, where reference stations have been monitored. A new sampling site was established in the Niantic River in 2000 within the only remaining viable eelgrass bed found in this estuary. Site relocation was necessitated by the 1999 dic-off of the eelgrass bed where the previous site had been located. This was the fifth time since 1985 that we have documented loss of an eelgrass bed in the Niantic River. The condition of the present bed is in question, as standing stock and shoot density estimates were low at this site in 2000 and 2001. Because the Niantic River is located well away from any influence of the MPS thermal plume, declines at NR sites cannot be attributed to MPS operation. The spatial pattern of decline among study populations in the vicinity of MPS suggests a relationship to nutrient enrichment. Niantic River eelgrass beds may be more susceptible to nutrient loading from fertilizers and domestic septic systems due to restricted tidal exchange of the estuary and proximity to expanding housing developments when compared to the JC and WP sites in Jordan Cove. Eelgrass declines in coastal embayments similar to the Niantic River in Rhode Island and Massachusetts have been directly linked to increased housing development within their watersheds.

## Rocky Intertidal Studies

Several important operational events during more thạn 20 years of rocky intertidal monitoring have resulted in identifiable ecological changes to the shore community near the MPS discharge. While measurable, these changes are not widespread, but remain restricted to approximately 150 m of shoreline on the east side of the power plant discharge to LIS. Thermal impacts to the shore community at Fox

Island were first observed in 1983, after the opening of the second quarry cut. Thermal addition to this site was modified when Unit 3 began commercial operation in 1986, and 3 -unit operating conditions over the next 11 years allowed for long-term successional community development under a relatively consistent thermal regime. This community exhibited some changes during the extended shutdown of all three reactors from March 1996 through June 1998. Results of post-shutdown studies (1999-2001) showed that the thermally adapted community at the Fox Island-Exposed (FE) site was highly resilient because, as units returned to service, the characteristic components of this community quickly recovered.
Qualitative algal sampling had documented seasonal shifts in annual species at FE during 3-unit operation, and following Unit 3 restart in 1998. These shifts included abbreviated season for coldwater species (e.g., Monostroma grevillei, Spongomorpha arcta and Dumontia contorta) and extended season for warm-water species (e.g., Grinnellia americana, Dasya baillouviana, and Bryopsis hypnoides). Seasonality of these species at FE during the recent shutdown period was more typical of other sites. Initial establishment of perennial populations of Graciliaria tikvahiae and Sargassum filipendula at FE was also detected through qualitative studies during early 3 -unit years. In addition, a new species not previously collected at any site (the red alga Hypnea musciformis) was identified at FE in 2001.
Dominant species abundance and distribution patterns at FE, established during 3 -unit operation, were more resilient to the return of ambient conditions. Thermal impacts had been most notable in the low intertidal during 3 -unit operation, due to tidal currents in the discharge area. The low intertidal Chondrus population and associated seasonal epiphytes at FE were replaced by a population of the opportunistic green alga Codium fragile supporting a perennial Polysiphonia spp. population and periodically heavy sets of the blue mussel Mytilus edulis. Winter declines in Polysiphonia abundance typical of other sites were observed at FE during the shutdown period, but otherwise, little change in low intertidal community composition was observed, relative to recent years when MPS was operating. Many characteristics of the impacted low intertidal community at FE (sparse, fluctuating populations of barnacles and Chondrus, heavy mussel sets) were probably related to the dominance of the Codium population, which persisted through the shutdown period, rather than direct thermal effects. This would explain the quick reversal of minor shutdown-related
changes following unit restarts. However, the FE Codium population has declined after the shut-down period, and has been largely replaced by expanding populations of Gelidium pusillum and Corallina officinalis.
High Ascophyllum nodosum growth was observed at all sites in 2000-2001, compared to historical means, with highest growth at Fox Island-New (FN) compared to other sites. This relationship among stations has been attributed to elevated temperatures from the MPS discharge. However, a similar relationship was observed during two recent growing seasons (1996-97 and 1997-98) while all three units were shutdown, but was not evident during the first year following Unit 3 (1998-99). Natural influences of other factors such as ambient temperature conditions, nutrients and light may play a more important role in determining Ascophyllum growing conditions at FN than does thermal plume incursion from the MPS discharge.
In addition to the localized changes noted for rocky intertidal communities in close proximity to the MPS discharge, the rocky interidal monitoring program has documented patterns and modifications unrelated to MPS operation. These include introduction and spread of an exotic red alga, Antithamnion pectinatum, a longterm region-wide increase in abundance of the common brown rockweed, Fucus vesiculosus, and evidence of the regional nature of barnacle recruitment success from year to year.

## Fish Ecology Studies

The objective of the fish ecology monitoring program at MPS is to determine whether operation of the electrical generating units has adversely affected the occurrence, distribution, and abundance of local fishes. Potential MPS impacts include entrainment of fish eggs and larvae through the condenser coolingwater system; impingement of juvenile and adult fish on intake screens, which has been mitigated by the installation of fish return sluiceways; and changes in distribution or abundance attributable to the thermal discharge. Trawl, scinc, and ichthyoplankton monitoring programs were established in 1976 to provide the basis for identifying taxa potentially affected, as well as information on long-term abundance trends used to measure changes in the local populations.
This report summarizes data collected in the monitoring programs from June 2000 through May 2001 (report year 2000-01). During the 2000-01 report period, MPS Units 2 and 3 were both operating most of the time and Unit 1 was shut down.
The potential effects of MPS were assessed by conducting detailed analyses on seven taxa most
susceptible to MPS operational impact from entrainment or thermal effects." Analyses of these species generally focused on comparing temporal trends over the past 25 years. No significant longterm trends were detected for juvenile and adult silversides (Menidia menidia and M. beryllina) collected by seine in Jordan Cove (JC), all life stages of grubby."(Myoxocephalus aenaeus), cunner (Tautogolabrus adspersus) 'eggs and larvae,' and tautog (Tautoga onitis) and sand lance (Ammodytes americanus) larvae. Atlantic menhaden (Brevoortia tyrañus) larvae showed a significant increasing trend in abundance during the past 25 years. A significant negative trend was observed for silversides at the Intake (IN) trawl station. Since the mid-1980s, cunner have become less abundant at IN, exhibiting a significant negative trend, which may be related to the removal of the Unit 3 rock cofferdam, a preferred habitat for this species. Despite the negative trend in abundance for cunner at trawl station IN, their abundance at JC this year was at an historic high. While tautog eggs continued to exhibit a declining trend, juvenile and adult tautog abundance was at a 25 -year high at station IN. The large numbers of tautog and cunner eggs entrained at MNPS did not appear to affect the spawning stocks of these two fishes because the proportion of juvenile recruits relative to adults has increased.

Densities of both bay anchovy (Anchoa mitchilli) eggs and larvac exhibited significant negative trends. This year the $\Delta$-mean density for anchovy eggs was the lowest recorded. However, the $\Delta$-mean density of larvae was within the range of abundance indices found during the past 5 years. Information from Rhode Island waters and Chesapeake Bay indicated that bay anchovy is experiencing a regional decline in abundance along the Atlantic coast. None of the long-term abundance trends determined for various life stages of selected species could be directly related to the operation of MPS.

## Winter Flounder Studies

The local Niantic River population of winter flounder (Pseudopleuronectes americanus) is potentially affected by the operation of MPS, particularly by entrainment of larvae through the cooling-water systems of the operating units. As a result, extensive studies of the life history and population dynamics of this important sport and commercial species have been undertaken since 1976. Each year, surveys of adult spawners are carried out in the Niantic River during late winter and early spring. Larval sampling is conducted at the plant discharges, in Niantic Bay,
and at three stations in the Niantic River. Age-0 juveniles are collected at two sites in the river. Winter flounder are also commonly taken in the yearround trawl monitoring program (TMP).

During the 2001 adult winter flounder spawning season, $\Delta$-mean trawl catch-per-unit-effort (CPUE) of fish larger than 15 cm in the Niantic River was 1.4 , the lowest value found in 26 years, although not significantly different from a CPUE of 1.6 found in 1996.. Abundance peaked in the early 1980s and decreased thereafter, most likely from stock and recruitment effects (i.e., decreased per-capita recruitment ät high stock sizes), a generally warming winter trend, and increased rates of exploitation. Niantic River CPUE was significantly correlated with several other winter flounder abundance indices from throughout Southern New England and current abundance remains low.
The Jolly stochastic model was applied to mark and recapture data to estimate the absolute abundance of the Niantic River adult spawning population. The abundance estimate for 2000 was 9.3 thousand for all winter flounder larger than 20 cm . This was considerably less than estimated population sizes during 1984-91, which ranged between about 33 and 80 thousand, but was similar to estimates made since 1995, which were between 5.5 and 8.5 thousand. Annual female spawner abundance estimates since 1976 ranged from a low of 2 thousand in 2001 to 75 thousand in 1982, with corresponding total egg production estimates from about 1.7 to 43.7 billion for the same years. Since 1995, nearly all annual sex ratios of spawners have been highly skewed ( $>2$ ) in favor of females, but most ratios were 1.7 or less from 1977 through 1994.

A post-spawning abundance survey was made in the river last year from mid-April through mid-May. Although less reliable because of non-random sampling, the CPUE of 11.8 for this survey was considerably higher than found during the regular spawning survey and indicated that more adult winter flounder were available for capture during late spring. It was not known whether or not most of these fish had been present during the spawning season in areas not sampled, or if they were from other stocks and entered the river after, spawning elsewhere. These fish were marked with a unique brand and recapture rates in 2001 indicated that most of these fish were probably from the Niantic River stock and also that abundance wás likely greater than indicated by the spawning survey CPUE.
Abundance of newly-hatched (Stage 1) winter flounder larvae was considerably below average in the Niantic River this year. However, despite recent low abundance of spawning females and egg
production, newly hatched larvae were still more numerous than expected in 5 of the last 6 years. This was attributed to higher egg survival, which may have been as much as four times greater than in earlier years and represented a potentially important compensatory mechanism. Given their relative scarcity in the Niantic River during 2001, larval winter flounder abundance in Niantic Bay was much higher than expected. Abundances of Stage 2 and 3 larvae were among the lowest observed since 1984 in the river, but densities were at relatively high levels in the bay. This suggested potentially high rates of flushing and higher survival in the bay and LIS.
Stage 4 larvae were also at record or near-record highs in the bay and at the larval station in the lower river, with the latter an indication of importation back into the river.
Rates of larval growth and development were positively correlated with water temperature, but other factors such as density and prey abundance probably also affected growth. Growth rate of larvae during 2001 in the river was about average, but in the bay growth rate was among the lowest of the 19 -year period. Larval mortality rate in 2001 was also the lowest observed in 18 years, helping to account for the high abundance of older larvae. Densitydependence was examined by comparing mortality and egg production estimates (a measure of yolk-sac larval stage abundance) at various monthly and seasonal water temperatures. Results of this analysis suggested that larval mortality decreased with decreasing egg production (i.e., density-dependency) and increasing April water temperatures (i.e., faster development).
Following larval metamorphosis and settlement, demersal age-0 young were sampled by beam trawl at two sites in the Niantic River. Relatively high abundance of newly settled young was found in early summer, reflecting the large numbers of Stage 4 larvae. However, this year-class experienced record low growth and near-record low survival. Thus, the 2001 year-class, which had the potential to be among one of the strongest produced since 1983, exhibited good, although not exceptional abundance at the end of summer.
The 2000-01 $\Delta$-mean CPUE calculated for young winter flounder taken during late fall and early winter at TMP stations did not reflect higher abundance of the 2000 year-class as indicated by beam trawl sampling during the summer of 2000 . Nevertheless, these two age-0 abundance indices were significantly correlated and the relative magnitude of these indices suggested that the 1988, 1992, and 1994 year-classes of winter flounder were relatively strong, whereas the 1993 and 1997 year-classes were weak. Regardless
of these values, however, fewer age-1 juveniles have been taken during each year of the Niantic River adult spawning population surveys since the early 1980s, with a CPUE for fish taken in the lower river navigational channel particularly low. This abundance decrease was much less when a CPUE was calculated for fish taken only in the upper portion of the river. The relative distribution of age-1 fish in Niantic River and Bay also may have changed over the years, which was attributed to environmental factors.
Young-of-the-year abundance indices were either not significantly correlated or were negatively correlated with the abundance of female adult spawners 3 to 5 years later. Conversely, positive correlations were found between age-1 abundance indices and these older fish. However, the forms of the significant relationships were unclear and none of the early life stages were considered to be a highly reliable predictor of potential future year-class strength. Unknown processes that occur after winter flounder become age-1 may be operating to produce fewer adult recruits from more abundant year-classes of juveniles. A stock and recruitment relationship was determined for the Niantic River winter flounder population using adult CPUE data. A February water temperature parameter significantly improved the model fit, but a depensatory parameter was nonsignificant.
The number of larvae entrained through the condenser cooling-water system at MPS is a measure of potential impact to winter flounder. Annual estimates of entrainment were related to both larval densities in Niantic Bay and plant operation. The 2000 and 2001 entrainment estimates of 331 and 376 million, respectively, were only exceeded by an estimate of 492 million in 1992. These totals reflected the high larval densities found in Niantic Bay during the past 2 years. As in previous years, Stage 3 larvae predominated (59\%) in entrainment collections. All entrained larvae are presumed to die for impact assessment purposes, although a limited entrainment survival study showed that some (5\%) older and larger larvae survived entrainment. The entrainment rate (annual abundance index divided by total flow) has varied since 1976 without trend, indicating that larval production and availability in Niantic Bay remained stable despite increased water use during the 1986-95 period of three-unit operation. Correlations between entrainment estimates and abundance indices of age-0 juveniles were positive, implying no entrainment effect and that the more larvae that were available for entrainment, the more that metamorphosed and settled in Niantic River and Bay. This was also illustrated by a comparison of annual entrainment and juvenile year-class strength,
which indicated that entrainment estimates served as an index of emerging year-class strength rather than being the most important factor in setting abundance.
The impact of larval entrainment on the Niantic River stock depends upon the fraction of the annual winter flounder production entrained each year (determined as equivalent eggs). Empirical massbalance calculations showed that a large number of entrained larvae likely come from a number of sources in LIS, including stocks associated with the Connecticut and Thames Rivers, as well as from the Niantic River. Estimates of the fraction of entrained larvae in 2001 from the Niantic River were similar for both the mass-balance model (21\%) and an independently conducted analysis of genetic stock identification (22\%). These remarkably similar estimates suggested that the mass-balance model provided accurate estimates of Niantic River larval stock entrainment. The Niantic River production loss estimate for 2001 was a relatively high $36.1 \%$ (18year geometric mean $=11.5 \%$ ), again likely reflecting a high flushing rate of larvae from the river. However, based on the apparent increase in egg survival noted in recent years, a factor which was not incorporated into the model, production loss estimates since 1995 may have been conservatively high.
A stochastic computer simulation model (SPDM) was used for long-term assessments of MPS impact. Annual female spawner biomass (lbs) was determined in simulations over a period extending from 1960, a decade before the operation of Unit 1, to 2060, 15 years after the projected shutdown of Unit 3 in 2045, which assumes that a 20 -year operating license renewal will be attained by MPS. Conditional larval entrainment mortality rates (termed ENT) from the mass-balance model were applied in the model. The long-term geometric mean value was used in SPDM projections going forward, and to account for model uncertainties, both high and low rates calculated by scaling the mean value by a factor of 1.5 and $1 / 1.5$, respectively, were also applied. Values of fishing mortality ( F ) provided by the Connecticut Department of Environmental Protection were used, with a recent mean rate of 0.74 applied going forward. Also, since the last application of the SPDM in an MPS Annual Report, the instantaneous natural mortality rate of age-1 winter flounder was reestimated and the model re-calibrated to match current Niantic River female spawner biomass.
In the SPDM simulations, five stochastic time-series were generated, including a theoretical unfished stock, the size of which depended only upon the dynamics of winter flounder reproduction, natural rate of increase, and environmental variability; a baseline stock affected by rates of fishing in addition
to the above; and three impacted stocks, which further added the effects of MPS (low, mean, and ENT rates going forward) to those of fishing and natural variation. Used initially in all simulations, an initial unfished stock size of 119,972 lbs represented the maximum spawning potential for the Niantic River stock and the critical stock size ( $25 \%$ of the calculated Maximum Spawning Potential) was determined as 29,993 lbs.

For the baseline projection, the exploited biomass was quickly reduced to about 30 thousand lbs by 1980 and declined further under high $F$ to a low of 5,630 lbs in 1994 , only $19 \%$ of the critical stock size. With some reduction in F beginning in 1999, the stock recovered somewhat, but remained between 12 and 13 thousand lbs through 2060. Under the mean ENT rate, the impacted stock also attained its lowest biomass of 4.3 thousand lbs in the mid-1990s. Absolute differences between the baseline and impacted stocks increased to about 6 thousand lbs in 2018. However, with the modeled retirements of Units 2 and 3, impacted stock sizes began to approach those of the baseline by 2060 . The simulations under the high and low ENT rates generally paralleled those of the mean rate, only differing in magnitude. Of note, the F rate used in the SPDM is critical and actual spawner abundance could depart greatly in the simulations if this or other simulated conditions were not matched in reality. Fishing reduces stock biomass at a much greater rate than larval entrainment by removing individuals, particularly larger ones, from a year-class every year as long as any fish remain. In contrast, entrainment removes fish only once in the lifetime of a generation and then early in life before many compensatory processes have occurred.
To provide some perspective on Niantic River winter flounder stock size estimates, annual exploitation rates were determined and used with annual recreational and commercial landings data of winter flounder for LIS. The calculated exploitable biomass of Niantic River winter flounder probably only represented about $2 \%$ of the entire winter flounder resource in LIS during the past two decades.

To date, efforts of regulatory agencies to control fishing mortality have not resulted in large increases in abundance for winter flounder stocks across the region or in the Niantic River. Even so, the remaining small adult spawning stock in the river continues to produce relatively large numbers of larvae and young fish, which are a likely result of population compensatory mechanisms. The effective retirement of Unit 1 in late 1995 was followed by an immediate reduction of about one-quarter of the MPS coolingwater flow, which has permanently lessened plant
impact, but has not resulted in stronger year-classes and subsequent enhanced recruitment to the spawning stock. Despite relatively good abundance of postentrainment immature winter flounder, significant recruitment to the adult spawning population has not occurred in recent years, due to as yet unknown factors removing these fish from the population. Environmental effects, including changes to the Niantic River and interactions with other species, especially during early life history, also are important factors likely affecting the winter flounder recruitment process.

## Introduction

## Reporting Requirements

This report summarizes results of ongoing environmental monitoring programs conducted in relation to the operation of Millstone Power Station (MPS). MPS can affect local marine biota in several ways: large organisms may be impinged on the traveling screens that protect the condenser cooling and service water pumps; smaller ones may be entrained through the condenser cooling-water system, which subjects them to various mechanical, thermal, and chemical effects; and marine communities in the discharge area may be subjected to thermal, chemical, and mechanical effects' resulting from the outflow of the cooling water. In addition, occasional maintenance dredging is done in the vicinity of the intake structures. The basis for the studies is the National Pollutant Discharge Elimination System (NPDES) permit (CT0003263), last issued by the Connecticut Department of Environmental Protection on December 14, 1992 to Northeast Nuclear Energy Company. The permit was transferred to Dominion Nuclear Connecticut, lnc. (DNC), effective on March 31, 2001. The regulations in the permit allow the MPS cooling water to be discharged into Long Island Sound (LIS) in accordance with Section 22a-430 of Chapter 446k of the Connecticut General Statutes and Section 301 of the Federal Clean Water Act, as amended. Paragraph 5 of the NPDES permit states that:

Thi permittee shall conduct or continue to conduct biological studies of the supplying and receiving waters, entrainment studies, and intake impingement monitoring. The studies shall include studies of intertidal and subtidal benithic communities, finfish communities and entrained plankton and shall include detailed studies of lobster populations and winter flounder populations.
In addition, paragraph 7 of the permit requires that:
On or before April 30, 1993 and annually thereafter, submit for review and approval of the Commissioner a detailed report of the ongoing biological studies required by paragraph 5 and as approved under. paragraph 6.
Furthermore, a decision and order of the Connecticut Siting Council (CSC) requires that the Council be informed of results of MPS environmental impact monitoring studies and any modifications made to these studies (paragraph 6 of the proceeding entitled "Docket No. 4, Certificate of Environmental

Compatibility and Public Need for an Electric Generating Facility Identified as 'Millstone Nuclear Power Station, Unit 3,' located in the Town of Waterford, Connecticut" and dated March 22, 1976). This report satisfies the requirements of the NPDES permit and of the CSC by updating and summarizing various studies conducted at MPS that were presented most recently in DNC (2001a).

## Study Area

MPS is situated on Millstone Point, about 8 km westsouthwest of New London on the Connecticut shore of LIS (Fig. 1). The property, covering an area of about 200 ha, is bounded to the west by Niantic Bay, to the east by Jordan Cove, and to the south by Twotree Island Channel. The MPS monitoring programs sample a study area of approximately $50 \mathrm{~km}^{2}$ that extends from the northern portions of the Niantic River and Jordan Cove to Giants Neck, 2 km south of Twotree Island, and 2 km east of White Point. Work takes place from the shoreline into areas as deep as 20 m southwest of $T$ wotree Island.
Strong tidal currents predominate in the vicinity of Millstone Point and influence the physical characteristics of the area. Average tidal flow through Twotree Island Channel is approximately $3,400 \mathrm{~m}^{3} \cdot \mathrm{sec}^{-1}$ and at maximum is about $8,500 \mathrm{~m}^{3} \cdot \mathrm{sec}^{-1}$ (NUSCO 1983). Current velocities are about 1 to 1.8 knots in the channel, slightly less ( 1 to 1.5 knots) near the plant and in Niantic Bay, and relatively weak in Jordan Cove and in the upper Niantic River. The currents are driven by semi-diurnal tides that have a mean and maximum range of 0.8 and 1.0 m , respectively. Thermal- and salinity-induced stratification may occur in regions unaffected by strong tidal currents. The greatest temperature variation has been observed in nearshore areas where water temperature can vary from -3 to $25^{\circ} \mathrm{C}$; salinity varies much less and typically ranges between 26 and $30 \%$. The bottom is generally composed of fine to medium sand throughout the area, but also includes some rock outcrops and muddy sand, especially near the shoreline. Strong winds, particularly from the southwest, can at times result in locally heavy seas (to 1.5 m or greater) near Millstone Point. Additional information on local hydrography and meteorology can be found in NUSCO (1983).


Fig. I. The study area for biological monitoring conducted to assess the effects of MPS operation.

## Millstone Power Station

The MPS complex consists of three nuclear power units; a detailed description of the station was given in NUSCO (1983). Unit 1 , a $660-\mathrm{MWe}$ boiling water reactor, began commercial operation on November 29, 1970 and on July 17, 1998 Northeast Utilities announced its retirement. Unit 2 is an $870-\mathrm{MWe}$ pressurized water reactor that began commercial operation in December 1975 and Unit 3 (1,150-MWe pressurized water reactor) commenced commercial operation on April 23, 1986. All units use oncethrough cooling water systems with rated flows of $28.0,36.1$, and $59.4 \mathrm{~m}^{3} \cdot \mathrm{sec}^{-1}$ for Units 1 through 3, respectively, including condenser and service water flow. Cooling water is drawn by pumps from depths of about $4.6-7.6 \mathrm{~m}$ below mean sea level within separate shoreline intakes located on Niantic Bay (Fig. 2 ). The intake structures, typical of many coastal power plants, have coarse bar racks ( 6.4 cm on center, $5.1-\mathrm{cm}$ gap) preceding vertical traveling screens to protect the plants from debris. Units 1 and 2 have always had $9.5-\mathrm{mm}$ mesh screens. Unit 3 originally had $4.8-\mathrm{mm}$ mesh screens, a combination of 9.5 - and $4.8-\mathrm{mm}$ mesh screens from early 1990 through summer 1992, and only $9.5-\mathrm{mm}$ mesh screens as of August 15, 1992. Fish return systems (sluiceways)
were installed at Unit I in December 1983, at Unit 2 in May 2000, and at Unit 3 during its construction to return aquatic organisms washed off the traveling screens back to LIS. Installation and operation of sluiceways have minimized the impact of impingement at MPS (NUSCO 1986, 1988a, 1994, DNC 2001b). A chronology of significant events associated with MPS construction and operation, including installation of devices designed to mitigate environmental effects and unit operational shutdowns exceeding 2 weeks, are found in Table 1. For reasons beyond the scope of this report, extended shutdowns occurred between late 1995 and early 1999, with Unit 3 returning to service in July 1998 and Unit 2 in May 1999 (Table 1). Some service water and condenser cooling water pumps were operated at each of these units during the period of extended shutdown, but only limited amounts of service water were withdrawn at Unit 1 after late 1995. Cooling-water use varied throughout the life of the station, depending upon plant operation, and total monthly flow (in millions of $\mathrm{m}^{3}$ ) from January 1976 through December 2001 is shown in Figure 3. During 2001, Unit 2 utilized $96.0 \%$ of its nominal maximum flow and Unit 3 totaled $88.6 \%$, which reflected the high capacity and sustained operation of both units this year, with the exception of the late winter refueling outage at Unit 3.

TABLE 1. Chronology of major construction and operation events at MPS through 2001.

| Date | Activity | Reference ${ }^{\text {a }}$ |
| :---: | :---: | :---: |
| December 1965 | Construction initiated for Unit I | NUSCO (1973) |
| November 1969 | Construction initiated for Unil 2 began | NUSCO (1973) |
| October 26, 1970 | Unit 1 initial criticality; produced first thermal eflluent | DNGL |
| November 29, 1970 | Unit 1 initial phase to grid | DNGL |
| December 28, 1970 | Unit I began commercial operation | DNGL |
| January 15, 1971 to February 22, 1971 | Unit 1 shutdown | DNGL |
| August-December 1972 | Surface boom at Unit 1 | NUSCO (1978) |
| November $1972{ }^{\text {i }}$ | Fish barrier installed at quarry cut | EL |
| September 3, 1972 to March 20, 1973 | Unit I shutdown | DNGL |
| November 1972 | Unit 2 coffer dam removed | NUSCO (1973) |
| April 18 to July 28, 1973 | Unit I shutdown. . $\therefore$ | DNGL |
| August-December 1973 | Surface boom at Unit 1 | NUSCO (1978) |
| July-December 1974 | Surface boom at Unit 1 | NUSCO (1978) |
| September 1 to November 5, 1974 | Unit 1 shutdown | DNGL |
| July-October 1975 | Surface boom at Unit 1 | NUSCO (1978) |
| July 1975 | Bottom boom installed at Unit 1 - | NUSCO (1978) |
| August 5, 1975 | Unit 3 coffer dam construction began | EL |
| September 10 to October 20, 1975 | Unit 1 shutdown | DNGL |
| October 7, 1975 | Unit 2 produced first efluent | EDAN |
| November 7, 1975 | Unit 2 initial criticality; produced first thermal eflluent | EDAN |
| November 13, 1975 | Unit 2 initial phase to grid $\quad:^{\text {a }}$ | DNGL |
| December 1975 | Unit 2 began commercial operation . | EL |
| March 19, 1976 | Unit 3 coffer dam construction finished | EL |
| June-October 1976 | Surface boom at Unit 2 | NUSCO (1978) |
| October 1 to December 2, 1976 | Unit 1 shutdown | DNGL |
| December 20, 1976 to January 20, 1977 | Unit 2 shutdown | DNGL |
| May 6 to June 25, 1977 | Unit 2 shutdown | DNGL |
| June-October 1977 | Surface boom at Unit 2 | NUSCO (1978) |
| November 20, 1977 to May I, 1978 | Unit 2 shutdown | DNGL |
| March 10 to April 15, 1978 | Unit I shutdown | DNGL |
| March 10 to May 21, 1979 | Unit 2 shutdown | DNGL |
| April 28 to June 27, 1979 | Unit 1 shutdown | DNGL |
| August 10 to 25, 1979 | Unit 2 shutdown | DNGL |
| November 1 to December 5, 1979 | Unit 2 shutdown | DNGL |
| May 7 to June 19, 1980 | Unit 2 shutdown | DNGL |
| June It to June 18, 1980 | Unit I shutdown | DNGL |
| August 15 to October 19, 1980 | Unit 2 shutdown | DNGL |
| October 3, 1980 to June 16, 1981 | Unit I shutdown | DNGL |
| January 2 to 19, 1981 | Unit 2 shutdown | DNGL |
| December 5, 1981 to March 15, 1982 | Unit 2 shutdown | DNGL |
| March 1981 | Bottom boom removed at Unit I ${ }^{\text {. }}$ | EL |
| September 10 to November 18, 1982 | Unit 1 shutdown | DNGL |
| March 2 to 18, 1983 | Unit 2 shutdown | DNGL |
| April-September 1983 | Unit 3 coffer dam removed, intake maintenance dredging | EL |
| May 28, 1983 to January 12, 1984 | Unit 2 shutdown | DNGL |
| December 1983 | Fish return system installed at the Unit 1 intake | EL |
| August 1983 | Second quarry cut opened | EL |
| April 13 to June 29, 1984 | Unit 1 shutdown | DNGL |
| February 15 to July 4, 1985 | Unit 2 shutdown | DNGL |
| June 1985 | Intake maintenance dredging | EL |
| September 28 to November 7, 1985 | Unit 2 shutdown | DNGL |
| October 25 to December 22, 1985 | Unit 1 shutdown | DNGL |
| November 1985 | Unit 3 produced first effluent | EDAN |
| February 12, 1986 | Unit 3 produced first thermal eflluent | EDAN |
| April 23, 1986 | Unit 3 began commercial operation | DNGL |
| July 25 to August 17, 1986 | Unit 3 shutdown | DNGL |
| September 20 to December 18, 1986 | Unit 2 shutdown | DNGL |
| December I to 15, 1986 | Unit 1 shutdown | DNGL |

TABLE I. (cont.).

| January 30 to February 16, 1987 | Unit 2 shutdown | DNGL |
| :---: | :---: | :---: |
| March 14 to April 10, 1987 | Unit 3 shutdown | DNGL |
| June 5 to August 17, 1987 | Unit 1 shutdown | DNGL |
| November 1, 1987 to February 17, 1988 | Unit 3 shutdown | DNGL |
| December 31, 1987 to February 20, 1988 | Unit 2 shutdown | DNGL |
| April 14 to May 1, 1988 | Unit 3 shutdown | DNGL |
| May 7-22, 1988 | Unit 2 shutdown | DNGL |
| October 23 to November 8, 1988 | Unit 3 shutdown | DNGL |
| February 4 to April 29, 1989 | Unit 2 shutdown | DNGL |
| April 8 to June 4, 1989 | Unit I shutdown | DNGL |
| May 12 to June 12, 1989 | Unit 3 shutdown | DNGL |
| October 21 to November 24, 1989 | Unit 2 shutdown | DNGL |
| March 30 to April 20, 1990 | Unit 3 shutdown; installation of some $9.5-\mathrm{mm}$ intake screen panels | DNGL; EL |
| May 8 to June 15, 1990 | Unit 2 shutdown | DNGL |
| September 14 to November 9, 1990 | Unit 2 shutdown | DNGL |
| February 2 to April 17, 1991 | Unit 3 shutdown; installation of new fish buckets and sprayers | DNGL; EL |
| April 7 to September 2, 1991 | Unit 1 shutdown | DNGL |
| April 23 to May 11, 1991 | Unit 2 shuldown | DNGL |
| May 26 to July 7, 1991 | Unit 2 shutdown | DNGL |
| July 25, 1991 to February 6, 1992 | Unit 3 shutdown; installation of new fish buckets and sprayers | DNGL; EL |
| August 7 to September 11, 1991 | Unit 2 shutdown | DNGL |
| October 1, 1991 to March 3, 1992 | Unit 1 shutdown | MOSR |
| November 6 to December 27, 1991 | Unit 2 shutdown | MOSR |
| January 28 to February 14, 1992 | Unit 2 shutdown | MOSR |
| March 22 to April 6, 1992 | Unit I shutdown | MOSR |
| May 16 to June 4, 1992 | Unit 3 shutdown; installation of new fish buckets and sprayers | MOSR; EL |
| May 29, 1992 to January 13, 1993 | Unit 2 shutdown | MOSR |
| July 4 to August 15, 1992 | Unit I shutdown | MOSR |
| August 15, 1992 | Completed installation of new fish buckets and sprayers at Unit 3 | EL |
| September 30 to November 4, 1992 | Unit 3 shutdown | MOSR |
| July 31 to November 10, 1993 | Unit 3 shutdown | MOSR |
| September 15 to October 10, 1993 | Unit 2 shutdown | MOSR |
| January 15 to May 23, 1994 | Unit I shutdown | MOSR |
| April 22 to June 18, 1994 | Unit 2 shutdown | MOSR |
| July 27 to September 3, 1994 | Unit 2 shutdown | MOSR |
| September 8-22, 1994 | Unit 3 shutdown | MIOSR |
| October 1, 1994 to August 4, 1995 | Unit 2 shutdown | MOSR |
| April 14 to June 7, 1995 | Unit 3 shutdown | MIOSR |
| November 30 to December 15, 1995 | Unit 3 shutdown | MOSR |
| November 4, 1995 | Unit I shutdown; retirement of unit announced July 17, 1998 | MOSR; EL |
| February 20, 1996 to May 11,1999 | Unit 2 shutdown | MOSR |
| March 30, 1996 to July 5, 1998 | Unit 3 shutdown | MOSR |
| December 11-29, 1998 | Unit 3 shutdown | MOSR |
| June 4, 1999 | Original Quarry Cut Fish Barriers replaced | EL |
| Aprit 30 to Sune 29, 1999 | Unit 3 shutdown | MOSR |
| April 22 to June 1, 2000 | Unit 2 shutdown | MOSR |
| May 23, 2000 | Commenced operation of Unit 2 fish retum sluiceway | EL |
| February 3 to March 30, 2001 | Unit 3 shutdown | MOSR |
| February 16 to April 2, 2002 | Unit 2 shutdown | MOSR |

[^1]

Fig. 2. The MPS site, showing the intake and discharge of each unit, the quarry, and the two quarry discharge cuts.


Fig. 3. Total monthly cooling-water use in millions of $\mathrm{m}^{3}$ at MPS from January 1976 through December 2001.

MPS cooling water can be heated in Units 1 (formerly), 2, and 3 from ambient temperature to a maximum of 13.9 (formerly), 12.7 , and $9.5^{\circ} \mathrm{C}$, respectively. Each unit has a separate discharge structure that releases the heated effluent into an abandoned granite quarry (ca. 3.5 ha surface area, maximum depth of approximately 30 m ). The cuts are equipped with fish barriers made up of $19-\mathrm{mm}$ metal grates, which serve to keep larger fish out of the quarry. The thermal discharge, which is typically about $10.5^{\circ} \mathrm{C}$ warmer than ambient under present station operation, exits the quarry through two channels (cuts), whereupon it mixes with LIS water (Fig. 2). All previous hydrothermal surveys conducted at MPS were described in detail in NUSCO (1988b). The thermal plume is warmest in the immediate vicinity of the cuts and the surface-oriented plume from three-unit operation was shown to cool to less than $2.2^{\circ} \mathrm{C}$ above ambient within about $1,100 \mathrm{~m}$ of the quarry (Fig. 4). Beyond this distance the plume is highly dynamic and varies mostly with tidal currents. Having a smaller volume, the plume from two-unit operation covers less area than did the former full three-unit operation shown in Figure 4 and calculations of its present extent are given in Adams (2001)

## Monitoring Programs

This report contains a separate section for each major monitoring program, some of which have been conducted without interruption or significant change since 1976. These long-term studies have provided the representative data and scientific bases necessary to assess potential biological impacts as a result of MPS construction and operation. The significance of changes found for various communities and populations beyond those that were expected to occur naturally were evaluated using best available methodologies. Programs discussed below include Winter Flounder Studies, Fish Ecology Studies, Lobster Studies, Eelgrass, Rocky Intertidal Studies, and Benthic Infauna. Reporting periods for each section vary and were predicated on biological considerations and processing time necessary for samples, as well as on regulatory requirements. In cases where the seasonal abundance of organisms differed from arbitrary annual reporting periods, the periods chosen were adjusted to best define the season of interest for a particular species or community.

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Low Slack Tide

Fig. 4. Locations of selected three-unit thermal plume isotherms $\left(1.5^{\circ} \mathrm{F}, 4^{\circ} \mathrm{F}, 6^{\circ} \mathrm{F}\right.$. and $\left.8^{\circ} \mathrm{F}\right)$ under various tidal conditions.


High Slack Tide

Fig. 4. (continued).

## Lobster Studies

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## Lobster Studies

## Introduction

The American lobster, Homarus americanus, is distributed in the Northwest Atlantic from Canada to Cape Hatteras and supports one of the most valuable commercial fisheries in the northeast United States ( $\$ 301.0$ million in 2000; NMFS landings data). For many years, fishery managers have been concerned that the intense exploitation of lobster throughout its range may impact long-term egg production and recruitment to coastal populations (Anthony and Caddy 1980; NFSC 1996; ASMFC 1997). As a result, a number of regulations have been implemented by state and federal agencies to minimize the chance of a population collapse due to recruitment failure by improving larval production and subsequent recruitment. Management measures include prohibition on possession of eggbearing (or v-notched) females, prohibition on spearing lobster, minimum legal size of 82.6 mm ( $31 / 4$, in carapace length), catch limit on lobster taken in trawl gear, and prohibition on possession of lobster parts. Additional measures implemented to control the coastwide escalation of fishing effort by regulating lobster traps used in the fishery include requirements for escape vents and biodegradable fasteners, maximum trap size restrictions, and limits on the number of traps used by fishermen.
The most recent stock assessment for American lobster indicated that instantaneous fishing mortality rates ( F ) in Long Island Sound (LIS) have exceeded 1.0 for the past decade and the number of traps used in the fishery has increased eightfold since the early 1980s (ASMFC 2000). Coinciding with the increase in fishing pressure, commercial landings of lobsters in Connecticut waters of LIS increased from 0.7 million pounds in 1979 to a record 5.6 million pounds in 1998 (Blake and Smith 1984; Smith et al. 1989; Crecco and Gottschall 1999). However, landings declined 70\% between 1998 and 2000 to only 1.7 million pounds. Landings declined even further by 2001 when 1.5 million pounds were harvested (CTDEP fishery statistics). The decline has been confirmed with fishery independent data collected in the CTDEP trawl survey which indicates that lobster abundance has fallen almost $60 \%$ since 1998 (Simpson et al. 2001). The recent decline in lobster abundance has been attributed to a significant lobster die-off in western LIS (LoBue 2001). Beginning in the fall of 1998 and throughout the summer of 1999, an unexplained highly invasive
disease emerged in lobsters harvested from western LIS (CTDEP 2000; LoBue and Gottschall 2000).
Lobstermen reported that more than half of the lobsters caught in traps were dead or died before arriving at port. Catch data compiled by CTDEP confirmed these observations, noting that commercial landings for all ports declined dramatically in the fall of 1999 (64$99 \%$ ) from the 1995-1998 average. Catch per trap haul in the commercial fishery during the fall 1999 declined $83 \%$ in western LIS and $39 \%$ in central LIS when compared to average CPUE during the period 19951998 (CTDEP 2000). At the same time, a widespread outbreak of shell disease syndrome was reported for lobsters in eastern LIS (DNC 2001a; Landers et al. 2001a) and other coastal areas of southern New England (Castro and Angell 2000). In January 2000, the U.S. Secretary of Commerce declared a failure of the commercial lobster fishery in LIS. As a result, the Long Island Sound Lobster Initiative (LISLI), was formed in 2001 by Sea Grant programs in New York and Connecticut along with CTDEP and NMFS. The LISLI is responsible for administering $\$ 6.6$ million in Federal funding for research investigations on causes and economic impacts of the die-off and shell disease in LIS lobsters. Funded researchers are currently investigating many possible factors on an ecosystemwide basis. These environmental, physiological, and biological stresses include: water quality conditions including elevated temperature and changes in salinity, environmental conditions such as storm events, pollution, lobster crowding, disease-causing organisms, and pesticides. At present, the status of the lobster population in LIS is uncertain and the long-term sustainability of the resource may be threatened.
Because of the regional economic importance of American lobster, the local population in the vicinity of Millstone Power Station (MPS) has been studied extensively since 1978 to determine if power plant operation has caused changes beyond those expected from natural variability and the high level of fishing. The potential impacts of power plant operations on the local population of lobsters include entrainment of larvae through the cooling water systems, impingement of juveniles and adults on the intake traveling screens, and effects of the heated effluent on juvenile and adult lobsters in the discharge area. The objectives of the lobster monitoring program at Millstone are to: 1) evaluate year-to-year, seasonal, and among-station changes in catch-per-unit-effort; 2) monitor lobster
population demographics including size structure, growth rates, sex ratios, female size at maturity, characteristics of egg-bearing females; and 3) monitor movements of lobsters in the vicinity of MPS. Beginning in 1984, studies were conducted during the hatching season to estimate the number of lobster larvae entrained through the cooling water systems. Impacts associated with recent plant operations on the local lobster population were assessed by comparing results of the 2001 study with those from 1978 to 2000. Emphasis has been placed on assessing long-term trends in the abundance and population characteristics of lobsters collected in the Millstone Point area. These results were compared, when appropriate, to studies conducted by other researchers in LIS and throughout the range of the American lobster.

## Materials and Methods

Lobsters were collected from May through October using pot gear. Four pot-trawls, each consisting of five double-entry wire pots ( $76 \times 51 \times 30 \mathrm{~cm} ; 2.5 \mathrm{~cm}^{2}$ mesh) equally spaced along a $50-75 \mathrm{~m}$ line buoyed at both ends, were fished in three rocky areas in the vicinity of MPS (NUSCO 1982, 1987a). Pots set in Jordan Cove (average depth 6 m ) were 500 m east of the Millstone discharge (Fig 1). The Intake station (average depth 5 m ) was 600 m west of the discharge near the power plant intake structures, and the Twotree station (average depth 12 m ) was located south of Millstone Point, about 2000 m offshore near Twotree Island. Pots were hauled on Monday, Wednesday, and Friday of each week, weather permitting, and twice per week on holiday weeks. On each sampling trip, surface and bottom water temperatures and salinities were recorded at each station. Lobsters were banded to restrain chelipeds, transported to the laboratory, and kept in a tank supplied with a continuous flow of seawater. Pots were rebaited and reset in the same area. On Fridays, all lobsters caught that week were examined and the following data recorded: sex, presence of eggs (berried), carapace length (CL) to the nearest 0.1 mm , crusher claw position, missing claws, and molt stage (Aiken 1973). The size at which females become sexually mature was determined (since 1981) by measuring the maximum outside width of the second abdominal segment of all females to the nearest 0.1 mm . Female size at sexual maturity was estimated by calculating the ratio of abdominal width to carapace length and plotting that ratio against carapace length (Skud and Perkins 1969; Krouse 1973). Lobsters were
tagged with a serially numbered international orange sphyrion tag (Scarratt and Elson 1965; Scarratt 1970), and released at the site of capture. Recaptured tagged lobsters and severely injured or newly molted (soft) lobsters were released after examination.

Entrainment studies were conducted since 1984 by sampling lobster larvae during the period of their occurrence (May through July) at one of the discharges of Units 1,2 , or 3 . Samples were collected with a 1.0 x 6.0 m conical plankton net of 1.0 mm mesh. The volume of cooling water sampled was estimated from the average readings of four General Occanic flowmeters located in the mouth of the net. The net was typically fished for 45-60 minutes and filtered about $4000 \mathrm{~m}^{3}$ of cooling water. From 1984 to 1993, eight lobster larvae entrainment samples (four day and four night) were collected each week. Based on a statistical review of the sampling frequency in 1994, the number of samples collected was reduced to six per week (three day and three night). Samples were transported to the laboratory where they were either immediately processed or placed in a 1.0 mm mesh sieve and held in tanks supplied with flowing seawater for later processing. All samples were processed within 24 hours by sorting in a white enamel pan. Lobster larvae were examined for movement and classified as live or dead. All larvae were also classified by developmental stage (I-IV) according to the criteria established by Herrick (1909). The abundance of larvae in entrainment samples was standardized as the number of larvae per unit-volume. The mean density of larvae from May through July was estimated by the mean of the assumed "delta" distribution, referred to as $\Delta$-mean (Pennington 1983; NUSCO 1988a). To estimate the total number of larvae entrained, the $\Delta$-mean density was multiplied by the total volume of water pumped through the plants during the sampling period.

Impingement studies were conducted at Unit I and 2 intakes from 1975 through 1987. Results summarized in NUSCO (1987a) included estimates of the total number of lobsters impinged, as well as their mean size, sex ratio, proportion of culls (missing claws), and survival of impinged lobsters. Possible impacts associated with impingement of lobsters at Units 1,2, and 3 were mitigated by installing fish return systems in the intakes, which return impinged organisms to LIS (NUSCO 1986a; 1987b; DN்C 2001b). Subsequently, NUSCO and the CT DEP agreed to discontinue impingement monitoring (NUSCO 1988b).


Fig. 1. Location of the Millstone Power Station (MPS), and the three lobster sampling stations, JC=Jordan Cove, $\mathrm{IN}=\mathrm{Intake}, \mathrm{TT}=\mathrm{T}$ wotree.

Catch-per-unit-effort (CPUE; i.e., the number of lobsters caught per pothaul) was used to describe the annual abundance of lobsters in the MPS area. Since the CPUE data are ratios, which are not additive and have an asymmetric distribution about the arithmetic mean, the geometric mean was computed to analyze trends in CPUE. The geometric mean is better suited ; for constructing asymmetric confidence intervals for log-normal data (McConnaughey and Conquest 1993). Annual geometric mean CPUEs were calculated for all lobster sizes. .
The annual abundance (CPUE) of legal-size lobsters in the MPS area was estimated by using the $\Delta$-mean. The $\Delta$-mean was a more appropriate statistic for describing the CPUE of legal-size lobster, since a large number of zero observations were present in the data. (i.e., many pots contain no legal-size lobsters). Both geometric means of all lobsters and $\Delta$-means of legalsize lobsters were used to compare annual variation in CPUE. In the following Results and Discussion.
section, the geometric mean abundance of all lobsters is called "mean total CPUE" while the $\Delta$-mean abundance of legal-size lobsters is referred to as "mean legal CPUE". The distribution-free, Mann-Kendall test (Hollander and Wolfe 1973) was used to determine the presence of significant trends in the time series of annual CPUE data, and of several other selected population characteristics (size composition, sex ratios, growth rates). Slopes of significant trends were calculated using Sen's estimator of the slope (Sen 1968).

The influence of water temperature on lobster molting was examined by estimating the time when lobster molts peaked each year and correlating the annual molt peaks with bottom water temperature. Molting peaks were derived using the inflection point of the Gompertz growth function (Draper and Smith 1981; Gendron 1989) fitted to data reflecting the cumulative percentage of molting lobsters at weekly


Fig. 2. Mean surface and bottom water temperatures from May through October at each of the lobster sampling stations from 1979 hrough 2001 and linear trends $( \pm 95 \%$ C.I.) of surface and bottom water temperature.
intervals during the molting season. The function used to describe annual molting peaks had the form:

$$
C_{t}=100 e^{-e^{-k(t-r)}}
$$

where $C_{t}=$ cumulative percentage of molting lobsters,
$\mathrm{t}=\quad$ time in weeks,
p $=$ inflection point scaled in weeks from May 1st,
$k=\quad$ shape parameter.
The derivative of the Gompertz function with respect to time yields a "molt frequency" function which describes the distribution of annual molts. Annual "molt frequencies" were then correlated with mean
bottom water temperature during May to investigate a possible relationship between water temperature and molting.

## Results and Discussion

## Water Temperature

Mean surface and bottom water temperatures measured from May through October at each station from 1979 to 2001 are presented in Fig. 2. Over the past 23 years, average surface water temperatures from May through October ranged between 16.0 and $19.6^{\circ} \mathrm{C}$ at Jordan Cove, $15.7^{\circ} \mathrm{C}$ and $18.2^{\circ} \mathrm{C}$ at Intake, and

TABLE 1. Catch statistics of lobsters caught in wire potsa from 1978 through 2001.

| Year | Total number caught | Number pots hauled | Geometric mean total CPUE | 95\% C.I. ${ }^{\text {. }}$ | $\begin{gathered} \Delta \text {-mean } \\ \text { legal } \\ \text { CPUE }^{\text {b }} \end{gathered}$ | 95\% C.I. | $\begin{gathered} \text { Commercial } \\ \text { catch } \\ \left(\mathrm{lbs} \times 10^{3}\right) \\ \hline \end{gathered}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1978 | 1824 | 1026 | 1.600 | 1.454-1.761 | 0.173 | 0.144-0.202 | - - |
| 1979 | 3259 | 2051 | 1.404 | 1.302-1.513 | 0.128 | 0.107-0.148 | 86.0 |
| 1980 | 2856 | 2116 | 1.103 | 0.997-1.221 | 0.109 | 0.092-0.126 | .81.1 |
| 1981 | 2236 | 2187 | 0.904 | 0.839-0.974 | 0.098 | 0.083-0.113 | 131.1 |
| 1982 | 9109 | 4340 | 2.006 | 1.925-2.089 | 0.165 | 0.144-0.186 | 136.5 |
| 1983 | 6376 | 4285 | 1.331 | 1.250-1.418 | 0.148 | 0.128-0.168 | 194.7 |
| 1984 | 7587 | 4550 | 1.607 | 1.540-1.677 | 0.159 | 0.140-0.179 | :251.7 |
| 1985 | 7014 | 4467 | 1.352 | 1.252-1.460 | 0.105 | 0.090-0.120 | 179.4 |
| 1986 | 7211 | 4243 | 1.585 | 1.501-1.673 | 0.086 | 0.074-0.097 | 157.3 |
| 1987 | 7280 | 4233 | 1.633 | 1.562-1.707 | 0.079 | 0.070-0.089 | 194.1 |
| 1988 | 8871 | 4367 | 1.929 | 1.846-2.015 | 0.079 | $0.068 \cdot 0.091$ | 217.8 |
| 1989 | 7950 | 4314 | 1.729 | 1.645-1.817 | 0.065 | 0.056-0.075 | 213.7 |
| 1990 | 7106 | 4350 | 1.531 | 1.455-1.610 | 0.076 | 0.065-0.087. | 327.0 |
| 1991 | 7597 | 4404 | 1.542 | 1.437-1.654 | 0.091 | 0.078-0.104 | 349.5 |
| 1992 | 11438 | 4427 | 2.457 | 2.352-2.565 | 0.085 | 0.075-0.095 | 446.9 |
| 1993 | 10195 | 4194 | 2.301 | 2.198-2.408 | 0.080 | 0.069-0.091 | 358.7 |
| 199.4 | 9849 | 4256 | 2.199 | 2.104-2.298 | 0.071 | 0.061-0.093 | 345.6 |
| 1995 | 6435 | 4317 | 1.261 | 1.152-1.380 | 0.080 | 0.068-0.093 | 308.8 |
| 1996 | 7531 | 4249 ' | 1.587 | 1.466-1.718 | 0.067 | 0.058-0.077 | 267.5 |
| 1997 | 9026 | 4293 | . 1.960 | 1.862-2.064 | 0.099 | 0.086-0.112 | 411.9 |
| 1998 | 10991 | 4380 | 2.315 | 2.189-2.448 | 0.125 | 0.108-0.141 | 495.4 |
| 1999 | 11216 | 4057 | 2.560 | 2.419-2.709 | 0.151 | 0.131-0.170 | 406.3 |
| 2000 | 8707 | 4192 | 1.849 | 1.721-1.986 | 0.092 | 0.078-0.106 | 247.3 |
| 2001 | 7268 | 4104 | 1.587 | 1.476-1.705 | 0.094 | 0.080-0.107 | 202.5 |

${ }^{2} 10$ wire pots fished at each station from August through October 1978, and from May through October 1979-81; 20 wire pots fished at each station from May through October 1982-2000.
${ }^{\text {b }}$ The minimum legal-size from 1978 to 1988 was $81.0 \mathrm{~mm}\left(3^{3} / 16 \mathrm{in}\right.$ ), minimum legal-size was increased in 1989 to $81.8 \mathrm{~mm}\left(3^{7 / 32}\right.$ in), and in 1990 to $82.6 \mathrm{~mm}(31 / 4 \mathrm{in})$.
${ }^{\text {c }}$ Annual commercial catch data from Area 1 eastern LIS (provided by CT DEP).
$15.2^{\circ} \mathrm{C}$ to $17.4^{\circ} \mathrm{C}$ at Twotree. Over the same period, the interannual variability in bottom water temperature was generally smaller than surface water temperature, with bottom water temperatures ranging between 15.0 and $17.5^{\circ} \mathrm{C}$ at Jordan Cove, 15.2 and $17.6^{\circ} \mathrm{C}$ at Intake, and $14.8^{\circ} \mathrm{C}$ and $17.2^{\circ} \mathrm{C}$ at Twotree. Surface water temperatures were notably higher than bottom water temperatures at the Jordan Cove station, which is located closest to the MPS discharge. Surface temperatures at this site were also slightly warmer., during the period 1986-1995 when 3 -units were operating. In comparison, surface and bottom water temperatures were most similar at the Twotree station, which is not influenced by the MPS thermal plume.
Long-term trends in the May to October water temperatures were examined" using SAS 'GLM Procedures (SAS 1990). Significant increasing trends in surface water temperatures were found at Intake ( $\mathrm{p}=0.012$, slope $=0.053^{\circ} \mathrm{C} /$ year) and Twötree ( $\mathrm{p}=0.011$, slope $=0.043^{\circ} \mathrm{C} /$ year). No significant trend was detected in surface water temperature at Jordan Cove, owing to the high degree of variability associated with changing levels of MPS operation (i.e., number of Units online from May to October). Bottom water
temperatures at each of the lobster sampling stations are not influenced by the MPS thermal plume and significant ( $p<0.05$ ) increasing trends were identified in bottom water temperature at each station over the past 23 years' (Jordan Cove slope $=0.052^{\circ} \mathrm{C}$ /year; Intake slope $=0.053^{\circ} \mathrm{C} /$ year; Twotree slope $=0.052^{\circ} \mathrm{C} /$ year).
Similar trends in long-term water temperature records. have been reported by other researchers in New England (Keller et al. 1999; Koeller 1999; Keller and Klein-MacPhee 2000; Wolff and Incze in press). Indications of warming trends are most evident during the winter months and appear to be partially correlated to local air temperature anomalies (Foertch 2000; Manning et al. 2001).

## Abundance and Catch-per-Unit-Effort

The total number of lobsters caught each year from May through October since 1978 ranged from 1,824 to 11,438 (Table 1). The total catch was lower from 1978 through 1981 (range $=1,824-3,259$ ) because only 10 wire traps were used at each station. : Beginning in 1982, 20 wire traps were used at each station and total catch fluctuated between 6,376 and 9,109 lobsters from

1982 to 1991. Catches peaked in $1992(11,438)$ and then declined by nearly $50 \%$ in $1995(6,435)$. For the next four years, catches increased each year and peaked again in $1999(11,216)$. Total catch declined about $22 \%$ in 2000 and again in 2001 ( $16 \%$ ) when 7,268 lobsters were caught in 4,104 pot-hauls. However, catch-per-pot averaged 1.587 lobsters during 2001, which was within the range of previous annual mean CPUEs ( 0.904 and 2.560 lobster/pot). Furthermore, the time series of total CPUE data has exhibited a significant increasing trend since 1978 (slope $=0.032$, $\mathrm{p}=0.014$ ). At present, the catastrophic die-off and decline in lobster abundance observed in western LIS has not occurred near MPS.
The catch of legal-size lobsters is highly dependent on the number of recruit-size lobsters one or two molts below minimum legal size. Since the majority of lobsters caught in our unvented traps are recruit-sized, annual fluctuations in total lobster CPUE correspond closely to changes observed in legal-size lobster abundance. For example, high total CPUEs in 1998 and 1999 corresponded to high legal CPUEs in those years. The 22\% decline in total CPUE from 1999 to 2000 corresponded to a $39 \%$ decline in legal-size lobster CPUE in 2000 ( 0.151 in 1999 to 0.092 in 2000; Table 1). However, despite the $16 \%$ decline in total CPUE observed from 2000 to 2001, legal CPUE this year was 0.094 , which was slightly higher than in 2000 and no significant long-term trend was found in the time series of annual $\Delta$-mean legal-size lobster CPUEs.
The trends observed in our indices of lobster abundance (total and legal CPUEs) closely followed trends in abundance of lobsters taken in the eastern LIS commercial fishery. Commercial catches fluctuated between $81,000 \mathrm{lbs}$ and $349,500 \mathrm{lbs}$ from 1979 to 1991 (Table 1). Catches peaked in 1992 (446,900 lbs) and then declined by $40 \%$ in 1996 ( $267,500 \mathrm{lbs}$ ).
Subsequently, catches increased and reached record levels of $495,400 \mathrm{lbs}$ in 1998. Following the record landings in 1998, commercial catches declined $18 \%$ in 1999 (406,300 lbs) and again in 2000 (39\%) when only $247,300 \mathrm{lbs}$ were taken. Commercial catch in eastern LIS declined further (18\%) in 2001 to about 202,500 lbs.
Annual CPUE values for all sizes of lobster and legal-size lobster are presented for each station in Figure 3. Total CPUE during 2001 was highest at Twotree (1.905), lowest at Intake (1.440), and intermediate at Jordan Cove (1.459). Over the past 24 years, total CPUE has significantly increased at Jordan Cove (slope $=0.033, p=0.026$ ) and Twotree (slope $=0.043, p=0.005$ ), but not at Intake. The

.Fig. 3. Mean total CPUE (geometric mean $\pm 95 \%$ C.I.) and mean legal CPUE ( 1 -mean $\pm 95 \%$ C.I.) of lobsters caught at each station from 1978 to 2001 (minimum legal size increased from 81.0 mm to 81.8 mm in 1989 and to 82.6 mm in 1990).

Twotree station also yielded the highest catch of legalsize lobsters during 2001 ( 0.110 ), followed by Intake (0.105) and Jordan Cove ( 0.067 ; Fig. 3). In general, when compared with the nearshore stations, Twotree has consistently yielded higher catches of legal-size lobsters. However, long term trend analysis indicates that catches of legal-size lobster have significantly declined at this station since 1978 (slope $=-0.004$, $\mathrm{p}=0.011$ ); whereas no trends in legal-size lobster CPUE were identified at Jordan Cove or Intake.

TABLE 2. Total number of lobsters and incidental catch of other species caught in traps.

|  |  |  |
| :--- | ---: | ---: |
|  | Range (1984-2000) | 2001 |
|  |  |  |
|  |  |  |
| Lobster | $6435-11438$ | 7268 |
| Rock, Jonah crab | $79-2033^{*}$ | $256^{*}$ |
| Spider crab | $1344-31480^{*}$ | $9316^{*}$ |
| Hermit crab | $192-721^{*}$ | 562 |
| Blue crab | $21-169$ | 50 |
| Winter flounder | $5-45^{*}$ | 9 |
| Summer flounder | $4-60^{*}$ | 40 |
| Skates | $14-94$ | 38 |
| Oyster toadfish | $5-76$ | 16 |
| Scup | $21-1039^{*}$ | 473 |
| Cunner | $41-239^{*}$ | 79 |
| Tautog | $39-250^{*}$ | 137 |
| Sea raven | $0-20$ | 1 |
| Whelks | $21-178^{*}$ | $29^{*}$ |

* Covariance analysis identified these catches as significant $(\mathrm{p}<0.05)$ factors affecting lobster CPUE.

In addition to lobsters, pot gear often catches other marine organisms including crustaceans, mollusks, and fish. The presence of these organisms in lobster pots can negatively influence the catch of lobster (NUSCO 1987a, 2001a). For example, the catchability of lobsters in pots fished near MPS was strongly depressed when spider crabs were abundant in pot gear (Table 2). On several occasions hundreds of spider crabs were caught in a single pot and completely blocked the entry funnels of the trap. Spider crabs can also indirectly influence lobster catchability by consuming bait. During 2001, spider crabs continued to be a dominant component of the total by-catch and significantly influenced lobster catches at the Intake station. Rock and Jonah crabs have been dominant components of the by-catch at Twotree and significantly influenced lobster catches at that site during 2001. At Jordan Cove, whelks were found to strongly influence lobster CPUE during 2001; this large gastropod also influenced lobster catches in previous study years. Results of our studies are similar to findings of other researchers in southern New England, which indicated that the incidental catches of rock crabs and spider crabs significantly affect lobster catch (Richards et al. 1983; Cobb et al. 1986; Richards and Cobb 1987). Behavioral interactions between crabs and European lobsters ( $H$. gammarus) were demonstrated to have substantial effects on the catch of both species in pot gear and operated in a densitydependent manner (Addison and Bannister 1998).
Another important factor which contributes to the inherent variability in the catch of lobsters in traps is the amount of time between trap-hauls or soaktime


Fig. 4. Relationship between catch-per-pot and soaktime during high (1999) and low (2001) years of lobster abundance.
(Saila et al. in press). Quite often commercial lobstermen will adjust the amount of time between pothauls to maximize their catch. In the LIS fishery, short soaktimes of 2-3 days are typical during summer "runs" as water temperature and lobster catchability increases; soaktimes are increased during winter months when water temperatures and lobster activity decreases. Long soaktimes of 7 days are more successful offshore than inshore, because of lower lobster density and a greater distance of attraction resulting from a larger foraging area (Miller 1995). In our study, traps are regularly checked on Mondays, Wednesdays, and Fridays from May through October, except during holiday weeks and during heavy sea conditions. As a result, there is little contrast in the number of days between pothauls. However, during 2001 and in some of the previous study years when lobster abundance was below average, the amount of time between pothauls significantly influenced the catch of lobsters (NUSCO 1986; 1990). The relationship between lobster catch-per-pot and soaktime during a high (1999) and low year (2001) of lobster abundance is illustrated in Figure 4. During 1999, when lobster CPUE was the highest observed in our studies and the effect of soaktime on lobster catch was not significant, catch-per-pot after a 2 day set averaged 2.7 lobsters. When lobster abundance was below average during 2001 and the effect of soaktime on lobster catch was significant, catch-per-pot averaged 1.7 lobsters after a 2 day set. In both years, catch-perpot peaked after a 3 day set averaging 3.0 and 1.9 lobsters in 1999 and 2001, respectively. After a 4 day set catch-per-pot decreased, but was equivalent (1.8 lobsters) in both high and low years of lobster abundance. In 1999, catch-per-pot increased again to 2.6 lobsters after a 5 day set, whereas catch decreased to 1.5 lobsters after 5 days in 2001. The causes of the
increased catch after a 5 day set in 1999 are unclear. Field observations indicate that it is unlikely related to the lobster bait, which is entirely consumed after 3 days. One possibility could be that the larger number of lobsters caught in years when abundance is high may attract other lobsters long after the bait has been consumed. Other factors may be related to changes in the commercial fishery such as trap selectivity (e.g., changes in escape vent dimensions) or trap saturation. Other researchers using underwater videos of baited lobster traps indicated that less than $6 \%$ of the lobsters that enter a trap are caught and more than $70 \%$ escape back through the entrance (Jury et al. 2001). These researchers also suggested that the efficiency of the traps and behavioral responses of lobsters to traps may be density dependent.

## Population Characteristics <br> Size Composition

Annual size frequency distributions of lobsters caught in $8-\mathrm{mm}$ size classes are illustrated in Figure 5. Although the catch of lobsters in all size classes declined during 2001, the pattern of size composition was similar to previous study years and continued to be dominated by pre-recruit ( $67-74 \mathrm{~mm}$ ) and recruit (7582 mm ) sized lobsters. A notable feature of the annual size-frequency distributions demonstrating the effects of the intense fishing pressure around MPS is the low number of lobsters in the $33-90 \mathrm{~mm}$ and $>90 \mathrm{~mm}$ size classes. Regardless of the number of lobsters in the recruit size class ( $75-82 \mathrm{~mm} \mathrm{CL}$ ), few lobsters in the $83-90 \mathrm{~mm}$ size class (full recruit) were caught in our traps because they were harvested by commercial lobstermen soon after molting to legal size. Lobsters larger than 90 mm CL were rarely present in the annual catch, which indicates that the LIS fishery is almost entirely ( $>90 \%$ ) dependent on new recruits.
Mean carapace length (CL) of all lobsters during 2001 was 71.0 mm , falling within the range of annual average sizes reported previously ( $69.5-72.2 \mathrm{~mm}$; Table 3). Percentage of legal-size lobsters ( $\geq 82.6 \mathrm{~mm}$ ) during 2001 was $4.9 \%$, within the range reported in previous studies when the legal size was $\geq 82.6 \mathrm{~mm}$ (1990-99; 3.1-5.7\%) and $\geq 81.0 \mathrm{~mm}$ (1978-88; 3.2$9.1 \%$ ). Since 1978, the percentage of legal-size lobsters in our catch has significantly declined (slope $=-$ $0.129, p=0.01$ ). The decline can be attributed to changes in fishery regulations (increases in minimum legal size in 1989 and 1990) and to the eightfold increase in the number of traps fished by Connecticut


Fig. 5. Number of lobsters caught in each $8-\mathrm{mm}$ carapace length category from (a) $<58 \mathrm{~mm}, 59-66 \mathrm{~mm}$, and $67-74 \mathrm{~mm}$ and (b) 75 $82 \mathrm{~mm}, 83-90 \mathrm{~mm}$, and $>90 \mathrm{~mm}$ ( 30 wood and 30 wire traps fished from 1978 to 1981; 60 wire traps fished from 1982 to 2001).
and New York fishermen in LIS since the early 1980s (Crecco and Gottschall 1999; ASMFC 2000).
The mean CL during 2001 was largest at Twotree ( 72.5 mm ), smallest at Jordan Cove ( 69.8 mm ), and intermediate at Intake ( 70.4 mm ; Table 4). Mean sizes at each of the three stations during 2001 were within the range of values reported from 1978 to 2000 (Twotree, $70.0-73.7 \mathrm{~mm}$; Jordan Cove, 68.8-71.8 mm; Intake $68.9-72.4 \mathrm{~mm}$ ). In contrast to previous study years when the percentage of legal-size lobsters was highest at Twotree, the percentage during 2001 was highest at Intake (5.6\%) followed by Twotree (4.9\%) and Jordan Cove (3.9\%). Since 1978 the percentage of legal size lobsters has significantly declined at Twotree (slope $=-0.237 ; p=0.008$ ); but not at Jordan Cove or Intake.

## Sex Ratios

The sex ratio of lobsters collected during 2001 was 0.57 females per male, compared to a range of 0.48 to 0.97 previously observed (Table 5). Since 1978, annual female to male sex ratios have exhibited a significant declining trend (slope $=-0.015, \mathrm{p}<0.0001$ ).

TABLE 3. Summary of lobster carapace length statistics for wire pot catches from May through October, 1978-2001.

|  | $\mathrm{N}^{2}$ | Carapace length (mm) Range | $\begin{gathered} \text { Mean } \\ \pm 95 \% \mathrm{Cl} \end{gathered}$ | Percentage of legal sizes ${ }^{\text {b }}$ |
| :---: | :---: | :---: | :---: | :---: |
| 1978 | 1508 | 53-111 | $71.4 \pm 0.33$ | 7.5 |
| 1979 | 2846 | 44-100 | $71.2 \pm 0.26$ | 7.6 |
| 1980 | 2531 | 40-96 | $70.7 \pm 0.27$ | 6.4 |
| 1981 | 1983 | 43-96 | $71.0 \pm 0.33$ | 8.8 |
| 1982 | 7835 | 45-103 | $70.8 \pm 0.15$ | 6.7 |
| 1983 | 5432 | 40-121 | $71.7 \pm 0.19$ | 9.1 |
| 1984 | 6156 | 45-107 | $71.8 \pm 0.18$ | 8.7 |
| 1985 | 5723 | 38-101 | $71.3 \pm 0.17$ | 5.9 |
| 1986 | 5961 | 36-107 | $70.1 \pm 0.17$ | 4.4 |
| 1987 | 5924 | 36-99 | $70.2 \pm 0.17$ | 3.9 |
| 1988 | 7144 | 21.97 | $69.5 \pm 0.16$ | 3.2 |
| 1989 | 6713 | 34-107 | . $69.9 \pm 0.17$ | 3.5 |
| 1990 | 6040 | 36-102 | $70.2 \pm 0.20$ | 4.9 |
| 1991 | 6449 | 31-101 | $70.2 \pm 0.20$ | 5.0 |
| 1992 | 9594 | 20-103 | $70.1 \pm 0.15$ | 3.3 |
| 1993 | 8487 | 30-102 | $70.8 \pm 0.15$ | 3.3 |
| 1994 | 7841 | 34-100 | $70.3 \pm 0.17$ | 3.1 |
| 1995 | 5472 | 37-101 | $71.9 \pm 0.20$ | 5.7 |
| 1996 | 6634 | 16-96 | $70.0 \pm 0.19$ | 3.8 |
| 1997 | 7528 | 35-100 | $71.5 \pm 0.17$ | 4.8 |
| 1998 | . 8958 | 25-99 | $71.3 \pm 0.16$ | 4.9 |
| 1999 | 9155 | 32-110 | $72.2 \pm 0.15$ | 5.6 |
| 2000 | 7151 | 46-110 | $71.1 \pm 0.17$ | 4.7 |
| 2001 | 5888 | 32-99 | $71.0 \pm 0.19$ | 4.9 |

${ }^{2}$ Recaptures not included.
${ }^{\mathrm{b}}$ The minimum legal size from 1978 to 1988 was $81.0 \mathrm{~mm} \mathrm{( } 3^{3} / 16$ in), minimum legal size was increased in 1989 to $81.8 \mathrm{~mm}\left(3^{7} / 32\right.$ in), and in 1990 , to $82.6 \mathrm{~mm}\left(3^{1 / 4} \mathrm{in}\right)$.

TABLE 4. Summary of lobster carapace length statistics for wire pot catches at each station from May through October, during the period from 1978 through 2000 and during 2001 studies.

|  | Mean carapace length (mm) ${ }^{2}$ | Percentage of legals ${ }^{\text {b }}$ |
| :---: | :---: | :---: |
| IORDAN COVE |  |  |
| 1978-2000 range | 68.8-71.8 | 2.7-8.2 |
| 2001 mean | 69.8 | 3.9 |
| INTAKE |  |  |
| 1978-2000 range | 68.9-72.4 | 2.8-9.2 |
| 2001 mean . | 70.4 | 5.6 |
| TWOTREE |  |  |
| 1978-2000 range | 70.0-73.7 | 3.1-14.2 |
| 2001 mean | 72.5 | 4.9 |
|  |  | - 1 |

${ }^{2}$ Recaptures not included.
${ }^{6}$ The minimum legal size from 1978 to 1988 was $81.0 \mathrm{~mm}\left({ }^{3} / 16\right.$ in), minimum legal size was increased in 1989 to 81.8 mm ( $3^{1 / 32}$ in), and in 1990 , to $82.6 \mathrm{~mm}(31 / 4 \mathrm{in})$.

During 2001,' female to male sex ratio was highest at Twotree (1.01), lowest at Jordan Cove (0.36) and intermediate at Intake (0.41). Values at each of the three stations were within the range of values reported

TABLE 5. Female to male sex ratios ${ }^{2}$ of lobsters caught in wire pots from May through October, 1978-2001.

|  | Jordan <br> Cove |  | Intake | Twotree | All <br> Stations |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1978 | 0.79 | 0.97 | 1.02 | 0.92 |  |
| 1979 | 0.68 | 0.83 | 1.15 | 0.82 |  |
| 1980 | 0.66 | 0.90 | 1.15 | 0.88 |  |
| 1981 | 0.70 | 0.71 | 1.19 | 0.86 |  |
| 1982 | 0.62 | 0.66 | 1.09 | 0.79 |  |
| 1983 | 0.72 | 0.67 | 1.25 | . |  |
| 1984 | 0.60 | 0.71 | 1.22 | 0.87 |  |
| 1985 | 0.70 | 0.67 | 1.38 | 0.82 |  |
| 1986 | 0.65 | 0.73 | 1.26 | 0.87 |  |
| 1987 | 0.71 | 0.63 | 1.24 | 0.88 |  |
| 1988 | 0.68 | 0.72 | 1.15 | 0.85 |  |
| 1989 | 0.64 | 0.65 | 1.08 | 0.79 |  |
| 1990 | 0.60 | 0.65 | 0.90 | 0.71 |  |
| 1991 | 0.51 | 0.57 | 1.13 | 0.74 |  |
| 1992 | 0.43 | i | 0.47 | 1.45 | 0.73 |
| 1993 | 0.47 | 0.59 | 1.59 | 0.84 |  |
| 1994 | 0.54 | 0.67 | 1.24 | 0.79 |  |
| 1995 | 0.53 | 0.61 | 0.93 | 0.71 |  |
| 1996 | 0.32 | 0.37 | 0.79 | 0.48 |  |
| 1997 | 0.35 | 0.48 | 0.77 | 0.52 |  |
| 1998 | 0.48 | 0.59 | 0.92 | 0.63 |  |
| 1999 | 0.49 | 0.60 | 0.79 | 0.62 |  |
| 2000 | 0.37 | 0.44 | 0.86 | 0.55 |  |
| 2001 | 0.36 | 0.41 | 1.01 | 0.57 |  |

${ }^{2}$ Recaptures not included.
over the past 24 years. The occurrence of more females at Twotree than at other stations has been consistent since 1975 (Keser et al. 1983; DNC 2001a). Significant declines in female to male ratios were noted from 1978 to 2001 at Jordan Cove and Intake (slope at both sites $=0.016 ; p<0.001$ ), but not at Twotree. The cause for the decline in female to male sex ratios is unclear, although other researchers working in southern New England and New York have observed similar declines in localized nearshore areas (K. Castro URI and K. Graulich NYSDEC, pers. comm.). Beginning in 1998, we recorded the sex of lobsters caught in our trawl monitoring program at Jordan Cove, Intake and Niantic River (see Fish Ecology section). These data also indicated a predominance of males in nearshore waters around MPS with female to male sex ratios of 0.72 in 1998, 0.74 in 1999, 0.47 in 2000, and 0.54 in 2001. Other lobster populations typically show a $1: 1$ 'sex' ratio (Templeman 1936; Ennis 1971, 1974; Stewart 1972; Krouse 1973; Thomas 1973; Briggs and Mushacke 1980; Lawton and Lavalli 1995), although there are some " reports" of male-" and female-dominated populations and seasonal assemblages (Cooper et al. 1975; Briggs and Mushacke'1979; Howell and Watson 1991, Howell et al. 1999). Changes in male and female
growth and maturity could cause differential susceptibility of the sexes to the trap fishery and lead to shifts in the sex ratio. In populations that are heavily fished, the sex ratios of individuals just above the legal size are expected to be skewed toward females because berried females are protected from harvest and therefore have lower mortality rates than males (Cobb 1995). Smith (1977) reported female to male sex ratios in the LIS commercial fishery ranging between 1.06 and 1.81 and more recently, sex ratios of lobsters caught in eastern LIS commercial traps were unusually higher, ranging between 2.61 and 6.29 females per male (Blake 1988). During routine trawl surveys conducted from 1986 to 1998 throughout LIS by the CTDEP, female to male sex ratios of lobsters one molt below legal-size averaged 1.44 during the spring and 0.96 during the fall (ASMFC 2000). In contrast, female to male sex ratios of legal lobsters caught in the CTDEP trawl surveys were dominated by males during the fall ( 0.64 ) and to a lesser degree during spring (0.96).

## Reproduction

Aiken and Waddy (1980) described a number of approaches to determine the size at which females become sexually mature. A simple technique is to examine the size distribution of egg-bearing females in the population. One limitation of this approach is that once females extrude eggs, they are protected from the fishery and the proportion of berried females above the minimum legal size can be overestimated. More accurate methods require sacrificing animals to examine the size and color of ovaries. For the past 21 years, we have used a technique first described by Templeman ( 1935,1944 ) who found that abdominal width measurements of females markedly increase at the first onset of sexual maturation. Female size at sexual maturity was estimated by calculating the ratio of abdominal width to carapace length (AW/CL) and plotting that ratio against carapace length (Skud and Perkins 1969; Krouse 1973). Based on measurements obtained from 66,000 non-berried females, mean AW/CL ratios were calculated for each $1-\mathrm{mm}$ CL and plotted against the carapace length of lobsters collected from 1981 to 2001 (Fig. 6). During 2001, females began to mature between 50 and 55 mm CL , and all females larger than 90 mm CL were mature. When comparing females caught in 2001 with those caught from 1981 to 2000, many of the individual $1-\mathrm{mm}$ mean values for 2001 were above the long-term mean.


Fig. 6. Morphometric relationship between the mean abdominal width to carapace length ratio ( $y$ ) and the carapace length ( $x$ ) of nonberried female lobsters during the period from 1981 to $2000(-)$ and during $2001(+++)$. Regression equation for each period: 1981-2000: $=0.61-\left(0.44^{*} 10^{-2}\right) x+\left(0.96^{*} 10^{-4}\right) x^{2}-\left(0.42^{*} 10^{-6}\right) x^{3}, r^{2}=.95$ 2001: $y=0.75-\left(1.07^{*} 10^{-2}\right) x+\left(1.88^{*} 10^{-4}\right) x^{2}-\left(0.83^{*} 10^{-4}\right) x^{3}, r^{2}=.95$


Fig. 7. Annual mean abdominal width to carapace length ratio of non-berried females from 1981 to 2001.

Closer examination of annual mean AW/CL ratios, revealed a shift in the relationship between female abdominal width and carapace length over the past 21 years. Annual mean $A W / C L$ values fluctuated between 0.610 and 0.620 from 1981 to 1987 and averaged 0.614 (Fig. 7). Beginning in 1988, mean values increased each year from 0.606 to 0.637 in 1993. From 1994 to 2001 annual $A W / C L$, values stabilized at values ranging between 0.627 and 0.639 , with an average of 0.632 . The discontinuity in the AW/CL ratio and the changes observed in the proportion and size distribution of berried females (see below) suggests a recent decrease in the size at sexual maturity of fenmale lobsters in LIS (Landers et al. 2001b).
The recent change observed in the time-series of morphometrics of non-berried female lobsters was followed by an increase in the abundance of eggbearing females. From 1978 through 1990, between 3.1 and $7.8 \%$ of females were egg-bearing (Table 6 ).

TABLE 6. Percentage of berried females ${ }^{2}$ caught at each station from May through October 1978-2001.

|  | Jordan <br> Cove | Intake | Twotree | All <br> Stations |
| :---: | :---: | :---: | :---: | :---: |
| 1978 | 3.5 | 3.1 | 7.4 | 4.7 |
| 1979 | 2.9 | 3.4 | 7.9 | 4.4 |
| 1980 | 2.9 | 2.4 | 6.0 | 3.9 |
| 1981 | 2.1 | 3.4 | 8.0 | 5.1 |
| 1982 | 1.3 | 0.9 | 7.2 | 3.1 |
| 1983 | 3.1 | 3.2 | 9.0 | 5.6 |
| 1984 | 4.3 | 3.2 | 12.7 | 7.5 |
| 1985 | 4.1 | 4.8 | 9.6 | 7.2 |
| 1986 | 4.2 | 2.6 | 9.7 | 6.1 |
| 1987 | 3.8 | 2.8 | 10.8 | 7.0 |
| 1988 | 4.1 | 2.8 | 7.3 | 5.1 |
| 1989 | 4.6 | 4.6 | 9.7 | 6.7 |
| 1990 | 3.7 | 4.6 | 14.2 | 7.8 |
| 1991 | 3.9 | 1.5 | 16.5 | 9.1 |
| 1992 | 4.2 | 2.4 | 27.3 | 15.3 |
| 1993 | 3.9 | 3.7 | 27.4 | 16.0 |
| 1994 | 8.4 | 5.8 | 25.1 | 15.0 |
| 1995 | 7.5 | 7.2 | 17.4 | 12.0 |
| 1996 | 4.3 | 4.0 | 17.4 | 10.4 |
| 1997 | 5.8 | 3.9 | 25.8 | 13.3 |
| 1998 | 8.3 | 6.0 | 27.9 | 13.5 |
| 1999 | 7.9 | 7.6 | 18.0 | 11.7 |
| 2000 | 2.8 | 6.4 | 18.5 | 11.0 |
| 2001 | 5.2 | 6.4 | 25.2 | 14.9 |
| 2 |  |  |  |  |

${ }^{\text {a }}$ Recaptures included

The percentage of berried females increased sharply to $16.0 \%$ in 1993 and remained at above-average levels through 2001, when $14.9 \%$ of females were eggbearing. More pronounced increases in berried female abundance occurred at Twotree, where the percentage of egg-bearing females has been highest since 1978 (Keser et al. 1983; DNC 2001a). Between 6.0 and $14.2 \%$ of females caught annually at Twotree were egg-bearing from 1978 to 1990, compared to between 16.5 and $27.9 \%$ annually from 1991 to 2001 (Table 6). Annual percentages of berried females at the nearshore Jordan Cove and Intake sites were also generally higher over the past decade (2.8-8.4\% and 1.5-7.6\%, respectively), than during the earlier study years from 1978 to 1990 (1.3-4.6\% and 0.9-4.8\%,respectively).
Following the recent decrease in female size at onset of sexual maturity, annual average carapace length of egg-bearing females has become smaller and the proportion of sublegal-size egg-bearing lobster has increased. The mean CL of berried females was 75.5 mm during 2001 and $93 \%$ were sublegal-size (Fig. 8). Similar average sizes and percentages of sublegal-size were observed in the past 10 years. During the early study years' from 1978 to 1987, egg-bearing females averaged $79.2 \mathrm{~mm}(\mathrm{CL})$ and $70 \%$ were sublegal-size. In recent study years from 1992 to 2001 , berried


Fig. 8. Percentage sublegal and mean carapace length ( $\pm 95 \%$ C.I.) of berried females collected from 1978 to 2001.
female average size decreased to 75.9 mm (CL) and $93 \%$ were below the legal size.
The size at sexual maturity for female lobsters varies widely across the species range (Aiken and Waddy 1980; Waddy et al. 1995). Maturation occurs at smaller sizes in shallow warmwater locations of southern New England and the Gulf of St. Lawrence and at larger sizes in deep coldwåter locations offshore and in the Gulf of Maine (Krouse 1973; Briggs and Mushacke 1979; Van Engel 1980; Aiken and Waddy 1986; Fogarty and Idoine 1988; Estrella and MćKiernan 1989; Blake 1994). Results from our studies over the past two decades showed that the onset of female lobster sexual maturity occurs between 50 and 60 mm (CL). Based on abdominal growth, our long-term study suggests a recent reduction in the size at which females become sexually mature. Additional eviderice supporting this finding includes an increase in the proportion of egg-bearing females and a decrease in their mean carapace length. It is presently unclear whether reductions in female size at maturity resulted from changes in environmental conditions or from selection pressures related to intense fishing pressure (Landers et al. 2001b). Average bottom seawater temperatures from 1979 to 2001 demonstrate a significant increasing trend (see below Molting and Growth) and, as mentioned earlier, the number of traps fished by Connecticut and New York fishers has increased almost eightfold in the past two decades (AFMSC 2000). Water temperatures can directly influence lobster maturation (Aiken and Waddy 1980), whereas high exploitation rates may be exerting a strong selection for earlier maturing lobsters. While the exact causes for the reduction in female size at maturity are presently unclear, the benefits of earlier maturation may be significant to our local lobster
population dynamics. Small size at maturity and subsequent egg production from sublegal-size females may explain why LIS lobster populations are resilient to high levels of fishing pressure.

## Molting and Growth

Lobster growth is a function of molt frequency and size (or weight) increase per molt. Water temperature is the most important factor regulating molting events (Aiken 1980). In our study area, the majority of molting lobsters were caught from late spring to early summer, although a secondary molt was observed in autumn of earlier study years (1978-1982) when sampling was conducted through November (Keser et al. 1983). The timing and frequency of lobster molts were examined using weekly cumulative percent-molt data. An example of these data are presented in Figure 9 a for lobsters caught during 2001 and during the period from 1980 to 2000 . The percentage of molting lobsters in the total catch peaked at about $12 \%$ during weeks 6 and 7 of the 2001 study compared to a peak of 7\% during previous studies (Fig 9b). Annual molting peaks were significantly ( $\mathrm{p}<0.05$ ) correlated with mean May bottom water temperatures. Molting occurred earlier in the years when May water temperatures were warmer than average. Conversely, peaks occurred later when water temperatures were colder than average. During 2001, molting peaked on June 15, when bottom water temperatures during May averaged $10.1^{\circ} \mathrm{C}$ (Fig. 10). In general, when bottom water temperatures during May averaged more than $10.0^{\circ} \mathrm{C}$, molting peaked during the first or second week of June. Molting peaks were delayed up to one month (e.g., July 12, 1994), when May temperatures averaged below $9.0^{\circ} \mathrm{C}$. Aiken and Waddy (1980) found that when water temperatures rise above $10^{\circ} \mathrm{C}$ lobsters quickly enter the premolt stage and progress to ecdysis. Early laboratory studies conducted by Templeman (1936) indicated that molting was delayed one week for every $1^{\circ} \mathrm{C}$ reduction in water temperature. In colder waters of Nova Scotia, differences in the timing of annual molts were linked to average bottom water temperatures in August (Tremblay and Eagles 1997).
Molt increments were deternined from lobster tagging studies by comparing CL measurements at the time of tagging with those from recaptured lobsters that had molted only once. Simple linear regressions of pre-molt (tag-size) and post-molt (recapture-size) sizes best describe growth for the size range of lobsters caught during our studies. Linear regression plots of


Fig. 9. a) Cumulative percentage of molting lobsters caught each week and b) molt frequency curves based on the Gompertz function fitted to data in a) of lobsters caught from May 1 (week I) through October 31 (week 26) during the period from 1980 to 2000 and during 2001.


Fig. 10. Relationship between the date of peak molting of lobsters collected near MPS (parameter t from the Gompertz function) and annual mean bottom water temperature during May (1980-2001).
growth for males and females caught in 2001 and those caught from 1979 through 2000 are shown in Figure 11; parameter estimates are provided in Table 7. During 2001, average molt increments of both sexes


Fig. 11. Relationship between carapace lengths at tagging and carapace lengths at recapture during the period from 1979 to 2000 and during 2001 at sites near MPS of both male and female lobsters.

TABLE 7. Simple linear regression equations describing the relationship between carapace lengths of male and female lobsters at tagging ( $\mathbf{x}$ ) and at recapture (y) in the vicinity of MPS from 1979 to 2000 and during 2001.

|  | MALES |  |  |  | FEMALES |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | N . | Growth model | $\mathrm{R}^{2}$ | Growth increment mean (mm) | N | Growth model | $\mathrm{R}^{2}$ | Growth increment mean ( mm ) |
| 1979-2000 | 1997 | $\mathrm{y}=18.934+0.846$ (x) | 0.72 | 8.45 | 2036 | $y=15.256+0.899(x)$ | 0.74 | 8.55 |
| 2001 | 46 | $y=12.785+0.918(x)$ | 0.84 | 7.17 | 23 | $y=10.069+0.945(\mathrm{x})$ | 0.95 | 6.39 |

were smaller than the average increments reported in previous studies. Males grew an average of 7.17 mm per molt during 2001, compared to an average of 8.45 mm from 1979 to 2000 . Average female incremental growth of 6.39 mm was smaller than males during 2001 and more than 2 mm smaller than the average female increment of 8.55 mm from 1979 to 2000 . When the three lobster collection stations were compared, molt increments and corresponding percentages of growth per molt during 2001 were largest at Twotree ( 7.38 $\mathrm{mm}, 10.5 \%$ ), and smaller at the nearshore Jordan Cove and Intake stations ( $6.70 \mathrm{~mm}, 10.4 \%$ and 6.56 mm , $9.9 \%$, respectively; Table 8). Although percent growth per molt was lower during 2001, average values of between 12.7 and $13.1 \%$ reported during the period 1979-2000 at the three stations were similar to others reported for LIS lobsters, which ranged from $11.6 \%$ to $15.8 \%$ for males and between $12.0 \%$ and $15.4 \%$ for

TABLE 8. Annual mean growth per molt (incremental and percentage) at each station based on tag and recapture studies conducted from 1979 through 2000 and during 2001.

|  | 1979-2000 |  | 2001 |  |
| :---: | :---: | :---: | :---: | :---: |
|  | $\qquad$ | \% | $\qquad$ | \% |
| Jordan Cove | 8.28 | 12.7 | 6.70 | 10.4 |
| Intake | 8.42 | 12.7 | 6.56 | 9.9 |
| Twotree | 8.75 | 13.1 | 7.38 | 10.5 |



Fig. 12 a) Annual mean incremental growth per molt of mate and female lobsters based on tag and recapture studies conducted from 1979 to 2001 and b) means and linear trend ( $\pm 95 \%$ C.l.) of bottom water temperature measured at 3 stations from May through October 1979 to 2001.
females (Stewart 1972; Briggs and Mushacke 1984; Blake 1994).
Male and female molt increments were highly correlated ( $\mathrm{r}=0.84, \mathrm{p}<0.0001$ ) and the time series of annual growh increments for both sexes (Fig. 12a) have exhibited a significant declining trend since 1979 (male slope $=-0.058, \mathrm{p}=0.006$; female slope $=-0.101$, $\mathrm{p}=0.001$ ). Average bottom seawater temperature measured from May through October at the three stations ranged between 15.0 and $17.4^{\circ} \mathrm{C}$ (Fig. 12b) and has exhibited a significant increasing trend (slope $=0.058, \mathrm{p}=0.004$ ). Despite this apparent inverse relationship, no significant correlation was found between temperature and the declining growh rates of both male and female lobsters in our study area. Molt increments were shown to be smaller in blue crabs raised in warmer water (Lefler 1972) and comparisons of molt increments from lobster tagging studies conducted in cooler offshore waters (Cooper and Uzmann 1971; Fogarty and Idoine 1988) with those in warmer areas (DNC 2001a) suggest this is also true for
adult lobsters. The declining trend in female growh may be related to earlier maturation, increased reproductive activity, and changes observed in berried female size structure observed over the past two decades as discussed in the previous section and by Landers et al. (2001b). Females may sacrifice somatic growth for development of ovaries and incubation and development of eggs. Finally, high exploitation rates may be exerting a strong selection for slower-growing earlier-maturing lobsters. Similar population responses have been demonstrated for spiny lobster populations under high fishing pressure (Polovina 1989; Chubb 1994) and for crayfish in controlled fishing experiments on closed populations (Momot 1998).

## Culls

The percentage of culls (i.e., lobsters missing one or both claws) was $11.9 \%$ of the total catch during 2001 and within the range of values reported since the study begin in 1978 (9.4-15.5\%; Table 9). The percentage of culls at the nearshore Jordan Cove and Intake stations was $15 \%$ during 2001, more than double the percentage found at Twotree ( $6.6 \%$ ). The pattern of higher claw-loss at the male-dominated nearshore sites has been consistent since the study began (Keser et al. 1983) and may be related to the fact that males are more aggressive, more active and more dominant than

TABLE 9. Percentage of culls (lobsters missing one or both claws) caught in wire pots at sites near MPS 1978-2001.

| caught in wire pots at sites near MPS $1978-2001$ |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Jordan <br> Cove | Intake | Twotrec | All <br> Stations |
| 1978 | 21.5 | 14.7 | 9.8 | 15.5 |
| 1979 | 17.3 | 17.8 | 88 | 15.5 |
| 1980 | 13.5 | 16.4 | 10.4 | 13.4 |
| 1981 | 13.4 | 16.7 | 7.1 | 12.1 |
| 1982 | 13.9 | 14.1 | 70 | 11.3 |
| 1983 | 14.6 | 15.3 | 8.2 | 12.4 |
| 1984 | 11.5 | 15.1 | 66 | 10.8 |
| 1985 | 15.1 | 13.9 | 7.2 | 11.1 |
| 1986 | 10.9 | 14.7 | 6.8 | 10.6 |
| 1987 | 11.9 | 14.7 | 6.2 | 10.3 |
| 1988 | 13.7 | 13.6 | 6.5 | 11.1 |
| 1989 | 14.4 | 14.3 | 8.8 | 12.2 |
| 1990 | 12.3 | 16.2 | 8.1 | 11.9 |
| 1991 | 14.5 | 14.0 | 8.2 | 11.8 |
| 1992 | 11.4 | 12.9 | 6.9 | 10.0 |
| 1993 | 11.2 | 12.6 | 7.7 | 10.1 |
| 1994 | 11.8 | 11.5 | 6.9 | 9.8 |
| 1995 | 14.2 | 15.5 | 7.8 | 11.9 |
| 1996 | 15.0 | 14.9 | 7.3 | 12.2 |
| 1997 | 15.1 | 12.8 | 6.2 | 11.2 |
| 1998 | 13.8 | 12.5 | 6.8 | 11.3 |
| 1999 | 12.9 | 12.0 | 7.4 | 10.7 |
| 2000 | 12.3 | 10.8 | 5.9 | 9.4 |
| 2001 | 15.0 | 15.0 | 6.6 | 11.9 |

TABLE 10. Lobster tag and recapture statistics from Millstone pots (May-Oct) and commercial pots (Jan.-Dec.) from 1978 to 2001.

${ }^{2}$ The minimum legal size from 1978 to 1988 was $81.0 \mathrm{~mm}(3 / 16 \mathrm{in})$, minimum legal size was increased in 1989 to $81.8 \mathrm{~mm}\left(3^{\mathrm{T}} / 32 \mathrm{in}\right)$, and in 1990 to 82.6 mm ( $31 / 4 \mathrm{in}$ ).
females (Atema and Voigt 1995). The escape vent regulation instituted in 1984 requires that commercial lobster pots contain an opening to allow escape of sublegal-size lobsters and reduce injury and mortality associated with overcrowded pots (Landers and Blake 1985; Landers and Keser 2000). Since 1984, claw-loss has averaged $10.9 \%$, which was lower than the average value of $12.7 \%$ reported before the implementation of escape vents. The benefits of incorporating escape vents in lobster traps have been noted by many researchers (Kiouse and Thomas 1975; Fair and Estrella 1976; Krouse 1978; Pecci et al. 1978; Fogarty and Borden 1980; Krouse et al. 1993). Fishery managers have recently implemented a number of additional escape vent requirements including larger vent size, use of biodegradable fasteners when installing vents, and required location of the vent within the trap (parlor section). These measures are expected to further improve lobster survival and recruitment by minimizing trap-related injury and mortality.

## Tagging Program

The total number of lobsters tagged during 2001 was 5,465 and within the range of values reported in 24 years of tagging studies ( $2,768-9,126$; Table 10). The percentage of lobsters tagged in 2001 that were
recaptured in Millstone pots during 2001 was $25.2 \%$ and was also within the range of values observed since the study began in 1978 (range $=14.4-26.2 \%$ ). The number of commercial lobstermen participating in the tag-recapture program declined dramatically last year; as a result, only $1.9 \%$ of the tagged lobsters were reported being caught in commercial pots, which was the lowest value reported in this study (previous range $=6.7-47.6 \%$ ). Many factors influence the percentage of tags returned annually by commercial lobstermen. Most notably; there is lag of about 1 year between the time when lobsters are tagged and the time when commercial lobstermen forward tag return information. The percentage recaptured in commercial traps during 2001 may increase as lobstermen submit their 2001 tag return information in 2002 . Some lobstermen may have left the industry due to the decline in lobster abundance over the past few years. In 1998 and 1999 when legal catches were high, lobstermen had little time to spend recording tag recovery information. . Additionally, the $\$ 2.00$ reward incentive for each tag return is minimal when compared to the price of their catch (pers. comm., several local lobstermen). Lastly, since the implementation of the escape vent regulation in 1984 and recent amendments to this regulation (i.e., larger vent size), fewer recaptures were made by commercial lobstermen.

Most tagged lobsters are sublegal and difficult to retain in commercial traps filted with escape vents. In contrast, the number of recaptures made by Millstone since 1984 increased, because our traps do not have escape vents and retain more tagged sublegal-size lobsters.
The mean CL of lobsters recaptured in Millstone traps was 73.6 mm during 2001 , which was within the range of average sizes reported previously (72.0-75.7 mm ; Table 10). In contrast, lobstermen consistently recaptured larger lobsters. During 2001, the mean CL of 82.6 mm was nearly 10 mm larger than in Millstone traps and among the largest average sizes reported in commercial gear (previous range $=75.5-82.9 \mathrm{~mm}$ ). The average size of tagged lobsters caught in commercial traps has significantly increased over the past 24 years (slope $=0.263 ; \mathrm{p}<0.001$ ). Shifts in the size of lobsters recaptured in Millstone and commercial traps were related to escape vent regulations (Landers and Keser 2000). Prior to the regulation, commercial lobsternen recaptured many of the tagged sublegal-size lobsters; with the regulation in force, many of the sublegal-size lobsters escaped from the vented commercial pots, but were retained in unvented Millstone pots. In eastem LIS, Landers and Blake (1985) noted a substantial reduction in the number of sublegal-size lobsters retained in vented pots, without a corresponding decrease in the catch of legal-size lobsters.
Since the tagging study began in 1978, commercial lobstermen have consistently caught a higher percentage of tagged legal-size lobsters than have Millstone researchers. During 2001, $49.0 \%$ of the recaptures in commercial pots were legal-size ( $\geq 82.6$ mm ) compared to only $6.8 \%$ in Millstone pots (Table 10). The 24 -year time series of percentage legal-size lobsters recaptured in Millstone pots has significantly declined (slope $=0.388 ; p<0.001$ ); no trend was identified in the percentage of legal-size lobsters in commercial traps. The decline in percentage of legalsize lobster recaptures in our traps is not surprising, given the fact that the number of traps fished by Connecticut and New York fishermen in LIS has increased eightfold since the early 1980s (ASMFC 2000). At such high levels of fishing effort, tagged legal-size lobsters are caught by commercial lobstermen soon after they are released and our chances of catching them again are small given the large number of commercial traps present in comparison to the small number of Millstone traps.

## Movement

Tag return information from the Millstone monitoring study and commercial lobstermen was used to assess the extent of lobster migrations in the MPS area.
During 2001, the majority of the lobsters recaptured in Millstone pots were caught at the same station at which they were released ( $99 \%$ at Jordan Cove, $95 \%$ at Twotree, and $92 \%$ at Intake). This pattern of shortrange movement among sublegal-size lobsters was also observed in previous years and in the recapture information provided by commercial lobstermen (Landers and Keser 1994). Since 1978, 92\% of the tagged lobsters recaptured in commercial pots were caught within 5 km of MPS (Fig. 13). Stewart (1972) demonstrated a strong homing behavior for the nearshore eastern LIS lobster population. Individual lobsters leave their burrows at night and return to the same shelters before dawn. This territorial and nocturnal behavior pattern limits their home range. The predominance of localized movement is typical for nearshore coastal lobster populations and agrees with results of other tagging studies conducted in eastem North America (Templeman 1940; Wilder and Murray 1958; Wilder 1963; Cooper 1970; Cooper et al. 1975; Fogarty et al. 1980; Krouse 1980, 1981; Campbell 1982; Ennis 1984; Watson et al. 1999).

Most of the lobsters recaptured more than 5 km from the study area had moved to the east or southeast. Since 1978, only 25 individuals were recaught in western LIS (Fig. 14). Over the past 24 years, more than 1,000 tagged lobsters were caught by commercial lobstermen in The Race, which suggests that this deep water channel between Long Island and Block Island is a migration route for lobsters that exit LIS. Once out of the Sound, lobsters moved casterly and were recaptured in Block Island (43) and Rhode Island Sounds (20). Some lobsters traveled more than 50 km to the east and were caught in Narragansett I3ay, RI (34), Buzzards Bay, MA (6), and in waters near Marha's Vineyard (10) and Nantucket Island (3; Fig. 15). Only two individuals were reported being caught in nearshore waters along the south shore of Long Island, compared to 29 lobsters which moved farther than 150 km to deeper offshore waters on the edge of the continental shelf, where they were caught in submarine-canyons (Hudson, Block, Atlantis, and Veatch). Similar exchanges between inshore and offshore lobster populations have been observed by other researchers (Saila and Flowers 1968; Uzmann et al. 1977; Cooper and Uzmam 1980; Campbell and


Fig. 13. Location and number of lobsters recaptured in the MPS area by commercial lobstermen from 1978 to 2001.


Fig. 14. Location and number of lobsters caught by commercial lobstermen farther than 5 km from MPS in the vicinity of Long Island and Block Island Sounds from 1978 to 2001.


Fig. 15. Number of tag returns at locations farther than 50 km from MPS from 1978 to 2001.



Fig. 16. Mean depth of capture location (a) and distance traveled each month (b) for lobsters caught in commercial traps from 1978 to 2001.

Stasko 1985, 1986; Estrella and Morrissey 1997). Based on our tagging studies, the number of lobsters captured offshore seems insignificant. However, fishing effort for lobsters along the continental shelf is low relative to coastal areas; therefore, the probability of capturing our tagged lobsters in this large area is concomitantly low.
The extent to which local lobsters undergo seasonal distribution shifts to offshore areas was further examined by comparing monthly changes in the average depth and distance traveled between release and recapture locations. For most months (Apr.-Nov.), the average depth of the areas where male and female lobsters were recaptured in commercial traps was similar to the depths, where lobsters were released (range $10-13 \mathrm{~m}$; Fig 16a). However, the average depths of capture location increased in March (15-20 m ) and December ( $14-24 \mathrm{~m}$ ), despite the fact that commercial lobstermen typically redeploy their gear to shallow nearshore waters during these winter months (pers. comm. several LIS lobstermen). Berried females exhibited the highest change in depth location. The average depth of the areas where berried females were caught in commercial traps was 13 m in November and
increased to 25 m in December; an opposite shift in depth occurred from March to April (15 to 10 m , respectively). Additional evidence for seasonal distribution shifts, particularly for egg-bearing females, was found in the average distance traveled between release and recapture locations (Fig. 16b). In June, berried females traveled an average of 2.2 km from the release sites; in July the distance traveled increased to 4.6 km . A second increase in distance traveled by berried females occurred from October to December ( 2.5 to 4.5 km ). These results suggest that lobsters, particularly egg-bearing females, move from shallow nearshore areas during the summer to deeper waters during the winter. Seasonal inshore-offshore migration of coastal lobster populations has been documented by many researchers (reviewed by Lawton and Lavalli 1995) and may be due to a number of factors including coastal physiography, increased turbulence during storms, and seasonal changes in catchability and water temperature (Wilder and Murray 1958; Cooper et al. 1975; Emis 1984). The pronounced seasonal migrations of berried females may be related to the need to optimize egg development and to meet the physiological demands of molting, mating and egg extrusion (Cooper and Uzmann 1971, 1980; Campbell 1986; Talbot and Helluy 1995). In addition, seasonal shallow-deep migrations by egg-bearing females may result in an intermixing of subpopulations at the time of hatching. Recent examinations of demographic linkages among nearshore and offshore lobster populations indicated that adult migrations and larval dispersal from offshore refugia to nearshore areas, or a combination of both mechanisms, may be responsible for the stability and resilience of highly exploited coastal populations of American and European lobster (Fogarty 1998; Sheehy et al. 1999).

## Entrainment

Only 58 lobster larvae were found in samples of MPS cooling water from May 21 to August 2, 2001, which represented the lowest number collected since 1984 when two or three units were operating (previous range 102-571). Stage composition during 2001 was 41 Stage I, 4 Stage II, 2 Stage III and II Stage IV, similar to previous studies when Stage I and Stage IV larvae predominated the collections (Fig. 17). Day and night samples were collected since $198+$ to examine diel variation in lobster larvae entrainment. During 2001, the density of lobster larvae in night samples was 0.315 per $1000 \mathrm{~m}^{3}$ and not significantly different from the


Fig. 17. Annual number of tohster larvae and their stage composition (Stage I-IV) collected in samples taken at the MPS discharges from 1984 through 2001.

TABLE 11. Annual $\Delta$-mean density of hobster larvae collected in day and night entrainment samples from 1984 through 2001.

| Year | Time of day | D-mean density ${ }^{3}$ | 95\% C.l. |
| :---: | :---: | :---: | :---: |
| 1984 | Day | 0.158 | 0.061-0.256 |
|  | Night | 0.737 | 0.138-1.336 |
| 1985 | Day | 0.390 | 0.0-0 820 |
|  | Night | 0620 | 0.290-0.951 |
| 1986 | Day | 0.324 | 0.063-0.585 |
|  | Night | $1.399^{\text {b }}$ | 0.556-2.2.42 |
| 1987 | Day | 0.791 | 0.040-1.542 |
|  | Night | 0667 | 0.205-1.129 |
| 1988 | Day | 0.727 | 0.0-1.653 |
|  | Night | 00.88 | $0271-1.106$ |
| 1989 | Day | 0.158 | 0087-0.229 |
|  | Night | $1.403^{\text {b }}$ | 0.537-2.269 |
| 1990 | Day | 0.341 | 0.101-0.581 |
|  | Night | $1.167^{\text {b }}$ | 0.569-1.765 |
| 1991 | Day | 0.287 | 0.131-0.442 |
|  | Night | $0.756^{\text {b }}$ | 0.502-1.010 |
| 1992 | Day | 1.299 | 0.043-2.555 |
|  | Night | 1.369 | 0.530-2.209 |
| 1993 | Day | 0.963 | 0.0-2.132 |
|  | Night | 1.168 | 0.0-2.433 |
| 1994 | Day | 0.268 | 0.085-0.452 |
|  | Night | $1.505^{6}$ | 0.706-2.303 |
| 1995 | Day | 0.594 | 0.0-1.310 |
|  | Night | 2.189 | 0.369-4.009 |
| 1996 | Day | 0.329 | 0.0-2.057 |
|  | Night | 0.242 | 0.0-1.215 |
| 1997 | Day | $0.2+4$ | 0.0-0.671 |
|  | Night | 0.122 | 0.0-0.281 |
| 1998 | Day | 0.500 | 0.0-1.257 |
|  | Night | 0.502 | 0.0-1.150 |
| 1999 | Day | 0.799 | 0.330-1.268 |
|  | Night | $4.497{ }^{\text {b }}$ | 1.281-7.713 |
| 2000 | Day | 04.45 | 0.101-0.788 |
|  | Night | 1034 | 0.0-2.074 |
| 2001 | Day | 0.290 | 0.067-0.513 |
|  | Nipht | 0.315 | 0.186-0.444 |

${ }^{2}$ Number per $1000 \mathrm{~m}^{3}$.
${ }^{6}$ Significant difference between day' and night densities based on 2 sample t-tests ( $\mathrm{p}<005$ ).

TABLE 12. Annual $\Delta$-mean density (number per $1000 \mathrm{~m}^{3}$ ) of lobster larvae in MPS entrainment samples during their season of occurrence and annual entrainment estimates with $95 \%$ C.I. from 1984 through 2001.

| Year | Time period included | Number of larvae | $\Delta$-mean density | 95\% C.I. | Cooling Vol. $\left(\mathrm{m}^{3} \times 10^{6}\right)$ | Entrainment estimate $\times 10^{3}$ | 95\% C.I. $\times 10^{3}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |  |
| 1984 | 21May-10Jul | 102 | 0.409 | 0.184-0.635 | 181.8 | 74.4 | 33.5-115.4 |
| - 1985 | - 15May-16Jul | 142 | 0.504 | 0.258-0.749 | 244.9 | 123.4 | 63.2-183.4 |
| $1986^{\text {a }}$ | 14May-14Jul | 232 | 0.857 | 0.418-1.297 | - 639.5 | 548.1 | 267.3-829.4 |
| 1987 | 18May-30Jul | 184 | 0.943 | 0.274-1.613 | . , 406.8 | 383.6 | 111.5-656.2 |
| 1988 | 16May-01Aug | 571 | 0.717 | 0.296-1.137 | 804.1 | 576.5 | 238.0-914.3 |
| 1989 | 22May-28Jul | 237 | 0.701 | 0.358-1.044 | 540.3 | 378.8 | 193.4-564.1 |
| 1990 | 14May-30Jul | 280 | 0.748 | 0.436-1.060 | $\because 747.9$ | 559.4 | 326.1-792.8 |
| 1991 | 07May-22Jul | 157 | 0.525 | 0.365-0.685 | - 541.6 | 284.3 | 197.7-371.0 |
| 1992 | 19May-14Jul | 625 | 1.334 | 0.652-2.016 | 449.5 | 599.6 | 293.1-906.2 |
| 1993 | 24May-25Jun | 218 | 1.081 | 0.273-1.889 | 346.1 | 374.1 | 94.5-653.8 |
| 1994 | 25May-04Aug | 257 | 0.908 | 0.445-1.371 | 715.4 | 649.6 | 318.4-980.8 |
| 1995 | 30May-21Jul | 254 | 1.385 | 0.470-2.300 | 476.1 | 659.4 | 223.8-1095.0 |
| 1996 | 06Jun-01Jul | 19 | 0.364 . | 0.194-0.535 | 53.0 | 19.3 | - 10.3-28.4 |
| 1997 | 30May-16Jul | 15 | 0.186 | 0-0.395 | 51.6 - | 9.6 | 0.0-20.4 |
| 1998 | 20May-23Ju! | 145 | 0.511 | 0.048-0.973 | 244.5 | 124.9 | 11.7-237.9 |
| 1999 | 17May-02Jul | 529 | 2.502 | 1.107-3.896 | 237.9 | 595.2 | 263.4-926.9 |
| 2000 | 17May-13Jul | 155 | 0.725 | 0.276-1.173 | 451.5 | 327.3 | 124.6-529.6 |
| 2001 | 21May-02Aug | 58 | 0.299 | 0.185-0.414 | 605.1 | '180.9 | 111.9-250.5: |

${ }^{2}$ Unit 3 began commercial operation.
density of 0.290 per $1000 \mathrm{~m}^{3}$ in day samples (Table 11). In six of the previous study years (1986, 1989, 1990, 1991, 1994, and 1999), significantly higher larval densities were observed in night samples; lobster larvae were never significantly more abundant in day samples. This observed diel variability was similar to results found elsewhere (review by Ennis 1995). Early laboratory studies on lobster larvae behavior demonstrated that Stage I larvae exhibit positive phototaxis and disperse from surface waters during darkness (Templeman 1937, 1939). In contrast, field súriveys conducted by Harding et al. (1987) in deeper, less turbid Canadian waters indicated that most Stage I larvae were collected at depths between 15 and 30 m during the day and were rarely found below 10 m at night. More recently, DiBacco and Pringle (1992) found significantly more Stage I larvae during nightime than during daytime in neuston tows made within a protected coastal embayment along the Nova Scotian Shelf. Since the MPS intakes are sited on the shoreline, and draw water from throughout the $8-10 \mathrm{~m}$ well-mixed water column, the'observed diel variability in our study may be influenced more by wind and tide generated water circulation than by vertical movement by larvae.
The $\Delta$-mean density of lobster larvae collected in entrainment samples during 2001 was 0.299 per 1000 $\mathrm{m}^{3}$, which was the lowest value reported in our entrainment studies when the MPS Units were operational (0.409-2.502; Table 12). An estimated

180,900 lobster larvae were entrained through the MPS cooling water system during 2001, which was within the range of previous studies when two or three units operated ( $74,400-659,400$ ). The lowest entrainment occurred in 1996 and 1997 when MPS units were offline ( 19,300 and 9,600 , respectively). Lobster larvae entrainment is directly related to both the annual larval density and the operational status of the MPS units during the hatching season. During years when lobster larvae abundance is high and units operate at full capacity, cooling-water demands are at a maximum and resulting entrainment estimates are higher. Conversely, entrainment estimates are low when lobster larvae abundance is low and one or more units are shutdown for maintenance or refueling.
Evaluating the effect of entrainment on lobster recruitment is difficult because of the high variability in lobster larvae abundance and stage composition (Bibb et al. 1983; Fogarty 1983; Lux et al. 1983; Blake 1984, 1988), the lack of reliable estimates of larval and postlarval survival rates (Phillips and Sastry 1980; Caddy and Campbell 1986; Cobb 1986; Blake 1991), and the uncertainty regarding post-settlement processes of early benthic phase lobsters controlling recruitment to the fishery (Hudon 1987; Incze and Wahle 1991; Wahle and Steneck 1991). Disagreement among researchers on the source and dispersion mechanism of lobster larvae and on egg and larval mortality has led to a range of survival estimates during the larval life history phase from less than $1 \%$ in Canadian waters (Scarratt

1964, 1973; Harding et al. 1982) to more than $50 \%$ in LIS (Lund and Stewart 1970; Blake 1991). Mechanisms of lobster larvae dispersal in coastal waters may be related to surface water circulation patterns (Fogart 1983). Surface currents regulated by the wind and tide converge and are visible on the surface waters as "slick" or "scum" lines. These convergence areas delineate zones of upwelling and downwelling and were reported to contain high densities of planktonic organisms including lobster larvae (Cobb et al. 1983; Blake 1988). Convergence areas were often seen in the MPS area stretching from near Twotree Island into Niantic Bay in 1988 and 1992 and could explain the large number of lobster larvae (all stages) collected in single samples of the cooling water in those years (DNC 2001a). Furthermore, based on the short duration of the first larval stage ( $3-5$ days), the source of Stage I larvae collected in the MPS cooling-water was probably from local spawners (e.g., from Twotree, where $25.2 \%$ of females were berried during 2001). The recent outbreak of shell disease in our area may be responsible for the lower abundance of larvae, if the disease affects egg-development or survival of newly hatched larvae. Stage IV larvac, however, are in the water column between 4 and 6 weeks, and, based on water circulation patterns in LIS, were unlikely to have originated locally. Lund and Stewart (1970) indicated that the large number of berried females found in western LIS compared to eastern LIS may be responsible for recruitment of Stage IV larvae throughout LIS. Stage IV larvae were found to exhibit directional swimming behavior and moved tens of kilometers from offshore spawning grounds (Cobb et al. 1989; Rooney and Cobb 1991; Katz et al. 1994). If lobster larvae found in our area originate from westem LIS, the recent lobster mortality event in western LIS could explain the lower abundance of lobster larvae during 2001. A similar 'source/sink' recruitment model was hypothesized by Fogarty (1998); he attributed the stability and resilience of overfished coastal lobster stocks to migration of adults and advection of larvae to coastal waters from less intensively fished areas far from shore on the edge of the continental shelf.

## Mortality and Shell Disease

Since 1984, we recorded the presence of dead or dying lobsters caught in our traps during each sampling trip. In addition, we qualitatively assessed external damage to the carapace and abdomen and noted the

TABLE 13. Incidence of dead or dying lobsters caught in traps from May through October 1984-2001.

| Year | Total <br> collected | Number <br> dead | $\%$ | Dead <br> males | Dead <br> females |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |
| 1984 | 7587 | 58 | 0.8 | 33 | $24(1)$ |
| 1985 | 7014 | 35 | 0.5 | 16 | 19 |
| 1986 | 7211 | 63 | 0.9 | 34 | 29 |
| 1987 | 7280 | 52 | 0.7 | 33 | 19 |
| 1988 | 8871 | 68 | 0.8 | 35 | 33 |
| 1989 | 7950 | 62 | 0.8 | 37 | 25 |
| 1990 | 7106 | 72 | 1.0 | 48 | $22(2)$ |
| 1991 | 7597 | 82 | 1.1 | 50 | $31(1)$ |
| 1992 | 11438 | 117 | 1.0 | 84 | 33 |
| 1993 | 10195 | 94 | 0.9 | 58 | $35(1)$ |
| 1994 | 9849 | 67 | 0.7 | 36 | 31 |
| 1995 | 6435 | 70 | 1.1 | 50 | 20 |
| 1996 | 7531 | 256 | 3.4 | 200 | $55(1)$ |
| 1997 | 9026 | 117 | 1.3 | 101 | 16 |
| 1998 | 10991 | 104 | 0.9 | 79 | 25 |
| 1999 | 11216 | 302 | 2.7 | 210 | $89(3)$ |
| 2000 | 8707 | 157 | 1.8 | 111 | $43(3)$ |
| 2001 | 7268 | 200 | 2.8 | 142 | $55(3)$ |
|  |  |  |  |  |  |
| 20 |  |  |  |  |  |

${ }^{2}$ Parenthetical values are number berried.
presence of shell disease (chitinoclasia). Two hundred ( $2.8 \%$ ) dead lobsters were observed in our catch during 2001, which was within the range of values observed in previous years ( $\mathrm{n}=35$ to $302,0.5-3.4 \%$; Table 13). In general, more males were found dead than females, especially from 1996 through 2001 when dead males ( $n=843$ ) outnumbered dead females ( $n=293$ ) by a factor of three. The cause of the recent increase in lobster mortality in our area and in western LIS is unclear. Pathologists have examined lobsters from western LIS at the request of NYSDEC and CTIDEP but have been unable to identify the cause of the die-offs. Comprehensive examinations of water quality parameters and bottom sediments failed to identify environmental conditions or toxins that could explain the lobster deaths. More recently, histologic examinations of lobsters revealed a systemic inflammatory disease affecting multiple tissues. The nervous system was markedly intlamed and the lesions were associated with a parasitic amoeboid protozoan (French et al. 2000, 2001). At present, the cause of death is hypothesized to be paramoebiasis. Most researchers agree that the die-off in western LIS was caused by multiple discase pathogens exacerbated by environmental stressors, such as increased seawater temperature, anoxia and contaminants. Stewart (1980) indicated that disease and parasitism are dominant sources of natural mortality in wild lobster populations. Mass mortality caused by fatal diseases such as gaffkemia (Snieszko and Taylor 1947) have been
reported for lobsters reared in high densities, but little is known about the prevalence and effects of pathogens on natural lobster populations (Martin and Hose 1995).
Lobster mortalities in the area around MPS may be related to shell disease, a more common affliction in marine and freshwater crustaceans. Shell disease is characterized as a deterioration of the exoskeleton by chitinoclastic microorganisms (Rosen 1970; Sinderman 1970). Gross signs of the disease are similar in all crustacean : species; the exoskeleton is pitted and marred with necrotic lesions and, although the disease is not immediately fatal, death may occur (Fisher et al. 1978). For lobsters, the unsightly appearance of the shell can greatly affect marketability. Chitin-digesting bacteria and fungi have been implicated as causative agents and include isolates of Vibrio spp., and several other Gram-negative bacilli (Hess 1937; Getchell 1989). From 1984 to 1997 only a few lobsters ( $\mathrm{n}=7$; $<0.1 \%$ ) were found with signs of shell disease. In the fall of 1998, 156 (6.5\%) lobsters had signs of minor shell disease. The incidence of shell disease increased markedly in 1999; with $24 \%$ and $38 \%$ of the total catch infected during September and October, respectively (Fig. 18a). In spring 2000, lobster biologists from southern New England and New York developed standardized procedures for monitoring the extent of shell disease in wild populations. The following index was established based on the percent shell coverage of disease symptoms (e.g., pitting, erosion, lesions) on the total surface area of the lobster: $0=$ no shell disease symptoms, $1=$ symptoms on $1-10 \%$ of shell surface, $2=$ symptoms on $11-50 \%$ of shell surface, and 3 =symptoms on $>50 \%$ of shell surface. During the past two years, the incidence of shell disease (all severity indices) closely corresponded to the molt cycle observed from May through October. Prior to the molt in May 2000 and 2001, 20\% to 30\% of the catch had shell disease (Fig. 18a). Following the major molt in July, only about $5 \%$ of the catch was afflicted in both years. The severity and percentage of diseased lobsters increased in August and September, reaching a peak in October both years when nearly $60 \%$ of the catch had symptoms of shell disease. In comparison, prevalence of shell disease in central and western LIS and in offshore canyon areas has never exceeded $5 \%$ (Landers et al. 2001a). In eastem LIS and other coastal areas of . southern New England, male and female lobsters of all sizes have been observed with shell disease, although larger sized individuals and egg-bearing females had higher incidence and severity .. of shell disease symptoms. Between $50 \%$ and $60 \%$ of the egg-bearing females had shell disease during 2000 and 2001,


Fig. 18. Percentage and severity of shell diseased lobsters collected in the vicinity of MPS from 1998 to 2001 (a) by month and (b) by sex.
respectively, which was considerably higher than the percentages observed in non-berried females and males (2000-2001 range $=12-21 \%$; Fig 18b). In nearshore waters of Rhode Island over $50 \%$ of the egg-bearing females observed in the trap fishery were infected with shell disease in 1999 (Castro and Angell 2000). This is most likely due to the molt cycle of large lobsters and egg-bearing females. Smaller lobsters molt more frequently ( $1-2$ times/yr or more) and shed their shells before severe shell disease symptoms occur. Large lobsters and egg-bearing females experience more shell deterioration because they may only molt every 2 years.
In addition to recording the presence and severity of shell disease on each lobster, we also noted the appearance of scars on the exoskeleton due to previous (i.e., premolt) infections of chitinoclasia. The occurrence of secondary infections began in June 2000 ( $16 \%$ ) following the onset of the major molt in May that year. The prevalence of secondary infections increased each month during 2000 and peaked in October when $53 \%$ of the catch had shell disease for a second time (Fig. 19). In May 2001, the levels of secondary infections were the same (53\%) as those observed the previous fall, but declined each month to


Fig. 19. Percentage of lobsters (by month and sex) infected with shell disease for a second time during 2000 and 2001 (based on presence of scars).
about 20\% in October 2001. At present, the causes of this unusual pattern of secondary infections (increase followed by a decrease) are unclear. One possibility is that the disease may be subsiding. Alternatively, in many cases secondary infections were found to occur on the same areas of the lobster shell; as bacteria consume these areas of the shell, previous disease-scars become obscured, which may have resulted in the declining pattern of secondary infections observed in 2001. Over the past two years, berried females had the highest percentage of secondary infections (29 and $45 \%$ ), followed by males ( 24 and $42 \%$ ), and nonberried females ( 14 and $37 \%$; Fig. 19). While these observations indicate that shell disease may not be immediately fatal, successive infections may cause additional physiological stress resulting in decreased locomotion and agonistic behavior which could lead to increased rates of predation.
A number of isolated outbreaks of shell disease have been reported in lobster populations along the New England coast in the past century. However, these reports were linited to impounded lobsters in the Gulf of Maine during the 1930s (Iless 1937; Taylor 1948) and to infrequent occurrences in wild lobsters in the 1980s along Massachusetts coastal waters and the New York Bight (Sinderman et al. 1989; Estrella 1991). The present epizootic is clearly different from any other cases reported for American lobster. The cause of the outbreak is unknown; it may be due to water quality degradation along the coast or to natural environmental factors such as warmer seawater temperature. However, it is surprising that the prevalence of shell disease was low in central and western LIS, two areas known to be more polluted with
domestic sewage and industrial contaminants than areas to the east. The etiology of shell disease may be enhanced by increased transmission due to crowding as a result of the sharply higher abundance of lobsters in recent years. Most alarming is the prevalence and severity of shell disease in egg-bearing female lobsters. The effect of the disease on lobster recruitment in southern New England may be significant if eggbearing females suffer higher natural mortality due to shell disease.

## Conclusions

The most notable changes in the population characteristics of local lobsters during 2001 were observed in the proportion of egg-bearing females and their size structure which may be related to changes in lobster growth and the size at which females become sexually mature. The benefits of earlier maturation may be significant to population dynamics. Small size at maturity and subsequent egg production from sublegal-sized females may explain why LIS lobsters are resilient to high levels of fishing pressure. Currently, the smaller size at sexual maturity allows females to spawn 1 or 2 times before reaching the legal size possibly sustaining current populations levels into the future. However, the long-term sustainability of the Connecticut fishery may now be threatened by the lobster die-off in western LIS and the widespread outbreak of shell disease in our area. The number of lobster larvae present in the samples of the cooling water was the lowest observed in 18 years of lobster larvae entrainment studies. The lobster mortality event in westem LIS could be responsible for the low number
of larvae collected in 2001, if larvae found in our area originate from the west. We found that berried females had the highest prevalence and severity of shell disease, but the effects of the disease on egg-development, hatching success and larval survival are unknown. The events of mass mortality in western LIS and the effects on recruitment in our area due to the recent outbreak of shell disease could be significant, if berried females suffer higher natural mortality. These effects are difficult to assess at this time, because lobsters require several years of growth before they are vuinerable to capture in our traps.
At present, the widespread die-off and recent catastrophic decline in lobster abundance observed in western LIS has not been observed in our study area in eastern LIS. Although catch-per-unit-effort of 1.587 lobster/pot during 2001 continued to decline from the record high CPUE of 2.560 observed in 1999, this value was within the range of annual CPUEs observed since 1978. . In addition, results from our studies over the past 24 years indicated a significant increasing trend in total CPUE, which is comprised mostly of recruit and prerecruit-sized lobsters (one and two molts below legal size). Furthermore, despite the fact that nearly all ( $>90 \%$ ) the lobsters larger than the minimum legal size are removed by fishing each year, no significant long-term trend in legal lobster CPUE was observed. These findings, combined with. the increasing trend in commercial lobster landings in our area of LIS, suggest that operation of the MPS cooling water system since the early 1970s has not caused a decrease in the local lobster population. Continued monitoring will help determine if operation of MPS has caused changes in the abundance and population characteristics of local lobsters beyond those expected from natural factors such as disease and the intense exploitation of this species in coastal waters.

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## Benthic Infauna

## Introduction

Marine benthic habitats in the vicinity of Millstone Power Station (MPS) and elsewhere support rich and diverse infaunal invertebrate communities. These communities are important as a source of food for numerous animal species, including lobsters and demersal fishes (Richards 1963; Moeller et al. 1985; Watzin 1986; Horn and Gibson 1988; Commito and Boncavage 1989; Franz and Tanacredi 1992; Commito et al. 1995). The natural activities of infauna, such as feeding, burrowing, and tube-building, also promote nutrient recycling from sediments to the water column (Goldhaber et al. 1977; Aller 1978; Gaston and Nasci 1988), and aid transfer of oxygenated water deeper into sediments. It is clear that the presence of a diverse benthic infaunal community is essential to the health of coastal marine ecosystems.
The close association of benthic communities with sediments, where most pollutants ultimately ${ }^{-}$ accumulate, also makes infaunal communities effective indicators of acute and chronic environmental conditions (Warwick 1986; Diaz and Schaffner 1990; Warwick et al. 1990; Somerfield et al. 1995; Zajac et al. 1998). Changes in benthic community structure and abundance that follow disturbance (Boesch and Rosenburg 1982; Young and Young 1982; Warwick et al. 1987, 1990; Gaston and Nasci 1988; Regnault et al. 1988; Rees and Eleftheriou 1989; NAESCO 1994; Prena 1995; Somerfield et al. 1995) provide a uscful baseline in the evaluation of impacts and recovery sequences that take place in benthic marine systems.
Coastal benthic communities are subject to a high degree of environmental variability (Holland 1985; Nichols 1985; Holland et al. 1987; Warwick 1988; Rees and Eleftheriou 1989; Boero 1994). The interaction of many physical and biological factors creating this natural variability, and their effects on the structural and functional ecology of benthic communities (Diaz and Schaffner 1990; Hutchings 1998; Zajac et al. 1998, 2000), hinders prediction of how these communities respond to disturbance. Longterm monitoring studies are essential to assess changes in marine benthic communities (Thrush et al. 1994; Prena 1995). Such studies are the principal means of documenting and characterizing changes in community structure and fluctuations in species abundance, which occur in response to climatic conditions (Boesch et al. 1976; Flint 1985; Jordan and Sutton 1985), to variations in biological factors (Levinton and Stewart 1982; Woodin 1982; Kneib 1988), and to human activities.

Studies of subtidal benthic infauna have identified impacts to communities that were attributed to Unit 3 intake construction (NUSCO 1987, 1988a) and to 3unit operations (NUSCO 1997), as well as to regional shifts in species composition and abundance that are assumed the result of natural events. The focus of this monitoring program is to measure infaunal species . composition and abundance, to identify spatial and temporal patterns in community structure and abundance, and to assess whether observed changes could possibly have been the result of construction and operation of MPS. This report presents results from the 2001 sampling year, and compares them to results summarized from previous years (1980-2000).

## Materials and Methods

Subtidal infaunal communities in the vicinity of MPS were sampled twice per year (June and September) from 1980 through 2001 at four stations (Fig. 1). The Giants Neck station (GN), located 6 km west of MPS, is outside the area potentially affected by power plant operations. Data from GN are used to identify possible region-wide shifts in infaunal community structure and composition occurring independently of power plant operations. The Intake station (IN), located 100 m seaward of MPS Unit 2 and Unit 3 intake structures, is exposed to scour produced by inflow of cooling water and the effects of periodic dredging. The Effluent station (EF), located approximately 100 m offshore from the station discharge into Long Island Sound, is exposed to increased water temperatures and scour, and to chemical or heavy metal additions to the cooling water disclarge. The Jordan Cove station (JC) is located 500 m east of MPS. The area encompassing this station experienced increases in surface water temperatures of 0.8 to $2.2^{\circ} \mathrm{C}$ above ambient during some tidal stages (primarily ebb tide) due to the 3 -unit thermal discharge of MPS (NUSCO 1988b).
At each station, ten replicate samples ( $0.0079 \mathrm{~m}^{2}$ each) were collected by SCUBA divers using a handheld coring device 10 cm in diameter $\times 5 \mathrm{~cm}$ decp. Each sample was placed in a 0.333 mm mesh Nitex bag and returned to the laboratory. Samples were fixed with $10 \%$ buffered formalin. After a minimum of 48 hours, organisms were floated from the sediments onto a 0.5 mm mesh sieve and preserved in a $70 \%$ ethanol solution with Rose Bengal added to facilitate sample processing. Samples were examined with dissecting microscopes (10x); organisms were sorted into major


Fig. 1. Map of the Millstone loint area showing the location of infaunal sampling stations (EF=Effucnt, GN=Giants Neck, IN=Intake, $\mathrm{JC}=$ Jordan Cove) established as part of the long-term monitoritg program for Millstone Power Station.

Groups (annelids, arthropods, molluses, and others) for later identification to the lowest practical taxon and counted. Oligochactes and rhynchocoels were each treated in aggregate because of the difficulties associated with identifying these organisms. Organisms too small to be quantitatively sampled by our methods (meiofauna; e.g., nematodes and the smaller ostracods, copepods, and forminifera) were not sorted. Grain size and sild/clay fraction were determined from a 3.5 cm diameter $\times 5 \mathrm{~cm}$ core, taken at the time of infaunal sampling. Sediment samples were analyzed using the dry sieving method described by Folk (1974).

## Data Analyses

## Sediments

Sediment sieve fractional weights were used to construct cumulative curves for 2001 and the period 1980-2000. June and September weights from each
sieve used for grain size analysis were pooled within the 1980-2000 period, with years serving as replicates. The curves were used to graphically illustrate differences in sediment composition between the 2001 period and previous years.

## Trend Analysis

The nonparametric (i.c., distribution-free) MannKendall test (Hollander and Wolfe 1973) was used to determine whether time-series data on community abundance, numbers of species, and selected taxa over the entire study period exhibited significant trends. Monthly (June and September) data were plotted against time; however, annual means of community and specitic population abundance were used as variables in the trend analysis.

## Community Analyses

Comparisons of annual collections at each station were made by calculating the Bray-Curtis similarity
index between each pair of years, using the formula (Clifford and Stephenson 1975):

$$
S_{j k}=\frac{\sum_{i=1}^{n} 2 \min \left(X_{i j}, X_{i k}\right)}{\sum_{i=1}^{n}\left(X_{i j}+X_{i k}\right)}
$$

where $S_{\mathrm{jk}}$ is the similarity index between year $j$ and year $k ; X_{\mathrm{ij}}$ is the $\log$ transformed $(\ln +1)$ abundance of taxon $i$ in year $j ; X_{i k}$ is the abundance in year $k$; and $n$ is the number of taxa in common, for which, on average, at least two individuals were found per year. A group-average-sorting, clustering algorithm was applied to the resulting similarity matrix (Lance and Williams 1967).

## Non Metric Multi-Dimensional Scaling (MDS)

The Non Metric Multi-Dimensional Scaling (MDS) application used here was introduced by Shepard (1962) and Kruskal (1964). MDS was applied to the same similarity matrix used in the classification and cluster analyses. This analysis was performed with the PRIMER suite of programs developed by the Plymouth Marine Laboratory (Warwick and Clarke 1991; Clarke 1993). The purpose of MDS is to construct a map or configuration of samples in a two-dimensional plot. There are no absolute similarity values associated with MDS. Interpretation of sample placement on a configuration/map is relative in nature, i.e., more similar samples are closer together. The iterative process of MDS seeks to position the elements in the configuration/map with the least amount of distortion (stress) derived from the original data matrix. Stress values less than 0.2 indicate a potentially useful 2 dimensional picture. Results of the cluster analyses are compared with the ordination plot as a check of the adequacy and mutual consistency of both representations.

## Results

## Sedimentary Environment

Sedimentary environments at infaunal sampling stations in the vicinity of MPS were analyzed for mean grain size and silt/clay content (Fig. 2). Sediment grain size means in 2001 were 0.34 mm and 0.85 mm (in June and September, respectively) at GN, 0.33 mm and
0.43 mm at $\mathrm{EF}, 0.18 \mathrm{~mm}$ and 0.30 mm at IN , and 0.14 mm and 0.20 mm at JC (Fig. 2). Mean grain sizes during 2001 were within the ranges of previous years at IN, EF and JC. Mean grain size at GN in September 2001 was the coarsest observed at any subtidal station over the study period, and was the result of the higher than normal proportion of broken blue mussel (Mytilus edulis) shell observed in the sample. Mean grain size increased at EF between 1985 and 1995 with a decreasing trend in subsequent years. Intake sediments have also shown a slight increase in grain size since 1989, while at JC, grain size has generally decreased since 1986. Grain size at GN has shown no consistent trend over the study period. Silt/clay contents of sediments collected in 2001 were highest at JC, $22.9 \%$ and $13.6 \%$ in June and September, respectively. At all other stations sediment silt/clay contents were less than $10 \%$ during 2001: EF ( $3.9 \%$ and $1.8 \%$ ); GN ( $7.4 \%$ and $0.4 \%$ ); IN ( $4.5 \%$ and $5.8 \%$ ) in June and September, respectively (Fig. 2). Silt/clay estimates at EF, IN, and JC in 2001 were within historical ranges. Sediment silt/clay content in September 2001 at GN was the lowest observed during since the study began in 1979 (also attributed to the high quantity of blue mussel shell).
Cumulative curves based on sediment sieve fraction weights (Fig. 3) were used to compare sediments collected during 2001 and the 1980-2000 period. Notable differences between cumulative curves from 2001 and the 1980-2000 period were observed at GN and JC, and were related to shifts in coarse grain fractions in 2001, i.e., an increase at GN and a decrease at JC. At EF and IN sediment curves illustrated only slight differences in grain-size distribution from previous years. EF sediments showed minor decreases in the coarse fractions, with an increase in the medium sand range. IN sediments in 2001 showed slight increases in the coarse and fine sand fractions compared to the 1980-2000 period.

## General Community Composition

Some useful descriptors of benthic community composition include total numbers of individuals and species, along with numbers of individuals and species within major taxonomic groups. Station totals for number of individual organisms collected in 2001 (Fig. 4) were highest at JC $(6,752)$, intermediate at GN $(5,265)$, and low at EF $(2,994)$ and IN $(2,859)$. These totals were within historical ranges for each station. Relationships among stations for total individuals in 2001 were consistent with those observed in most previous years. For example, during the 22 study years, total abundance has been highest at JC for 18 years, and lowest at IN for 16 years.





Fig. 2. Mean grain size (mun) and silvclay (\%) of sediments at MNPS infaunal stations for June and September from 1980 to $200 t$.


Fig. 3. Cumulative curves based on fractional weights of sediment collected during the 1980-2011 period and during 2001.


Fig. 4. Total annual abundance of individual organisms collected during 1980-2001 at MPS infaunal stations.

Polychaetes comprised more than $65 \%$ of the individuals collected at three of the four stations in 2001: 4,810 polychaetes were collected at JC, 4,052 at GN, and 1,871 at $\mathrm{IN}^{( }$(Fig. 5). At EF, the 1,748 polychaetes collected in 2001 comprised $58 \%$ of the total individuals. Oligochactes were the second most abundant group at all sites in 2001: annual counts at JC ( $1, .453$ ), EF $(1,018), \mathrm{GN}(991)$ and IN (717) were all within historical ranges.
Arthropods were the third-ranked taxonomic group in 2001 at EF, GN and IN ( 91,115 and 183 individuals, respectively), and ranked fourth in total abundance at JC (120). Arthropod abundances in 2001 were at historical lows at EF (91) and GN (115). At JC and IN, 2001 arthropod abundances (120 and 183, respectively) were at the low end of the historical ranges. Molluscs ranked fourth in abundance at EF (77), GN (64) and IN (68) and third at JC (304). At all stations, mollusc abundances in 2001 were relatively low but within historical ranges. The extremely high abundance of Mytilus edulis at EF observed last year (i.e., over 3,000 individuals) was not repeated in 2001.

Annual total numbers of species in 2001 (Fig.6) were highest at JC (105), followed by IN (97), EF (96), and GN (94); these values were within the range of previous study years at all stations. Similar to total abundance discussed above, most of the species identified were polychaetes: 64 at $\mathrm{JC}, 60$ at $\mathrm{GN}, 57$ at IN and 53 at EF (Fig. 6). Polychacte species numbers for all stations were within their historical ranges. Numbers of arthropod species in 2001 were highest at EF and IN (24) followed by GN and JC (20), with all station values within historical ranges. The relationship of numbers of arthropod species among stations has remained relatively stable over years, with IN generally
having highest arthropod species numbers and JC most often the lowest. Numbers of mollusc species in 2001 was similar among EF (16), IN (16) and GN (14); the number of mollusc species was highest at JC (21).

## Faunal Abundance

Mean faunal densities in June and September 2001 collections, respectively, were 181 and 105 individuals per core at IN, 181 and 119 at EF, 313 and 214 at GN, and 300 and 375 at JC (Fig. 7). Faunal densities were within historic ranges established since 1980 at all stations. Analyses of long-term trends in mean community abundance indicated a significant increase in number of individuals collected at IN since 1980. There were no significant trends in mean density evident at any other station during the 1980-2001 study period.

## Numbers of Species

Mean numbers of species per core in 2001 June and September (respectively) were 29 and 19 at EF, 29 and 21 at GN, 29 and 16 at IN, and 32 and 28 at JC (Fig. 8). All 2001 means were within the range of means observed at each station established for previous study years. Trend analysis revealed a significant long-term increase in annual mean species numbers per core over the entire study period (1980-2001) at IN. There was a significant decreasing long-term trend at EF. There was no long-term trend in annual mean number of species per core at GN or JC.


Fig. 5. Total abundance of individuals by major taxonomic group at the MISS infamal stations during 1980-2001. Molluse abundance at EF in $20 \times 10$ ( $3,3.44$ individuals) was outside the range of typical values and was not plotted proportionally. See Community Composition Section for more discussion.





Fig. 6. Total number of species within major taxonomic groups, and overall total collected, at the infaunal stations during 1980-2001.


Fig. 7. Mean faunal density (per core) at Millstone subtidal stations in June and September for every gear from 1980 through 2001 . Vertical line marked with (*) indicates Unit 3 restart in July 1998.


Fig. 8 Mean number of species (per core) at subtidal stations in June and September for every year from 1980 through 2001. Vertical line marked with (*) indicates Unit 3 restart in July 1998

## Community Dominance

Relative abundance of common taxa provides insight into community dominance and annual variation, and long-term infaunal community structure. The relative abundances of infaunal taxa were calculated for each station during 2001 and compared to means over the previous 21 years (19802001; Table 1). While many taxa were collected in 2001, only four (representatives of the class Oligochaeta, and the polychaetes Aricidea catherinae, Tharyx spp. and Polycirrus eximius) had relative abundances of $>10 \%$ at one or more stations. The live top-ranked taxa at each station in 2001 accounted for $64 \%$ or more of the total individuals, and were in descending order: Oligochaeta, and the polychaetes Polycirrus eximius, Parapionosyllis longicirrata, Protodorvillea gaspeensis and Tharyx spp. at EF: Tharyx spp., Oligochacta, Aricidea catherinae, $P$. cximins, and $P$. gaspeconsis at $\mathrm{GN} ; d$. catherinac, Oligochaeta, and the polychactes Exogone hebes. Tharyx spp., and $P$. gaspecnsis at IN; and, $A$. catherinue, Oligochaeta, and the polychactes Scoletoma temuis, Thary: spp., and $P^{\prime}$. cximius at JC. In 2001, five taxa were among the top ten dominants at all sites: Oligochacta, Polycirrus eximias, Protodorvillea gaspeensis, Tharyx spp. and Polydora socialis.
Variability in relative abundance levels and shifts in rankings among dominant taxa from year-to-year have been typical of benthic communities at each monitoring site, and are often related to short-term pulses in abundance of particular taxa. Some of these pulses occur at more than one site, suggesting regional factors. $A$ good example of this phenomenon is the regional inerease in relative abundance of the Polydora socialis in 2001. This species was not among the overall study period dominants at any station, and has rarely been among the ammal dominam taxa. Multi-site increases in relative abundance (compared to the overall study period) of Polycirrus eximios (all sites), Tharyx spp. (GN, IN, and JC) and Parapionosyllis longicirrata (EF, IN, and JC) were also observed in 2001. Other species have exhibited periodic regional pulses in past years two or more times during the study period, including Mediomastus ambiseta and Prionospio steentrupi, but were not among community dominants in 2001. Similarly, the amphipod Leptocheirus pinguis accounted for $4.3 \%$ of the total organisms collected at both IN and JC over the 1980-2001 study period, ranking sixth and fifih, respectively, but relative abundance was $<1 \%$ at both stations in 2001.
Other shifts in relative abundance in 2001 were not consistent among sites or were site-specific. For
example, when compared to the 1980-200t period, Aricidea catherinae abundance in 2001 was high at IN and JC, but low at GN. Oligochaete relative abundance at IN also showed a large increase in 2001 (25.6\%) when compared to the overall study period (12.3\%). At the other sites, 2001 oligochacte abundance was lower than (EF) or was similar to (JC and GN) the 1980-2001 period. Some site-specific shifts in relative abundance moted during 2001 include increases of Polydora caulleryi at EF, and decreases of Pygospio clegans, Capitella spp., and Ampelisca verilli at IN.

## Dominant Taxa

Seven infaunal taxa have been selected as representative of sites affected by construction and operation of MPS. These taxa are useful biomonitoring tools because they are also indicators of natural environmental stress (e.g., storms). Trends in their abundance were examined using the same techniques as those applied to time-series of overall community abundance and numbers of species.

Oligochaetes - Representatives of the class Oligochata are among the dominant taxa throughout the study period at all stations, accounting for 1.3$43 \%$ of the organisms collected since 1980. During 2001, oligochacte abundance during June and September, respectively, averaged 55 and $47 /$ core at EF, 63 and $36 /$ core at $(3 N, 41$ and $32 / \mathrm{core}$ at 1 N , and 104 and $42 /$ eore at JC. At all stations, oligochate densities in 2001 were within the ranges of densities for previous study years (Fig. 9a-d). Oligochacte abundances were higher in the June than Septermber at all stations. Long-term (1980-2001) trend amalysis of oligochaete abundance revealed that the increase noted at IN was significant (Fig. 9e). No significant long-term trends were detected at all other stations.

Aricidea catherinae - The polychacte A. catherinae was among the top ten dominant taxa at all stations except EF in 2001 (Table 1). Average densities in June and September 2001 (Fig. 9e-h) were highest at JC (39 and $143 /$ core, respectively) and IN (47 and 37/core, respectively). Lower average densities occurred at EI: ( 2 and $4 / \mathrm{core}$ ) and GN ( 24 and $20 / \mathrm{core}$ ). Average densities all all stations in 2001 were within historical ranges.

Long-term (1980-2001) trend amalysis indicated a significant increase in abundance of Aricidea cotherinae at IN and JC, and a significant decreasing trend at GN. Aricidea abundances at EF have been

TABLE 1. Relative abundance (\%) and coefficient of variability (C.V.) of each of the ten most abundant taxa collected at Millstone infaunal monitoring stations during 2001 and for the overall study period (1980-2001).

|  | 2001 | 1980-2001 | 1980-2001 |
| :---: | :---: | :---: | :---: |
| EFFLUENT | \% | \% | C.V. |
| Oligochaeta | 34.0 | 43.1 | 6.9 |
| Polycirrus eximius | 10.2 | 6.3 | 26.4 |
| Parapionosyllis longicirrata | 8.6 | 2.8 | 30.6 |
| Protodorvillea gaspeensis | 7.3 | 5.1 | 9.5 |
| Tharyx spp. | 4.5 | 4.7 | 40.4 |
| Polydora caulleryi | 4.1 | 0.9 | 33.7 |
| Neanthes acuminata | 2.7 | 0.7 | 24.7 |
| Polydora socialis | 2.3 | 0.4 | 29.1 |
| Capitella spp. | 1.9 | 0.8 | 21.9 |
| Microphthalmus aberrans | 1.9 | 0.6 | 16.5 |
| Mytilus edulis | 0.7 | 2.7 | 84.2 |
| Mediomastus ambiseta | 0.5 | 2.7 | 36.7 |
| Tellina agilis | 0.6 | 2.2 | 16.3 |
| Rhynchocoela | 1.8 | 1.9 | 22.0 |
| Aricidea catherinae | 1.7 | 1.8 | 17.5 |

## GIANTS NECK

| Tharyx spp. |  | 38.5 | 22.9 | 9.4 |
| :---: | :---: | :---: | :---: | :---: |
| Oligochaeta |  | 18.8 | 20.6 | 6.3 |
| Aricidea catherinae |  | 8.4 | 12.1 | 14.7 |
| Polycirrus eximius |  | 6.9 | 2.9 | 16.2 |
| Protodorvillea gaspeensis |  | 3.6 | 3.0 | 7.8 |
| Polydora socialis |  | 3.5 | 0.6 | 31.8 |
| Exogone dispar |  | 3.0 | 2.3 | 10.6 |
| Polydora caulleryi |  | 2.2 | 1.2 | 22.8 |
| Ampharete americana |  | 1.2 | 0.5 | 19.2 |
| Exogone hebes |  | 1.1 | 0.3 | 24.3 |
| Scoletoma tenuis |  | 1.1 | 1.8 | 14.5 |
| Mediomastus ambiseta |  | 0.6 | 7.1 | 24.4 |
| Prionospio steenstrupi |  | 0.7 | 4.4 | 55.2 |
| Ampelisca vadorum |  | 0.3 | 1.7 | 41.7 |

C.V. $=($ Standard Error/Mean $) \times 100$

|  | 2001 | 1980-2001 | 1980-2001 |
| :---: | :---: | :---: | :---: |
| INTAKE | \% | $\%$ | C.V. |
| Aricidea catherinae | 29.7 | 12.3 | 23.5 |
| Oligochacta | 25.6 | 12.6 | 15.2 |
| Exogone hebes | 8.1 | 5.0 | 16.1 |
| Tharyx spp. | 7.4 | 3.9 | 16.5 |
| Protodorrillea gaspeensis | 3.7 | 2.5 | 17.2 |
| Parapionosyllis longicirrata | 2.7 | 0.5 | 29.0 |
| Polydora socialis | 2.1 | 0.4 | 32.7 |
| Polycirrus eximias | 1.7 | 0.5 | 28.7 |
| Owenia fusiformis | 1.1 | 0.8 | 58.8 |
| Euclymene spp. | 1.) | 0.4 | 28.6 |
| Mediomastus ambiseta | 1.0 | 8.6 | 32.0 |
| Prionospio stecnstrupi | 0.3 | 5.0 | 69.7 |
| Leptocheirus pinguis | 0.8 | 4.3 | 70.9 |
| Pygospio elegans | 0.5 | 3.4 | 43.4 |
| Capitella spp. | 0.5 | 3.2 | 22.7 |
| Ampelisca verrilli | 0.3 | 2.9 | 28.3 |

## JORDAN COVE

| Aricidea catherinae | 27.0) | 19.1 | 1.4.3 |
| :---: | :---: | :---: | :---: |
| Oligochacta | 21.5 | 23.9 | 9.4 |
| Scoletoma temuis | 9.8 | 5.0 | 10.3 |
| Tharye spp. | 6.6 | 4.2 | 12.6 |
| Polycirrus eximias | 4.7 | 3.8 | 18.4 |
| Exogone heloes | 3.8 | 1.2 | 25.8 |
| Nucula anmulata | 2.7 | 1.7 | 14.8 |
| Parapionsyllis longicirrata | 2.3 | 1.2 | 28.8 |
| Polydora socialis | 2.1 | 0.3 | 42.2 |
| Protodorvillea gaspeensis | 2.0 | 0.6 | 18.5 |
| Mediomastus ambiseta | 1.7 | 14.4 | 26.8 |
| Leptochcirus pinguis | 0.8 | 4.3 | 32.5 |
| Prionospio steconstrupi | 0.1 | 4.2 | 50.1 |
| Capitella spp. | 1.1 | 1.4 | 16.1 |

C.V. $=($ Standard Error/Mean $) \times 100$




Fig. 9. Mean density of selected taxa in June and September and linear regression of the 1980-2001 period. Vertical line marked with (*) indicates Unit 3 restart in July 1998.


Fig. 9. (comt.)
low throughout the study period, with no apparent trend.

Mediomastus ambiseta - Although not among the ten numerically abundant taxa at any station in 2001 (Table 1), Mediomastus has exhibited periodic pulses in abundance that have placed it among the historical dominants, particularly at JC. Average June and September densities in 2001 were among the lowest values recorded at all stations, and never exceeded 6 individuals/core. Highest M. ambiseta density was at JC (6/core in June and September) and typically averaged between 1-2/core at all other stations (Fig. $9 \mathrm{i}-1$ ). Although low, mean densities during 2001 were within the range of values observed at all stations since 1980. There were no significant long-term trends in abundance at any station over the study period (1980-2001).

Tharyx spp. - Tharyx spp. ranked first in relative abundance in 2001 at GN (38.5\%), fourth at JC (6.6\%) and IN (7.4\%), and fifth at EF (4.5\%) (Tab. 1). Densities during June and September 2001 (Fig. $9 \mathrm{~m}-\mathrm{p}$ ) were highest at GN (114 and 89/core), with lower densities observed at JC (21 and 23/core), IN ( 12 and 9/core), and EF (2 and 11/core). Analysis of long-term data (1980-2001) indicated a significant increasing trend at GN, IN and JC. There was no trend in Tharyx spp. density at EF.

Polycirrus cximius - The polychacte P. eximius was among the dominant taxa during 2001 at EF, GN and JC, ranking second, fourth, and fifth, respectively (Tab. 1). June and September mean densities in 2001 were 14 and $17 /$ core at EF, 17 and 20/core at GN and 4 and $28 /$ core at EF (Fig. $9 q$-s). High annual variability in density has been characteristic of $P$. eximius over the study period at EF and JC, with pulses of abundance observed both before and after Unit 3 startup in 1986 and following the MPS restart in 1998. Densities at GN have been lower and appear to exhibit a gradual increasing tendency. Pulses occurred, but on a smaller scale and coinciding with those observed at EF and JC. The erratic nature of population density data over the entire study period (1980-2001) at each station resulted in no significant long-term trends.

Protodorvillea gaspeensis - P. gaspeensis was among the top ten dominant taxa in 2001 at all stations (Table 1). - P. gaspeensis rankings during 2001 were fourth at EF, fifth at GN and IN, and tenth at JC. June and September 2001 densities were 12 and $10 /$ core at EF, 10 and 9/core at GN, 8 and 3/core at IN, and 6 and 7/core at JC. These densities were all within each station's historical range (Fig. 9t-w).

Trend analysis on the entire time-series (1980-2001) indicated a significant increasing trend in $P$. gaspeensis density at IN and JC. No significant longterm trends were detected at either EF or GN.

Nucula annulata - The bivalve mollusc $N$. annulata was among the infaunal community dominants only at JC in 2001, ranking seventh in relative abundance (Table 1). This species was also among the overall study period dominants at JC, ranking ninth. Data from IN are also included because this species has exhibited increases in abundance concurrent with construction and operation of Unit 3 during the 1984-1990 period. Mean densities of $N$. annulata in June and September 2001 were 7 and 11/core at JC, and 1/core at IN (Fig. $9 x-y)$. These density estimates were within the range of estimates from previous years. Trend analysis of the 22 -year time-series revealed a significant increasing trend in $N$. annulata density at JC, but no significant trend at IN.

Parapionosyllis longicirrata - P. longicirrata was among the dominant taxa at EF and JC in 2001, ranking third and eighth in relative abundance, respectively (Table 1). Abundance of $P$. longicirrata, based on mean density during June and September 2001, respectively, were 23 and $3 /$ core at EF, and 9 and 7/core at JC. Densities at both stations appear to have decreased in 2001, relative to high levels noted in recent past years; however, analyses of the 22 -year time-series indicate a significant increasing trend at both EF and JC.

## Multivariate Analyses

Several infaunal taxa exhibited temporal trends in abundance associated with construction and operation activities at MPS. Trends in the abundances of other taxa were common throughout the greater MPS area, but were unrelated to the power plant. Each species represents one component of a complex community, affected by many biological and physical factors. To provide a more complete characterization of local infaunal communities, multivariate Bray-Curtis similarity indices were calculated for each pair of annual collections, using all species whose abundance averaged at least two individuals per year. Relationships of annual collections based on comparisons of these indices' are illustrated as a clustering dendrogram and as a two-dimensional representation of a multi-dimensional scaling plot for each station (Fig. 10).


Fig. 9. (cont.)


Fig. 9. (cont.)


Fig. 9. (cont.)

## Cluster Analysis

The clustering dendrogram based on EF data shows a similarity among collections from the 22 study years of approximately $65 \%$ (Fig. 10at). Five groups of annual collections were apparent at the $70 \%$ or greater similarity level. Group I consisted of collections from 1980-1985, and these years distinguished themselves from all or most other groups with high densities of the polychacte taxa Polycirrus eximius, Tharyx spp., Eumida sanguinea, and Exogone hebes, and the molluse Tellina agilis. The next three years (1986-1988) composed Group II, and were characterized by lower abundances of $P$.
eximios, Tharnx spp. and dricidea catherinae, and high abundances of oligochaetes, the blue mussel Mytilus edulis, and the anemone Maliphanella luciae. Group II collection years coincided with the period of scour-related sediment changes associated with Unit 3 stat-up and early operational years. Group III included annual collections from 1993 through 1996. This group was characterized by moderate abundances of a number of polychaete species (c.g., A. catherinae, Parapionossllis longicirrata, and Protodonvillea gaspeensis), and low densities of Mcdiomastus ambiscta and $P$. eximius. Representatives from the class Rhynchocoela were most abundant during this period. Group IV


Fig. 9. (cont.)





Fig. 9. (cont.)
consisted of annual collections made from 1997 through 1999, a period which encompasses the extended shutdown of MPS. The EF infaunal community during these years was characterized by high abundances of the polychaetes $P$. longicirrata, M. ambiseta, A. catherinae, P.eximius and Pygospio elegans, and the amphipod Ampelisca vadorum. Collections from the last two years (2000 and 2001) along with those from the period 1989-1992 were included in Group V. This group was characterized primarily by high abundances of $P$. eximius, $P$ : gaspeensis, and Prionospio steenstrupi, in addition to the highest abundance of Mytilus edulis and lowest abundance of the M. ambiseta.
Overall similarity among annual collections at the reference site. (GN) was highest of any station at $>70 \%$ (Fig. 10b). Group I years (1981-1982) were characterized by high densities of Aricidea catherinae and the amphipod Gammarus lawrencianus and low abundances of oligochaetes, Mediomastus ambiseta, Exogone dispar, and the amphipod Ampelisca vadorum. Collections from 1983 through 1986 and 1980 compose Group II, and were characterized by high densities of $M$. ambiseta, Polydora caulleryi, and the amphipod Phoxocephalus holbolli and low abundances of oligochaetes. Group III (1987-1991) was distinguished from other groups by high abundances of the amphipods Leptochierus pinguis and A. vadorum, concurrent with low abundances of Tharyx spp. Annual collections from 1992-1994 formed Group IV. During these years, the abundance of the polychaetes Prionospio steenstrupi and M. ambiseta along with $A$. vadorum were at the highest of any grouping. Group V contained annual collections from 1995-1999 and the infaunal community during these years was characterized by moderate abundances of most taxa relative to other groupings. The most recent years (2000-2001) formed Group VI which was characterized by high abundances of Tharyx spp. Polycirrus eximius, $P$. gaspeensis, Polydora socialis and P. caulleryi, and the lowest abundance of M. ambiseta.
High infaunal community variability resulted in relatively low overall similarity ( $<60 \%$ ) among the four groups of annual collections identified at IN (Fig. 10c). Initial study years included in Group I (1980-1983) had collections with generally low abundances of most infaunal taxa, with the exception of moderate numbers of Aricidea catherinae and oligochaetes, and high abundance of maldanids relative to other groups. Unit 3 intake dredging, cofferdam removal and startup all occurred during the next four study years (1984-1987) that comprise Group II. This group is distinguished from others by high abundance of amphipods (Leptocheirus pinguis, Ampelisca verrilli," Ampelisca vadorum, and

Ampelisca abdita)' and the polychaetes Mediomastus ambiseta, Owenia fusiformis and Polydora cornuta, and low abundance of A. catherinae and oligochactes. Abundances of all amphipods declined during Group III study years (1988-1991), as populations of $A$. catherinae and oligochactes rebounded and highest abundances were observed for the polychaetes Exogone hebes, Capitella spp., and Polydora quadrilobata, and the mollusc Nucula annulata. Collections from the last 10 years (1992-2001) formed Group IV. High abundances of oligochaetes and a number of polychaete species (e.g., A. catherinae, M. ambiseta, Prionospio steenstrupi, -Pygospio elegans, Tharyx spp., and Protodorvillea gaspeensis) were observed during this period.
.. Collections from JC clustered into three groups at about a $65 \%$ similarity level (Fig. 10d). Group I collection years (1980-1986) were characterized by the highest density of oligochactes and the polychaetes Mediomastus ambiseta and Polydora caulleryi, and low abundance of Aricidea catherinae, Prionospio steenstrupi, Leptocheirus pinguis, Nucula annulata and juveniles of the polychaete family Maldanidae. Group II (1987-1993) collections were distinguished by higher densities of $P$. steenstrupi, $L$ pinguis, Polycirrus eximius and N. annulata, and by lower oligochaete and Mediomastus ambiseta densities. Group III was made up of collections from 1994 through 2001. This group of 8 years had highest densities of the polychactes A. catherinae, Mediomastus ambiseta, and Tharyx spp., and $P$. eximius densities. Several other taxa (e.g., Maldanidae, Protodorvillea gaspecensis, Exogone hebes and Parapionsyllis longicirrata) were also abundant during this period.

## Multi-Dimensional Scaling (MDS)

The MDS plots included in Figure 10 illustrate temporal community shifts and groupings of annual collections that are generally consistent with their associated clustering dendrograms. However, these two-dimensional plots are more amenable to multiple among-year comparisons, are more sensitive to less :abundant taxa, and reveal the degree of community shifts based on distance between points representing annual collections. For example, at EF , there are three island groupings that correspond to pre-Unit 3 years ( $1980-1985$ ), the initial three years following Unit 3 start-up, (1986-1988), and a loose grouping of all other years including the period of extended shutdown (1997-1999). Examination of spatial relationships among individual points in the MDS plot for EF reveals that the largest year to year shift occurred between 1985 and 1986, when increased water flow from Unit 3 start-up scoured the discharge


Fig. 10. Clustering dendrogram of natural log-transformed density data of annual infanal commonities and list of taxa (mean density/year) associated with major group divisions in classification and cluster analyses and Multidimensional Sealing two dimensional plots for the years 1980-2001.


| Taxon | Group I 1981-1982 | $\begin{gathered} \text { Group II } \\ 1983-1986 \\ 1980 \\ \hline \end{gathered}$ | Group III 1987-1991 | $\begin{aligned} & \text { Group IV } \\ & \text { 1992-1994 } \end{aligned}$ | $\begin{aligned} & \text { Group V } \\ & \text { 1995-1999 } \end{aligned}$ | $\begin{gathered} \text { Group VI } \\ 2000-2001 \end{gathered}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |
| Tharyx spp. | 1011 | 1322 | 740 | 1396 | 1325 | 1729 |
| Oligochaeta | 963 | 985 | 1090 | 1139 | 1096 | 1041 |
| Aricidea catherinae | 1677 | 834 | 917 | 328 | 720 | 372 |
| Mediomastus ambiseta | 35 | 715 | 263 | 400 | 342 | 72 |
| Prionospiosteenstrupi | 160 | 77 | 316 | 1306 | 323 | 24 |
| Polycirrus eximius | 66 | 146 | 142 | 75 | 171 | 405 |
| Protodorvillea gaspeensis | 143 | 182 | 201 | 143 | 167 | 204 |
| Exogone dispar | 48 | 135 | 114 | 145 | 123 | 147 |
| Ampelisca vadorum | 6 | 29 | 218 | 246 | 115 | 124 |
| Scoletoma tenuis | 147 | 137 | 152 | 81 | 91 | 10 |
| Phoxocephalus holbolli | 101 | 138 | 86 | 46 | 77 | 8 |
| Leptocheirus pinguis | 14 | 39 | 297 | 8 | 69 | 36 |
| Polydora caulleryi | 53 | 111 | 51 | 16 | 68 | 153 |
| Polydora socialis | 5 | 20 | 16 | 46 | 46 | 176 |
| Capitella spp. | 38 | 44 | 62 | 50 | 50 | 24 |
| Mitrella lunata | 30 | 27 | 15 | 90 | 37 | 45 |
| Rhynchococla | 23 | 42 | 36 | 50 | 35 | 34 |
| Polydora quadrilobata | 16 | 38 | 68 | 57 | 34 | 6 |
| Ampharete americana | 11 | 14 | 27 | 60 | 34 | 65 |
| Harmothoe imbricata | 20 | 35 | 25 | 61 | 33 | 11 |
| Gammarus lawrencianus | 110 | 54 | 14 | 0 | 33 | 17 |
| Pagurus acadianus | 26 | 28 | 55 | 28 | 32 | 14 |



Fig 10 (con't.)

| taxon | Group I | Group II | Group III | Group IV |
| :---: | :---: | :---: | :---: | :---: |
|  | $1980-1983$ | $1984-1987$ | $1988-1991$ | $1992-2001$ |


| Aricidea catherinat | 1.57 | 30 | 180 | 671 |
| :---: | :---: | :---: | :---: | :---: |
| Leptochicrus pinguis | 13 | 894 | 46 | 67 |
| Oligochaeta | 1.58 | 72 | 216 | 574 |
| Mediomastus ambiseta | 13 |  | 24 | 380 |
| Prionospio steenstrupi | 27 | 42 | 89 | 359 |
| Eadgone helos | 71 | 19 | 185 | 151 |
| Pygospior elagrans | 14 | 16 | 60 | 266 |
| Tharyespp. | 66 | 53 | 83 | 1.18 |
| C'apitella spp. | 73 | 61 | 1.51 | 38 |
| Nucula annalata | 14 | 12610 | 1.45 | 27 |
| Amprelasca verrilli | 61 | 1711 | 37 | 21 |
| Ampelasca vadoriam | 8 | 14.5 | 12 | 11.3 |
| Tellina agilis | 56 | 48 | 8.4 | 70 |
| Ampelisca abdita | 9 | 18.5 | 5 | 1) |
| Probodorvillea graspeensis | 2.5 | 6 | 46 | 122 |
| Mahdandac | 146 | 1 | 13 | 3.3 |
| Unciola irrorata | 8 | 90 | 31 | 44 |
| Polydura atuadrildoatas | 24 | 1.5 | 105 | 14 |
| Owenia fastormis | 4 | 112 | 26 | 14 |
| P'olydera cornata | 4 | 112 | 18 | 11 |
| Giammaras lawrencianas | 52 | 3 | 43 | 15 |
| l'asurus lonsicarpus | 2 | 97 | 0 | 0 |
| Spiophantes bombyr | 18 | 14 | 25 | 19 |
| Shicrophtalmus aberrans | 4 | 21 | 25 | 20 |



Fig IO. (con't.)

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| Taxon | $\begin{gathered} \text { Group I } \\ 1980-1986 \\ \hline \end{gathered}$ |  | $\begin{gathered} \text { Group II } \\ 1987-1993 \\ \hline \end{gathered}$ |  | $\begin{gathered} \text { Group III } \\ \text { 1994-2001 } \end{gathered}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Oligochaeta | 2365 |  | 1400 |  | 1456 |
| Aricidea catherinue | 795 |  | 1443 |  | 2385 |
| Mediomastus ambiseta | 1707 |  | 649 | . | 1636 |
| Prionospio steenstrupi | 65 |  | 973 |  | 132 |
| Leptocheirus pinguis | 53 |  | 680 |  | 358 |
| Scoletomatenuis | 349 |  | 414 |  | 311 |
| Tharyx spp. | 282 | $\cdots$. | 227 |  | 517 |
| Polycirrus eximius | 273 |  | -. 457 |  | 190 |
| Nucula annulata | 38 |  | . 194 |  | 145 |
| Capitella spp. | 102 |  | 134 |  | 74 |
| Tellina agilis | 117 | ; | 75 |  | 118 |
| Exogone hebes | 36 | , | $\because 42$ |  | 183 |
| Parapionosyllis longicirrata | 48 |  | - 28. |  | 186 |
| Microphthalmus aberrans | 44 |  | 87 |  | 90 |
| Maldanidae | 2 |  | 26 |  | 172 |
| Polydora caullergi | 140 | - | 12 |  | 13 |
| Protodorvillea gaspeensis | 13 |  | $\therefore 28$ | : | 92 |
| Phyllodace mucosa | 33 |  | - 36 |  | 56 |
| Mitrella lunata | 52 |  | . 38 |  | 34 |
| Rhynchocoela | 46 |  | 50 |  | 27 |



Fig. 10 ( con't $^{\prime}$ )
area and resulted in sudden sedimentological changes in the discharge area and infaunal shifts described for Group Il of the dendrogram. The largest grouping represents remaining years (1989-2001), when an altered sediment regime was maintained by chronic discharge llow conditions. This stabilization allowed rebounds of certain populations impacted by initial scouring effects, including Polycirrus eximius and Aricidea catherinae, and a decline in oligochacte abundance to levels similar to pre-scouring years.
Annual collections in the GN MDS plot show litle evidence of distinct groupings (Fig. 10b). The orientation of annual collections reflect subtle, natural changes in community composition described in the clustering dendrogram. A gradual temporal community shift is apparent in the MDS plot attributed primarily to long-term increases of Tharyx spp. Exogone dispar, Protodorvillea gaspeensis, Polycirrus eximitus and oligochactes, and a concomitant decrease of Aricidea catherinae.
The MDS plot of annual IN collections (Fig IIc), illustrates high variability in community structure during the period from 1980-1990. This period was marked by substantial community changes as a result of disturbances from Unit 3 intake construction and dredging activities, as discussed in the previous section. The greatest community shift, based on the MDS plot, occurred between 1983 and 1984, when Unit 3 intake colferdam removal and dredging ativities were conducted. The years inmediately following these activities (1984-1987) form a grouping in close proximity to each other with a low similarity to other annual collections, consistent with Group Il of the cluster dendrogram. Colletions from 1988-1990 were considered a transition pathway with taxa such as Aricidea catherintae showing increases over previous years and lower abundances of amphipods. The MDS plot shows a difference in the association of the 1991 collection from that of the cluster analysis due in parn to the increases in $A$. catherinac. All remaining years (1091-2001) form a distinct, relatively cohesive island and demonstrate the post-disturbance community, with a trend toward more consistent community structure indicative of stability and recovery.

Scouring in the discharge area after Unit 3 start-up resulted in silt redistribution that had a strong effeet on the JC infaunal community, as described above. The magnitude of community changes from year to year at JC are illustrated in the MDS plot (Fig. 10d), and were greatest between 1986 and 1987 when large increases in sediment siltelay content were observed (Fig. 2). Two island groupings shown in the MDS plot (1980-1985 (less 1981) and 1987-2001) reffect the shift in the JC community state caused by siltation. Some of these shifts were first ohserved
during the second half of 1986, which is a distinct outlier in the MDS plot.

## Discussion

Monitoring studies of benthic infaunal communities inhabiting marine sediments in the MPS vicinity have documented both long-term trends and short-term changes in community and population level parameters and sedimentary characteristics. This 22year database has provided evidence of impacts from acute and chronic disturbance events related to MPS construction and operation, along with assessment of the degree to which these disturbances have affected or continue to affect each station in relation to factors unrelated to station operation. Infaunal communities exhibiting evidence of MPS inpacts were observed at the stations in the immediate vicinity of MPS (IN, EI and JC). Community changes at these sites were closely associated with changes in sediment composition related 10 power plant-induced disturbance. The infaunal community at GN reference site was unaffected by plant operation and provided a baseline from which variability associated with natural envirommental cycles and disturbances could be assessed.
The GN station, which is not intluenced by MPS has exhibited changes in sediments and community structure over the entire stady period. Sediment mean grain sife was most variable at this site, but silt/clay content was least variable and no long-term trends were apparent for ejher parameter. White the GN infaunal community has generally been domimated by three taxa over the study period (Tharys spp., oligochactes and Aricidea cotherinate), some longlerm trends in abundance of these taxa cominued through the 200). Tharyx spp. abundance has continued to increase over the 22 -year study period. while A. catherinac, abundant in early monitoring years, declined. Some species at (aN have exhibited relatively high year-to-year fluctuations. In many cases, these fluctuations proved useful in validating area-wide shifts in species abundance and community structure in response to matural causes and unrelated 10 MIPS operation. For example, there was a pulse in abundance of the polychate Polydora socialis at all stations in 2001, indicating a regional phenomenon. Other similar population shifts inchude several areawide increases of the opportunistic polyehacte Mediomastus ambiseta (NUSCO 1989, 1996, 1999) and large increases in abundance of the amphipods Leptocheirus pinguis and Ampelisce spp., and of the polychactes Prionospio stcenstrupi in 1992 and Pygospio elegans in 1998 (NUSCO 1993, 1999). These increases could not be explained by changes in
site-specific sedimentary or regional abiotic factors (NUSCO 1989, 1993, 1999). . However, because these changes were all observed at reference site GN, their cause was determined to be independent of power plant construction or operation.
Physical disturbances initiated community changes at IN and JC in the period between 1980-1987. Prior impacts were noted from 1983 to 1985 at IN, and resulted from dredging and cofferdam removal during Unit 3 construction (NUSCO 1987). Sedimentary characteristics at IN, :(primarily sil//clay :content levels) have stabilized and become more consistent between sampling periods. - Evidence of some infaunal community stabilization and recovery was also apparent. In particular, numbers of individuals and species richness at IN have increased over the study period, as have abundances of organisms typically more common in early study years or at other sites, such as Aricidea catherinae and Tharyx spp.
These trends, along with concomitant decreases in abundance of Nucula annulata and other opportunistic species (e.g., the amphipods Leptocheirus pinguis and Ampelisca spp.), indicate some degree of recovery at IN. Recovery at IN is not complete, however, as other organisms that established post-impact community dominance, such as oligochaetes and Protodorvillea gaspeensis, maintained or increased their degree of dominance in recent years and may persist indefinitely. Long-term recovery following disturbance is typical of marine benthic communities (Kaplan et al. 1974; Swartz et al. 1980; Nichols 1985; Berge 1990; Zajac et al. 1998).

Silt deposition at JC attributed to sediment scouring from the MPS discharge area following Unit 3 startup in 1986 resulted in increased silt/clay content and abrupt infaunal community changes. The silt deposition and concurrent large infaunal community changes occurred over a short period at the beginning of the 3 -unit operational period.
The altered infaunal community has exhibited changes in recent years moving toward pre-impact community structure. For example, A. catherinae and Tharyx spp. abundances both advanced to levels observed prior to 1986 within a few years of Unit 3 start-up. However, continued trends toward recovery noted in previous years (e.g., reduced silt/clay content during 1994 and 1995, and rebounding abundances of ; oligochaetes and $P$. eximius through 1993; NUSCO 1994, 1995) were short-term habitat or community changes not reflected in 2001 data. Additionally, the opportunistic mollusc Nucula annulata maintained a population at JC through 2001 that remains well above pre-Unit 3 levels. This biological evidence of disturbance is consistent with trends in sediment
characteristics; elevated silt/clay levels in sediments at JC persisted through 2001. These observations are similar to those of other researchers studying the effects of siltation on benthic infaunal communities (Rhoads and Young 1970; Jumars and Fauchald 1977; Turk and Risk 1981; Maurer et al. 1986; Emerson 1989; Brey 1991; Currie and Parry 1998).
The benthic infaunal site that still experiences active MPS operational impact processes is the EF station, located in the path of the discharge 100 m from shore. Current scour from the MPS discharge modifed sediments and the infaunal community at EF, immediately after Unit 3 startup in 1986. Relatively coarse sediment with low sil/clay levels were characteristic of the 3 -unit benthic habitat at EF .
While sediment characteristics remain different from those observed prior to 1986, the altered sedimentary environment at $E F$, and the infaunal community it supports, stabilized under the new environmental conditions created by the 3 -unit discharge. Even though Unit 1 has been permanently retired, the discharge from Units 2 and 3 represents almost $80 \%$ of the 3 -unit flow. The relative stabilization of the sediments at EF allowed for rebounds of Tharyx spp. and Aricidea catherinae, taxa common during 2 -unit operation. In addition, oligochaete and Protodorvillea gaspeensis abundances have generally decreased from high abundances during the early 3 unit period. Oligochaete abundances at EF during the last two years (2000 and 2001) were among the lowest observed during the study period, possibly related to the heavy mussel set observed in the discharge area in 2000 (also noted on nearby rocky shores; see Rocky Intertidal section of this report).
High mussel settlement is likely the result of conditions created by the MPS discharge, (current velocities provided more food and substrate was amenable to mussel settlement) as no similar settlement was observed on sediments at the other monitoring sites. High mussel cover on sediments has been shown to have a strong influcnce on infaunal. community structure and composition (Ragnarsson and :Raffaelli 1999). Other indications of sediment stabilization at EF are population increases consistent with regional pulses of populations discussed above (e.g., Mediomastus ambiseta, Polycirrus eximius, and Polydora socialis). During earlier years of 3 -unit operation, because natural regional factors were overridden by plant discharge effects, region-wide pulses in the above taxa had not been observed at EF.
Changes observed in community structure at impacted stations could define permanent shifts in the dynamics of benthic communities in the vicinity of MPS. Shifts in benthic community structure at EF were similar to those caused by commerical scallop dredging in Port Phillip, Australia, which were noted
over a 20-year period (Curric and Parry 1998). However, establishing which components of benthic community structure are due to natural cycles or maninduced disturbance is difficult because the length of natural cycles is unpredictable (Gray and Christie 1983). Additional monitoring will determine if these trends are indicative of current operating conditions (Units 2 and 3 only), or are simply short-term fluctuations related to other factors.

## Conclusions

Benthic habitats and their associated infaunal communities at IN, JC and EF Juring 2001 continue to exhibit patterns in community structure typical of disturbance related to construction and operation of Unit 3. While sediment grain size exhibited the highest variability at the unimpacted site GN, silt/clay content and infaunal community structure at this site exhibited highest stability over the study period relative to the oller monitoring sites. Some indications of increased stability and limited community recovery have been observed in recent ${ }^{-}$ years at stations impacted by short-term episodic disturbance events (e.g., dredging and construction activities at IN , and siltation at JC). This recovery is ongoing but progression is slow. Community parameters of species composition and population abundance show similarities to the pre-disturbance period, but also clearly show differences related to past disturbance at both stations. Data from the recent two-year shutdown period (1996-1998) and from the years following restart provided additional evidence that the MIPS discharge has been a dominant factor in structuring both the sedimentary environment and infamal community in the immediate vicinity of the discharge. In the absence of discharge effects, the EF community exhibited detectable shifts in some populations that had not been seen during 3 -unit operation. Aside from these population changes, overall recovery of the community and sedimentary enviromment during this period was limited, indicating that post-operational recovery will be slow. Reversal of most trends ohserved during the shutdown period were noted soon after MIPS restart, owing to full operation of Units 2 and 3 for much of the last three years.

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## Eelgrass

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## Introduction

Eelgrass (Zostera marina L.) is the dominant marine angiosperm in temperate coastal regions of the Northern Hemisphere (Setchell 1935; Thayer et al. 1984). Extensive eelgrass meadows found in shallow estuaries and lagoons provide multi-level ecological benefits that make them key components, to many coastal marine systems. High eelgrass primary productivity is utilized directly by many consumers including numerous waterfowl, fish and invertebrate grazers (see review by Valentine and Heck 1999), but much of the production enters the foodweb through the detrital pathway (Thayer et al. 1984). Increased habitat complexity within eelgrass meadows enhances primary and secondary production and species diversity in coastal ecosystems. Eelgrass meadows support productive epiphytic communities (Thayer et al. 1984; Nelson and Waaland 1997), and provide feeding and nursery grounds for many species, including commercially important finfish and invertebrates (Orth 1973; Heck et. al. 1989, 1995; Gotceitas et al. 1997; Mattila et al. 1999; Short et al. 2001).

Historical reports from eastern Long Island Sound (LIS) of eelgrass populations (e.g., Marshall 1947, 1960, 1994; Koch and Beer 1996) describe fluctuations in abundance and distribution over the last century similar to those observed elsewhere in the western North Atlantic Ocean (Tutin 1942; Rasmussen 1977; Orth and Moore 1983; Short et al. 1996). Populations along the coastline of Connecticut were almost completely eliminated by 1933. Some recovery occurred by the late 1950 s, and by the early 1970s celgrass beds in southeastern Connecticut were again extensive (Knight and Lawton 1974; McGill 1974; Marshall 1994). .: Since the early 1980s, however, eelgrass populations in Connecticut and elsewhere have declined (e.g., Orth and Moore 1983; Short 1988; Koch and Beer 1996; Short et al. 1996). Because of the ecological importance and high historical variability of eelgrass, and the prediction that the 3-unit thermal plume from Millstone Power Station (MPS) could reach to the nearby populations in Jordan Cove (Knight and Lawton 1974; NUSCO 1988), the present study to monitor local eelgrass populations was initiated in 1985. Objectives of the present study are to identify temporal patterns" of eelgrass population characteristics in the vicinity of MPS and to determine the extent to which changes in these patterns are the result of natural variability or MPS operation.

## Materials and Methods

Three eelgrass study sites in the vicinity of MPS were sampled during 2001 (White Point-WP, Jordan Cove-JC, and Niantic River-NR; Fig. 1). The WP and JC stations, located 1.6 km and 0.5 km east of the power plant discharge, respectively, were within the area potentially influenced by the 3 -unit thermal plume (NUSCO 1988). The NR sites, located about 3 km from Millstone Point, are unaffected by power plant operation (Fig. 1). Water depths (at mean low water) were 2.5 m at WP, 1.5 m at NR and 1.1 m at JC. The WP and JC sites have been sampled since 1985. A new sampling site was established in the Niantic River in 2000 (NR5) due to bed die-off at the 1999 site (NR4). Location changes and periods of sampling for all Niantic River sites are described in Figure 1.
Samples were collected monthly at each site from June through September, the period of maximum standing stock and plant density. At each station, 16 samples were collected by SCUBA divers from randomly placed quadrats ( $25 \times 25 \mathrm{~cm}, 0.0625 \mathrm{~m}^{2}$ ) within a 10 m radius of the station marker. The upright shoots from plants within each quadrat were harvested, placed in a 0.333 mm mesh bag, returned to the laboratory and analyzed. A 3.5 cm diameter x 5 cm deep core was taken concurrently with eelgrass samples for analysis of sedimentary characteristics at each station. Water temperature at the JC site was measured by an, encased thermistor-recorder, suspended just above the sediment surface. Temperature measurements have been recorded in Jordan Cove since 1991.
All shoots collected were counted in the laboratory and the longest blade of each shoot (up to 20 per sample) was measured to the nearest centimeter. The number of reproductive shoots in each sample was used to estimate the percentage of reproductive shoots in the population. Shoots were rinsed in freshwater to remove invertebrates and epiphytes. Eelgrass standing stock was estimated as the weight of the shoots taken from each quadrat. From 1985 to 1987, shoots were weighed, then dried in an oven at $80^{\circ} \mathrm{C}$ to constant weight. Dry weights from 1988 to 2001 were estimated from the wet-weight/dry-weight relationship obtained above.
Linear regression methods were used to examine long-term trends in eelgrass shoot density, shoot length and standing stock. These methods were applied to all the data,' except for outliers identified by the SAS (1999) univariate procedure, and removed prior to analysis. This analysis was not


Fig. 1. Map of the Millstone Point area, showing the location of eclgrass sampling stations: JC=Jordan Cove, NR=Niantic River ( $1=$ sampled 1985.Junc 1986 and 1993-1994, $2=$ sampled July-5 eptember 1986, $3=$ sampled 1987-1992. 4=sampled 1995-1999. $5=$ sampled $20(0)-2(0) 1)$, WP=White Point.
conducted on Niantic River data due to the lack of a continuous time-series at any given sampling location resulting from periodic sampling bed die-off.

Mean sediment grain size and silt/clay content were determined using the dry sieving method (Folk 1974). Sediment samples were heated to $500^{\circ} \mathrm{C}$ for 24 h to determine organic content, estimated as the difference between dry-weight and ash-weight. Both silvelay and organic content were recorded as a percentage of the total sediment sample weight.

## Results

## Temperature

Average daily seawater temperatures at the MPS intakes and discharge, and at the JC sampling station, during June through September 2001 are presented in Figure 2. With both Units 2 and 3 operating at full power, eflluent temperatures were as high as $10-11^{\circ} \mathrm{C}$ above intake and JC temperatures during this period. Peak daily average temperature occurred on August

31 at all three sites $\left(22.5^{\circ} \mathrm{C}\right.$ at the intake, $23.0^{\circ} \mathrm{C}$ at JC , and $33.4^{\circ} \mathrm{C}$ at the discharge). Daily average seawater temperatures at the JC eelgrass station were generally warmer than ambient temperatures by 1-2 ${ }^{\circ} \mathrm{C}$ through August. At times, seawater temperatures were up to $4-5^{\circ} \mathrm{C}$ warmer at JC during afternoons on sunny days.

## Sediments

Monthly measures (June-September) of mean grain size, silt/elay and organic content have been used to describe eelgrass monitoring sites since 1985 (Fig. 3). Sediments at stations nearest MPS (JC and WP) have shown a high degree of stability relative to the variable nature of those at NR. Much of the variability at NR has been attributed to frequent relocation of the sampling site, the patchy distribution of eelgrass within the Niantic River, and subsequent exposure to water currents and sellement of sediment finc fractions.

Sediments collected during 2001 were coarser at JC (mean grain size $0.24-0.28 \mathrm{~mm}$ ) than those at WP (0.13-0.16 mm) or NRS (0.06-0.1.4 mm). Sediment


Fig. 3. Mean grain size, organic content and sil//clay content of sediments at MPS eelgrass stations, Jordan Cove (JC), Niantic River (NR) and White Point (WP), sampled during the period June-September from 1985 through 2001.


NIAJJTIC RIVER


WHI"E POINT


Fig. 4. Monthly mean shoot density ( $\pm 95 \%$ C.I.) and annual mean density at ecigrass sites.
organic content in 2001 was higher at NR5 (5.97.9\%) than at WP (1.5-3.2\%) or JC (0.9-2.1\%). Silt/clay content in 2001 was highest at NR5 (21.9$44.9 \%$ ), intermediate at WP (5.6-9.9\%), and lowest at JC (2.3-4.1\%). Sediment parameter estimates in 2001 were within historical ranges established for each station since 1985.

## Shoot Density

Annual mean shoot density (shoots $\mathrm{m}^{-2}$ ) among eelgrass stations in 2001 was similar to previous years; highest at JC (714), intermediate at WP (200) and lowest at NRS (139; Fig. 4). While shoot densities at all stations in 2001 were within historical ranges, mean annual density at NR5 remained among the lowest observed at any Niantic River site since 1985. Monthly shoot densities in 2001 ranged from 655 (July) to 817 (June) at JC, 113 (September) to 281 (August) at WP, and 107 (September) to 208 (July) at NR5. Monthly mean densities at all stations were within the ranges of previous years. Linear regression analysis of shoot density over the entire time-series indicated no significant trend at JC or WP. Although the regression analysis was not performed, there was an obvious decline in shoot density in the Niantic River.

## Shoot Length

Annual mean shoot lengths for 2001 were longest at WP ( 90 cm ), shortest at NR5 ( 46 cm ), and intermediate at JC ( 52 cm ; Fig. 5). Annual means at all stations were within historical ranges observed since 1985. Maximum monthly shoot lengths in 2001 were highest in July and August at JC ( 61 cm ), in June and August at NR5 ( 49 cm ), and in June at WP ( 105 cm ). Linear regression of shoot length data over the entire time-series indicated significantly decreasing slopes at JC ( $\mathrm{n}=1083$; slope $=-0.08 \mathrm{~cm}$ month ${ }^{-1}$; $p<0.0001$ ) and WP ( $n=1064$; slope $=-0.08$ cm month ${ }^{-1} ; p<0.0001$ ).

## Standing Stock

Eelgrass standing stock (g dry wt $\mathrm{m}^{-2}$ ) annual mean for 2001 was higher at JC (215) than at WP (154) or NR5 (43; Fig. 6). These annual standing stock estimates were within the historic ranges. Monthly standing stock estimates in 2001 ranged from 164 (August) to 277 (July) at JC, from 68 (September) to 221 (August) at WP, and from 23 (September) to 62 (July) at NR5. Linear regression analysis of standing stock data over the entire time-series indicated a significantly decreasing trend at JC ( $\mathrm{n}=1015$; slope $=$ $-0.60 \mathrm{~g} \mathrm{~m}^{-2}$ month $^{-1} ; \mathrm{p}<0.001$ ) and WP ( $\mathrm{n}=998$;
slope $=-0.17 \mathrm{~g} \mathrm{~m}^{-2}$ month $^{-1} ; p=0.013$ ). A decline in standing stock estimates was also apparent in the Niantic River over the study period; however, linear regression was not performed on data from these populations due to the lack of a continuous timeseries at any given sampling location resulting from periodic sampling bed die-off.

## Seed-Bearing Shoots

Annual and monthly numbers of seed-bearing shoots collected at each station are presented in Figure 7. Total seed-bearing shoot abundance in 2001 was highest at JC (70) followed by NR5 (42) and WP (27). Seed bearing shoots in 2001 were found in June through August at all stations, with the highest number of shoots occurring in June ( 41 at JC, 25 at NR, and 19 at WP). While some seed-bearing shoots were collected in August 2001, there has been a general regional trend over time that seed-bearing shoots have occurred less frequently during late summer. Seed-bearing shoots were consistently observed in samples collected from all sites in August and September prior to 1991, but were rare in those months in subsequent years.

## Discussion

Eelgrass populations monitored near MPS exhibited considerable within- and among-year variability in population parameters at all three study locations. While relatively minor, but statistically significant, declines in eelgrass shoot length and biomass were observed at sites nearest the MPS thermal plume (JC and WP), these eelgrass populations persisted as healthy beds over the course of this study. In contrast, eelgrass beds in the Niantic River (NR 1-5), located well outside the thermal plume area, experienced the largest declines in population characteristics of any site.
Researchers have shown that increased water temperatures (e.g., from heated power plant effluents) eliminated eelgrass from nearby areas (Phillips 1974; Thayer et al. 1984). Similarly, decline of the seagrasses Thalassia testudinum and Halodule wrightii in Florida (Roessler and Zieman 1969; Wood et al. 1969; Zieman 1970; Roessler 1971; Blake et al. 1976; Thorhaug et al. 1979) and the marsh grass Spartina alterniflora in Maine (Keser et al. 1978) were directly attributed to temperature increases from power plant effluents. However, effluent-related water temperature increases experienced by populations in the above-cited studies were considerably higher than those observed in this study.



WHITE FOINT


Fig. 5. Monthly mean shoot length ( $\pm 95 \%$ C.I.) and amual mean leng'h at eclgrass sites



WHITE POINT


Fig. 6. Monthly mean standing stock biomass ( $\mathbf{9 5 \%}$ C.I.) and annual mean biomass at eelgrass sites


Fig. 7. Total number of seed-tearing shoots collected monthly and annually at MPS eelgrass sites.

Daily temperature fluctuations of up to $5^{\circ} \mathrm{C}$ were measured at our JC study site during mid-day in summer, and were most pronounced on sunny days at low tide. This diurnal pattern during summer
suggests that increases in seawater temperature at this site is primarily attributed to solar warming. Any thermal input to JC from the MPS cooling water discharge ( $<l^{\circ} \mathrm{C}$; NUSCO 1988) is difficult to
distinguish from natural fluctuations, suggesting very limited thermal effects.
This study provides evidence of long-term trends in eelgrass population characteristics from a region that has experienced widespread declines over the last two decades. Similar trends of eelgrass populations have been documented elsewhere in eastern North America, including the Chesapeake Bay (e.g., Orth 1976; Orth and Moore 1983, 1986; Moore et al. 1996, 1997; Moore and Wetzel 2000) and coastal embayments of Rhode Island (Harlin and ThorneMiller 1981; Thorne-Miller and Harlin 1984; Short et al. 1996), Massachusetts (Dexter 1947, 1985; Roman and Able 1988; Short and Burdick 1996) and New Hampshire (Riggs and Fralick 1975; Short et al. 1986). This study complements the few reports available on temporal trends of eelgrass in LIS (Dexter 1946; Marshall 1947, 1960, 1994; Knight and Lawton 1974; Koch and Beer 1996). These reports indicate that eelgrass populations in LIS have fluctuated since the widespread die-off in the 1930s, similar to long-term trends reported elsewhere in eastern North America. Several reports note that eelgrass populations in the vicinity of MPS had experienced considerable recovery by the 1970s (Knight and Lawton 1974; Marshall 1994), with a gradual decline beginning in the 1980s (Vozarik et al. 2000; DNC 2001). Eelgrass distribution along the north shore of LIS once reached as far west as Westchester County, NY, but now is limited to the easternmost third of the Connecticut coastline (Koch and Beer 1996; Randall et al. 1999). Our results show that this west-to-east trend in eelgrass decline may be progressing. Given that much of the shoreline and watershed areas of LIS are experiencing the type of expanding industrial and residential development thought to be detrimental to eelgrass elsewhere (e.g., Orth and Moore 1983; Thayer et al. 1984; Short et al. 1996), there is need to better document changes in eelgrass populations, to more definitively identify and possibly reduce sources of suggested impact responsible for observed declines.
It is often not clear how trends and fluctuations in eelgrass population characteristics observed in this study are related to physical :factors such as, temperature or sedimentary characteristics. We observed abbreviated or shifted period of occurrence of seed-bearing plants, which were more commonly found at all stations throughout the June-September during the period 1985-89 than during later years, when they occurred primarily in June and July. Analysis of long-term temperature data from the MPS intakes revealed a significant warming trend in seawater temperatures over the period 1976-2000, and this trend was particularly pronounced during the winter/spring (January-June; Foertch 2000). The winter/spring temperature rise may have resulted in
earlier onset of both the optimum temperature range for 'eelgrass reproduction' ( $9-15^{\circ} \mathrm{C}$; Setchell 1929; Thayer et al. 1984) and of more stressful temperatures above $15-20^{\circ} \mathrm{C}$ when eelgrass ceases to produce seeds (Burkholder and Doheny 1968; Orth and Moore 1983). Seawater temperature rise may have contributed to declines observed in other population parameters as well.
Two short-term population declines in shoot density and standing stock biomass were directly associated with localized population overgrowth. In July 1991, a bloom of the filamentous green alga Cladophora spp. covered the entire WP eelgrass bed. As a result of this bloom, shoot density declined and the sparse remaining plants appeared pale yellow in August and September. Studies in Oregon, USA (Kentula and McIntire 1986) and Hampshire, UK (den Hartog 1994) found similar demise of eelgrass following green algal blooms of Enteromorpha prolifera and $E$. radiata, respectively. Loss of plants from NR3 in 1992 was attributed to fouling by blue mussels (Mytilus edulis) which covered blades so heavily that they sank to the bottom where they were completely overgrown. Reusch et al. (1994) suggested that blue mussels can be beneficial to eelgrass through biodeposition, water nutrient enhancement, and turbidity reduction. However, mussels in their study were rarely observed attached to blades or rhizomes, but rather to each other forming an understory mat below the eelgrass canopy. The NR3 bed had not recovered from this event through 2001.
Extensive die-off of most of the remaining eelgrass in the Niantic River, including sampling bed for site NR4, occurred between July and August 1999. This die-off coincided with a large increase in sediment silt/clay content at NR4. The cause of this sedimentary change was not determined. Turbidity and high siltation rates over short periods have been shown to be detrimental to seagrasses (Moore et al. 1996, 1997; Vermaat et al. 1996). In addition, we observed thick mats (up to 25 cm ) of the red macroalga Agardhiella subulata covering the sediment surface and lower portions of eelgrass blades at NR4 preceding the die-off.: This habitat change : in the Niantic River was also reported by Goldberg et al. (2000), who noted evidence of hypoxia and migration of scallops out of the NR4 eelgrass bed. This condition has been demonstrated by Hauxwell et al. ( 2001 ) and others (see review of McGlatherty 2001) to be detrimental to eelgrass by limiting light and creating anoxia and high ammonia levels within the mat through algal decomposition and remineralization.
The spatial pattern of decline among study populations in the vicinity of MPS suggests a relationship to nutrient enrichment. The location exhibiting highest variability in eelgrass population
parameters monitored in this study is the Niantic River, where a patchy population of transient beds has been observed since 1985. This population was extensive during the 1970s (McGill 1974; Marshall 1994), but nearly nonexistent by 1999 (this study and Vozarik et al. 2000). Marshall (1994) and Short (1988) suggested that the decline of eelgrass in the Niantic River was due to a combination of poor water quality and the presence of Labyrinthula, a marine slime mold implicated as the causative factor in eelgrass wasting disease. The Niantic River study populations experience the least tidal flushing and are nearest to freshwater input sources and dense housing developments, when compared to the other two study populations near MPS. Thus, Niantic River celgrass beds may be more susceptible to nutrient loading from fertilizers and domestic septic systems than beds in Jordan Cove. As with many areas in the northeast U.S., considerable housing development has occurred along the Niantic River shoreline and within its watershed over the past 20 years. The JC site receives more tidal flushing and is farther removed from coastal nutrient input sources, and eclgrass population declines there were quite minor by comparison. The study population most removed from these sources and lotated along an open coastline with unrestricted tidal llow, WP, showed the least change of the three study areas over the $17-\mathrm{yr}$ study period.
Studies conducted in Ninigret Pond, RI and Waquoit Bay, MA linked both long- and short-term declines in eelgrass populations to increased housing development in the watershed (Short and Burdick 1996; Short et al. 1996; Bowen and Valiela 2001). Specifically, increased surface run-off and high groundwater discharge of nutrients from on-site septic systems were implicated as causal factors in declines of eelgrass population parameters in these studies. In another study in Ninigret Pond, Thorne-Miller and Harlin (1984) found a negative correlation between eelgrass production-to-biomass ratio ( $\mathrm{P} / \mathrm{B}$ ) and distance from the breachway to the Atlantic Ocean, suggesting a positive effect of ocean water on eelgrass production. These coastal systems are similar to the Niantic River in that they are enclosed embayments where environmental conditions are strongly intluenced by local landuse practices. While municipal sewerage has been installed in some parts of the Niantic River watershed (e.g., the eastern shore of the River), most of the developed portions still rely on on-site septic systems. In addition, old septic systems no longer in use may still be sources of nutrient input to the River, especially after periods of heavy precipitation. Moore et al. (1997) attributed losses and lack of recovery of eelgrass in an upriver section of the York River in Chesapeake Bay to seasonal pulses of higher turbidity relative to
downriver sites where turbidity was generally lower and eelgrass has persisted.

Biological disturbances may also have contributed to eelgrass losses in the Niantic River. Increased numbers of waterfowl known to reduce eelgrass coverage (Brent geese Branta bernicla, and swans Cygnus spp.; Valentine and Heck 1999) have been noted grazing on celgrass in the Niantic River (pers. obs.). In addition, large increases in green crabs (Carcinus maenas) have been documented in the River over the last 20 yrs (DNC 2001); their burrowing activities have been shown to damage celgrass (Davis et al. 1998).

## Conclusions

Eelgrass beds at the two monitoring sites nearest MPS (JC and WP) exhibited periodic fluctuations in shoot density and standing stock biomass, but generally support healthy populations, based on parameters monitored, throughout the 17-year study period, including 2001 when most parameters estimates were above average. These two populations are considered potentially impacted by the MPS thermal plume, but temperature monitoring has not provided evidence of a power plant influence. Rather, observed variability in water temperatures is indicative of natural solar warming and hydrodynamic conditions in Jordan Cove. These natural factors are particularly influential at JC, which is the most shallow of the study sites, and is immediately adjacent to extensive shallow sand flats vulnerable to solar warming. In addition io temperature data, there does not appear to be any relationship between WP and JC poputation fluctuations and power plant discharge flow and heat output. With Unit 1 permanently retired, the likelihood of thermal plume incursion at JC and WP in the future is further reduced.

Population variability has been highest during the study period in the Niantic River, where reference stations have been monitored. Since 1985, these studies have documented complete die-off of five separate eelgrass beds within the river. During 2001, population condition at NRS was generally poor. The exact cause of the long-term decline of eelgrass in the Niantic River is unknown. Because the Niantic River is located well away from any influence of the MPS thermal plume, declines there have been, and continue to be, related to other environmental factors such as nutrient input from domestic septic systems. disease, or increased turbidity.

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# Rocky Intertidal Studies 

## Introduction

Shore habitats in New England, including those near the discharge point of Millstone Power Station (MPS), are often comprised of hard substrata in the form of boulders and exposed bedrock ledge, which support rich and diverse communities of attached algae and animals. These communities are important and productive components of coastal ecosystems. Given their ecological importance and vulnerability to thermal plume impacts, studies of rocky shore communities are often included in ecological monitoring programs designed to assess the impacts of shore-sited power plants (Vadas et al. 1976, 1978; Wilce et al. 1978; NAESCO 1994; NAI 1999; NUSCO 1999; DNC 2001).

Rocky intertidal studies at MPS are part of a comprehensive environmental monitoring program whose primary objective is to determine whether temporal and spatial differences (e.g., in abundance, distribution or species composition) among communities at several sites in the Millstone Point area can be attributed to construction and operation of MPS. To achieve this objective, studies were designed and implemented to identify attached algal and animal species found on nearby rocky shores, to describe temporal and spatial patterns of occurrence and abundance of these organisms, and to identify physical and biological factors that induce variability in such communities. This research includes qualitative algal sampling, abundance (percentage cover) measurements of intertidal organisms, and growth and mortality studies of the brown macroalga, Ascophyllum nodosum. The following report discusses results of sampling and analysis in the most recent study year and compares these results to those of previous years, with particular attention to events related to construction and operation of MPS that may have altered environmental conditions on nearby shores.

## Materials and Methods

## Qualitative Algal Sampling

'Qualitative algal collections were made during oddnumbered months at four rocky intertidal stations (Fig. 1). These stations are, in order of most to least exposed to prevailing winds and storm forces: Fox IslandExposed (FE), Millstone Point (MP), White Point
(WP), and Giants Neck (GN). The MP station was added in September 1981; FE, WP and GN have been sampled since March 1979. A year of qualitative sampling is determined to be from March to the following January, both months inclusive. In other words, the latest year of qualitative algal data (2001) comprises collections from March 2001 to January 2002.

The FE station, approximately 100 m east of the MPS discharges, is directly exposed to the thermal plume during part of the tidal cycle; MP and WP are 300 and 1700 m from the discharges, respectively, and potentially impacted by the plume. The GN station is about 6.5 km west of Millstone Point and unaffected by MPS operation.

- Qualitative collections were used to characterize the attached flora at each site during each sampling period. Algal samples were identified fresh or after short-term freezing. Voucher specimens were made using various methods: in saturated NaCl brine, as dried herbarium mounts, or as microscope slide preparations.
The qualitative species list includes all attached, macroscopic algal species recorded from MPS sampling stations. Excluded from these lists are diverse diatom taxa, cyanobacteria and some crustose, endophytic or endozooic algal species. These elements of the microbiota are present but difficult to consistently collect, and, for many species, to identify as components of a large-scale environmental program. Also included in our lists are taxa that may be conspecific or subspecific forms, or alternate life history stages of erect macroalgae. For simplicity, we refer to each of these entities as a species throughout this report. Except where noted, nomenclature follows that of South and Tittley (1986), as updated by Villalard-Bohnsack (1995) and Sears (1998).


## Abundance Measurement

Abundance of rocky intertidal organisms was expressed as a percentage of substratum cover. At each qualitative collection station, five permanent strip transects 0.5 m wide were established perpendicular to the water-line, extending from Mean High Water to Mean Low Water levels. Each transect was subdivided into $0.5 \mathrm{~m} \times 0.5 \mathrm{~m}$ quadrats and was non-destructively sampled six times per year, in odd numbered months. The latest year for abundance measurement data is the same as that described for qualitative algal sampling data. The total number of quadrats in each transect


Fig. 1. Location of the MPS rocky intertidal sampling sites: $G N=G i a n t s$ Neck, MP=Millstone Point, FE=Fox Island-Exposed, WP=White Point.


Fig. 2. Detail map of the MPS vicinity: FN=Fox Island-New Ascophyllum site ( 1985 -present); MP and FE as in Figure 1.
depended on the slope of the transect. The percentage of substratum cover of all organisms and remaining free space in each quadrat was subjectively determined.

Understory organisms, i.e., species that were partially or totally obscured by the canopy layer, were assigned a percentage value that approximately corresponded to their actual substratum coverage. Each quadrat was assigned to a zone based on its tidal height: Zone 1 (high intertidal), Zone 2 (mid intertidal), or Zone 3 (low intertidal).

## Ascophyllum nodosum Studies

Growth and mortality of Ascophyllum nodosum, a perennial brown alga, were studied at two reference stations (GN and WP mentioned previously; Fig. 1) and a potentially impacted station (FN, about 150 m from the quarry discharges, northeast of the Fox IslandExposed sampling site; Fig. 2). Ascophyllum populations at GN and WP have been monitored since 1979, and those at FN since 1985. Ascophyllum had been monitored earlier, at a site ca. 75 m east of the original Millstone quarry cut (FO), from 1979 to 1984. This Ascophyllum population was eliminated in the summer of 1984 by exposure to elevated temperatures
from the thermal plume discharged through two quarry cuts (NUSCO 1987).
Upright shoots, or fronds, of Ascophyllum were measured , monthly, after onset of new vesicle formation, from April to the following April. At each station, fifty fronds were marked at their bases with a numbered plastic tag, and five apices on each individual were marked with colored cable ties. Linear growth was determined by measurements made from the top of the most recently formed vesicle to the apex of the developing axis, or apices if branching had occurred.. Monthly measurement of tagged plants began in June; in April and May, vesicles were not yet sufficiently large to be tagged, and tips were measured on randomly chosen individuals. Tags lost to thallus breakage were not replaced, and the pattern of loss was used as a measure of mortality. Loss of the entire frond was assumed when both the base tag and tip tags were missing. Tip survival was based on the number of remaining tip tags.

## Data Analysis

Analysis of qualitative algal collections includes a calculation of a frequency of occurrence index, based on the percentage of collections in which each species was found out of all possible collections (e.g., at a station, in a month, during a year). This index was used to calculate similarities among annual collections, using the Bray-Curtis formula (Clifford and Stephenson 1975):

$$
S_{j k}=\frac{\sum_{i=1}^{n} 2 \min \left(X_{i j}, X_{i k}\right)}{\sum_{i=1}^{n}\left(X_{i j}+X_{i k}\right)}
$$

where $S_{j k}$ is the similarity index between collections $j$ and $k ; X_{\mathrm{ij}}$ is the frequency of occurrence index for species $i$ in collection $j$; $X_{i k}$ is the index in collection $k$; and $n$ is the number of species in common. A flexiblesorting ( $\beta=-0.25$ ), clustering algorithm was applied to the resulting similarity matrix (Lance and Williams 1967). .. Additionally, multi-dimensional scaling ordination (MDS) was used to create a 2 -dimensional representation of comparisons of annual collections. This analysis was performed using a group-averaging algorithm in the PRIMER suite of programs developed by the Plymouth :Marine Laboratory (Warwick and Clarke 1991; Clarke 1993).
Quantitative analyses included determination of abundance of intertidal organisms as percentage of substratum covered by each taxon. Substratum not
occupied by macrobiota was classed as free space. Cover values of selected species were plotted against time." Similarities of communities (represented as annual collections at each station) were calculated using the Bray-Curtis coefficient formula cited above, substituting untransformed percentages for frequency of occurrence indices. Comparison of station/year collections was done using the same MDS techniques described above for qualitative algal analyses.
:A Gompertz growth curve was fitted to Ascophyllum length data using non-linear "regression methods (Draper and Smith 1981). The Gompertz function form used (Gendron 1989) has three parameters, related by the formula:

$$
L_{t}=\alpha e^{-e^{-t(t-t \cdot t)}}
$$

where $L_{1}$ is the predicted length at time $t, \alpha$ is the asymptotic length (estimate of length at the end of the growing season), $k$ is the rate of decrease of specific growth (shape parameter), and $t_{0}$ denotes the time at which the inflection point occurs (time when length is increasing most rapidly). The $\alpha$ parameter was compared among stations and between periods using 2sample $t$-tests ( $\mathrm{p}=0.05$ ) based on the asymptotic standard errors of the parameter estimates. The first derivative of the Gompertz function represents an instantaneous growth rate, with a maximum value occurring at $t_{0}$. Growth data representing the latest growing season (2000-2001) were plotted for all stations together and for each station separately, with summaries of previous years' data. Ascophyllum mortality was presented as loss of fronds and tips over time.

## Results and Discussion

## Qualitative Algal Studies

Water temperature is frequently implicated as a critical environmental factor in determining macroalgal spécies occurrence and distribution (Hoek 1982, 1984; Breeman 1988; Lüning 1990), and is often an important regulatory cue for algal life cycles (Lüning 1980; Swenarton 1997). Macroalgal communities in the vicinity of MPS are exposed to elevated water temperatures resulting from the thermal effluent discharge, and therefore, alterations of spatial and temporal patterns of species occurrence are likely. The current qualitative algal sampling program is used to monitor these patterns by applying various floristic analyses to data compiled from periodic algal collections.

Qualitative algal sampling results are presented in Table 1 as percent frequency of species occurrence by month, by station. The total number of species identified in 2002 was 102. This total was within the range of annual totals for previous study years (81111). One new taxon, the red alga Hypnea musciformis, was added to our species list in 2001; and owing to continued nomeclatural ambiguity, we follow Sears (1998) in renaming Scytosiphon simplicissimus and Ceramium nodulosum back to S. lomentaria and C. rubrum, respectively. Since 1979,153 algal species have been collected in odd-numbered months at the current four sampling sites; this compares with 161 species reported in the last year that nine stations were sampled (until 1995; see NUSCO 1996). Of the 'lost' species, only Laminaria digitata had occurred as more than a trace component of our flora. $L$ digitata had been relatively common, but only at the Twotree Island (TT) sampling site, where sampling has been suspended since 1995.
Because elevated temperatures mäy cause temporal shifts in occurrence of seasonal species, identification of components of the flora which exhibit natural seasonality provide a baseline from which power plantinduced changes can be assessed. A characteristic suite of species typical of cold-water period (January-May) collections in the Millstone area includes Dumontia contorta, Polysiphonia stricta, Spongonema tomentosum, Desmarestia viridis, Halosiphon tomentosus, Ulothrix flacca, Urospora penicilliformis, Monostroma grevillei, Protomonostroma undulatum and Spongomorpha arcta (Table 1). An equally distinctive group of species characteristic of warmwater (July-November) collections includes Champia parvula, Lomentaria baileyana, Aglaothamnion roseum, Ceramium diaphanum, Grinnellia americana, Dasya baillouviana, Polysiphonia harveyi, Hincksia mitchelliae, Enteromorpha, clathrata, Chaetomorpha linum, Bryopsis plumosa and B. hypnoides.
Shifts in natural occurrence patterns related to thermal plume exposure (i.e., decreased occurrence of cold-water species resulting from an abbreviated season, or increased occurrence over an extended season for species with warm-water affinities) can be detected by comparing species frequencies at stations potentially exposed to the thermal plume to other stations beyond its influence. The only station where such shifts have been and continue to be evident is the study site nearest the discharge, FE. For example, two cold-water red algae, Dumontia contorta and Polysiphonia stricta, were common components of the local winter/spring flora and were found in collections at unimpacted stations (GN, MP and WP) an average of $35 \%$ and $33 \%$ of the time, respectively (Table 1). However at FE, these two species were only found in
$9 \%$ and $14 \%$ of all collections, and most of these specimens were found prior to thermal plume effects caused by the opening of the second quarry cut in 1983, ie., in group fel described below. Other coldwater species (Desmarestia viridis, Monostroma grevillei, Protomonostroma undulatum and Spongomorpha arcta) occurred occasionally at FE, but much less frequently than at the other three stations. It is important to note that all these cold-water species were found at FE during the 1996-98 station shutdown period, presumably because of the ambient temperature conditions in the absence of cooling water discharge. By contrast, a number of warm-water seasonal species were more common at FE than at other sites. For example, Aglaothamnion roseum and Hincksia. mitchelliae occurred in $22 \%$ and $41 \%$, respectively, of the collections at FE, but in only $6 \%$ and $17 \%$ of collections at unimpacted stations, respectively. Other warm-water seasonals found with higher frequency at FE included Grinnellia americana, Dasya baillouviana, and Bryopsis hypnoides. These species were less common at FE during the shutdown period, which we attribute to the lack of thermal addition to FE discussed above.

Several perennial species exhibited shifts in occurrence patterns at FE. We documented the establishment of populations of species with geographical distributions which extend into warm temperate and tropical regions, and are therefore tolerant of elevated temperature regimes at FE (e.g., Gracilaria tikvahiae, Agardhiella subulata, Sargassum filipendula, and this year Hypnea musciformis; Taylor 1957; Lüning 1990). Similarly, some species near the southern limit of their normal geographical ranges, such as Mastocarpus stellatus and Polysiphonia lanosa, experienced elimination at FE either directly, from exposure the MPS thermal plume, or indirectly, from reduction of suitable substrata.

Cluster analyses, based on annual collections at each station, also reflect both site-specific and area-wide changes to the algal flora. For instance, groupings of collections at GN, MP and WP (Fig. 3a-c) separate into early and later sampling years, with the point of separation around 1986-88. This separation was influenced by the increasing contribution in recent years of species like Antithamnion pectinatum, an introduced species (Verlaque and Riouall 1989; Villalard-Bohnsack 1995) first observed in the MPS area in 1986 and now common at all study sites (Foertch et al. 1995), and Gelidium pusillum, which has been considerably more abundant during 3 -unit operation at GN and WP. Even with such floristic separation; all annual collections at GN, MP and WP clustered at greater than $50-60 \%$ similarity, indicating a

TABLE I. Qualitative algal collections (Mar. 1979 - Jan. 2002) by month, and by station. Values represent number of times found, as a percentage of possible times found. A dash before a species name indicates that it was collected in the latest report year. Taxa enclosed in quotes are, or may be, conspecific or subspecific forms or alternate life-history stages; see text for details. The FE, by group columns refer to the dendrogram groupings in Fig. 3; the unimpacted station average is the mean of GN, MP and WP (' T '=present, but $<1 \%$ ).

|  | by month |  |  |  |  |  | by station |  |  |  |  |  | FE, by group |  |  |  |  | unimpact.$\frac{\text { sta. avg. }}{6}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Rhodophyta | J |  | M | $\underline{J}$ | $\underline{S}$ | N |  | GN M | MP |  |  |  |  |  |  |  |  |  |
| Stylonema alsidii | 4 | 2 | 0 | 7 | 20 | 6 | 8 | 8 | 0 | 9 | 0 | 33 | 8 |  |  |  | 0 |  |
| -Erythrotrichopeltis ciliaris | 25 | 11 | 11 | 12 | 29 | 30 | 26 | 25 | 5 | 21 | 13 | 75 | 25 | 58 | 0 | 26 | 0 | 17 |
| -Erythrotrichia carnea | 10 | 4 | 3 | 3 | 13 | 8 | 9 | 12 | 3 | 4 | 0 | 0 | 0 | 0 | 17 | 15 | 17 | 6 |
| Erythrocladia subintegra | 1 | 0 | 0 | 0 | 1 | 3 | 2 | 0 | 2 | 0 | 0 | 8 | 0 | 0 | 0 | 4 | 0 | 1 |
| Erythropeltis discigera | 2 | 1 | 0 | 1 | 4 | 7. | 7 | 1 | 1 | 1 | 0 | 8 | 33 | 8 | 0 | 4 | 8 | 1 |
| -Bangia atropurpurea . | 74 | 82 | 35 | 10 | 27 | 51 | 44 | 44 | 50 | 48 | 38 | 33 | 25 | 50 | 50 | 50 | 50 | 47 |
| -Porphyra leucosticta | 72 | 80 | 62 | 18 | 10 | 30 | 44 | 46 | 53 | 40 | 29 | 33 | 33 | 67 | 42 | 46 | 67 | 46 |
| -Porphyra carolinensis | 7 | 3 | 2 | 1 | 3 | 4 | 9 | 0 | 4 | 1 | 0 | 0 | 0 | 0 | 0 | 4 | 83 | 2 |
| -Porphyra umbilicalis | 58 | 76 | 85 | 43 | 26 | 39 | 62 | 43 | 63 | 51 | 71 | 33 | 33 | 100 | 67 | 70 | 25 | 53 |
| -Porphyra linearis .. | 13 | 8 | 3 | 0 | 0 | 0 | 2 | 1 | 11 | 4 | 0 | 0 | 0 | 0 | 17 | 0 | 8 | 5 |
| Audouinella purpurea | 3 | 0 | 2 | 1 | 1 | 2 | 4 | 1 | 1 | 1 | 25 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| -Audouinella secundata | 33 | 21 | 21 | 17 | 19 | 12 | 22 | 21 | 23 | 17 | 21 | 33 | 8 | 8 | 17 | 28 | 25 | 20 |
| Audouinella daviesii | 6 | 2 | 2 | 4 | 3 | 4 | 5 | 4 | 2 | 4 | 4 | 0 | 0 | 8 | 0 | 4 | 25 | 3 |
| -Audouinella saviana | 8 | 13 | 15 | 4 | 11 | 15 | 14 | 14. | 7 | 9 | 4 | 25 | 25 | 17 | 0 | 15 | 17 | 10 |
| Audouinella sp. | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| Audouinella dasyae | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| -Gelidium pusillum | 38 | 31 | 27 | 33 | 34. | 42, | 43 | 64 | 1 | 25 | 0 | 0 | 0 | 0 | 92 | 67 | 100 | 30 |
| Nemalion helminthoides | 0 | 0 | 0 | 6 | 0 | 0 | 0 | 0 | 4 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| -Bonnemaisonia hamifera | 1 | 4 | 9 | 12 | 0 | 1 | 0 | 1 | 1 | 17 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 6 |
| Trailliella intricata' | 0 | 0 | 1 | 0 | 0 | 0 | 0. | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| -Agardhiella subulata | 25 | 18 | 12 | 20 | 27 | 28 | 64 | 4 | 3 | 14 | 17 | 25 | 75 | 67 | 50 | 87 | 92 | 7 |
| Polyides rotundus | 3 | 2 | 6 | 12 | 8 | 7 | 7 | 2 | 5 | 12 | 8 | 0 | 0 | 0 | 17 | 7 | 8 | 6 |
| -Cystoclonium purpureum | 73 | 64 | 73 | 44 | 20. | 54 | 24 | 66 | 62 | 68 | 79 | 42 | 0 | 0 | 33 | 6 | 17 | 65 |
| -Gracilaria tikvahiae | 15 | 7 | 2 | 2. | 13 | 11 | 32 | 0 | 0 | 1 | 0 | 8 | 0 | 25 | 33 | 50 | 75 | T |
| -Ahnfeltia plicata | 37 | 38 | 35 | 42 | 29 | 34 | 31 | 12 | 55 | 47 | 92 | 67 | 0 | 0 | 50 | 9 | 17 | 38 |
| -Phyllophora pseudoceranoides | 20 | 10 | 4 | 9 | 6 | 11 | 5 | 9 | 6 | 20 | 17 | 0 | 0 | 0 | 8 | 4 | 0 | 12 |
| -Coccotylus truncatus | 8 | 13 | 10 | 6 | 7. | 10 | 2 | 5 | 6 | 22 | 8 | 0 | 0 | 8 | 0 | 0 | 0 | 11 |
| -Chondrus crispus | 97 | 97 | 97 | 98 | 97 | 97 | 88 | 1001 | 100 | 100 | 100 | 75 | 0 | 83 | 100 | 100 | 100 | 100 |
| -Mastocarpus stellatus | 57 | 54 | 52 | 53 | 52 | 63 | 6 | 32 | 98 | 91 | 21 | 25 | 0 | 0 | 0 | 0 | 0 | 73 |
| -Hypnea musciformis | 1 | 0 | 0 | 0 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 25 | 0 |
| Rhodophysema georgii | 0 | 0 | 1. | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| -Corallina officinalis | 75 | 72 | 71 | 71 | 73 | 74 | 97 | 4 | 99 | 93 | 96 | 100 | 92 | 100 | 100 | 96 | 100 | 66 |
| -Dumontia contorta | 22 | 67 | 70 | 7 | 1 | 1 | 9 | 43 | 28 | 33 | 33 | 25 | 0 | 0 | 8 | 0 | 0 | 35 |
| Gloiosiphonia capillaris | 1 | 2 | 7 | 0 | 0 | 0 | 5 | 0 | 1 | 1 | 4 | 0 | 0 | 0 | 17 | 7 | 0 | 1 |
| -Chorcocolax polysiphoniae | 9 | 13 | 7 | 8 | 3 | 3 |  | 20. | 7 | 1 | 8 | 0 | 0 | 0 | 0 | 0 | 0 | 9 |
| Hildenbrandia rubra | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| -Palmaria palmata | 18 | 28 | 27 | 25 | 7 | 12 | 6 | 21. | 23 | 28 | 13 | 8 | 0 | 8 | 0 | 4 | 8 | 24 |
| -Champia parvula | 28 | 11 | 6 | 58 | 76 | 63 | 40 | 32 | 33 | 57 | 21 | 58 | 33 | 50 | 25 | 43 | 58 | 40 |
| -Lomentaria baileyana | 2 | 0 | 0 | 8 | 47 | 7., | 14 | 14 | 1 | 12 | 8 | 33 | 25 | 25 | 0 | 11 | 17 | 9 |
| -Lomentaria clavellosa | 7 | 11 | 9 | 1 | 2. | 3. | 1 | 6 | 3 | 12 | 0 | 8 | 0 | 0 | 0 | 0 | 0 | 7 |
| Lomentaria orcadensis | 1 | 1 | 1 | 0 | 6 | 0 | 1 | 1 | 0 | 4 | 0 | 0 | 0 | 0 | 0 | 2 | 8 | 1 |
| -Antithamnion cruciatum | 25 | 2 | 9 | 45 | 38 | 38 | 19, | 25 | 19 | 41 | 33 | 25 | 33 | 17 | 0 | 17 | 0 | 28 |
| -Antithamnion pectinatum | 61 | 33 | 25 | 47 | 63. | 62 | 49 | 36 | 66 | 45 | 0 | 0 | 0 | 58 | 42 | 85 | 75 | 49 |
| Callithamnion corymbosum | 0 | 0 | 0 | 0 | 2. | 2 | 1 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 1 |
| -Aglaothamnion roseum | 4 | 2 | 1 | 8 | 26 | 18 | 22 | 6 | 5 | 6 | 33 | 0 | 8 | 33 | 0. | 33 | 0 | 6 |
| -Callithamnion tetragonum | 36 | 19 | 10 | 11 | 18 | 30 | 15 | 18 | 29 | 22 | 63 | 42 | 0 | 8 | 0 | 0 | 0 | 23 |
| -Aglaothamnion byssoides | 0 | 0 | 0 | 6 | 0 | 0 | 1. | 2 | 0 | 0 | 0 | 0 | 0 | 0 |  | 2 | 8 | 1 |
| -'Callithamnion baileyi' | 21 | 9 | 4 | 24 | 22' | 37. | 4 | 26 | 32 | 19 | 0 | 0 | 0 | 0 | 8. | 7 | 0 | 26 |
| -Ceramium deslongchampii | 2 | 0 | 1 | 1 | 7. | 3 | 0 | 7 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 |
| -Ceramium diaphanum | 1 | 0 | 2 | 30 | 42 | 11 | 10 | 12 | 11 | 24 | 4 | 0 | 8 | 0 | 8 | 13 | 33 | 16 |
| -Ceramium rubrum | 83 | 83 | 85 | 90 | 83 | 85 | 71 | 91 | 83 | 95 | 100 | 92 | 42 | 83 | 50 | 74 | 17 | 90 |
| Ceramium fastigiatum | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| -Spermothamnion repens | 54 | 34 | 25 | 45 | 48 | 60 | 12 | 54 | 43 | 67 | 21 | 17 | 0 | 0 | 8 | 15 | 8 | 55 |

TABLE I. (cont.)
Rhodophyta
-Spyridia filamentosa
Scagelia pylaisaei
Griffithsia globulifera
-Grinnellia americanum
Phycodrys rubens
-Dasya baillouviana
Chondria sedifolia
-Chondria baileyana
Chondria capillaris
-Polysiphonia denudata
-Polysiphonia harveyi
-Polysiphonia lanosa
-Polysiphonia nigra
-Polysiphonia fucoides
-Polysiphonia stricta
-Polysiphonia elongata
Polysiphonia fibrillosa
-Polysiphonia flexicaulis
-Rhodomela confervoides

| Phaeophyta |
| :--- |
| -Ectocarpus fasciculatus |
| -Ectocarpus siliculosis |
| Ectocarpus sp. |
| -Hincksia granulosa |
| -Hincksia mitchelliae |
| -Pilayella littoralis |
| -Spongonema tomentosum |
| -Acinetospora sp. |
| -Ralfsia verrucosa |
| -Elachista fucicola |
| Halothrix lumbricalis |
| -Leathesia difformis |
| -Chordaria flagelliformis |
| Sphaerotrichia divaricata |
| Eudesme virescens |
| -Pogotrichum filiforme |
| Punctaria tenuissima |
| Phaeosaccion collinsii |
| -Punctaria latifolia |
| -Punctaria plantaginea |
| -Petalonia fascia |
| -Scytosiphon lomentaria |
| Desmarestia aculeata |
| -Desmarestia viridis |
| Chorda filum |
| Halosiphon tomentosus |
| -Laminaria longicruris |
| -Laminaria saccharina |
| -Sphacelaria cirrosa |
| Sphacelaria rigidula |
| -Ascophyllum nodosum |
| -Fucus distichus s edentatus |
| Fucus distichus s evanescens |
| -Fucus spiralis |
| -Fucus vesiculosus |
| -Sargassum filipendula |



TABLE I. (cont.)
Chlorophyta
-Ulothrix flacca
-Urospora penicilliformis
Urospora wormskjoldii
'Urospora collabens'
Acrochaete viridis
-Monostroma grevillei
-Protomonostroma pulchrum
Monostroma oxysperma
-Spongomorpha arcta
-Spongomorpha aeruginosa
'Codiolum gregarium'
Capsosiphon fulvescens
Capsosiphon groenlandicum
-Blidingia minima
Blidingia marginata
-Enteromorpha clathrata
-Enteromorpha flexuosa
-Enteromorpha intestinalis
-Enteromorpha linza
-Enteromorpha prolifera
Enteromorpha torta
Enteromorpha ralfsii
Percursaria percursa
-Ulva lactuca
-Prasiola stipitata
-Chaetomorpha linum
Chaetomorpha melagonium
-Chaetomorpha aerea
-Cladophora albida
-'Cladophora flexuosa'
'Cladophora glaucescens'
Cladophora laetevirens
-Cladophora refracta'
-Cladophora sericea
'Cladophora crystallina'
-Cladophora hutchinsiae
-Cladophora rupestris
-Cladophora ruchingeri
-Rhizoclonium riparium
'Rhizoclonium keneri'
'Rhizoclonium tortuosum'
-Bryopsis plumosa
-'Bryopsis hypnoides'
Derbesia marina
-Codium fragile
-


| FE, by group |  |  |  |  |  |  | unimpact. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | fe4 | fe5 |  |  | sta. avg. |
| 54 | 17 | 8 | 33 |  |  | 17 | 29 |
| 38 | 17 | 33 | 67 | 33 | 22 | 25 | 33 |
| . 13 | 0 | 8 | 58 | 25 | 19 | 0 | 7 |
| 0 | 8 | 17 | 8 | 0 | 0 | 0 | 1 |
| 0 | 0 | 0 : | 0 | 0 | 0 | 0 | . 1 |
| 21 | 25 | 17 | 0 | 8 | 2 | 0 | 19 |
| 33 | 25 | 0 | 17 | 25 | 7 | 0 | 30 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| 33 | 0 | 0 | 25 | 25 | 4 | 0 | 23 |
| 13 | 0 | 8 | 0 | 8 | 0 | 0 | 13 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| 4 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| 75 | 67 | 67 | 75 | 83 : | 91 | 75 | 69 |
| 8 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| 25 | 33 | 33 | 17 | 0 | 6 | 8 | 14 |
| 63 | 50 | 75 | 92 | 58 | 851 | 100 | 54 |
| 38 | 25 | 67 | 8 | 0 | 13 | 8 | 30 |
| 75 | 50 | 42 | 921 | 100 | 961 | 100 | 79 |
| 25 | 33 | 25 | 17 | 8 | 6 | 0 | 23 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| 4 | 0 | 0 | 0 | 0 | 2 | 0 | 2 |
| 0 | 0 | 0 : | 0 | 0 | 0 | 0 | 1 |
| 100 | 100 | 67 | 100 | 100 | 85 | 92 | 95 |
| 0 | 0 | 0 | 0 | 0 | 2 | 0 | 21 |
| 88 | 67 | 8 | 17 | 17 | 13 | 17 | 60 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| 63 | 75 | 58 | 83 | 83 | 781 | 100 | 50 |
| 13 | 0 | 0 | 0 | 0 | 2 | 8 | 5 |
| 0 | 25. | 42 | 33 | 25 | 13 | 17 | 21 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| 25 | 8 | 0 | 17 | 0 | 6 | 0 | 7 |
| 46 | 33 | 58 | 25 | 33 | 26 | 50 | 30 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | T |
| 8 | 25 | 0 | 0 | 0 | 4 | 8 | 7 |
| 4 | 0 | 0 | 0 | 0 | 15 | 25 | 6 |
| 0 | 0 | 8 | 8 | 8 | 24 | 25 | 6 |
| 25 | 17 | 8 | 0 | 0 | 7 | 0 | 19 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| 4 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 21 | 17 | 33 | 25 |  | 17 | 8 | 4 |
| 0 | 17 | 0 | 8 | 0 | 24 | 8 | 7 |
| 33 | 17 | 0 | 8 | 0 | 0 | 0 | 1 |
| 100 | 100 | 100 | 100 | 100 | 981 | 100 | 88 |



Fig. 3. Clustering dendrogram of percent similarity of qualitative algal collections, by year, at rocky intertidal sampling stations. Numbered groups at FE are further characterized in Table 1.
high degree of consistency in the year-to-year algal flora at these sites.
In contrast, the overall similarity of annual collections at FE was only about $35 \%$, with seven groupings apparent at about the $70 \%$ similarity level (Fig. 3d; Table 1). The first group represents collections made during 2 -unit 1 -cut operational years (1979-82), when the flora at FE was similar to that observed at other unimpacted stations (Table 1). Temperature conditions were severely altered when the second quarry cut was opened in 1983, and account for the relatively low algal diversity at FE, and increased occurrence of some of warm-water species discussed above (e.g., Lomentaria baileyana, Champia parvula, and Hincksia mitchelliae) in 1983 and 1984 (Group 2). Algal diversity remained low in Group 3 years (1985 and 1986), when dominant perennials (Chondrus crispus, Ascophyllum nodosum and Ahnfeltia plicata) and their associated epiphytes (e.g., Ceramium rubrum, Protomonostroma undulatum, Elachista fucicola, Cystoclonium purpureum and Polysiphonia :lanosa) were eliminated, while opportunistic species, including Codium fragile, Polysiphonia harveyi and Enteromorpha intestinalis became more common. Another floristic shift occurred during the early years of 3 -unit operation (1987 and 1988; Group 4). Increased water flow out of the quarry cuts from the Unit 3 discharge reduced temperature extremes at FE , allowing for reestablishment of some species (e.g., Chondrus crispus, Polysiphonia fucoides, Elachista fucicola, and Spongomorpha arcta), and initial colonization by new populations (Sargassum filipendula and Gracilaria tikvahiae). This group was further distinguished by the initial appearance Antithamnion pectinatum, which was found throughout the study area during this period.

More consistent water temperature regimes occurred over the next eight years of the 3 -unit operating period (1989-1996) which are included in Group 6. This' relative consistency allowed for some "further successional development of what has become a unique flora at FE, characterized by perennial populations of; Sargassum filipendula and Gracilaria tikvahiae and shifts in seasonal species occurrence patterns described above. It will be interesting to note whether the recently observed Hypnea musciformis population will persist. Similar floristic shifts have been observed by other researchers studying attached algae near thermal effluents (Vadas et al. 1976; Wilce et al. 1978; Schneider 1981). However, this algal community has exhibited a degree of resiliency to more than two years. of plant shutdown (1996-1998), as Group 6 also includes 1999, the first year when both Units 2 and 3 operated after the shutdown. Detectable changes to the FE algal flora did occur during the shutdown period, as
indicated by the separate grouping of 1997 and 1998 (Group 5). Some examples include reduced occurrence of established perennial populations of Sargassum and Gracilaria, brief reoccurrence of cold-water annuals discussed above, and reduced occurrence of warmwater annuals relative to previous years during 3 -unit operation. The close linkage of Group 7 years (2000 and 2001) to Group 6 indicates these changes were transient, as many characteristics of the thermal algal community returned following the restart of Units 2 and 3.
Relationships among all station/year collections were examined using multi-dimensional scaling (MDS) techniques. The resulting MDS plot (Fig. 4) revealed two distinct clusters. The largest cluster is comprised of all annual collections made at unimpacted sites (GN, MP and WP) along with FE collections made prior to observable thermal impacts (1979-1983). Initial years of thermal impacts at FE (1984-1986) appear as spatially distinct outliers in the MDS plot, and represent early successional years during the most severe thermal loading after the opening of the second quarry cut. The unique algal community that developed at FE after Unit 3 start-up in 1986, including the years of shutdown and reactor restarts, are represented by annual collections included in a smaller cluster. Years comprising this cluster are the same as those included in Groups 5-7 of the dendrogram discussed above (1987-2001). The temporal course of algal community development under various thermal regimes following Unit 3 start-up can be tracked in this cluster. This course demonstrates the establishment of a new environmental domain at FE during 1987-2001 (sensu Bradbury et al. 1984) that exhibits some degree of resilience to fluctuation in thermal regimes, such as ambient conditions during station shutdown from 1996 to 1998.

## Abundance Measurement

Thermal impacts also change abundance and distribution of existing species; such changes could be undetected based solely on qualitative sampling described in the previous section. Therefore, more quantitative assessments of distribution and abundance patterns of dominant intertidal organisms, based on percentage substratum cover of some seaweeds discussed above, and also of several invertebrates, were conducted using permanently marked transects. This study was designed to sample species abundance over an area sufficiently large as to accurately describe large-scale patterns of abundance in each intertidal zone (high, mid and low) at each sampling site. Among-station differences in abundance patterns could then be related to site-specific physical and biological


Fig. 4. Two-dimensional MDS plot for annual qualitative algal collections from rocky intertidal transects at $\mathrm{FE}, \mathrm{GN}$, and WP (1979-2001). and MP (1982-2001).
controlling mechanisms including, for stations near the MPS discharge, exposure to elevated temperature regimes. Abundance patterns of ecologically important intertidal organisms follow, along with analyses of overall community structure.

## Barnacles

Barnacles (primarily Semibalanus balanoides) can occupy a large proportion of intertidal rock surfaces on local shores, relative to other sessile invertebrates. They also exhibit spatial and temporal patterns of abundance that can be related to specific environmental factors. Barnacle abundance is generally highest in the mid-intertidal zone, where habitat conditions are optimal relative to upper and lower zone conditions. Barnacle abundance in the upper intertidal zone is limited by shorter immersion time, reducing time for larval settlement and feeding, and creating more physical stress from desiccation and temperature extremes (Connell 1961; Grant 1977; Gaines and Roughgarden 1985; Connolly and Roughgarden 1999). Longer immersion time of low intertidal surfaces dramatically increases larval supply and, coupled with extensive algal canopy, improves survival conditions (Leonard 1999), but higher predation rates and interspecific competition for space (lack of available
substratum) substantially limits abundance (Connell 1961; Underwood and Denley 1984; Minchinton and Scheibling 1991, 1993; Bertness et al. 1999).
Seasonally, barnacles exhibit an annual pattern of abundance marked by reproduction and settlement in early spring, rapid growth and surface cover increases in summer, and decreased abundance through autumn and winter due to a combination of overcrowding, predation and physical disturbance (Connell 1961; Menge 1976; Hughes and Griffiths 1988; Bertness 1989; NUSCO 1993; Bertness et al. 1998). In addition, long-term monitoring studies in Plymouth, U.K. summarized by Southward (1991) have revealed trends in barnacle abundance and distribution over periods of $10-25$ years that may be related to sea water temperature, solar (sunspot) cycles, or changing weather patterns and other effects of global climate shift.
Barnacle abundance patterns described above, both seasonal and spatial, were observed at all Millstone study sites over the 23 -year study period (Fig. 5).
Maximum barnacle cover in the high intertidal (Zone 1) during 2001 ranged from $3 \%$ (GN) to $29 \%$ (MP). Minimum coverage in Zone 1 ranged from $\leq 2 \%$ ai GN and WP to $8 \%$ at MP. In the mid intertidal (Zone 2), maximum barnacle cover in 2001 was lowest at FE ( $12 \%$ ) and highest at GN (67\%); minimum cover in


Fig. 5. Abundance of barnacles in each zone, and of predatory snails in Zone 3, of undisturbed transects, from March 1979 through January 2002.
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Fig. 5. (cont.)

Zone 2 was lowest at $\mathrm{FE}(<1 \%)$ and highest at GN ( $14 \%$ ). Low intertidal (Zone 3) maxima during 2001 ranged from $6 \%$ (FE) to $33 \%$ (MP). The annual minimum in Zone 3 in 2001 was $0 \%$ at FE and MP, $<1 \%$ at WP, and $4 \%$ at GN. Peak abundance of predatory snails (primarily the oyster drill, Urosalpinx cinera) at all study sites occurred in July or September 2001, and ranged from $0.2 \%$ (GN) to $3.3 \%$ (WP).

While predictable annual barnacle abundance cycles have been observed at each site over the study period, recruitment and peak abundance levels have been more variable from year to year. Peak abundance estimates in 2001 were low at most sites (excluding GN) and have been low regionally for three of the past four sampling years, with 2000 being the exception. These and other studies (e.g., Underwood and Denley 1984; Minchinton and Scheibling 1991; Miron et al. 1999) show a strong relationship between larval settlement and peak abundance levels. In 2000 and most study years, seasonal cycles were characterized by heavy set of cyprid larvae in spring, followed by growth to a summer peak in percent cover, as described above. The specific factors influencing barnacle recruitment from year to year are unclear, but the regional nature of recruitment success suggests natural, possibly climatic, environmental factors unrelated to power plant operation.
Natural regional environmental factors were also important in determining barnacle population structure at FE, particularly during the recent shutdown period (1996-98). Barnacle abundance was generally high during this period in all three zones, when compared to years during 3 -unit operation. However, the substantial influence of MPS operation on barnacles at FE again became apparent with plant restart. Since 1998, peak barnacle abundance at FE has been low, with near complete elimination of barnacles in Zones 2 and 3 each summer. This sharp a decline had not been observed in previous 3 -unit operational years, and may be due to a combination. of thermal effects and regionally poor barnacle settlement and recruitment discussed above. Clearly, regional conditions for barnacle recruitment were better in 2000, and seasonal patterns of abundance at FE were more consistent with those observed during most 3 -unit operational years (i.e., 1988-1995). Similarly, low peak barnacle abundances at MP and WP in 1998-99 and 2001 noted above were also evident at FE. The thermal regime at FE under current 2-unit operation (Units 2 and 3; temperatures $6-7^{\circ} \mathrm{C}$ above ambient) is not as stressful as that observed after the opening of the second quarry cut in 1983, and prior to initial Unit 3 start-up in 1986. The 2-unit (Units 1 and 2) 2-cut discharge during this period elevated temperatures by $9-11^{\circ} \mathrm{C}$ at FE and caused complete elimination of barnacles in Zones 2
and 3 by October (Fig. 5; NUSCO 1992). Because of reduced temperature extremes under current operating conditions, barnacle survival was observed in Zones 1 and 2 at FE through the summers of 2000 and 2001. However, these current conditions appear more stressful than those during 3 -unit, 2 -cut operation (1986-1995), and barnacle abundance has been low at FE since MPS restart in 1998. It is likely that low abundance is related to other community changes at FE in recent years discussed below.

## Fucus

Fucoids are major contributors to seaweed biomass in many parts of the world, including arctic, temperate, tropical and antarctic waters (Lüning 1990). Ecological studies of fucoids began in earnest at the turn of the century; however, most progress in understanding the demography and community ecology of these plants has been made since the early 1970s (Chapman 1995). Recent research has shown that Fucus canopy can be a major influence on abundance of other algac, sessile invertebrates, snails, and on the recruitment, regeneration and growth/self thinning of Fucus itself (McCook and Chapman 1997; Jenkins et al.1999). On local shores near MPS, Fucus vesiculosus forms an extensive canopy over barnacles in the mid intertidal zone, and also occurs in high and low intertidal zones. Other species of Fucus included in our abundance estimates are found occasionally at our study sites, but contribute relatively little in terms of percent substratum coverage. These species include $F$. distichus subsp. edentatus, $F$. distichus subsp. evanescens (both occur mostly subtidally) and $F$. spiralis, which occurs in the high intertidal.
Fucus populations in the MPS area exhibit distribution patterns and seasonal abundance cycles similar to those reported elsewhere in the North Atlantic Ocean (Lubchenco 1980, 1983; Topinka et al. 1981; Creed et al. 1996; Johnson et al. 1998; Karez and Chapman 1998). At most MPS study sites, Fucus abundance typically peaks annually in late summer or autumn, reflecting high recruitment and growth rates prior to and during this period (Fig. 6). Peak abundance in Zone 1 during 2001 was greatest at FE ( $72 \%$ ) and lowest at GN ( $<1 \%$ ). Fucus abundance was highest in Zone 2, relative to upper and lower zones, at all stations except FE (owing to the high Fucus cover in Zone 1). Fucus abundance reached its highest at MP; peaking at 66\%; Zone 2 abundance peak in 2001 was lowest at WP (52\%). In Zone 3, maximum Fucus cover during 2001 was greatest at GN (31\%); the lowest abundance peak occurred at FE (5\%). Adult Fucus is relatively resistant to herbivory, but newlysettled germlings may be vulnerable. Grazing snails


Fig. 6. Abundance of Fucus in each zone, and of grazing snails in Zone 3, of undisturbed transects, from March 1979 through January 2002.


Fig. 6. (cont.)
(primarily the common periwinkle Littorina littorea) were most abundant at GN and WP with maximum cover estimates in Zone 3 reaching 5\% at both sites. Grazing snails were rare or absent from most low intertidal collections at MP in 2001, and were not observed at all in Zone 3 quadrats at FE since 1998.
Fucus abundance patterns varied among study populations, reflecting environmental conditions unique to each site. In general, Fucus is most abundant on moderately exposed shores, such as those at FE and MP. Fucus abundance is limited at highly exposed sites by physical stress from wave shock, while at more sheltered sites, like WP and GN, these species are often outcompeted for space by another fucoid, Ascophyllum nodosum (Schonbeck and Norton 1978, 1980; Keser and Larson 1984). Vertical distribution patterns of intertidal Fucus are generally determined by desiccation rate gradients controlled by the degree of wave exposure, as well as slope of available substratum (Johnson et aL. 1998). More detailed description of the role these natural, site-specific characteristics play in determining Fucus zonation patterns at each study site is provided in previous reports (e.g., NUSCO 1992, 1993).

Since 1983, MPS operation has had a measurable impact on the Fucus abundance at FE. Prior to that, the two operating units discharging through a single quarry cut resulted in only a small thermal addition $\left(<2^{\circ} \mathrm{C}\right)$ to the shoreline at FE , which had no detectable effects to the intertidal community there (NUSCO 1992). Dramatic community changes were observed after the opening of the second quarry cut, particularly during the summer of 1984 with two units operating. The perennial Fucus population occupying all three intertidal zones was eliminated by temperature increases of up to $13^{\circ} \mathrm{C}$. As discussed previously in the barnacle section of this report, Unit 3 start-up reduced thermal stress to the adjacent shoreline, as increased discharge velocity carried the zone of extreme temperature conditions beyond the FE study site during most of the tidal cycle. As a result, perennial Fucus populations returned to mid and upper intertidal zones at FE, because they were exposed to air at low tide, the time of maximum thermal incursion. However, Fucus in Zone 3 was exposed to temperature increases of 8$9^{\circ} \mathrm{C}$ for several hours during each tidal cycle; this regime would totally eliminate Fucus by September of most 3 -unit operational years. A similar regime was apparent at FE from 1999 to 2001 as Zone 3 Fucus was eliminated in each summer with both Units 2 and 3 operating.

Ambient temperature conditions during the extended 3-unit shutdown from 1996 to 1998 allowed Fucus plants to survive in Zone 3 through summer during both 1996 and 1997, similar to pre-impact years of 1 -
cut operation. Even after Unit 3 restart in June 1998, some Fucus survived the seasonal maximum temperatures of late summer. Summer survival of low intertidal Fucus at FE had been observed in only two 3unit years (1993 and 1995) when a thermal discharge was present.

Fucus exhibited long-term abundance cycles at study sites more distant from the discharge than FE; these are likely unrelated to MPS operation. Most notable of these is the protracted decline/recovery cycle at MP discussed in previous reports (e.g., NUSCO 1996, 1998). Fucus abundance in all zones at MP appears to have plateaued at historic high levels established in recent years. We have no explanation for this longterm abundance cycle at MP. Proximity of MP to the MPS discharge (ca. 300 m to the east) and the moderate temperature increases measured directly there ( $2-3^{\circ} \mathrm{C}$ above ambient during slack tides; NUSCO 1994) suggest the possibility of a power plant impact. A direct thermal impact is unlikely, however, as the present Fucus population at FE recovered relatively rapidly after Unit 3 start-up, even under much greater temperature extremes than those at MP. The increase in Fucus abundance at MP in recent years could be related to lower abundance of grazing snails since 1993. It is unclear whether this decline in snail abundance at MP is related to the thermal plume or other factors. However, it is interesting to note that grazing snail abundance at MP in July 2000 was almost $7 \%$ cover, and concurrent peak summer Fucus cover was the lowest in six years. Data from WP and GN suggest that the increasing trend at MP may be part of an area-wide trend, as a gradual increase in Fucus abundance has become apparent over the last 15-20 years in Zones 2 and 3 at both stations.

## Chondrus and common epiphytes

The common red alga Chondrus crispus is among the most abundant low intertidal organisms on New England rocky shores (Menge 1976; Lubchenco 1980; Lubchenco and Menge 1983) including those near MPS. As discussed in previous sections, low intertidal habitat near the MPS discharge is more susceptible to thermal impacts than are higher zones. Therefore, documentation of abundance patterns of Chondrus and its associated epiphytes is critical to our ccological monitoring program. Perennial stands of Chondrus exclude many other species from Zone 3, including Fucus vesiculosus discussed above (Lubchenco 1980). Two seasonally abundant algal taxa coexist as epiphytes on Chondrus (i.e., Monostroma spp. (including Protomonostroma) and Polysiphonia spp.) instead of competing directly for primary space.

Spatial and temporal distribution patterns of these algae in the Millstone area are discussed below.
Extensive, well-established Chondrus populations are documented at three of the four study sites (all but FE) during the study period. Given its perennial habit, Chondrus abundance fluctuates little seasonally or from year to year at these three sites. During 2001, percent cover estimates were consistent with historical levels, peaking at $53 \%$ at $\mathrm{GN}, 82 \%$ at MP, and $76 \%$ at MP (Fig. 7).
The Chondrus population at FE had abundance levels similar to sites mentioned above prior to 1984 (40$75 \%$ ), but has since been reduced to scattered individual plants, with abundance estimates rarely exceeding $3 \%$. The extensive Chondrus population at FE was eliminated in 1984 by elevated water temperatures from the 2 -cut 2 -unit discharge (NUSCO 1987). Since that time, only a few scattered Chondrus plants have been observed in upper Zone 3 study quadrats. These plants are present during cooler months, but their upright portions are typically eliminated each summer. This summer decline was attributed to elevated water temperatures from the 2 -cut 3 -unit discharge. However, a similar decline was observed in both 1996 and 1997 during plant shutdown, suggesting other mechanisms, possibly seasonal overgrowth by Codium fragile and blue mussels (Mytilus edulis). These two species, and their influence on low intertidal community structure, will be discussed further in the next section. Regardless of the causal mechanisms, the pattern of summer disappearance of Chondrus, established under 3-unit operating conditions, was observed in each of the summers after the MPS shutdown period ended, including 2001.
Common epiphytes that exhibit warm-water and coldwater seasonality in the local low intertidal zone have shown temporal shifts in abundance in response to altered temperature regimes, typical of those at FE during MPS operation (NUSCO 1997). Polysiphonia spp. (mostly P. harveyi) are common warm-water epiphytes on Chondrus, Ascophyllum and Codium; they may also grow attached to rock. The annual abundance cycle of Polysiphonia spp. is characterized by a late summer peak, with cover declining to near $0 \%$ by winter at most study sites (Fig. 7). Peak abundance during 2000 was lowest at GN (2\%) and highest at FE (86\%). The annual cycle in Polysiphonia spp. abundance has been consistent at all stations except at FE throughout the study period. Elevated temperature regimes at FE since the opening of the second quarry cut (1983) produced favorable conditions for these species by extending the season of occurrence and increasing the levels of peak abundance, as well as by decreasing the abundance of grazing snails (Fig. 6).

These temperature regimes at FE have also allowed Polysiphonia spp. to persist through cold water months, when such species are typically absent from other sites, including FE prior to 1983. The return of ambient temperature conditions at FE following the 3 -unit shutdown has resulted in a winter/spring decline in Polysiphonia abundance similar to that observed at the other rocky shore stations, and at FE prior to the opening of the second quarry cut. Following MPS restart (July 1998), Polysiphonia persisted through the 1999 cold-water season at FE, with abundance never dropping below $4 \%$ cover. However, during the late winter of 2000 and 2001, Polysiphonia abundance declined to $<1 \%$ even though two units were operating during that time. Polysiphonia is often observed as an epiphyte on the often dominant Codium fragile population at FE. Relatively low Codium abundance since 1998 (see next section) may explain the concomitant and atypical winter decline of Polysiphonia.
The annual abundance cycle of Monostroma spp. (M. grevillei and Protomonostroma undulatum) is out of phase with that described for Polysiphonia spp., i.e., peak abundance is observed during cold water months (late winter/early spring) and these species disappear during warm-water months (July-December; Table 1, Fig. 7). This annual abundance cycle occurred consistently over the study period at all study sites except FE. Peak Monostroma abundance in 2001 was $11 \%$ at GN, $13 \%$ at MP, and $12 \%$ at WP. Since 1984 and excluding the recent extended shutdown period, Monostroma has been observed in FE study transects only rarely, and its cover has never exceeded $1 \%$. Prior to 1984, peak annual Monostroma cover at FE was similar to other exposed sites, ranging from $17 \%$ to $48 \%$. Monostroma occurred in low abundance ( $<1 \%$ cover) in FE study quadrats during the three cold water periods of 1996-98 shutdown, but has not been observed since the plants were restarted. While Monostroma is clearly temperature-limited during thermal discharge, other environmental factors may also inhibit at FE , such as limited suitable substratum, e.g., a host species (Chondrus crispus) and bare rock.

## Additional Taxa at Fox Island-Exposed

A number of other low intertidal taxa at FE have become important to the monitoring program because they have exhibited localized shifts in abundance. Population shifts of these taxa reflect acute effects of power plant operational changes "and long-term successional development of the thermally altered community in the nearfield discharge area. Therefore, time-series of abundance of these taxa are presented in Figure 8 and discussed below to provide a more

Fig. 7. Abundance of Chondrus and major epiphytes in Zone 3 of undisturbed transects, from March 1979 through January 2002.


Fig. 7. (cont.)
110 Monitoring Studies, 2001


Fig. 8. Abundance of additional taxa of interest in Zone 3, of undisturbed transects at FE, from March 1979 through January 2002.
comprehensive description of ecologically significant changes to the intertidal community at FE.
Codium fragile - This siphonaceous green alga is now a common component of the algal flora of eastern Long Island Sound, and is frequently found at all MPS monitoring stations (Table 1). Codium is an introduced species, first collected on the northeast coast of North America in 1957 at nearby East Marion, Long Island, NY (Bouck and Morgan 1957). This opportunistic or 'weed' species quickly colonized other areas of New England, including Connecticut. Optimal light requirements of Codium are comparable to those of other low intertidal algae such as Chondruis (Mathieson and Burns 1971). However, Codium is competitively inferior to Chondrus (through competitive exclusion), as demonstrated by low Codium abundance $(<5 \%$ : cover) at sites where extensive Chondrus canopy is found (DNC 2001). Codium is considered a. community dominant only at FE, with maximum covèr often approaching $90 \%$ since 1984 . Elimination of the FE Chondrus population in 1984 by elevated temperatures allowed Codium to develop a perennial population during the 3 -unit operational period, which persisted through the $1996-98$ shutdown period. However, since 2000, Codium cover at FE has declined precipitously, and remained $<1 \%$ in Zone 3 throughout 2001. This decline has coincided with expansion of populations of Gelidium pusillum and Corallina officinalis, discussed below.
Mytilus edulis - Blue mussels were observed periodically at most study sites (DNC 2001), and have often been among the dominant low intertidal taxa, particularly at FE (Fig. 8). Temporal mussel abundance patterns were altered at FE after Unit 3 startup in 1986. Prior to 1986, mussel abundance at FE never exceeded $5 \%$, but during 3 -unit operation, annual abundance peaks often reached $20-30 \%$ and in one instance (1994) exceeded $90 \%$. Mussel cover in Zone 3 at FE in 2001 peaked at $32 \%$. The higher settlement of mussels at FE since 1986 was previously attributed to hydrodynamic characteristics of the 3 -unit thermal plume, but may also be related to indirect effects such as the ability of mussels to settle in the extensive low intertidal Codium canopy. Following settlement, initial higher growth and subsequent high moriality were attributed to elevated water temperatures at FE : Thermal plume incursion at FE produced optimum temperature conditions for growth ( $10^{\circ}$ to $20^{\circ} \mathrm{C}$; Seed and Suchanek 1992) coinciding with spring and early summer phytoplankton blooms. However, thermal incursion proved detrimental by late summer, as, temperatures approached and often exceeded $27^{\circ} \mathrm{C}$, the maximum temperature for adult survival (Gonzales and Yevich 1976). High mussel cover was observed in 2000 and 2001 even with the sharp decline in Codium
abundance, indicating that replacement species in Zone 3 (ë.g., Gelidium pusillum discussed below) provide suitable substratum for settlement.
Antithamnion pectinatum - A more recent introduction to the Millstone area low intertidal zone, Antithamnion pectinatum (Foertch et al. 1995), occurs commonly as an epiphyte on Chondrus, Corallina and Ulva. This species is common in warm temperate waters of the western Pacific, but has also been identified as a recent introduction to the Mediterranean Sea (Verlaque and Riouall 1989) and the Azores (Athanasiadis and Tittley 1994). Since its introduction to the Millstone area in 1987, A. pectinatum has been found at all sampling sites, including FE near the MPS discharge (DNC 2001), and in Rhode Island (VillalardBohnsack 1995). Because this species is present at all of our study sites, including FE, its occurrence is considered an area-wide phenomenon unrelated to MPS. In the past several years," A. pectinatum has become increasingly common at FE, with peak cover estimates at time exceeding 30\%: In 2001, maximum cover of this species FE was just over $4 \%$. The generally higher abundance of A. pectinatum in recent years is likely due to expansion of the low intertidal host species Corallina officinalis coupled with the return of elevated temperature regime after the 199698 MPS shut-down period.
Other species - Abundance time-series of three perennial macroalgae (Gelidium pusillum, Corallina officinalis, and Sargassum filipendula) are presented in Figure 8 for the first time, because their populations have exhibited gradual increases since the late 1980 s, and are now considered community dominants at FE. All three species have biogeographical ranges that included Long Island Sound, but also extend into tropical waters (Taylor 1957). Qualitative sampling indicates that Gelidium occurrence has increased regionally since 1986 (Table 1), and has now established an extensive turf in the low intertidal at FE at times exceeding $60 \%$ cover. Reduced competition for space with Codium in recent years has allowed for population expansion. The Corallina population at FE also appears to have benefitted from reduced Codium abundance at FE , as Zone 3 cover has approached 20\% in recent years. Sargassum was first observed in 1986 at FE, and has occurred periodically primarily during warm water' months, but has rarely been collected at other sampling sites (Table 1). Declines noted in winter at FE are attributed to loss of upright thalli, presumably a response to colder water temperatures. The Sargassum population at FE appears dependent on thermal input from MPS, as few plants were observed during the 1996-98 shutdown period. After this period, extended operation of Units 2 and 3 provided beneficial conditions for the "FE' population as

Sargassum abundance exceeded $30 \%$ cover at times, the highest level observed since the study began.

## Community Analysis

Over one hundred macroalgal and invertebrate taxa occur in local rocky shore communities. Abundance and distribution of these taxa are influenced by complex interactions between physical processes (e.g., tidal height, exposure to waves, water temperature) and biological processes (e.g., inter- and intraspecific competition for light, space and nutrients, grazing and predation (including that by taxa not normally considered intertidal organisms, such as fish and shorebirds), growth and reproductive cycles). Characterization of these communities may be descriptive; abundance of populations that are stable or predictably variable may be represented as time-series of percentage of substratum coverage, as described in previous sections. However, comparisons among stations, or among years at a given station, may also be made using multivariate techniques, similar to those described in the Qualitative Algal section, using the abundance of all taxa found in the transects, even those that are rare or unpredictable in their occurrence.
Bray-Curtis similarity matrices, using annual average abundances of all taxa found in mid and low intertidal zones at each station, are illustrated as clustering dendrograms (Fig. 9). Annual samples at FE (Fig. 9a) were much more dissimilar than those at other stations; at the $40 \%$ similarity level, years grouped into three distinct clusters. Group I (1979-83) is comprised of early study years prior to observable effects from the opening of the second quarry cut. This community was characterized by high Chondrus, Fucus and barnacle coverage, with an appreciable amount of available free space (rock); it was similar to communities found at nearby unimpacted sites. Group II comprises 1984-86, when the opening of the second cut substantially altered the community at FE. Fucus and Chondrus abundance declined considerably, and these species were replaced with Codium and ephemeral green algae (Enteromorpha spp.). The remaining 15 study years make up Group III (1987-2001), which is further divided into three subgroupings at the $55 \%$ similarity level. Group IIIa is comprised of the 3 -unit operating period from 1987 to 1996 and is characterized by the return of Fucus to Zone 2, expansion of the Codium population in Zone 3, and less bare rock. Group IIIb (1994 and 1997-1998) is distinguished from Group IIIa by lower Codium cover, high cover of Mytilus and more diatoms. The last three years (1999-2001) are included in Group IIIc, and are characterized by low abundance of Codium and barnacles and high
abundance of Polysiphonia harveyi, Corallina officinalis, and Gelidium pusillum.
The high among-year similarity illustrated by the dendrograms for the other three rocky intertidal monitoring stations reflects the more stable environmental conditions at these sites. Annual collections at GN formed three distinct groupings at the $70 \%$ similarity level (Fig. 9b). Groúp I (1982 and 1983) was distinguished from Groups II (1979-1981, 1984-1991, and 2000) and III (1992-1999, 2001) by lower abundances of barmacles and Fucus, and more bare rock. At MP, three groupings were noted at the $60 \%$ similarity level (Fig. 9c): Group I (1982-1987), Group II (1988-1994) and Group III (1995-2000). A pronounced increasing trend in Fucus abundance at MP discussed earlier, along with more subtle increases in ephemeral green algae (Ulva lactuca and Enteromorpha linza), Polysiphonia harveyi, and Mytilus and a decline in available rock over the years were the major community changes that explain among-group dissimilarities. In addition, Group III at MP was further distinguished from other groups by higher Chondrus cover. Similar temporal trends in Fucus abundance and available rock factored strongly in groupings determined by cluster analysis at WP (Fig. 9d).
To provide a comparison of community states at FE to those observed at unimpacted stations, multidimensional scaling techniques were applied to a BrayCurtis similarity matrix comprised of station/year collections from all stations. These techniques have proven useful elsewhere for understanding spatial and temporal patterns of complex assemblages on rocky shores (Dye 1998; Underwood and Chapman 1998). The resulting MDS plot (Fig. 10) has a low stress level (0.09), indicating a good representation of the ordination in two dimensions (Clarke 1993), and reveals two distinct groupings with several outliers. Most collections fall into a relatively tight grouping, which comprises all annual collections at the unimpacted stations, along with annual collections from FE prior to the opening of the second quarry cut (19791983). The compactness of this cluster, with considerable overlap among collections, suggest that stations unaffected by MPS form a distinct community state relative to impacted areas. The changing community states at FE are illustrated by the relationships among the remaining collections in this MDS plot and these relationships are similar to those in the clustering dendrogram discussed above (Fig. 9). Annual collections at FE from 1984 to 1986 are spatially distinct, due to dramatic community changes at FE following the opening of the second quarry cut (see subgroup II in the dendrogram). The other grouping in Figure 10 includes FE annual collections


| taxon | Group I | Group II | Groú III | Group IIIa | Groun IIIb | Group IIIc |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: |
| Codium fragile | 0.95 | 24.98 | 26.25 | 36.05 | 17.97 | 5.14 |
| Fucus vesiculosus | 24.33 | 2.76 | .24 .40 | 25.45 | 23.65 | 21.96 |
| Semibalanus balanoides | 18.94 | 15.29 | 8.84 | 10.78 | 8.52 | 3.36 |
| Polysiphonia harveyi | 5.89 | 6.03 | 15.32 | 12.61 | 9.73 | 29.04 |
| Mytilus edulis | 0.34 | 2.55 | 13.58 | 7.61 | 33.75 | 11.34 |
| rock | 16.91 | 16.04 | 3.25 | 3.48 | 3.26 | 2.54 |
| Chondrus crispus | 30.28 | 4.88 | 0.78 | 0.48 | 0.93 | 1.51 |
| Enteromorpa linza | 1.23 | 6.31 | 8.10 | 9.06 | 6.75 | 6.55 |
| Enteromorpha flexuosa | 0.92 | 14.54 | .4 .64 | 4.92 | 3.78 | 4.65 |
| Ulvalactuca | 3.09 | 3.00 | 5.27 | 5.95 | 5.06 | 3.46 |
| diatoms | 0.11 | 1.41 | 5.14 | 2.85 | 9.52 | 7.61 |
| Corallina officinalis | 0.56 | 0.36 | 2.53 | 0.89 | 1.82 | 8.16 |
| Gelidium pusillum | 0.00 | 0.00 | 3.70 | 0.77 | 1.19 | 15.00 |

Fig. 9. Clustering dendrogram of percent similarity of undisturbed communities, by year, at rocky intertidal sampling stations: a) Fox Island-Exposed, b) Giants Neck, c) Millstone Point, and d) White Point. Each dendrogram is followed by a listing of taxa whose mean percent substratum coverage (by station) was at least $5 \%$ in at least one of the groupings determined by community analysis; group numbers correspond to those in the dendrograms.


| taxon | Group I | Group II | Group III |
| :--- | ---: | :---: | :---: |
| Semibalanus balanoides | 23.66 | 35.21 | 32.34 |
| Fucus vesiculosus | 13.90 | 24.18 | 35.88 |
| Chondrus crispus | 24.35 | 24.91 | 26.70 |
| rock | 28.75 | 19.16 | 13.58 |
| Ascophyllum nodosum | 5.33 | 5.78 | 5.76 |



| taxon | Group I | Groun II | Group III |
| :--- | ---: | ---: | ---: |
| Chondrus crispus | 36.49 | 35.97 | 44.68 |
| Semibalanus balanoides | 24.40 | 26.38 | 26.65 |
| Fucus vesiculosus | 5.35 | 13.96 | 41.56 |
| rock | 22.38 | 12.55 | 6.07 |
| Enteromorpha linza | 1.15 | 5.69 | 7.36 |
| Polysiphonia harveyi | 1.09 | 4.97 | 6.80 |
| Mytilus edulis | 0.77 | 3.69 | 8.14 |
| Ulva lactuca | 1.10 | 3.96 | 7.14 |



Fig. 9. (cont.)


Fig. 10. Two-dimensional MDS plot for annual percent cover data from rocky intertidal transects at FE, GN, and WP (1979-2001), and MP (1982-2001).
from 1988 to 2001, years of development and maintenance of the thermal community established under 3 -unit operation. Recent years (i.e., 2000 and 2001) are becoming increasingly dissimilar from other FE collection years, due to the large decline in Codium, and increases of Mytilus, Corallina, and Gelidium discussed for Group IIIc in Figure 9, changes particularly obvious in 2001.

## Ascophyllum nodosum Studies

Populations of the rockweed Ascophyllum nodosum form dense, nearly monospecific, stands on sheltered rocky shores locally, including some areas near the MPS outfall. Monthly monitoring of population growh and mortality has been conducted at three locations since 1979 to assess possible effects of MPS operation. Ascophyllum exhibits easily quantifiable growth responses to even slight changes in temperature, which makes this species a critical biomonitoring tool for studies of the ecological effects of thermal effluents. This attribute makes Ascophyllum a key species within; the MPS ecological monitoring program and these studies, as elsewhere (e.g., Vadas et al. 1976, Wilce et al. 1976), document the value of this species as a sensitive indicator of local environmental conditions. Review of phenological, ecological and applied
monitoring studies of Ascophyllum was presented in NUSCO (1993). Results of 2000-2001 Ascophyllum growth and mortality studies, compared with results from previous years, are presented below.

## Growth

Annual Ascophyllum growth is described using a Gompertz growth model (Gendron 1989) fitted to monthly Ascophyllum tip length data (Fig. 11). Parameters of this model provide useful indicators of Ascophyllum population growth characteristics. The $\alpha$ parameter of the model, used as an estimate of total. annual growth, was highest at FN in 2000-2001 (118.9 $\mathrm{mm})$, with lower estimates at $\mathrm{GN}(112.6 \mathrm{~mm})$ and WP ( 108.7 mm ; Fig. 11a). Ascophyllum annual growth estimates at $\mathrm{FN} \cdot$ in 2000-2001 was significantly higher ( $\mathrm{P}<0.05$ ) than the estimates at GN and WP; the difference between annual growth at GN and WP. was not significant. Growth rates were determined by extracting the first derivative of the Gompertz growth model and ploting results on the same set of axes (Fig. 11b). The inflection point, a parameter of the Gompertz model which identifies the time of maximum , growth rate, occurred earliest in 2000-2001 at FN (5 July; . $19.7 \mathrm{~mm} / \mathrm{mo}$ ), followed by WP ( 18 July; 18.6 $\mathrm{mm} / \mathrm{mo}$ ) and GN ( 22 July; $17.6 \mathrm{~mm} / \mathrm{mo}$ ).


Fig. It. Ascophyllum growth during 2000-2001: a) the Gompertz growth model fitted to monthly tip length data (ertor bars represent monthly mean lengths $\pm 2$ SE), and b) instantaneous growth rates based on the first derivative of the Gompertz model, including inflection points.

The annual growth estimate at FN during 2000-2001 was not significantly different from the 15 -year (19852000) mean of all previous years ( 115.0 mm ; Fig. 12). The inflection point for previous years' data at FN was 17 days later than during 2000-2001, with a lower peak growth rate ( $22 \mathrm{July} ; 16.1 \mathrm{~mm} / \mathrm{mo}$ ). At GN, growth in 2000-2001 was significantly higher than the mean of the previous 21 years ( 95.5 mm ). The growth curve for the previous years at GN had a later inflection point ( 27 July) and a lower peak growth rate ( $14.2 \mathrm{~mm} / \mathrm{mo}$ ) than during 2000-2001. Growth during 2000-2001 at WP was significantly higher than the mean of previous 21 years $(88.2 \mathrm{~mm})$. The peak growth rate for previous years ( 12.8 mm ) was lower and occurred later ( 30 July) than during the current study year.
Relationships of Ascophyllum growth characteristics among the three populations monitored in 2000-2001 were similar to many previous years, particularly to those during 3 -unit operation. Highest growth frequently occurred at the site nearest the discharge (FN), lowest growth at WP and intermediate growth at GN. In addition, earlier and higher peak growth rate at FN compared to WP and GN was also frequently noted in previous years. The MPS thermal plume was thought to be a dominant influence on Ascophyllum


Fig. 12. Ascophyilum growth and growth rates during 2000-
 including inflection points. Error bars represent monthly mean lengths $\pm 2 \mathrm{SE}$.
growth at FN until recent years. During the first year of monitoring at FN (1985-86), with only one unit operating and temperature increases of only about $1^{\circ} \mathrm{C}$ for 1-2 hours each tidal cycle, growth was not significantly different from GN or WP. Higher temperature increases (up to $3-4^{\circ} \mathrm{C}$ for $3-4$ hours each tidal cycle) resulted from Unit 3 startup during the 1986-87 growing season. This level of thermal input was believed to create favorable conditions for Ascophyllum growth by: 1) extending the period of "normal" or "ambient" peak growing conditions for local populations ( $18-21^{\circ} \mathrm{C}$; Kanwisher 1966; Chock and Mathieson 1979); 2) more closely synchronizing
these periods of optimal growing temperatures with the period of maximum daily solar irradiance (June); and 3) elevating temperatures in late summer above normal maxima but below stress levels ( $22-25^{\circ} \mathrm{C}$ ), increasing plant respiration and growth rates without exceeding photosynthate production (Brinkhuis et al. 1976; Stromgren 1977. 1981; Vadas et al. 1978). During subsequent 3 -unit years, there appeared to be a relationship between the thermal load produced by the power plant (affected by the number and duration of unit outages) and the degree of growth enhancement at : FN (NUSCO 1992). A similar relationship was observed at our original experimental population (FO) closer to the discharge from 1979 to 1983, prior to thermal effluent-related elimination after the opening of the second quarry cut (NUSCO 1992).
However, higher growth continued at FN through.. 1996-98 MPS shutdown period when no thermal effluent was present, suggesting that natural factors at FN were more influential than previously thought. Possible natural factors included solar warming of water from nearby sandflats and water circulation patterns in Jordan Cove (NUSCO 1999, 2000). Further confounding our hypothesis of power plant-related growth enhancement at FN were atypical relationships among monitored populations during 1998-99 while a thermal effluent was present (i.e., growth at FN was significantly lower than growth at GN, and not significantly different : from growth at WP). , High growth at GN and WP in 1998-99 cannot be explained by water temperature, as ambient temperatures were well within historical ranges (NUSCO 1999). Atypically high growth relative to previous years was again observed at GN and WP during 2000-2001, and was also noted at FN. Other natural environmental factors must contribute to variability in growth, especially at GN, which is unaffected by the MPS thermal plume. Thermal effects have also never been observed at WP, even during years of 3 -unit operation. The lack of a clear relationship between seawater. temperature and Ascophyllum growth at all three sample sites provides evidence that other regional or site-specific environmental factors, possibly nutrients or light, account for much of the spatial and temporal variability in growth observed since 1985.

## Mortality

Ascophyllum populations endure considerable stress from a variety of biotic and abiotic factors in the intertidal zone (NUSCO 1992), often resulting in breakage and loss (mortality) of the upright shoots or fronds. Thalli compensate for this loss of biomass through growth of suppressed fronds, described as meristem banks (Cousens 1986; Vadas et al. 1990) that


Fig. 13. Ascophyllum mortality, as number of remaining tagged plants, at each station.
were previously covered and shaded by a welldeveloped main shoot canopy. Population mortality is monitored in this study by examining patterns of frond base tag loss (referred to as plant loss; Fig. 13) and apical tag loss (tip loss; Fig. 14). . Plant losses at FN and WP during 2000-2001 were equal ( $46 \%$ ), and both lower than their respective historic means ( $61 \%$ and 57\%). Plant loss at GN in 2000-2001 (48\%) was lower than the historical mean of $57 \%$. Patterns of tip loss at each station were similar to those described above for plant loss. Tip loss estimates at FN and WP during


Fig. 14. Ascophyllum mortality, as number of remaining tagged tips, at each station.

1999-2000 were also equal to each other ( $69 \%$ ), and also both lower than means from previous study years ( $81 \%$ and $73 \%$, respectively). Tip loss at GN in 2000$2001(72 \%)$ was lower than the historic mean ( $76 \%$ ).
Ascophyllum mortality trends observed throughout this study reveal no evidence of power plant impact, with the exception of elimination at the original experimental study site (FO) in 1984 (NUSCO 1992). Our current sampling site nearest the discharge (FN) has generally had higher mortality rates than reference sites, with some exceptions including the last two
sampling years. However, these higher mortality rates do not appear to be related to proximity to the discharge, but rather to the higher degree of population exposure to wind- and wave-induced stress at FE , compared to the more sheltered reference sites. An area-wide seasonal pattern of mortality has been observed ihroughout our studies, which further implicates wave-induced stress as a major cause of mortality. Throughout the study, mortality rates were highest during the months of August through November, when strong storms and high energy waves were frequent. Many studies elsewhere point to the strong relationship between mortality and degree of site exposure to prevailing winds and storms (Baardseth 1955, 1970; Jones and Demetropoulos 1968; Vadas et al. 1976, 1978; Wilce et aL 1978; Cousens 1982, 1986; Vadas and Wright 1986).
The Ascophyllum population at our original impacted site (FO) has shown few signs of recovery following power plant-induced elimination 1984. As discussed in previous reports (NUSCO 1996, 1997), some individual thalli have settled, grown and persisted at FO during 3-unit operation and more recent years; however, no significant recovery has occurred to date. In addition, no observable recruitment of Ascophyllum occurred during more than two years of the recent station shutdown. Given its documented poor recruitment under natural conditions (Vadas et al. 1990), particularly at more exposed sites like FO (Bertness et al. 1999), it appears likely that postoperational recovery of this population will be slow.

## Conclusions

More than 20 years of ecological monitoring studies have documented that impacts associated with operation of MPS to rocky intertidal habitats remain restricted to approximately 150 m of shoreline on the east side of the discharge to Long Island Sound. This area, which includes the Fox Island study sites, has been exposed to the thermal effluent for most years since the opening of the second quarry cut in 1983, and has supported a unique community of seaweeds and invertebrates that has undergone a degree of successional development over the last 18 years. This community has exhibited resilience to recent operational events (e.g., more than two years of plant shutdown, Unit 3 restart in 1998, Unit 2 restart in 1999), when only minor and transient community changes were observed.
The intertidal community at FE that developed under modified thermal regimes in the discharge area was characterized by shifts in occurrence and abundance of many intertidal species at FE, e.g., absence or
abbreviated season for species with cold-water affinity (Chondrus, Monostroma, Dumontia), and presence or extended season of occurrence for species with warmwater affinity (Codium, Sargassum, Gracilaria). In 2001, a new species to include in the latter category, Hypnea musciformis, established à population at FE that appears to be expanding. Most of these community characteristics remained evident during the shutdown period from 1996 to 1998. Sublle community changes attributed to the return of ambient temperature conditions to FE included: , 1) higher frequency, of some cold water annuals (e.g., Monostroma spp., Spongomorpha spp. and Polysiphonia stricta) and lower frequency of warmwater annuals (e.g.,' Bryopsis spp., Polysiphonia harveyi and Grinnellia americanum); 2) higher mid and low intertidal abundance of barnacles and Fucus throughout the year; and 3 ), small increases in abundance of Chondrus in the low intertidal. However, these responses to ambient temperature conditions were reversed following MPS restart.
High Ascophyllum growth was observed at all sites in 2000-2001 compared to historical means, with highest growth at FN compared to other sites. This. relationship among stations has been attributed to elevated temperatures from the MPS discharge. However, a similar relationship was observed during two recent growing seasons (1996-97 and 1997-98) while all three units were shutdown, but was not evident during the first year following Unit 3 restart (1998-99). Natural influences of other factors such as ambient temperature conditions, nutrients and light may play a more important role in determining Ascophyllum growing conditions at FN than does thermal plume incursion from the MPS discharge.
These studies were also sufficiently sensitive to detect important regional changes to rocky shore communities unrelated to MPS operation.. These included the introduction and spread of an exotic red alga Antithamnion pectinatum, a species native to the Pacific Ocean and not previously reported in the North Atlantic, and an increase in abundance of Fucus over the last 15 years. In addition, these studies also revealed evidence of the regional nature of barnacle recruitment success from year to year.
In summary, the current rocky intertidal monitoring program has provided sufficient characterization of local rocky shore communities to detect and document ecologically significant changes, both related and unrelated to MPS operation. More important, this, long-term monitoring database provides a useful foundation from which further changes related to MPS operation or other factors can be assessed.

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## Fish Ecology Studies

## Introduction

Long Island Sound supports a diverse assemblage of fish species, including year-round residents, seasonally migratory fishes, and rarely seen transient species. Several of these species support important commercial and sport fisheries worth millions of dollars each year in Connecticut (Sampson 1981; Blake and Smith 1984).
The objective of the fish ecology monitoring program at Millstone Power Station (MPS) is to determine whether operation of the electrical generating units has adversely affected the occurrence, distribution, and abundance of local fishes. Potential MPS impacts include entrainment of early life history stages through the condenser cooling-water system, impingement of juvenile and adult fish on intake screens, and changes in distribution or abundance attributable to the thermal discharge. Numbers of fish eggs and larvae entrained have been reliably estimated each year since 1976. However, effects of entrainment mortality are more difficult to assess as fish population dynamics are influenced by many biological processes, such as compensatory mortality, density-dependent growth, age at maturity, fecundity, population age composition and size structure, and stock size and range.
Estimates of larger fish impinged on traveling screens can be readily made, but in the case of eggs and larvae, the implications of fish removal are difficult to assess as adult populations are concurrently affected by natural and fishing mortality rates. At MPS, however, the impact of fish impingement has largely been mitigated by the installation and operation of fish return sluiceways (NUSCO 1986, 1988c, 1994b; NNECO 1999; DNC 2001b).
Changes in the thermal regime of local waters due to MPS operation have been documented (NUSCO 1988b; see the Introduction section to this report). Should water temperatures exceed species tolerance level, individuals may be forced to move from the area, abandoning potentially important spawning or nursery grounds. Chronic exposure to increased temperature regimes may also affect the seasonality of spawning, growth and development of individuals, and possibly even local stock population dynamics.
Several monitoring programs were established to provide information for the assessment of impacts
from MPS operation on local fish assemblages. The programs include sampling of ichthyoplankton (fish eggs and larvae) at the power station and in nearby waters, demersal fishes by trawl, and shore-zone species by seine. These monitoring programs have been modified several times as a result of periodic evaluations (NUSCO 1987, 1994a, 1995). Data collected provided a basis for identifying taxa potentially affected by plant operation, as well as information on long-term abundance trends used to measure changes in local populations.

Fishes potentially impacted by MPS were selected either because of their susceptibility to entrainment or because of their distribution in relation to the thermal plume. Selected taxa include the American sand lance (Ammodytes americanus), anchovies (Anchoa spp.), Atlantic menhaden (Brevoortia tyrannus), silversides (Menidia spp.), grubby (Myoxocephalus aenaeus), cunner (Tautogolabrus adspersus), and tautog (Tautoga onitis). • During the past several years, increased emphasis has been placed on the tautog. This species supports one of the principal sport and commercial fisheries of LIS (Smith et al. 1989; ASMFC 1996) and its numbers have been declining in many areas since 1984, likely from overfishing of this slow-growing and long-lived fish (Simpson et al. 1995).
This report summarizes data collected from the monitoring programs during June 2000 through May 2001. These data are compared to findings presented in DNC (2001a). MPS Units 2 and 3 were both operating during most of this report period. Unit 3 was shut down from February 3 to March 30, 2001 and Unit 2 operated throughout the year. Unit 1 was shut down permanently on November 4, 1995, which resulted in less impact on all life stages of fish because entrainment and impingement rates are directly related to the amount of cooling-water used. The potential effects of MPS are assessed below for each selected fish taxon.

## Matcrials and Methods

Results of year-round sampling are presented using a 12 -month period that extends from June of one year through May of the following year. Because of occasional overlap in the occurrence of a species during the May-June transitional period, speciesspecific analyses are based on actual periods of occurrence instead of being constrained to a May 31
endpoint. When the season of occurrence for a species crossed a calendar year, the annual period was termed a report year (e.g., 2000-01). When a species was primarily collected only within a calendar year, the annual period was presented as a specific year (e.g., 2001). The materials and methods that follow correspond to the most recent sampling periods that have been completed.

## Ichthyoplankton Program

Sampling frequency of ichthyoplankton entrained through the MPS cooling-water system varied seasonally during 2000-01. Both day and night samples were collected twice a week during June through August, once a week in September and February, and three times a week during March through May. Only one daytime sample per week was collected during October through January. Samples were collected at either the Unit 2 or 3 discharge (station EN, Fig. 1).
To collect samples from the plant discharge, a $1.0 \times$ $3.6-\mathrm{m}$ conical plankton net with $335-\mu \mathrm{m}$ mesh was
deployed using a gantry system. Four General Oceanic flowmeters (Model 2030) were mounted in the mouth of the net and positioned to account for horizontal and vertical flow variations. Sample volume (about $200 \mathrm{~m}^{3}$, except during periods of high plankton or detritus concentrations, when volume was reduced) was determined from the average readings of the four flowmeters. Under normal plant operation, the net was usually deployed for 3 to 4 minutes. All ichthyoplankton collections were preserved in $10 \%$ formalin.

Ichthyoplankton samples were split in the laboratory using a NOAA-Bourne splitter (Botelho and Donnelly 1978); fish eggs and larvae were removed from the samples with the aid of a dissecting microscope. Successive splits were completely sorted until at least 50 larvae (and 50 eggs for samples processed for eggs) were found, or until one-half of the sample had been examined. Larvae were identified to the lowest practical taxon and enumerated in all samples, except for June samples, when only two (one day and one night) samples per week were typically examined. Tautog,


Fig. I. Location of current trawl, seine and ichthyoplankton sampling stations.
cunner, and anchovy (bay anchovy, A. mitchilli and striped anchovy, A. hepsetus) eggs were identified and enumerated in all samples collected from April through August. Tautog and cunner. (Family Labridae) have eggs of similar appearance and were distinguished on the basis of a weekly bimodal distribution of egg diameters (Williams 1967). All ichthyoplankton densities were reported as a number per $500 \mathrm{~m}^{3}$ of water filtered.

## Trawl Program

Triplicate bottom tows were made to collect demersal fish using a $9.1-\mathrm{m}$ otter trawl with a $0.6-\mathrm{cm}$ codend liner every other week throughout the year at three stations: Niantic River (NR), Jordan Cove (JC), and Intake (IN) (Fig. 1). A typical standard tow was 0.69 km in length, but if the trawl net became loaded with macroalgae and detritus, tow distances were shortened and catches standardized to 0.69 km by proportionally adjusting the catch; this only occurred at the NR or JC sites. Catch was expressed as the number of fish per standardized tow (CPUE). Up to 50 randomly chosen individuals of certain selected species per station were measured (total length) to the nearest mm . Catch of tautog in lobster pots (see Lobster Studies section for sampling methods) was used to supplement the trawl abundance data for this species.

## Seine Program

Shore-zone fish were sampled using a $9.1 \times 1.2-\mathrm{m}$ knotless nylon seine net of $0.6-\mathrm{cm}$ mesh. Triplicate shore-zone hauls (standard distance of 30 m ) were made parallel to the shoreline at Jordan Cove (JC): biweekly from May through November (Fig. 1). Collections were made during a period 2 hours before and 1 hour after high tide. Fish from each haul were identified to the lowest possible taxon, counted, and the total length of up to 50 randomly selected individuals of each species from each replicate were measured to the nearest mm total length. Catch was expressed as number of fish per. haul.

## Data Analyses <br> Abundance Estimates

A $\Delta$-mean was used as an index of abundance of juvenile and adult fish collected in the trawl and
seine programs, and of fish eggs and larvae in the ichthyoplankton program. The $\Delta$-mean is the best estimator of the mean for abundance data that approximates the lognormal distribution and contains numerous zeros (Pennington 1983, 1986). Calculation of this index and its variance estimate was described in detail in NUSCO (1988a). Because of varying sampling frequencies, the $\Delta$-mean indices of ichthyoplankton taxa were weighted by the largest number of samples collected in a week to standardize data across weeks and years. With species that occurred seasonally, data to calculate each $\Delta$-mean were restricted by month to reduce the number of zero values in the distribution tails, which extend beyond the occurrence seasonal boundaries. A nonparametric, distribution-free "Mann-Kendall test (Hollander and Wolfe 1973) was used to determine the direction of change if an annual $\Delta$-mean timeseries represented a significant ( $p \leq 0.05$ ) trend. Sen's (1968) nonparametric estimator of the slope was used to describe the rate of change of significant trends.: This approach to trend analysis was suggested by Gilbert (1989) as being particularly well-suited for analysis of environmental monitoring data, because no distributional assumptions are required and small sample sizes are acceptable. Graphical methods are useful aids to formal tests for trend analysis (Velleman and Hoaglin 1981). In particular, methods for smoothing time series of abundance data help visualize short-term patterns often obscured by high variability. Moving averages were used to smooth large abundance fluctuations and plotted against time to provide a visual aid to the statistical trend analyses. The time interval or "window" for the moving average was about $20 \%$ of the time series length as suggested by Gilbert (1989). Longer intervals provide a greater smoothing effect, but they may also obscure short-term patterns which can be of interest in our relatively short time series of abundance data.

## Entrainment Estimates

Entrainment estimates of dominant ichthyoplankton were calculated from daily density estimates at station EN. These estimates were based on the parameters of a Gompertz function fitted to the entrainment data. The distribution of egg and larval abundances over time is usually skewed because their densities increase rapidly to a maximum and then decline slowly. The cumulative density over time from this type of distribution resembles a
sigmoid-shaped curve, for which the inflection point occurs at the time of peak abundance. The Gompertz function (Draper and Smith 1981) was used to describe the cumulative egg and larval abundance distribution. Thus, the inflection point was not constrained to be the mid-point of the sigmoid curve as is the case in the frequently used logistic and probit curves. The particular form of the Gompertz function used (Gendron 1989) was:

$$
\begin{equation*}
C_{t}=A \times \exp (-\exp [-k \times\{t-p\}]) \tag{1}
\end{equation*}
$$

where
$C_{t}=$ cumulative density at time $t$
$t=$ time in days from the date when the eggs
or larvae first occur
$A=$ total or asymptotic cumulative density
$p=$ inflection point in days since first occurrence date
$\kappa=$ shape parameter.
The origin of the time scale was set to the date when the eggs or larvae generally first appeared in the waters off MPS. Least-squares estimates, standard errors, and asymptotic $95 \%$ confidence intervals of the $A, p$, and $\kappa$ parameters were obtained by fitting the above equation to the cumulative abundance data using nonlinear regression methods (Proc NLIN; SAS Institute Inc. 1990). The cumulative data were obtained as the running sums of the weekly geometric means of the abundance data per unit volume. A geometric mean of weekly densities was used in analyses because the data generally followed a lognormal distribution (McConnaughey and Conquest 1993) and weekly sampling frequencies varied.
A "density" function was derived algebraically by calculating the first derivative of the Gompertz function (Eq. 1) with respect to time. This density function, which directly describes abundance over time (abundance curve), has the form:

$$
\begin{equation*}
\mathrm{d}_{t}=A^{\cdot} \times k \times \exp (-\exp [-k \times\{t-p\}]-k \times[t-p]) \tag{2}
\end{equation*}
$$

where $A^{\prime}$ equals $7 \times A$ because the cumulative densities were based on weekly ( 7 -day period) geometric means, $d_{t}$ is density on day $t$ and all the other parameters are as described in Equation 1. Daily entrainment was estimated by multiplying these daily densities $d_{t}$ by the daily volume of cooling water that passed through MPS. Annual entrainment estimates were determined by summing all daily estimates during the period of occurrence.

# Results and Discussion 

Species Composition and Relative Annual Abundance

A total of 121 fish taxa was recorded as eggs, larvae, juveniles, or adults from collections made in the trawl, seine, and ichthyoplankton programs as part of the Fish Ecology monitoring studies at MPS from June 1976 through May 2001. This total includes fishes collected at present and former sampling stations during this 25 -year period, with 111 taxa taken by trawl, 49 by seine, and 58 enumerated in ichthyoplankton samples (Appendix I). Anchovies (mostly bay anchovy) accounted for almost half ( $46 \%$ ) of the larvae collected at the MPS discharges (station EN) from June 1976 through May 2001, although they only accounted for $5 \%$ of the larvae this year. Overall, winter flounder (Pseudopleuronectes americanus) comprised another $15 \%$ and thirteen other taxa made up most of the remainder (Table 1). Although Atlantic menhaden (Brevoortia tyrannus) accounted for almost $8 \%$ of all larvae entrained, since June 1976, its larvae comprised $63 \%$ of the total larval entrainment during 2000-01. Cunner, tautog, and anchovies accounted for nearly $87 \%$ of the eggs collected. Silversides (Atlantic silverside, M. menidia, and inland silverside, M. beryllina) dominated ( $80 \%$ ) the seine catch at station JC (Appendix II); another $10 \%$ were killifishes (striped killifish, Fundulus majalis, and mummichog, $F$. heteroclitus), with fourspine stickleback (Apeltes quadracus) and Atlantic menhaden each making up $4 \%$ of the catch. There

TABLE 1. Taxonomic composition of ichthyoplankton collected at EN (as a percentage of the total) from June 1976 through May 2001 for larvae and May 1979 through September 2000 for eggs.

| Taxon | Larvae | Eggs |
| :--- | :---: | :---: |
| Anchoa spp. | 45.7 | 4.9 |
| Pseudopleuronectes americanus | 14.9 |  |
| Brevoortia tyrannus | 7.7 |  |
| Ammodytes americanus | 7.4 |  |
| Myoxocephalus aenaeus | 5.9 |  |
| Pholis gunnellus | 2.8 |  |
| Tautogolabrus adspersus | 2.4 | 54.0 |
| Tautoga onitis | 2.1 | 27.6 |
| Enchelyopus cimbrius | 1.5 |  |
| Ulvaria subbifurcata | 1.1 |  |
| Liparis spp. | 1.1 |  |
| Clupea harengus | 1.0 |  |
| Syngnathus fuscus | 0.9 |  |
| Scophthalmus aquosus | 0.7 |  |
| Peprilus triacanthus | 0.7 |  |
|  |  |  |

was a dramatic increase in the numbers of Atlantic menhaden in the seine catch, with the catch during the past 3 years accounting for $72 \%$ of the 25 -year total catch. Eight taxa accounted for about $80 \%$ of the total catch at the three trawl stations (Appendices III-V). These were the winter flounder (42\%), scup (Stenotomus chrysops; 11\%), silversides (mostly Atlantic silverside; 7\%), windowpane (Scophthalmus aquosus; 5\%), grubby (4\%), skates (mostly little skate, Raja erinacea; also the winter skate, R. ocellata, and clearnose skate, R. eglanteria; 3\%), anchovies (mostly bay anchovy; 3\%) and cunner (3\%). Eight species had higher abundance at least one station during 2000-01 than during any other year of the 25 -year time period. These were the tautog at two stations (NR and JC), scup, and smallmouth flounder (Etropus microstomus) at NR; northern puffer (Sphoeroides maculatus), rock gunnel (Pholis gunnellus), and striped cusk-eel (Ophidion marginatum) at JC, and summer flounder (Paralichithys dentatus) and lumpfish (Cyclopterus lumpus) at IN (Appendices III-V). The total catch of fish over the 25 -year period was similar between IN $(123,639)$ and NR $(120,230)$, with the catch at each of these stations roughly twice the total of

71,208 fish at JC. Temporal changes in the composition of dominant taxa (cited above) collected in the trawl and ichthyoplankton programs during the 25year period were compared using $\Delta$-means. Change's in the composition of seine catches were not examined because silversides have always dominated the catch and these changes will be discussed below. In trawl sampling, winter flounder and scup had the largest annual $\triangle$-mean CPUE during each report year, with silversides, grubby, windowpane, and skates also relatively numerous (Table 2). The 2000$01 \Delta$-mean CPUE for winter flounder (7.7) was the second lowest recorded, with only 1997-98 having a lower $\Delta$-mean CPUE. Alternatively, the $\Delta$-mean CPUE for scup (34.3) was the third highest in the 25year period. The 2000-01 $\Delta$-mean CPUE for silversides ( 0.8 ) and grubby ( 0.5 ) were relatively low with silversides having the second lowest and grubby the lowest recorded. The $\Delta$-means of 1.4 for skates and 2.8 for windowpane were within the range of historic catches (Table 2).
Each species and life stage of ichthyoplankton collected at EN occurred during specific time periods. Therefore, $\Delta$-mean densities ( $\mathrm{no} . / 500 \mathrm{~m}^{3}$ ) were computed from data taken during standardized

TABLE 2. The annual $\triangle$-meana ${ }^{4}$ CPUE (no. $/ 0.69 \mathrm{~km}$ ) of the most abundant fish collected by trawl at JC, IN, and NR for each report year from June 1976 through May 2001.


[^2]periods of occurrence for each taxon. Because of seasonal occurrence and sample processing, some species have means determined through 2001 and others through 2000 (Table 3). Cunner eggs were always the most abundant of the fish eggs collected.

Tautog had the second most abundant eggs entrained. A large decline in anchovy egg abundances occurred after 1984, with the 2000 $\Delta$-mean density ( $<1.0$ ) being the lowest.

TABLE 3. The annual $\Delta$-mean density ( $\mathrm{no} / 500 \mathrm{~m}^{3}$ ) of the most abundant fish eggs and larvae collected at EN for each year from June 1976 through May 2000.

| Taxon | 1976 | 1977 | 1978 | 1979 | 1980 | 1981 | 1982 | 1983 | 1984. | 1985 | 1986 | 1987 | 1988 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| EGGS* |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T. adspersus | - | - | - | 8,348 | 8,379 | 7,326 | 7,874 | 7,580 | 6,707 | 12,842 | 2,579 | 5,017 | 5,388 |
| T. onitis | - | - | - | 1,648 | 3,741 | 2,501 | 3,561 | 2,372 | 1,817 | 4,027 | 2,833 | 2,972 | 2,211 |
| Anchoa spp. | - | - | - | 1,558 | 999 | 769 | 499 | 2,415 | 3,631 | 118 | 586 | 64 | 32 |
| larvae |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Anchoa spp. | 1,152 | 931 | 483 | 2,168 | 2,430 | 5,768 | 816 | 1,421 | 302 | 1,102 | 1,244 | 126 | 359 |
| P. americamus | - | 106 | 143 | 114 | 285 | 129 | 233 | 297 | 210 | 180 | 87 | 109 | 116 |
| A. americanus ${ }^{\text {c }}$ | - | 94 | 318 | 119 | 111 | 136 | 21 | 27 | 18 | 9 | 3 | 13 | 41 |
| B. tyrannus | 5 | 3 | 3 | 1 | 2 | 1 | 9 | 18 | 2 | 38 | 2 | 2 | 5 |
| M. aenaeus | - | 41 | 38 | 36 | 38 | 107 | 72 | 68 | 50 | 68 | 34 | 29 | 95 |
| P. gunnellus | $\stackrel{-}{-}$ | 13 | 13 | 16 | 58 | 58 | 27 | 13 | 14 | 14 | 22 | 4 | 26 |
| T. adspersus | 29 | 58 | 1 | 13 | 58 | 78 | 31 | 49 | 4 | 12 | 4 | 5 | 9 |
| T. onitis | 37 | 36 | 1 | 11 | 46 | 83 | 44 | 33 | 3 | 15 | 3 | 7 | 17 |
| E. cimbrius | 2 | 8 | 6 | 8 | 6 | 1 | 6 | 13 | 5 | 8 | 8 | 12 | 45 |
| Liparis spp. | 27 | 30 | 10 | 16 | 22 | 5 | 13 | 8 | 36 | 1 | 4 | 42 | 18 |
| U. subbifurcata | - | 5 | 9 | 14 | 14 | 16 | 17 | 6 | 4 | 60 | 7 | 9 | 23 |
| $S$ S. fuscus | 3 | 7 | 4 | 9 | 8 | 13 | 7 | 9 | 9 | 5 | 4 | 6 | 7 |
| S. aquosus | 9 | 11 | 1 | 5 | 5 | 5 | 2 | 13 | 3 | 1 | 4 | 3 | 5 |
| P. triacanthus | 14 | 3 | 1 | 2 | 11 | 17 | 9 | 9 | 1 | 2 | 3 | <1 | 9 |
| C. harengus | - | 1 | 1 | 1 | <1 | 6 | 1 | 0 | 1 | $<1$ | 2 | 1 | 14 |
| Taxon | 1989 | 1990 | 1991 | 1992 | 1993 | 1994 | 1995 | 1996 | 1997 | 1998 | 1999 | 2000 | 2001 |
| EGGS ${ }^{\text {a }}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |
| T. adspersus | 6,255 | 7,269 | 6,987 | 2,776 | 4,535 | 8,722 | 4,266 | 8,801 | 3,610 | 1,459 | 3,530 | 8,245 | - |
| T. onitis | 3,373 | 1,942 | 2,040 | 1,189 | 1,394 | 1,350 | 1,807 | 2,323 | 587 | 897 | 1,373 | 2,322 | - |
| Anchoa spp. | 32 | 89 | 317 | 62 | 329 | 234 | 118 | 36 | 5 | 114 | 1 | <1 | - |
| LARVAE ${ }^{\text {b }}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Anchoa spp. | 619 | 1,122 | 799 | 178 | 203 | 475 | 181 | 175 | 131 | 106 | 129 | 65 | - |
| P. americanus | 203 | 106 | 99 | 381 | 21 | 142 | 224 | 81 | 255 | 137 | 182 | 328 | 296 |
| A. americanus ${ }^{\text {c }}$ | 31 | 24 | 7 | 18 | 28 | 43 | 63 | 18 | 11 | 28 | 13 | 53 | 12 |
| B. tyrannus | 47 | 16 | 81 | 37 | 8 | 44 | 56 | 145 | 23 | 28 | 58 | 293 | - |
| M. aenaeus | 63 | 30 | 24 | 58 | 34 | 48 | 43 | 85 | 140 | 55 | 39 | 45 | 188 |
| P. gunnellus |  | 6 | 3 | 15 |  | 28 | 17 | 41 | 22 | 42 | 12 | 22 | 21 |
| T. adspersus | 14 | 68 | 209 | 8 | 10 | 25 | 12 | 12 | 7 | 21 | 15 | 17 | - |
| T. onitis | 15 | 33 | 99 | 13 | 6 | 12 | 8 | 18 | 2 | 14 | 59 | 12 | - |
| E. cimbrius | 31 | 37 | 98 | 5 | 18 | 9 | 8 | 2 | 10 | 9 | 4 | 2 | - |
| Liparis spp. | 12 | 3 | 23 | 14 | 12 | 5 | <1 | 46 | 2 | 2 | 0 | 4 | - |
| U. subbifurcata | 41 | 51 | 34 | 28 | 2 | 18 | 8 | <1 | 18 | 1 | 10 | 1 | 1 |
| S. firscus | 5 | 3 | 5 | 3 | 6 | 4 | 6 | 2 | 2 | 2 | 4 | 3 | - |
| S. aquosus | 3 | 4 | 12 | 2 | 2 | 3 | 1 | 4 | 2 | 1 | 2 | 3 | - |
| P. triacanthus | 5 | 29 | 10 | 2 | 2 | 5 | 3 | 1 | 2 | 1 | 2 | 4 | - |
| C. harengus | 1 | 1 | 2 | 9 | 7 | 4 | 4 | 33 | 11 | 7 | 13 | 0.5 | 1 |

[^3]Abundance of most of the 15 dominant larval taxa was within the range of values found in previous years. Atlantic menhaden and grubby larval $\Delta$-mean densities were the highest ever recorded. However, larval anchovy $\Delta$-mean densities were at an historic low. On occasion, larvae of other species, particularly those of cunner, tautog, rock gunnel (Pholis gunnellus), fourbeard rockling (Enchelyopus cimbrius), and radiated shanny (Ulvaria subbifurcata) were relatively abundant.

## Entrainment Estimates

Entrainment of fish eggs and larvae in the condenser-cooling water system represents a direct impact from the operation of MPS. Most fish eggs collected at station EN were from cunner, tautog, and anchovies and the most numerous larvae were anchovies, winter flounder, American sand lance, grubby, and Atlantic menhaden (Table 1). The annual numbers of eggs and larvae entrained were related to their abundance at station EN and plant operations (i.c., cooling-water usage). MPS Units 2
and 3 were both operating during most of this report period. Unit 3 was shut down from February 3 to March 30, 2001 and Unit 2 operated throughout the year. Unit I was shut down permanently and is being decommissioned. The 2000 entrainment estimates of both cunner ( 4,802 million) and tautog eggs ( 2,149 million) were within the historic ranges (Table 4). Anchovy egg entrainment was at a historic low ( $<1$ million) in 3 of the last 4 years.
The 2000 entrainment estimate of 68 million larval anchovies was less than the long-term average of 433 million (Table 5), this was due to the low density of anchovy larvae this year. The 2001 entrainment estimate of 12 million American sand lance larvae was within the range of previous estimates. Entrainment estimates of grubby larvae in 2001 (178 million) and Atlantic menhaden larvae in 2000 (474 million) were the highest recorded. This was due to their high densities during these years (see Table 3). Entrainment of winter flounder larvae totaled 376 million in 2001 and this loss is evaluated in the Winter Flounder Studies section of this report.

TABLE 4. Estimated number of cunner, tautog, and anchovy eggs entrained each year from 1979 through 2000 at MPS and the volume of cooling water on which the entrainment estimates were based.

| Year | Cunner |  | Tautog |  | Anchovies |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $\begin{aligned} & \text { No. entrained } \\ & \left(\times 10^{\prime \prime}\right) \end{aligned}$ | $\begin{aligned} & \text { Volume }\left(\mathrm{m}^{3}\right)^{\prime} \\ & \left(\times 10^{\prime \prime}\right) \end{aligned}$ | $\begin{aligned} & \text { No. entrained } \\ & \left(\times 10^{\prime}\right) \end{aligned}$ | $\begin{aligned} & \text { Volume }\left(m^{3}\right)^{2} \\ & \left(\times 10^{6}\right) \end{aligned}$ | $\begin{aligned} & \text { No. entrained } \\ & \left(\times 10^{\prime}\right) \end{aligned}$ | $\begin{aligned} & \text { Volume }\left(m^{3}\right)^{2} \\ & \left(\times 10^{6}\right) \end{aligned}$ |
| 1979 | 1,053 | 684 | 448 | 684 | 324 | 578 |
| 1980 | 1,660 | 762 | 969 | 762 | 87 | 604 |
| 1981 | 1,547 | 769 | 1,398 | 769 | 287 | 673 |
| 1982 | 2,078 | 808 | 1,253 | 808 | 210 | 680 |
| 1983 | 1,899 | 753 | 1,019 | 753 | 371 : | 601 |
| 1984 | 2.135 | 779 | 1,323 | 779 | 883 | 659 |
| 1985 | 2,814 | 790 | 1,720 | 790 | 27 | 675 |
| 1986 | 2,855 | 1,772 | 3,750 | 1,772 | 522 | 1,484 |
| 1987 | 4,090 | 1,687 | 3,597 | $\because 1,687$ | 31 | 1,402 |
| 1988 | 4,294 | 1,843 | 2,693 | 1,843 | 15 | 1,558 |
| 1989 | 4.307 | 1,547 | 3,002 | 1,547 | 5 | 1,327 |
| 1990 | 3,634 | 1,724 | 2,101 | 1,724 | 27 | 1,465 |
| 1991 | 4.117 | 1,198 | 1,521 | 1,198 | 105 | 978 |
| 1992 | 2.648 | 1,484 | 1,338 | 1,484 | 18 | 1,216 |
| 1993 | 5,421 | 1,655 | 2,062 | 1,655 | 228 | 1,357 |
| 1994 | 6,146 | 1,627 | 2,069 | - 1,627 | 177 | 1,389 |
| 1995 | 5,527 | 1,536 | 2,562 | 1,536 | 30 : | 1,327 |
| 1996 | 872 | $\cdots 264$ | . 313 | 264 | 4 | 168 |
| 1997 | 569 | 212 | - 111 | 212 | $<1$ | 161 |
| 1998 | 581 | 745 | 496 | 745 | 53 | 644 |
| 1999 | 1,959 | 1,200 | 1,168 | $\therefore 1,200$ | <1 | 1,004 |
| 2000 | - 4,802 | : 1,402 | 2,149 | - $\therefore 1,1,402$ | $<1$ | 1,173 |

[^4]TABLE 5. Estimated number of anchovy, winter flounder, American sand lance and grubby larvae entrained each year from 1976 through 2001 at MPS and the volume of cooling water on which the entrainment estimates were based.

| Year | Anchovies |  | Winter Flounder |  | American sand lance ${ }^{\text {a }}$ |  | Grubby |  | Atlantic menhaden |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | No. entrained ( $\times 10^{6}$ ) | Volume $\begin{gathered} \left(\mathrm{m}^{3}\right)^{b} \\ \left(\times 10^{6}\right) \end{gathered}$ | No. entrained ( $\times 10^{6}$ ) | $\begin{aligned} & \text { Volume } \\ & \left(\mathrm{m}^{3}\right)^{6} \\ & \left(\times 10^{6}\right) \end{aligned}$ | No. entrained (x106) | $\begin{aligned} & \text { Volume } \\ & \left(\mathrm{m}^{3}\right)^{b} \\ & \left(\times 10^{6}\right) \end{aligned}$ | No. entrained ( $\times 10^{6}$ ) | $\begin{aligned} & \text { Volume } \\ & \left(\mathrm{m}^{3}\right)^{\mathrm{b}} \\ & \left(\times 10^{6}\right) \end{aligned}$ | No. entrained ( $\times 10^{6}$ ) | $\begin{aligned} & \text { Volume } \\ & \left(\mathrm{m}^{3}\right)^{6} \\ & \left(\times 10^{6}\right) \end{aligned}$ |
| 1976 | 378 | 591 | 121 | 629 | 16 | 800 | 12 | 644 | 3 | 890 |
| 1977 | 414 | 549 | 29 | 443 | 80 | 888 | 31 | 667 | 2 | 857 |
| 1978 | 161 | 631 | 80 | 390 | 176 | 710 | 11 | 477 | 3 | 1,035 |
| 1979 | 805 | 533 | 44 | 342 | 111 | 810 | 20 | 564 | $<1$ | 895 |
| 1980 | 877 | 486 | 168 | 562 | 112 | 974 | 32 | 742 | 2 | 700 |
| 1981 | 1,448 | 610 | 45 | 373 | 75 | 627 | 42 | 418 | 2 | 999 |
| 1982 | 449 | 525 | 164 | 638 | 27 | 927 | 48 | 650 | 14 | 859 |
| 1983 | 613 | 463 | 211 | 541 | 30 | 996 | 55 | 745 | 20 | 753 |
| 1984 | 167 | 581 | 84 | 508 | 18 | 877 | 39 | 673 | 4 | 989 |
| 1985 | 690 | 579 | 80 | 469 | 8 | 900 | 35 | 627 | 44 | 816 |
| 1986 | 1,093 | 1,206 | 123 | 1,064 | 4 | 1,766 | 54 | 1,341 | 5 | 1,776 |
| 1987 | 117 | 1,114 | 165 | 1,193 | 35 | 1,962 | 52 | 1,453 | 2 | 1,785 |
| 1988 | 383 | 1,284 | 184 | 1,173 | 86 | 1,813 | 112 | 1,299 | 7 | 2,009 |
| 1989 | 530 | 1,158 | 167 | 889 | 44 | 1,708 | 68 | 1,198 | 208 | 1,885 |
| 1990 | 978 | 1,221 | 133 | 1,174 | 45 | 2,004 | 47 | 1,415 | 37 | 1,939 |
| 1991 | 451 | 754 | 116 | 750 | 7 | 1,572 | 31 | 1,090 | 56 | 1,085 |
| 1992 | 151 | 979 | 492 | 1,075 | 22 | 1,526 | 73 | 1,157 | 52 | 1,601 |
| 1993 | 197 | 1,050 | 42 | 1,387 | 48 | 2,011 | 52 | 1,429 | 28 | 1,800 |
| 1994 | 509 | 1,193 | 173 | 920 | 65 | 1,819 | 56 | 1,203 | 70 | 1,923 |
| 1995 | 175 | 1,198 | 214 | 1,006 | 89 | 1,783 | 58 | 1,221 | 91 | 1,860 |
| 1996 | 24 | 133 | 51 | 472 | 18 | 1,056 | 41 | 997 | 23 | 211 |
| 1997 | 17 | 125 | 76 | 173 | 3 | 227 | 28 | 180 | 5 | 213 |
| 1998 | 63 | 571 | 84 | 358 | 11 | 474 | 22 | 348 | 35 | 944 |
| 1999 | 136 | 905 | 146 | 748 | 14 | 1,112 | 49 | 844 | 140 | 1,532 |
| 2000 | 68 | 1,008 | 331 | 1,003 | 88 | 1,613 | 47 | 1,131 | 474 | 1,647 |
| 2001 | - ${ }^{\text {c }}$ | - | 376 | 963 | 12 | 1,513 | 178 | 1,024 | - | - |

- Includes data from December of the previous calendar year.
b Volume was determined from the condenser cooling water flow at MPS during the season of occurrence for each taxa.
c Not calculated because larvae occur after the end of the report period (May 2001).


## Impingement

Impingement impacts at the MPS were substantially abated due to the shutdown of MPS Unit 1 ( $23 \%$ cooling-water reduction) and the installation of aquatic organism returns at Units 2 and 3 (NUSCO 1988c, 1994b; DNC 2001b). A return system was included as part of the intake structure during the construction of Unit 3 ; in early 2000 a sluiceway was backfitted at Unit 2. A 1-year study to evaluate its operation and effectiveness was completed in July 2001. A report of the results submitted to the CTDEP concluded that the system worked as designed and successfully returned impinged marine organisms back to Long Island Sound (DNC 2001b).

## Selection of Potentially Impacted Taxa

Additional data analyses were completed for selected taxa that were identified as potentially impacted, either because of their prevalence in entrainment samples or because of possible influence by the thermal discharge. Taxa potentially vulnerable to entrainment include American sand lance, anchovies, Atlantic menhaden, grubby, cunner, tautog, and winter flounder. Silversides were also selected for further analysis because their distribution in Jordan Cove may be affected by the MPS thermal discharge. The $\Delta$-mean densities (no. $/ 500 \mathrm{~m}^{3}$ ) of eggs and larvae, $\Delta$-mean CPUE in trawl ( $\mathrm{no} . / 0.69 \mathrm{~km}$ ), and seine CPUE ( $\mathrm{no} . / 30 \mathrm{~m}$ ) were calculated for abundant life stages of these selected taxa. Information on the winter flounder is presented in a separate section of this report (see Winter Flounder Studies) and is not included among the fishes discussed below.

## American sand lance

The American sand lance is a schooling fish common in estuaries and in coastal inshore waters from Labrador to Chesapeake Bay (Richards 1982; Nizinski et al. 1990). Sand lance have a life span of 5 to 9 years, but populations are typically dominated by the first three age groups (Reay 1970). Sexual maturation occurs at age-1 or 2 with adults spawning once a year, predominantly between November and March (Richards 1963, 1982; Scott and Scott 1988; Westin et al. 1979; Grosslein and Azarovitch 1982). Eggs are demersal and adhesive (Fritzsche 1978; Smigielski et al. 1984). Embryonic and larval development is lengthy (Smigielski et al. 1984).
American sand lance were collected in all three Fish Ecology sampling programs, although relatively few juveniles and adults were taken by seine or trawl (Appendices II-V). Most sand lance were collected : as larvae in winter and spring at station EN. Abundance appeared to be considerably less in February than during January or March through May. Larval abundance peaked in the late 1970s and early 1980s, followed by a rapid decline in 1982 (Table 6; Fig. 2). The $\Delta$-mean density of larvae at EN during 2001 was 12 , which was within the range of the historic data, although among the lowest of the timeseries. There was no significant trend ( $p=0.07$ ) detected over the 25 -year time series based on the Mann-Kendall test.
Sand lance larvae are the fourth most abundant larval taxon collected at station EN (Table 1). Annual entrainment estimates, which were based on abundance and related to cooling-water flow during the larval season, ranged from 3 to 176 million (Table 5). The 2001 entrainment estimate of 12 million larvae and the MPS cooling-water flow during the period of occurrence were comparable to the historic range (Table 5).
Declines in sand lance abundance during the 1980s were also apparent in other areas of the Northwest Atlantic Ocean. Larval densities in LIS over a 32 year period (1951-83) were highest in 1965-66 and 1978-79; the latter peak was also evident throughout the entire range of American sand lance (Monteleone et al. 1987). This high abundance persisted throughout the Northwest Atlantic until 1981 and the decline that followed appeared to be inversely correlated with that of Atlantic herring (Clupea harengus) and Atlantic mackerel (Scomber scombrus) (Nizinski et al. 1990). These two fishes prey heavily upon sand lance, and following their decreases from overfishing in the 1970s sand lance'

TABLE 6. The annual $\Delta$-mean ${ }^{2}$ density ( $\mathrm{no} . / 500 \mathrm{~m}^{3}$ ) and $95 \%$ confidence interval of American sand lance larvae collected at EN from June 1976 through May 2001.


- Data seasonally restricted to December-May and year given for $\Delta$-means is that of the latter 5 months.
increased in abundance (Sherman et al. 1981; Monteleone et al. 1987). In more recent years, both of these predators have again become more abundant and sand lance abundance hás decreased (Stephenson and Kornfield 1990; Smith and Morse 1993; NMFS 1995). Given the large abundance changes of American sand lance along the Atlantic coast, effects of MPS operation are difficult to ascertain, but are likely small in comparison to the large-scale natural


Fig. 2. Annual $\Delta$-mean densities (data points) with the 5-. year moving average (line) of American sand lance larvae at station EN from 1977 through 2001.
fluctuations typically associated with this species. The sand lance is a short-lived species and, as such, any changes in abundance from the MPS outages should have been readily detected. No changes were evident in 2001 and larval abundance was within its historic range.

## Anchovies

The bay anchovy, one of the most abundant fishes found along the Atlantic Coast (McHugh 1967), is usually the dominant summer ichthyoplankton species found within its range (Leak and Houde 1987). This species ranges from Mexico to Cape Cod and occasionally into the Gulf of Maine (Hildebrand 1943; Bigelow and Schroeder 1953; Grosslein and Azarovitch 1982). Bay anchovies are common in nearshore and estuarine waters during warmer months, but move offshore in winter (Vouglitois et al. 1987). Chesapeake Bay bay anchovies were found to have little genetic variation, indicating a lack of stock structure, likely due to enormous population size and considerable movements and mixing of stocks (Morgan et al. 1995). This homogeneity serves to lessen the effects of any localized impacts. Although the striped anchovy also occurs from Uruguay to Nova Scotia, its occurrence north of the Chesapeake Bay is variable and the striped anchovy is usually found farther offshore than the bay anchovy (Hoese and Moore 1977; Smith 1985). The eggs of the two species can be readily distinguished and since 1979, when eggs were first identified to species in MPS samples, about $96 \%$ of the anchovy eggs collected at station EN were determined to be bay anchovy. Therefore, most of the anchovies collected in the Fish Ecology programs were likely bay anchovy, even if only identified to genus.
The bay anchovy can mature at 2.5 to 3 months and individuals spawn repeatedly during the summer (Luo and Musick 1991). In LIS, spawning takes place at depths of 20 m or less from May through September, with a peak during June and July (Wheatland 1956; Richards 1959). Spawning appears to be correlated with high zooplankton abundances (Castro and Cowen 1991; Peebles et al. 1996) and warm water temperatures (Zastrow et al. 1991). Eggs are pelagic and hatch in about 24 hours at water temperatures of $27^{\circ} \mathrm{C}$ (Kuntz 1914). Mortality rates of eggs and larvae are relatively high (Leak and Houde 1987; Houde et al. 1994; Dorsey et al. 1996), particularly from predation by ctenophores and jellyfish (Govoni and Olney 1991; Purcell et al.
1994). Juveniles and adults are important forage for many recreationally and commercially important fishes (Vouglitois et al. 1987) and also have high mortality rates (Newberger and Houde 1995).
Anchovies were collected in all three programs, but rarely by seine and only sporadically by trawl (Appendices II-V). Juvenile anchovies resulting from the summer spawn were typically captured by trawl from August through October. Anchovy abundance in some years varied by two orders of magnitude and the majority were collected in only a few years of sampling at the three trawl stations (Appendices III-V).
Anchovies dominated larval collections and their eggs ranked third in abundance (Table 1). Eggs and larvae occurred in collections made from June through September. Annual egg and larval abundances were significantly correlated (Spearman's rank-order correlation coefficient $\mathrm{r}=0.71$; $\mathrm{p}=$ 0.001 ). In 2000, the $\Delta$-mean density of $<1.0$ for anchovy eggs was the lowest recorded as was the $\Delta$ mean density of 65 of larvae (Table 7; Fig. 3). Other incidences of large annual changes in bay anchovy

TABLE 7. The annual $\Delta$-mean ${ }^{2}$ density (no $/ 500 \mathrm{~m}^{3}$ ) and $95 \%$ confidence interval of anchovy eggs and larvae collected at EN from June 1976 through May 2000.

| Year | $\Delta$-mean density <br> of eggs | $\Delta$-mean density <br> of larvae |
| :---: | :---: | :---: |
| 1976 |  | $1,152 \pm 419$ |
| 1977 |  | $931 \pm 408$ |
| 1978 | $1,558 \pm 807$ | $483 \pm 206$ |
| 1979 | $999 \pm 689$ | $2,168 \pm 908$ |
| 1980 | $769 \pm 273$ | $2,430 \pm 1,249$ |
| 1981 | $499 \pm 202$ | $816 \pm 3,326$ |
| 1982 | $2,415 \pm 2,130$ | $1,421 \pm 530$ |
| 1983 | $3,631 \pm 3,528$ | $302 \pm 165$ |
| 1984 | $118 \pm 72$ | $1,102 \pm 453$ |
| 1985 | $586 \pm 366$ | $1,244 \pm 893$ |
| 1986 | $64 \pm 19$ | $126 \pm 69$ |
| 1987 | $32 \pm 26$ | $359 \pm 216$ |
| 1988 | $32 \pm 29$ | $619 \pm 416$ |
| 1989 | $89 \pm 88$ | $1,122 \pm 853$ |
| 1990 | $317 \pm 306$ | $799 \pm 801$ |
| 1991 | $62 \pm 70$ | $178 \pm 80$ |
| 1992 | $329 \pm 260$ | $203 \pm 103$ |
| 1993 | $234 \pm 204$ | $475 \pm 410$ |
| 1994 | $118 \pm 154$ | $181 \pm 117$ |
| 1995 | $36 \pm 29$ | $175 \pm 196$ |
| 1996 | $5 \pm 5$ | $131 \pm 90$ |
| 1997 | $114 \pm 123$ | $106 \pm 80$ |
| 1998 | $1 \pm 1$ | $129 \pm 135$ |
| 1999 | $<1 \pm 1$ | $65 \pm 39$ |
| 2000 |  |  |
|  |  |  |

[^5]

Fig. 3. Annual $\Delta$-mean densities (data points) with moving averages (line) of 4 years for anchovy eggs (1979-2000) and 5 years for anchovy larvae (1976-2000) at station EN.
egg abundance were observed in LIS during 1952-55 (Richards 1959) and in Barnegat Bay, NJ during 1976-81 (Vouglitois et al. 1987). At MPS, egg and larval densities decreased by an order of magnitude since the early to mid-1980s and have remained relatively low, except for larvae in 1990. Results of a Mann-Kendall test indicated that densities of both eggs and larvae have had significant negative trends (eggs $p=0.001$, slope $=-43$; larvae $p=0.001$, slope $=-50.4$ ) for the entire time-series.
The entrainment of eggs and larvae by MPS probably represents the greatest direct operational impact on anchovies that spawn in the Millstone area. In 2000, the entrainment estimate of eggs ( $<1$ million) was similar to the 1999 estimate. Both estimates were comparable to the 1997 estimate when the units were shut down and cooling-water usage was an order of magnitude lower. The larval entrainment estimate ( 68 million) was higher than the 1996, 1997 and 1998 estimates, but was much lower than in the past (Table 5). Densities of both anchovy eggs and larvae had significant negative trends. Bay anchovies appear to be experiencing a regional decline in abundance. A sharp drop in abundance was measured over the past decade in Narragansett Bay, Rhode Island (Tim Lynch, RI DEM, pers.
comm.). The Maryland DNR Juvenile Finfish Seining Survey data also indicated that the bay anchovy population declined dramatically in Chesapeake Bay after 1993 (Price 1999). The decrease in anchovy eggs and larvae seen in recent years is probably reflective of regional decreases in abundance. The fact that there is little genetic variation among the abundant anchovy stocks along the Atlantic Coast serves to lessen any localized impacts of localized fluctuations in abundance.

## Atlantic menhaden

The Atlantic menhaden is a pelagic species that filter-feeds phytoplankton, zooplankton and detritus (Lewis and Peters 1994). Atlantic menhaden have been reported from Nova Scotia to southeastern Florida, but is most common from the Gulf of Maine to northern Florida (Reintjes 1969; Ahrenholz 1991). It is a coastal migratory species considered to be estuarine-dependent, with young-of-the-year found in coastal bays, rivers, and sounds, often penetrating to upstream limits of saline water (Reintjes and Pacheco 1966). The Atlantic menhaden supports the largest commercial fishery along the Atlantic coast with landings primarily used for fish meal, oil, and solubles (Vaughan and Smith 1988; Ahrenholtz et al. 1987). This species also serves as forage for many marine fishes, birds, and mammals. Although it has a life span of up to 10-12 years, the current fishery is highly dependent upon pre-spawning fish; most females mature during their second year of life at about 180 mm in length (Lewis et al. 1987). Based on its movements and distribution, the Atlantic menhaden is treated as a single stock for management, although Epperly (1989) found some evidence for northern and southern subpopulations that were based on meristic and biochemical differences.
Atlantic menhaden exhibit a distinct annual cycle of movements (Nicholson 1971, 1978; Dryfoos et al. 1973; Kroger and Guthrie 1973). More localized movements have been attributed to responses to gradients in the densities of their phytoplankon prey (Friedland et al. 1989, 1996). Fish are found throughout most of the range during summer, although the population is generally stratified by age and size, with the oldest and largest individuals found farthest north. Adult fish begin to move southwards in September, followed by juveniles later in fall and early winter. Atlantic menhaden overwinter in waters off the Carolinas and southward and begin relatively rapid movements northward in March and
early April and re-occupy their summer range by about June. Spawning takes place at night (Ferraro 1980b), and can occur during any month of the year, but the presence of larvae and juveniles in specific waters is largely associated with the annual movement/reproductive cycle. Most spawning takes place in the ocean where early larval growth and development occurs. Some spawning also takes place in major estuaries (Ahrenholz 1991). Spawning was reported to occur within LIS and adjacent waters from late spring through early fall (Dietrich 1979; Ferraro 1981; Powell and Phonlor 1986). The bimodal distribution of larvae found in the MPS samples, however, suggested two distinct spawning periods in LIS, a larger one in the summer and a smaller event during fall. Ferraro (1981) also reported a similar annual bimodal spawning periodicity for Atlantic menhaden in the Peconic Bays, NY. Allantic menhaden found offshore of New England and New York waters begin spawning in September and spawning activity increases as fish move progressively south in October and November, peaking off North Carolina in winter. Eggs and larvae are tolerant of a relatively wide range of salinities and temperatures (Ferraro 1980a). Small larvae are moved landward by Ekman transport (Nelson et al. 1977), and further development occurs in nearshore coastal and estuarine waters, which are sought out by juveniles. Larval and juvenile movements (including diel vertical migration) have been found to be affected by environmental cues, such as light (Forward et al. 1993, 1996) and temperature (Friedland and Haas 1988; De Vries et al. 1995) with increased swimming activity and movement towards surface waters occurring at night.
Atlantic menhaden were taken in all three monitoring programs, but were collected irregularly by trawl or seine. Most ( $90 \%$ ) Atlantic menhaden were collected in 6 out of 25 years of sampling at the trawl stations (Appendices III-V). The majority (53\%) of the Atlantic menhaden collected by trawl were found at NR. The Atlantic menhaden collected by seine at JC were mostly ( $>70 \%$ ) taken in the last three years (Appendix II). The shore-zone seine catch of Atlantic menhaden was composed entirely of young-of-the year, reflecting the landward migration of juveniles seeking coastal and estuarine areas as nursery zones as previously described by Nelson et al. (1977).
Some eggs of Atlantic menhaden were identified in entrainment collections taken from April 1979 through September 1992, but they were never abundant and made up less than $0.1 \%$ of all fish eggs
taken (NUSCO 1994a). This suggested that most spawning in LIS did not occur near MPS, even though the Atlantic menhaden was the third most abundant larval taxon collected (Table 1). The $\Delta$-mean density of 293 in 2000 for Atlantic menhaden was the highest recorded and larvae were twice as abundant as the second highest year of 1996 (Table 8). Larval densities have increased since 1988, with many density estimates during recent years above the moving average, especially in 2000 (Fig. 4). Based on a Mann-Kendall test, a significant ( $p=0.001$; slope $=2.5$ ) increasing trend was detected during the entire operational period of 19762000.

The foremost operational impact by MPS to Atlantic menhaden is the entrainment of larvae. In 2000, the entrainment estimate of 474 million larvae was the highest recorded (Table 5). The large increase in Atlantic menhaden abundance over the past decade indicated that MPS has had minimal or no impact on their numbers, with abundance in northern waters likely increasing because of reduced fishing mortality along the Atlantic coast.

TABLE 8. The annual $\Delta$-mean ${ }^{\text {a }}$ density (no. $/ 500 \mathrm{~m}^{\mathbf{l}}$ ) and $95 \%$ contidence interval of Atlantic menhaden larvae collected at EN from June 1976 through May 2000.

| Year | S-Inean density |
| :---: | :---: |
|  |  |
| 1976 | $5 \pm 1$ |
| 1977 | $3 \pm 1$ |
| 1978 | $3 \pm 1$ |
| 1979 | $1 \pm 1$ |
| 1980 | $2 \pm 1$ |
| 1981 | $1 \pm 0.4$ |
| 1982 | $9 \pm 3$ |
| 1983 | $18 \pm 10$ |
| 1984 | $2 \pm 1$ |
| 1985 | $38 \pm 22$ |
| 1986 | $2 \pm 1$ |
| 1987 | $2 \pm 1$ |
| 1988 | $5 \pm 4$ |
| 1989 | $47 \pm 64$ |
| 1990 | $16 \pm 12$ |
| 1991 | $81 \pm 140$ |
| 1992 | $37 \pm 28$ |
| 1993 | $8 \pm 4$ |
| 1994 | $44 \pm 76$ |
| 1995 | $56 \pm 60$ |
| 1996 | $145 \pm 431$ |
| 1997 | $23 \pm 18$ |
| 1998 | $28 \pm 60$ |
| 1999 | $58 \pm 63$ |
| 2000 | $293 \pm 373$ |
|  |  |

[^6]

Fig. 4. Annual $\Delta$-mean densities (data points) and the 5 year moving average (line) of Atlantic menhaden larvae at station EN from 1976 through 2000.

## Silversides

The Atlantic and inland silversides are sympatric along most of the east coast of North America and reside in bays, estuaries, and salt marshes. The Atlantic silverside ranges from the Gulf of St. Lawrence to northern Florida (Conover 1992) and the inland silverside from Cape Cod to South Carolina (Johnson 1975). Both species are abundant, but in general, the Atlantic silverside is more numerous than the inland silverside, except in low salinity waters (Bengtson 1984, 1985). Both species are important trophic links between zooplankton and larger piscivorous fishes and birds. Most silversides mature as yearlings and only live 1 to 2 years. Spawning begins at water temperatures of $9-12^{\circ} \mathrm{C}$ and occurs during the day at high tide on a semilunar cycle (Middaugh 1981; Conover and Ross 1982; Jessop 1983; Conover and Kynard 1984). Adhesive eggs are laid in shallow water on vegetation (Conover and Kynard 1984). Larvae are planktonic, but remain near spawning areas. Sex is indeterminate until fish reach 8 to 21 mm in length and sex ratio is affected by prevailing water temperature during development (Conover and Kynard 1981; Conover and Fleisher 1986). Growth of young is rapid and mean lengths of Atlantic silverside can exceed 90 mm by November with fish from early spawns larger at any given age than late-spawned fish (Conover 1979; Bengtson et al. 1987). Atlantic silversides migrate offshore during : winter, but remain in waters within 40 km of shore and in depths of less than 50 m (Conover and Murawski 1982). In contrast, inland silversides have small home ranges (Hoff 1972). Siversides typically suffer high (>99\%) overwinter mortality (Conover and Ross 1982; Conover 1992).

Nearly all silversides collected in the trawl monitoring program were Atlantic silverside, with less than $0.1 \%$ identified as the inland silverside since 1981, when the two species were first distinguished in the MPS sampling. More than $80 \%$ of the silversides collected by seine were Atlantic silverside, although relative proportions varied from year to year. Collectively, silversides dominated the seine catch ( $80 \%$ ) -at JC (Appendix II) and were taken by trawl ( $7 \%$ of the trawl catch) in winter (Appendices III-V), but were rarely ( $<0.1 \%$ of larvae collected since 1976) found in ichthyoplankton collections.

Silversides were taken at the three trawl stations from October through February after individuals left the shore areas to overwinter in deeper waters. Annual trawl abundances fluctuated considerably (Table 9). Few silversides were taken at IN in 200001. At JC and NR, the $\Delta$-mean CPUE was within the historical ranges. The 2000-01 $\Delta$-mean CPUE was below the moving average at all three stations (Fig. 5 ). A significant negative trend was found at IN ( $p=$ 0.001 , slope $=-0.4$ ) over the 25 -year period, but no trend was found at NR or JC, based on the MannKendall test.

TABLE 9. The annual $\triangle$-mean ${ }^{1}$ CPUE (no. $/ 0.69 \mathrm{~km}$ ) and $95 \%$ confidence interval of Atlantic silverside collected by trawl at three selected stations from June 1976 through May 2001.

| Year | $\cdot \mathrm{IN}$ | JC | NR |
| :---: | :---: | :---: | :---: |
| $1976-77$ | $15 \pm 16$ | $13 \pm 20$ | $77 \pm 283$ |
| $1977-78$ | $29 \pm 92$ | $6 \pm 612$ | $10 \pm 21$ |
| $1978-79$ | $60 \pm 105$ | $9 \pm 8$ | $2 \pm 1$ |
| $1979-80$ | $42 \pm 276$ | $6 \pm 17$ | $4 \pm 6$ |
| $1980-81$ | $8 \pm 17$ | $4 \pm 5$ | $3 \pm 4$ |
| $1981-82$ | $6 \pm 9$ | $1 \pm 0$ | $6 \pm 8$ |
| $1982-83$ | $2 \pm 4$ | $1 \pm 2$ | $12 \pm 5$ |
| $1983-84$ | $2 \pm 4$ | $4 \pm 1$ | $1 \pm 6$ |
| $1984-85$ | $2 \pm 6$ | $5 \pm 11$ | $1 \pm 1$ |
| $1985-86$ | $7 \pm 8$ | $6 \pm 8$ | $3 \pm 6$ |
| $1986-87$ | $5 \pm 3$ | $8 \pm 7$ | $110 \pm 222$ |
| $1987-88$ | $3 \pm 5$ | $2 \pm 2$ | $15 \pm 27$ |
| $1988-89$ | $2 \pm 1$ | $1 \pm 0$ | $25 \pm 14$ |
| $1989-90$ | $1 \pm 1$ | $2 \pm 2$ | $12 \pm 20$ |
| $1990-91$ | $1 \pm 0$ | $1 \pm 0$ | $17 \pm 11$ |
| $1991-92$ | $12 \pm 10$ | $2 \pm 1$ | $19 \pm 7$ |
| $1992-93$ | $115 \pm 156$ | $7 \pm 2$ | $24 \pm 15$ |
| $1993-94$ | $5 \pm 7$ | $3 \pm 2$ | $3 \pm 2$ |
| $1994-95$ | $1 \pm 1$ | $4 \pm 3$ | $2 \pm 2$ |
| $1995-96$ | $2 \pm 1$ | $2 \pm 2$ | $1 \pm 1$ |
| $1996-97$ | $4 \pm 2$ | $0.4 \pm 0.1$ | $13 \pm 8$ |
| $1997-98$ | $1 \pm 1$ | $0.5 \pm 0.5$ | $2 \pm 5$ |
| $1998-99$ | $1 \pm 1$ | $21 \pm 65$ | $7 \pm 11$ |
| $1999-00$ | $3 \pm 3$ | $8 \pm 7$ | $7 \pm 4$ |
| $2000-01$ | $<1 \pm 0$ | $1 \pm 2$ | $3 \pm 7$ |
|  |  |  |  |

2. Data seasonally restricted to November-February at IN and NR and October-January at JC.


Fig. 5. Annual $\Delta$-mean CPUE (data points) and the 5 -year moving average (line) of Atlantic silversides taken by trawl at stations NR, IN, and JC from 1976-77 through 2000-01.

The $\Delta$-mean CPUE for Atlantic and inland silversides taken by seine during 2000 ( 135 and 9 , respectively) was within the previously observed range of values (Table 10). In 2000, the abundance of Atlantic silverside exceeded that of the inland silverside (Fig. 6). The 2000 annual $\Delta$-mean CPUE value for Atlantic silverside was greater than the moving average, however, inland silverside abundance was less than its moving average (Fig. 7). Abundance data of both species showed no significant trends for the 20 years tested based on the Mann-Kendall test.

To determine if any changes in size distributions have occurred, the length-frequencies for seine and trawl catches (expressed as percentages) were examined comparing the current year with historic data (Fig. 8). More $60-80 \mathrm{~mm}$ Atlantic silversides

TABLE 10. The annual $\Delta$-meana CPUE ( $n$ o 30 m ) and $95 \%$ confidence interval of Atlantic silverside and inland silverside collected by seine at JC from 1981 through 2000.

| Year | Atlantic silverside | Inland silverside |
| :---: | :---: | :---: |
|  | $152 \pm 251$ | $3 \pm 3$ |
| 1981 | $111 \pm \pm 162$ | $6 \pm 16$ |
| 1982 | $397 \pm 598$ | $88 \pm 243$ |
| 1983 | $29 \pm 24$ | $3 \pm 2$ |
| 1984 | $19 \pm 12$ | $4 \pm 8$ |
| 1985 | $172 \pm 385$ | $14 \pm 21$ |
| 1986 | $109 \pm 90$ | $3 \pm 2$ |
| 1987 | $96 \pm 108$ | $27 \pm 54$ |
| 1988 | $70 \pm 93$ | $14 \pm 16$ |
| 1989 | $83 \pm 80$ | $133 \pm 234$ |
| 1990 | $38 \pm 11$ | $74 \pm 37$ |
| 1991 | $78 \pm 55$ | $43 \pm 27$ |
| 1992 | $60 \pm 73$ | $5 \pm 5$ |
| 1993 | $37 \pm 28$ | $63 \pm 64$ |
| 1994 | $87 \pm 73$ | $60 \pm 114$ |
| 1995 | $78 \pm 60$ | $27 \pm 23$ |
| 1996 | $93 \pm 78$ | $5 \pm 37$ |
| 1997 | $108 \pm 79$ | $4 \pm 3$ |
| 1998 | $135 \pm 201$ | $9 \pm 13$ |
| 1999 |  |  |
| 2000 |  |  |
|  |  |  |

and smaller ( $<40 \mathrm{~mm}$ ) inland silversides were observed in seine samples this year when compared to the historic length range. The silversides caught by trawl were similar in size as those found in previous years, although there was a somewhat higher proportion of $80-100 \mathrm{~mm}$ fish than in the past.
The primary impact of the operation of MPS on silversides may be the incursion of the thermal plume into the shore-zone area, potentially causing disruption in spawning activity and influencing the distribution of juveniles and adults. The thermal


Fig. 6. Annual $\Delta$-mean CPUE ( $n o . / 30 \mathrm{~m}$ ) of Atlantic silverside (dashed line) and inland silverside (solid line) taken by seine at station JC from 1981 through 2000.


Fig. 7. Annual $\Delta$-mean CPUE (data points) and the 5 -year moving average (line) of Atlantic silverside and inland silverside taken by seine at station JC from 1981 through 2000. (Note that the vertical scales differ among the graphs).
increase is only $0.8^{\circ} \mathrm{C}$ at the JC station (NUSCO 1988b). Elevated summer temperatures in Jordan Cove appear to be more directly related to solar heating of the shallow sand flats (such as at the JC seine site) than to the MPS thermal plume (see the Eelgrass section of this report). Therefore, it appears unlikely that a small increase in water temperature could affect the reproductive success of a species that ranges as far south as northern Florida.

## Grubby

The grubby is a demersal fish found in shallow waters along the Atlantic coast from the Gulf of St. Lawrence to New Jersey. It tolerates a wide range of temperature and salinity (Bigelow and Schroeder 1953). Individuals reside in protected shallow water on mud or sand bottoms, peat reefs, and in eelgrass beds (Ennis 1969; Lazzari et al.1989) and occur throughout the year near MPS. Similar to the winter flounder, grubby produce blood plasma antifreeze proteins and can remain active in very low water temperatures (Reisman et al. 1987). Female grubby reach maturity within one year (Lazzari et al. 1989). Grubby spawn throughout the winter and have a demersal, adhesive egg with an incubation time of 40


Fig. 8. Length-frequency distribution ( $20-\mathrm{mm}$ length intervals) of Atlantic silverside and inland silverside taken by seine at station JC and Allantic silverside taken by trawl at stations NR, IN, and JC from June 1976 through May 1999 and from June 2000 through May 2001.
to 44 days at a water temperature of $4.6-6^{\circ} \mathrm{C}$ (Lund and Marcy 1975; Lazzari et al. 1989). Richards (1959) reported larvae present in LIS from February through April and Laroche (1982) noted that they are more abundant near the bottom than at the surface. The grubby is a small fish and has no sport or commercial value. Given its protective spines and cryptic coloration, it probably also has limited forage value. Grubby prey upon many small fishes and benthic invertebrates (Lazzari et al. 1989; Levin 1991).

The grubby was the fifth-most abundant larval fish collected at EN, and primarily occurred in samples from February through May. Grubby accounted for $5.9 \%$ of all larvae collected from June 1976 through May 2001 (Table 1). The $\Delta$-mean density of larvae
for 2001 of $188 / 500 \mathrm{~m}^{3}$ was the highest recorded (Table 11) and well above the 5 -year moving average (Fig. 9). Because of fluctuating annual abundance, a Mann-Kendall test revealed no significant temporal trend in larval abundance. The entrainment estimate of 178 million larvae in 2001 also the highest recorded because of the high density (Table 5).
Predominantly a shallow-water fish, the grubby was the fifth-most abundant fish taken by trawl at the three stations with $59 \%$ of the total collected at NR (Appendices III-V). In 2000-01, catches at all three trawl stations were below the moving average, with the catch at IN being the lowest recorded (Table 12; Fig. 10). No significant trends in abundance were found. Smaller grubbies were generally taken by trawl in 2000-01 based on the percent lengthfrequency distributions (Fig. 11).

Entrainment of grubby larvae is the primary direct plant impact on the resident grubby population. However, larval grubby abundance was at a historic high. This unexploited species has been among the most stable of the fishes residing near MPS. Because

TABLE 11. The annual $\Delta$-mean² density (no. $/ 500 \mathrm{~m}^{3}$ ) and $95 \%$ conlidence interval of grubby larvae collected at EN from June 1976 through May 2001.

| Year | $\Delta$-mean density |
| :---: | :---: |
| 1977 | $41 \pm 9$ |
| 1978 | $38 \pm 9$ |
| 1979 | $36 \pm 7$ |
| 1980 | $38 \pm 7$ |
| 1981 | $107 \pm 27$ |
| 1982 | $72 \pm 13$ |
| 1983 | $68 \pm 19$ |
| 1984 | $50 \pm 15$ |
| 1985 | $68 \pm 23$ |
| 1986 | $34 \pm 10$ |
| 1987 | $29 \pm 7$ |
| 1988 | $95 \pm 35$ |
| 1989 | $63 \pm 18$ |
| 1990 | $30 \pm 8$ |
| 1991 | $24 \pm 6$ |
| 1992 | $58 \pm 17$ |
| 1993 | $34 \pm 9$ |
| 1994 | $48 \pm 16$ |
| 1995 | $43 \pm 15$ |
| 1996 | $85 \pm 37$ |
| 1997 | $140 \pm 60$ |
| 1998 | $55 \pm 19$ |
| 1999 | $39 \pm 24$ |
| 2000 | $45 \pm 19$ |
| 2001 | $188 \pm 89$ |
|  |  |

[^7]

Fig. 9. Annual $\Delta$-mean densities (data points) with a 5 year moving average (line) of grubby larvae at station EN from 1977 through 2001.
the grubby is a short-lived species that matures in 1 year, any changes in abundance resulting from MPS would have been apparent. Since grubby abundance has been stable, it is likely that the plant has had little or no effect on the population.

TABLE 12. The annual $\Delta$-mean* CPUE (no. 0.69 km ) and $95 \%$ confidence interval of grubby collected by trawl at three selected stations from June 1976 through May 2001.

| Year | NR | JC | IN |
| :--- | ---: | ---: | ---: |
| $1976-77$ | $0.9 \pm 0.3$ | $0.6 \pm 0.2$ | $0.6 \pm 0.1$ |
| $1977-78$ | $0.5 \pm 0.1$ | $2.2 \pm 0.5$ | $1.1 \pm 0.2$ |
| $1978-79$ | $1.2 \pm 0.2$ | $2.0 \pm 0.6$ | $0.7 \pm 0.2$ |
| $1979-80$ | $3.3 \pm 0.9$ | $0.7 \pm 0.1$ | $0.9 \pm 0.2$ |
| $1980-81$ | $3.8 \pm 1.1$ | $\mathrm{I} .1 \pm 0.2$ | $2.1 \pm 0.6$ |
| $1981-82$ | $7.5 \pm 2.5$ | $1.0 \pm 0.2$ | $2.3 \pm 0.6$ |
| $1982-83$ | $11.7 \pm 2.7$ | $1.4 \pm 0.2$ | $2.2 \pm 0.5$ |
| $1983-84$ | $4.1 \pm 0.8$ | $1.7 \pm 0.3$ | $1.7 \pm 0.3$ |
| $1984-85$ | $5.9 \pm 1.2$ | $1.6 \pm 0.3$ | $0.9 \pm 0.2$ |
| $1985-86$ | $2.3 \pm 0.5$ | $1.4 \pm 0.3$ | $0.7 \pm 0.1$ |
| $1986-87$ | $7.2 \pm 2.3$ | $1.1 \pm 0.2$ | $0.9 \pm 0.2$ |
| $1987-88$ | $3.7 \pm 1.2$ | $1.2 \pm 0.2$ | $1.1 \pm 0.2$ |
| $1988-89$ | $10.5 \pm 2.3$ | $1.0 \pm 0.1$ | $1.4 \pm 0.3$ |
| $1989-90$ | $3.6 \pm 2.0$ | $0.4 \pm 0.1$ | $1.0 \pm 0.3$ |
| $1990-91$ | $8.0 \pm 2.0$ | $0.4 \pm 0.1$ | $0.8 \pm 0.2$ |
| $1991-92$ | $3.4 \pm 0.5$ | $0.5 \pm 0.1$ | $1.0 \pm 0.2$ |
| $1992-93$ | $6.2 \pm 2.0$ | $1.4 \pm 0.3$ | $1.9 \pm 0.3$ |
| $1993-94$ | $2.2 \pm 3.0$ | $0.7 \pm 0.5$ | $1.9 \pm 3.8$ |
| $1994-95$ | $3.7 \pm 1.6$ | $2.9 \pm 1.1$ | $1.6 \pm 0.6$ |
| $1995-96$ | $1.9 \pm 1.0$ | $0.7 \pm 0.3$ | $0.7 \pm 0.2$ |
| $1996-97$ | $5.5 \pm 1.3$ | $0.7 \pm 0.1$ | $2.3 \pm 0.5$ |
| $1997-98$ | $3.7 \pm 1.6$ | $0.9 \pm 0.3$ | $2.7 \pm 1.1$ |
| $1998-99$ | $3.3 \pm 1.0$ | $1.8 \pm 0.6$ | $1.3 \pm 0.5$ |
| $1999-00$ | $0.8 \pm 0.2$ | $1.2 \pm 0.4$ | $0.9 \pm 0.4$ |
| $2000-01$ | $0.7 \pm 0.3$ | $0.6 \pm 0.3$ | $<0.1 \pm 0$ |
|  |  |  |  |
|  |  |  |  |

[^8]

Fig. 10. Annual $\Delta$-mean CPUE (data points) with a 5 -year moving average (line) of grubby taken by trawl at stations NR, IN, and JC from 1976-77 through 2000-01. (Note that the vertical scales differ among the graphs).


Fig. 11. Length-frequency distribution ( $10-\mathrm{mm}$ length ${ }^{\prime}$ intervals) of grubby taken by trawl at stations NR, IN, and ' JC from June 1976 through May 2000 and from June 2000 through May 2000.

## Cunner

The cunner occurs from Newfoundland to Chesapeake Bay (Scott and Scott 1988), and prefers structural habitats, such as rocks, pilings, eelgrass or mussel beds, and macroalgae. Cunner are inactive at night (Bradbury et al. 1997) and become torpid when water temperatures fall below $5-8^{\circ} \mathrm{C}$ (Green and Farwell 1971; Olla et al. 1975; Dew 1976). Individual cunner maintain highly localized home ranges (Green 1975; Olla et al. 1975; Gleason and Recksiek 1988), may establish defended territories (Pottle and Green 1979b), and most do not undertake extensive movements (Green and Farwell 1971; Olla et al. 1979; Lawton et al. 1996). Most cunner live 5 to 6 years, with a maximum age of about 10 , which is less than one-third of the life span of the closely related tautog (Dew 1976; Regan et al. 1982).
Cunner mature at age- 1 to 2 and spawn during May through September from afternoon into the evening (Johansen 1925; Dew 1976; Pottle and Green 1979a; Green et al. 1985). Larger ( $>20 \mathrm{~cm}$ ) males may spawn with individual females, whereas smaller (80180 mm ) males and females spawn in large ( $30-$ $150+$ ) groups (Pottle et al. 1981). Lawton et al. (1996) reported all cunner larger than 65 mm observed in western Cape Cod Bay to be mature. The pelagic eggs hatch in 2 to 6 days, depending upon water temperature (Williams 1967; Dew 1976). Williams et al. (1973) noted that only about $5 \%$ of cunner eggs survive to hatching. Newly-hatched larvae are 2 to 3 mm in length, metamorphose by 10 mm , and settle into preferred habitats (Miller 1958; Levin 1991). Tupper and Boutilier (1997) reported that settlement was not affected by habitat type or adult density, but post-settlement survival, recruitment success, and adult densities were positively correlated with habitat complexity. They concluded that habitat-mediated post-settlement processes played an important role in cunner population dynamics.
The cunner has little commercial value and is generally not sought after by sport fisherman, although numerous individuals are often caught while fishing for other species (MacLeod 1995). Regionally, declining trends in adult cunner abundance have been observed in LIS (CTDEP 1998; Smith et al. 1989), Cape Cod Bay (Lawton et al. 1994), and Mount Hope Bay (MRI 1994).

In the MPS area, cunner eggs and larvae are present primarily from May through July. Among the early developmental stages collected at station EN, cunner eggs were the most abundant of all egg taxa, whereas larvae were less common, ranking only seventh
overall (Table 1). The $2000 \Delta$-mean density of eggs was $8,245 / 500 \mathrm{~m}^{3}$ and was within the range of historic data (Table 13), as was the egg entrainment estimate of 4,802 million (Table 4). The 2000 annual $\Delta$-mean egg density was above the moving average (Fig. 12), while the corresponding larval $\Delta$-mean density of $17 / 500 \mathrm{~m}^{3}$ was within the historic range and approximately equal to the moving average (Table 13; Fig. 12). Based on the Mann-Kendall test, no significant temporal trends were detected for cunner eggs and larvae.
Cunner were rarely taken by seine at JC (Appendix II), but juveniles and adults were caught by trawl, mostly from spring through summer. More cunner were taken at IN $(4,270)$ than at JC $(3,866)$, with relatively few (602) found at NR (Appendices III-V). However, during the past 3 years, many more cunner were collected at JC $(1,760)$ than at IN (112). The $2000-01$ annual $\Delta$-mean CPUE values for IN (0.9) remained below catches made in the late 1970s and early 1980s, whereas the $2000 \Delta$-mean CPUE value for JC (13.9) was the highest recorded there (Table 14). Annual $\Delta$-mean CPUE at JC was well above the

TABLE 13. The annual $\Delta$-mean density (no. $/ 500 \mathrm{~m}^{3}$ ) and $95 \%$ confidence interval of cunner eggs and larvae collected at EN from June 1976 through May 2000.

| Year | $\Delta$-mean density <br> of eggs | $\Delta$-mean density <br> of larvae |
| :--- | :---: | :---: |
| 1976 |  | $29 \pm 14$ |
| 1977 |  | $58 \pm 28$ |
| 1978 |  | $1 \pm 0$ |
| 1979 | $8,349 \pm 4,601$ | $13 \pm 5$ |
| 1980 | $8,379 \pm 3,788$ | $58 \pm 19$ |
| 1981 | $7,326 \pm 3,890$ | $78 \pm 36$ |
| 1982 | $7,874 \pm 2,359$ | $31 \pm 14$ |
| 1983 | $7,580 \pm 6,106$ | $49 \pm 26$ |
| 1984 | $6,707 \pm 4,494$ | $4 \pm 2$ |
| 1985 | $12,842 \pm 6,156$ | $12 \pm 10$ |
| 1986 | $2,579 \pm 1,460$ | $4 \pm 1$ |
| 1987 | $5,017 \pm 3,536$ | $5 \pm 3$ |
| 1988 | $5,388 \pm 3,608$ | $9 \pm 4$ |
| 1989 | $6,255 \pm 5,302$ | $14 \pm 12$ |
| 1990 | $7,269 \pm 7,198$ | $68 \pm 61$ |
| 1991 | $6,987 \pm 5,482$ | $209 \pm 157$ |
| 1992 | $2,776 \pm 1,654$ | $8 \pm 4$ |
| 1993 | $4,535 \pm 2,612$ | $10 \pm 6$ |
| 1994 | $8,722 \pm 9,644$ | $25 \pm 18$ |
| 1995 | $4,266 \pm 3,222$ | $12 \pm 9$ |
| 1996 | $8,801 \pm 6,043$ | $12 \pm 6$ |
| 1997 | $3,610 \pm 3,157$ | $7 \pm 4$ |
| 1998 | $1,458 \pm 6,313$ | $21 \pm 18$ |
| 1999 | $3,530 \pm 3,044$ | $15 \pm 11$ |
| 2000 | $8,245 \pm 8752$ | $17 \pm 14$ |
|  |  |  |
|  |  |  |
| Data seasonally restricted to May-August for eggs and June- |  |  |
| August for larvac. |  |  |
| 9 |  |  |



Fig. 12. Annual $\Delta$-mean densities (data points) with moving averages (line) of 4 years for cunner eggs (19792000) and 5 years for cunner larvae (1976-00) at station EN. (Note that the vertical scales differ between the graphs).
moving average, while catches at IN just above the moving average (Fig. 13). Based on the MannKendall test no significant trend in abundance was detected at JC. However, catches exhibited a significant decreasing trend at $\operatorname{IN}(p<0.001$; slope $=-0.3)$. Contributing to the decrease at IN was the mid-1983 removal of a cofferdam that was in place during the construction of the Unit 3 intake structure. This rock cofferdam provided good habitat for cunner and may have increased their availability to sampling by trawl at the nearby IN station (NUSCO 1997). The MannKendall test was again executed for annual catches (1984-2001) at $\operatorname{IN}$ after the coffer dam was removed and no trend ( $p=0.8$; slope $=0$ ) was detected.
Ages of cunner were assigned based on an agelength key provided by Serchuk (1972) to determine an age-frequency distribution of fish collected by trawl. Percent length-frequency distributions were determined for past years (1976-99) and 2000-01 (Fig. 14). The size distributions in 2000-01 differed from the historic data with almost $80 \%$ of the cunner caught during 2000-01 being young-of-the-year. This trend (higher percentage of small fish) has been consistent over the past decade.

TABLE 14. The annual $\triangle$-mean ${ }^{\text {a }}$ CPUE (no. 0.69 km ) and $95 \%$ confidence interval of cunner collected by trawl at two selected stations from June 1976 through May 2000.

| Year | IN | JC |
| :---: | :---: | :---: |
|  |  |  |
| 1976 | $26.0 \pm 19.0$ | $4.0 \pm 2.0$ |
| 1977 | $24.0 \pm 23.0$ | $3.0 \pm 1.0$ |
| 1978 | $6.0 \pm 3.7$ | $3.0 \pm 1.4$ |
| 1979 | $29.0 \pm 23.0$ | $9.0 \pm 5.0$ |
| 1980 | $23.0 \pm 16.0$ | $6.0 \pm 2.0$ |
| 1981 | $12.0 \pm 10.0$ | $5.0 \pm 2.2$ |
| 1982 | $5.0 \pm 3.0$ | $4.0 \pm 2.0$ |
| 1983 | $3.0 \pm 1.3$ | $4.0 \pm 2.0$ |
| 1984 | $2.0 \pm 1.0$ | $2.0 \pm 1.0$ |
| 1985 | $1.0 \pm 0.6$ | $1.0 \pm 0.5$ |
| 1986 | $0.1 \pm 0.2$ | $0.5 \pm 0.4$ |
| 1987 | $0.2 \pm 0.2$ | $0.4 \pm 0.2$ |
| 1988 | $0.3 \pm 0.1$ | $3.0 \pm 3.4$ |
| 1989 | $0.9 \pm 0.4$ | $0.8 \pm 0.4$ |
| 1990 | $0.4 \pm 0.1$ | $0.9 \pm 0.2$ |
| 1991 | $0.4 \pm 0.1$ | $2.3 \pm 0.7$ |
| 1992 | $1.0 \pm 0.7$ | $1.4 \pm 0.5$ |
| 1993 | $0.1 \pm 1.1$ | $1.4 \pm 0.7$ |
| 1994 | $0.4 \pm 0.1$ | $0.8 \pm 0.5$ |
| 1995 | $0.8 \pm 0.4$ | $1.3 \pm 1.1$ |
| 1996 | $0.3 \pm 0.2$ | $0.2 \pm 0.2$ |
| 1997 | $0.5 \pm 0.3$ | $0.9 \pm 0.6$ |
| 1998 | $0.7 \pm 0.4$ | $7.2 \pm 5.2$ |
| 1999 | $0.4 \pm 0.2$ | $11.2 \pm 10.0$ |
| 2000 | $0.9 \pm 0.8$ | $13.9 \pm 10.0$ |
|  |  |  |

* Data seasonally restricted to May-August at (N and MaySeptember at JC.

The entrainment of eggs represents the greatest potential impact on the cunner population in the vicinity of MPS. Both egg abundance and larval density in 2000 were within the historic range, and young-of-the-year $\triangle$-mean CPUE at the JC trawl stations was at a historic high.

## Tautog

The tautog ranges from New Brunswick to South Carolina, but is most common from Cape Cod to the Delaware Capes (Bigelow and Schroeder 1953). Tautog are active during the day, but are quiescent during night (Olla et al. 1974). Adult tautog prefer rocky areas and similar reef-like habitats near shore from spring through fall. Juveniles are typically found in eelgrass beds and among macroalgae in coves and estuaries (Tracy 1910; Bigelow and Schroeder 1953; Wheatland 1956; Cooper 1965; Briggs and O'Conner 1971; Sogard and Able 1991; Hostetter and Munroe 1993; Dorf and Powell 1997). Young tautog are more restricted in their movements than adults and remain in close association with


Fig. 13. Annual $\triangle$-mean CPUE (data points) with the 5 year moving average (line) of cunner taken by trawl at stations IN and JC from 1976 through 2000. (Note that the vertical scales differ among the graphs).
shelter throughout the year (Olla et al. 1978). During winter, adults move to deeper ( $25-55 \mathrm{~m}$ ) water, while juveniles remain inshore to overwinter in a torpid state (Cooper 1965; Olla et al. 1974). Tautog are long-lived with maximum age reported for males of 34 years and 22 years for females (Chenoweth 1963; Cooper 1964). Adult growth rates have been estimated for several regions ranging from


Fig. 14. Length-frequency distribution by length (mm) and age (determined from age-length key of Serchuk 1972) of cunner taken by trawl at stations IN and JC from June 1976 through May 2000 and from June 2000 through May 2001.

Narragansett Bay to Virginia (Cooper 1964; Simpson 1989; Hostetter and Munroe 1993).
Male tautog mature when 2 to 3 years old and females at age-3 to 4; fecundity at size and age was reported by Chenoweth (1963). Adults return to nearshore waters in spring prior to spawning, with a high proportion of fish returning to the same spawning area each year (Cooper 1965). Spawning occurs during afternoon or early evening hours from mid-May until mid-August in LIS (Wheatland 1956; Chenoweth 1963; Olla and Samet 1977, 1978). The pelagic eggs hatch in 42 to 45 hours at $22^{\circ} \mathrm{C}$ (Williams 1967; Fritzsche 1978) and larvae are capable of feeding by 52 hours post-hatching (Schoedinger and Epifanio 1997). The pelagic larval stage lasts about 3 weeks and individuals settle on the bottom when they reach a size of about 17 mm (Sogard et al. 1992; Dorf 1994). Estimated growth rate during pre-settlement is about 0.75 mm per day and during post-settlement is about 0.5 mm per day (Sogard et al. 1992; Dorf 1994). In laboratory experiments, Laurence (1973) found that larval tautog may encounter potential energy deficits at higher ( $>19^{\circ} \mathrm{C}$ ) temperatures and Schoedinger and Epifanio (1997) found that prey densities strongly affected larval growth and development. Size at the end of the first growth season in Narragansett Bay (about 50 mm total length; Dorf 1994) was less than that found in a southerm New Jersey estuary ( 75 mm standard length; Sogard et al. 1992) and this was attributed to a longer growing season in southern waters.
Tautog were collected primarily as eggs in the ichthyoplankton entrainment program. Since 1979, eggs ranked second in abundance from collections at station EN (Table 1). Tautog and its sympatric species, cunner, have similar early life history characteristics, and annual $\Delta$-mean densities of their eggs were correlated (Spearman's rank-order correlation coefficient $r=0.47 ; p=0.03$ ). The $2000 \Delta-$ mean density for tautog eggs of $2,322 / 500 \mathrm{~m}^{3}$ was within the historic range and along with a similar value found in 1996, was the highest density seen since 1989 (Table 15). The $2000 \Delta$-mean was above the 5 -year moving average line (Fig. 15), although a significant negative trend $(p=0.006$, slope $=-88)$ was found over the 25 -year period according to a Mann-Kendall test. The 2000 annual entrainment estimate of 2.1 billion also fell within the historic range and was the highest estimate since 1995 (Table 4).

In contrast to eggs, tautog larvae were not a predominant taxon, ranking eighth in entrainment since 1976 (Table 1). Larval abundance in 2000 was

TABLE 15. The annual $\Delta$-mean² density (no. $/ 500 \mathrm{~m}^{\mathbf{3}}$ ) and $95 \%$ confidence interval of tautog eggs and larvae collected at EN from June 1976 through May 2000.

| Year | $\Delta$-mean density <br> of eggs | $\Delta$-mean density <br> of larvae |
| :---: | :---: | :---: |
| 1976 |  | $37 \pm 16$ |
| 1977 | $1,648 \pm 566$ | $36 \pm 17$ |
| 1978 | $3,741 \pm 1,482$ | $1 \pm 1$ |
| 1979 | $2,501 \pm 604$ | $11 \pm 5$ |
| 1980 | $3,561 \pm 1,400$ | $86 \pm 18$ |
| 1981 | $2,372 \pm 994$ | $44 \pm 36$ |
| 1982 | $1,817 \pm 504$ | $33 \pm 21$ |
| 1983 | $4,027 \pm 2,424$ | $3 \pm 2$ |
| 1984 | $2,833 \pm 1,212$ | $15 \pm 12$ |
| 1985 | $2,972 \pm 1,232$ | $3 \pm 2$ |
| 1986 | $2,211 \pm 906$ | $7 \pm 3$ |
| 1987 | $3,373 \pm 2,002$ | $17 \pm 10$ |
| 1988 | $1,942 \pm 978$ | $15 \pm 7$ |
| 1989 | $2,040 \pm 1,052$ | $33 \pm 28$ |
| 1990 | $1,189 \pm 462$ | $99 \pm 51$ |
| 1991 | $1,394 \pm 582$ | $13 \pm 4$ |
| 1992 | $1,350 \pm 658$ | $6 \pm 3$ |
| 1993 | $2,807 \pm 798$ | $12 \pm 8$ |
| 1994 | $583 \pm 2,032$ | $8 \pm 4$ |
| 1995 | $897 \pm 423$ | $18 \pm 16$ |
| 1996 | $1,373 \pm 636$ | $2 \pm 1$ |
| 1997 | $2,322 \pm 1336$ | $14 \pm 13$ |
| 1998 |  | $58 \pm 41$ |
| 1999 | $12 \pm<1$ |  |
| 2000 |  |  |

- Data seasonally restricted to May-August for eggs and JuneAugust for larvae.
just below the 5 -year moving average (Fig. 15). No trends in larval abundance were found. Annual abundances of tautog and cunner larvae were significantly correlated (Spearman's rank-order correlation coefficient $\mathrm{r}=0.84 ; \mathrm{p}<0.001$ ), indicating common processes that affected the abundance of larvae for both of these wrasses.
A preliminary study to examine initial and latent survival of entrained tautog and cunner eggs was conducted on four occasions during the summer of 2001 using a $0.5-\mathrm{m}$ plankton net having $0.333-\mathrm{mm}$ mesh. To collect eggs, the net was hauled vertically in the Unit 3 discharge and also in Niantic Bay near the MPS intakes to provide control samples on June 18, June 25, July 9, and July 23, 2001. Net contents were emptied into a bucket with seawater of the same temperature as the collection site and sorted in the laboratory. Sample processing was typically completed within 3 hours from the time of collection. Collected eggs were assessed for viability and live specimens were incubated in chambers with flowing seawater for up to 60 hours to assess latent mortality. Identification to species was deternined by examining hatched larvae at the completion of the


Fig. 15. Annual $\Delta$-mean densities (data points) with moving averages (line) of 4 years for tautog eggs (19792000) and 5 years for tautog larvae ( $1976-2000$ ) at station EN. (Note that the vertical scales differ between the graphs).
incubation period. Initial viability of wrasse eggs was consistently higher at the intake site than at the discharge site on all sample dates. The survival data is summarized in Table 16. On average, $53 \%$ of the eggs collected at the intake site were alive at time of collection compared to only $20 \%$ alive at the discharge site. Latent mortality was higher for
entrained eggs with only $4 \%$ of eggs surviving to hatch while approximately $20 \%$ of the eggs collected at the intake site hatched. Tautog larvae were more prevalent than cunner on all sampling dates, comprising between $60 \%$ and $77 \%$ of hatched larvae. Few samples were collected during this pilot study and more extensive sampling is planned for 2002.
Juvenile and adult tautog were sampled in two monitoring programs at MPS, trawl and lobster pot monitoring. Data were analyzed on an annual basis to compare catches from the two programs. Tautog were caught infrequently by trawl because they prefer rocky or reef habitats and are less vulnerable to this sampling; gear; annual $\Delta$-mean CPUE could not be calculated because of too many zero values. As an alternative, the annual sum of catches at the trawl stations were used as an index of abundance (Table 17). No significant trends were found for the combined catch of tautog at the three inshore trawl stations during the 26 -year period. However, the total catch of tautog at $\operatorname{IN}$ was at a historic high in 2001, and ranked third at JC and sixth at NR.
Tautog are routinely found in pots used in the lobster population monitoring program (see the Lobster Studies section for details). Since 1988, these fish were counted and measured to provide another index of tautog abundance. Total annual (May-October) catches were examined at each of the three lobster monitoring program stations (Jordan Cove, designated herein as JC; Intake, IN; and Twotree, TT) (Table 18). In the lobster pot program, no significant trends were found for the three stations combined, at IN, or TT. However, there was a significant increasing trend (slope $=6.4 ; \mathrm{p}<0.007$ ) in the catch of tautog at JC.

TABLE 16. Summary of wrasse egg entrainment survival study conducted in 2001.

| Date sampled | Collection site | ```Total no. eggs examined``` | No. viable 0-hours | No. hatched 12-hours | No. hatched 36-hours | No. hatched 60-hours | Initial \% viable | hatched | \% tautog |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 18 June | Intake | 173 | 90 | 0 | -2 | 33 | 52.0 | 20.2 | 60.3 |
|  | Discharge | 438 | 110 | . 0 | 8 | 5 | 25.1 | 3.0 | 39.7 |
| 25 June | Intake | 227 | 66 | 15 | 12 | 3 | 29.1 | 13.2 | 77.3 |
|  | Discharge | 437 | 77 | 18 | 7 | 2 | 17.6 | 6.2 | 22.7 |
| 9 July | Intake | 220 | 188 | 1 | 59 | 2 | 85.5 | 28.2 | 69.4 |
|  | Discharge | 583 | 192 | 1 | 7 | I | 32.9 | 1.5 | 30.6 |
| 23 July | Intake | 15 | 7 | 0 | 0 | 0 | 46.7 | 0 | n/a |
|  | Discharge | 262 | 10 | 0 | 0 | 0 | 3.8 | 0 | $n / \mathbf{a}$ |

TABLE 17. Total annual catch of tautog collected by trawl at three selected stations from 1976 through 2001.

| Year | NR | JC | IN | Total |
| ---: | ---: | ---: | ---: | ---: |
|  |  |  |  |  |
| 1976 | 46 | 113 | 76 | 195 |
| 1977 | 15 | 59 | 70 | 198 |
| 1978 | 27 | 56 | 83 | 169 |
| 1979 | 47 | 20 | 46 | 173 |
| 1980 | 25 | 24 | 28 | 91 |
| 1981 | 126 | 35 | 52 | 178 |
| 1982 | 80 | 19 | 40 | 167 |
| 1983 | 31 | 16 | 46 | 90 |
| 1984 | 5 | 27 | 47 | 67 |
| 1985 | 25 | 58 | 25 | 99 |
| 1986 | 100 | 33 | 13 | 183 |
| 1987 | 26 | 31 | 37 | 72 |
| 1988 | 50 | 23 | 25 | 118 |
| 1989 | 36 | 34 | 17 | 84 |
| 1990 | 89 | 44 | 13 | 140 |
| 1991 | 67 | 90 | 8 | 124 |
| 1992 | 22 | 15 | 51 | 26 |
| 1993 | 12 | 19 | 13 | 120 |
| 1994 | 129 | 73 | 28 | 92 |
| 1995 | 40 | 47 | 22 | 230 |
| 1996 | 46 | 26 | 24 | 109 |
| 1997 | 125 | 71 | 12 | 86 |
| 1998 | 138 | 27 | 208 |  |
| 1999 | 238 | 138 | 45 | 430 |
| 2000 | 324 | 261 | 119 |  |
| 2001 | 117 | 119 | 341 |  |
|  |  |  |  |  |
|  |  |  |  |  |

Length-frequency distributions of tautog caught by trawl and those found in the lobster pots were compared (Fig. 16). Ages were assigned to length categories based on age-length information for LIS (Simpson 1989). Young tautog, ages 1 and 2, accounted for a high proportion of the tautog caught in the trawl program, while those found in pots were dominated by fish of ages 3,4 and 5 , during which both males and females become mature. Lobster pots select for certain size-classes of tautog because

TABLE 18. Total annual catch of tautog collected in the lobster monitoring program at selected stations from 1988 through 2001.

| Year | IN | JC | TT | Total |
| :--- | :--- | ---: | ---: | ---: |
| 1988 | 47 | 15 | 40 | 102 |
| 1989 | 25 | 22 | 20 | 67 |
| 1990 | 27 | 8 | 11 | 46 |
| 1991 | 48 | 7 | 27 | 82 |
| 1992 | 32 | 11 | 21 | 64 |
| 1993 | 64 | 12 | 26 | 102 |
| 1994 | 29 | 8 | 43 | 80 |
| 1995 | 14 | 18 | 10 | 42 |
| 1996 | 23 | 134 | 63 | 210 |
| 1997 | 12 | 27 | 16 | 55 |
| 1998 | 91 | 80 | 64 | 235 |
| 1999 | 97 | 86 | 28 | 211 |
| 2000 | 85 | 80 | 48 | 213 |
| 2001 | 23 | 106 | 8 | 137 |
|  |  |  |  |  |



Fig. 16. Length-frequency distribution by length (mm) and age (determined from age-length key of Simpson 1989) of tautog taken by trawl and lobster from 1988 through 2001.
the $2.5 \mathrm{~cm}^{2}$ wire mesh do not retain smaller individuals when pots are hauled. Also, the $15-\mathrm{cm}$ diameter of the funnel entrances restricts the entry of most larger individuals.
The two monitoring programs sample different segments of the tautog population. The trawl stations are located in preferred juvenile habitat containing eelgrass beds and macroalgae bottoms in coves and estuaries (Sogard and Able 1991; Hostetter and Munroe 1993; Dorf and Powell 1997). Adult tautog favor rocky areas and similar reef-like habitats near shore from spring through fall (Tracy 1910; Bigelow and Schroeder 1953; Wheatland 1956; Cooper 1964; Briggs and O'Conner 1971) which are sampled by lobster pots. The trawl program provides an index of juvenile tautog in the MPS area, while the lobster pot catches provide a reliable index of newly recruited adults. It is interesting to note that in Figure 17, during some years, peaks in trawl catches preceded those in the lobster pot catches. Most notable was an increase in the 1995 trawl catch followed by a 1996 increase in lobster pot catches. More recently, this pattern did not hold as there were very high catches of juvenile tautog in trawl catches, but lobster pot catches have not yet reflected this abundance.
The greatest direct impact of MPS on tautog stocks is the entrainment of eggs. Tautog eggs are spawned during early evening, the pelagic eggs are dispersed rapidly from spawning sites by tidal transport, and hatch in less than 48 hours. Egg mortality is high immediately following spawning, probably from predation. Since 1993, special studies have been conducted on tautog eggs at MPS to assess the potential entrainment impact on tautog stocks. These tautog egg studies have included diel abundance changes in entrainment samples, spatial distribution in eastern LIS, daily spawning periodicity, and estimation of natural egg mortality. These studies were summarized in NUSCO (1997, 1998, 1999, 2000).


Fig. 17. Total annual catch of tautog in trawl and lobster pot collections from 1988 through 2001.

Recent work to determine seasonal fecundity estimates based on histological examination of tautog from Virginia waters demonstrated that tautog females are capable of spawning numerous times within a year. White (1996) determined that an individual female tautog spawned, on average, every 1.14 days during their spawning period and, based on this spawning frequency, estimated the total reproductive output for age-3 through 9 female tautog to range between 168,000 and $11,053,00$ eggs. Chenoweth (1963) reported maximum egg production per unit of ovary weight for fish 7 to 9 years old with production declining in fish age-16 and older. We attempted to verify seasonal fecundity estimates for tautog in Connecticut waters by sponsoring a study that was initiated during the summer of 2000 and continued in 2001. This research at the Avery Point campus of the University of Connecticut includes long-term observations of spawning male and female tautog in a laboratory holding facility as well as histological examination of gonads of field-collected fish. Results of this tautog reproductive biology study will be submitted following the completion of the research.

Based on trawl catches, juvenile tautog abundance has increased in recent years. In addition, numbers of newly recruited mature adults have not declined since the late 1980s, as indicated by the selective catch of 3- to 5-year-old tautog in lobster pots in the Millstone area. Therefore, changes in the relative proportion of juveniles and adults were probably unrelated to entrainment losses. In addition, the decline in juvenile and adult tautog abundance in LIS that began in the mid-1980s (Simpson et al. 1995) coincided with the decreasing trend in eggs collected at EN. If the decrease in adults was caused by entrainment losses, then the reduction in egg abundance should have lagged the decline of juveniles by several years because females do not mature until age-3 or 4 . Therefore, the lower
abundance of tautog eggs was probably due to a decline in the abundance of spawning adults from fishing mortality rather than from the operation of MPS. During the 1990s, the instantaneous fishing mortality rate for tautog was estimated at about 0.54 (annual fishing mortality of $42 \%$ ) and various survey biomass indices declined by more than half from the previous decade (ASMFC 1996). At present, tautog stocks are overfished and because of the long life and slow growth of this species, abundance should remain depressed until fishing mortality is reduced to less than half of current levels.' Recent increases in juvenile abundance may indicate that management measures may be working. However, due to the strong correlation found between the abundance of cunner and tautog larvae, environmental and biotic factors are likely also important in determining yearclass strength.

## Conclusions

Potential MPS :impacts on local fish populations include entrainment of eggs and larvae, impingement of juvenile and adults, and distributional changes as a result of the thermal discharge. During the 2000-01 report period, MPS Units 2 and 3 were both operating most of the time and Unit 1 was shutdown. Cooling-water volume was reduced as a result of the decommissioning of Unit 1 , which resulted in less entrainment of fish eggs and larvae and impingement of juveniles and adults, as entrainment and impingement rates are directly related to the amount of cooling-water used. Impingement impacts were further reduced at MPS with the installation of an aquatic organism return sluiceway in early 2000 at Unit 2. A 1-year study to evaluate its operation and effectiveness was completed in July 2001 and it was concluded that the . system worked as designed and successfully returned impinged marine organisms to Long Island Sound.
Detailed analyses were conducted on seven taxa that were most susceptible to MPS operational impact due to entrainment or effects of the thermal discharge. Analyses of these species generally focused on comparing temporal trends over the past two and one-half decades. No significant long-term trends were ' detected for juvenile and adult silversides by seines at JC, all life stages of grubby, cunner eggs and larvae, and tautog and sand lance larvae. Atlantic menhaden larvae showed a significantly increasing trend in abundance during the past 25 years. A significant negative trend was observed for silversides at the IN trawl station. Since
the mid-1980s, cunner have become less abundant at IN , exhibiting a significant negative trend, which may be related to removal of the Unit 3 rock cofferdam, a preferred habitat for this species. Despite the negative trend in abundance for cunner at trawl station IN, tautog abundance was at a historic high at station $\mathbb{I N}$, as was cunner abundance at station JC. Tautog eggs exhibited a negative trend in abundance. The large numbers of tautog and cunner eggs entrained at MPS did not appear to affect the future spawning stocks of these two fishes because the proportion of juvenile recruits relative to adults has increased.

Densities of both anchovy eggs and larvae showed significant negative trends. This year the $\Delta$-mean density for anchovy eggs was the lowest recorded. However, the $\Delta$-mean density of larvae was within the range of abundances found during the past 5 years. The bay anchovy appears to be experiencing a regional decline in abundance. A sharp drop in abundance was measured over the past decade in Narragansett Bay and populations declined dramatically in Chesapeake Bay after 1993. The bay anchovy is an important forage species for striped bass and recent increases in striped bass abundance along the Atlantic Coast may have contributed to the reduced numbers of bay anchovy.

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## APPENDICES

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APPENDIX I. List of fishes collected in the Fish Ecology sampling programs (June 1979-May 2001; all stations).


APPENDIX I. (continued).

| Scientific name | Common name | Trawl | Seine | Ichthyoplankton |
| :---: | :---: | :---: | :---: | :---: |
| Monacanthus hispidus | planehead filefish | * |  |  |
| Monocanthus spp. | filefish | * |  |  |
| Morone americana | white perch | - |  | * |
| Morone saxatilis | striped bass | * | * |  |
| Mugil cephalus | striped mullet | * | * | * |
| Mugil curema | white mullet |  | * |  |
| Mullus auratus | red goatfish | * |  |  |
| Mustelis canis | smooth dogfish | * |  |  |
| Myliobatis freminvillei | bullnose ray | * |  |  |
| Myoxocephalus aenaeus | grubby | * | * | * |
| Myoxocephalus octodecemspinosus | longhorn sculpin | * |  | * |
| Myoxocephalus spp. | sculpin | * |  |  |
| Ophidiidae | cusk-cels | * |  |  |
| Ophidion marginatum | striped cusk-eel | * | * | * |
| Ophidion welshi | crested cusk-eel |  |  |  |
| Opsanus tau | oyster toadfish |  |  |  |
| Osmerus mordax | rainbow smelt | * | * | * |
| Paralichthys dentatus | summer tlounder | * |  | + |
| Paralichthys oblongus | fourspot flounder | * |  | - |
| Peprilus triacanthus | butterfish | * | * | * |
| Petromyzon marinus | sea lamprey | * |  |  |
| Pholis gunnellus | rock gunnel | * | * | * |
| Pollachius virens | pollock | * |  | * |
| Pomatomus saltatrix | bluefish | * | * |  |
| Priacanthus arenatus | bigeye |  |  |  |
| Priacanthus cruentatus | glasseye snapper |  |  |  |
| Pristigenys alta | short bigeye | * |  |  |
| Prionotus carolinus | northern searobin | * | * | * |
| Prionotus evolans | striped searobin | * | * | * |
| Pseudopleuronectes americanus | winter flounder | * | * | * |
| Pungifius pungitius | ninespine stickleback | * | * | * |
| Raja eglanteria | clearnose skate | * |  |  |
| Raja erinacea | little skate |  |  |  |
| Raja ocellata | winter skate |  |  |  |
| Salmo truta | brown trout | * |  |  |
| Sciaenidae | drums |  |  | * |
| Scophthalmus aquosus | windowpane | * | * | * |
| Scomber scombrus | Atlantic mackerel | * |  | * |
| Scyliorhimus retifer | chain dogfish | * |  |  |
| Selar crumenopthalmus | bigeye scad | * |  |  |
| Selene setapinnis | Atlantic moonfish | + |  |  |
| Selene vomer | lookdown | , | * |  |
| Synodus foetens | inshore lizardfish | * |  |  |
| Sphyraena borealis | northern sennet | * |  |  |
| Sphoeroides maculatus | northern puffer | * | * | * |
| Squalus acanthias | spiny dogfish | * |  |  |
| Stenotomus chrysops | scup | * |  | * |
| Strongylura marina | Atlantic needlefish |  | * |  |
| Syngnathus fuscus | northern pipefish | - | * | - |
| Tautogolabrus adspersus | cunner | * | * | * |
| Tautoga onitis | tautog | * | * | * |
| Trachinotus falcatus | permit | * | * |  |
| Trachurus lathami | rouglı scad | * |  |  |
| Trachinocephalus myops | snakefish | * |  |  |
| Trinectes maculatus | logehoker | * |  |  |
| Ulvaria subbifurcata | radiated shanny | * |  | * |
| Upeneus parvus | dwarf goatfish | * |  |  |
| Urophycis chuss | red hake | * |  |  |
| Urophycis regia | spotied hake | * |  |  |
| Urophycis tenuis | white hake | * |  |  |
| Urophycis spp. | hake | * | * | * |

APPENDIX IL. Toal number of samples collected and number of fish caught by seine at station IC during each repor year from June 1976 through May 2000.

| Yezz | 76-77 | 77.78 | 78.79 | 79.80 | 80.81 | 81-82 | 82.83 | 83-84 | 84.85 | 85.86 | 86-87 | 87-88 | 88.89 | 89.90 | 90-91 | 91-92 | 92-93 | 93.94 | 94.95 | 95-98 | 96-97 | 97.98 | 98.99 | 99-00 | 00-01 | Toul |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Number of samples | 24 | 24 | 24 | 24 | 24 | 24 | 33 | 42 | 54 | 36 | 36 | s0 | 60 | 60 | 60 | 63 | 45 | 45 | 42 | 42 | 45 | 42 | 42 | 48 | 48 | 951 |
| Taxon ${ }^{\text {a }}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Menidia spp. | 37,264 | 15,962 | 505 | 443 | 5,834 | 1,587 | 2,006 | 7,667 | 1.056 | 484 | 3.103 | 4.174 | 3,548 | 2.369 | 5,611 | 3.585 | 4,504 | 2.234 | 2.922 | 4,443 | 6.196 | 7,999 | 3.629 | 3.990 | 4,191 | 135,308 |
| Fundulus spp. | 1,634 | 714 | 706 | 472 | 515 | 308 | 640 | 667 | 1.312 | 759 | 80 | 364 | 2,294 | 639 | 831 | 1,152 | 310 | 300 | 1.635 | 609 | 117 | 117 | 56 | 139 | 63 | 16,449 |
| B. Drannus | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 3 | 2 | 25 | 510 | 171 | 2 | 47 | 14 | 881 | 23 | 381 | 2.341 | 1,798 | 1,193 | 7.393 |
| A. quadrocus | 463 | 592 | 237 | 264 | 48 | 93 | 88 | 1.827 | 169 | 104 | 29 | 92 | 144 | 301. | 125 | 1.076 | 55 | 36 | 29 | 9 | 48 | 13 | 23 | 7 | 27 | 6.184 |
| C. variegatus | 42 | 284 | 33 | 16 | 7 | 33 | 133 | 27 | 25 | 23 | 1 | 2 | 10 | 2 | 14 | 1.169 | 8 | 0 | 53 | 11 | 2 | 3 | 0 | 4 | 1 | 1.907 |
| P. satarix | 1 | 0 | 1 | 1 | 0 | 1 | 135 | 0 | 2 | 0 | 0 | 1 | 1 | 3 | 797 | 2 | 0 | 10 | 4 | 3 | 1 | 2 | 1 | 9 | 6 | 981 |
| P. pungitius | 2 | 0 | 6 | 1 | 3 | 2 | 3 | 295 | 7 | 3 | 8 | 2 | 12 | 3 | 4 | 4 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 357 |
| G. aculeams | 8 | 141 | 13 | 2 | 2 | 2 | 2 | 49 | 5 | 3 | 14 | 2 | 32 | 0 | 0 | 2 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 1 | 0 | 280 |
| S. puscus | 1 | 1 | 6 | 1 | 1 | 2 | 12 | 9 | 9 | 5 | 1 | 8 | 12 | 5 | 8 | 8 | 5 | 2 | 13 | 12 | 7 | 39 | 32 | 12 | 34 | 245 |
| Gadidae | 0 | 0 | 9 | 0 | 20 | 12 | 11 | 6 | 0 | 2 | 3 | 0 | 1 | 1 | 0 | 0 | 1 | 0 | 1 | 0 | 6 | 14 | 0 | 12 | 10 | 109 |
| M. cephalus | 0 | 0 | 3 | 1 | 41 | 1 | - 4 | 4 | 1 | 0 | 0 | 4 | 39 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 100 |
| M. currema | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 9 | 0 | 0 | 0 | 42 | 1 | 9 | 1 | 0 | 1 | 30 | 0 | 1 | 1 | 1 | 0 | 0 | 97 |
| C. harengus | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 30 | 0 | 6 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 5 | 22 | 3 | 74 |
| Lpana | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 1 | 0 | 15 | 10 | 2 | 0 | 32 | 3 | 0 | 0 | 1 | 0 | 1 | 1 | 0 | 1 | 70 |
| P. americanus | 1 | 1 | 0 | 0 | 6 | 0 | 0 | 2 | 4 | 7 | 4 | 0 | 0 | 3 | 10 | 3 | 2 | 0 | 3 | 0 | 5 | 0 | 10 | 2 | 2 | 65 |
| T.oniris : | 0 | 0 | : 0 | 0 | $\cdots 0$ | 0 | 4 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 7 | 1 | 0 | 0 | 1 | 4 | 0 | 1 | 2 | 11 | 11 | 45 |
| C. wheartandi ${ }^{-1}$ | 0 | 0 | $\therefore 0$ | 0 | 0 | 0 | 4 | 2 | 6 | 11 | 2 | 6 | 1 | 3 | 0 | 1 | 0 | 2 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 41 |
| T. filcarus* | 0 | 0 | 1 | 0 | $\because$ | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 22 | 6 | 0 | 0 | 0 | 0 | 0. | 2 | 0 | 0 | -1 | 0 | 0 | 33 |
| A. nostrata | 9 | 4 | '12 | 2 | 2 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 31 |
| M. aenaeus | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 2 | 2 | 0 | 1 | 0 | 0 | 0 | 1 | 2 | 3 | 1 | 0 | 2 | 0 | : 1 | 9 | 26 |
| C. hippos | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 1 | 3 | 0 | 1 | 1 | 2 | 0 | 0 | 9 | 0 | 1 | 0 | 23 |
| Anchoa spp. | 0 | 0 | 0 | 0 | 2 | 0 | 7 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 2 | 0 | 1 | 0 | 2 | 0 | 0 | 1 | 1 | 0 | 21 |
| O. mordar | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 18 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | $\because 0$ | 0 | 0 | 0 | 20 |
| T. odspersius | 0 | 0 | 2 | 0 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 | 0 | 3 | 14 |
| A. pseuldiarensus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 | 0 | 0 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 1 | 0 | 11 |
| L.apodus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 7 | 0 | 0 | 0 | 0 | 0 | 0 | 7 |
| A. americamus | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | . | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 6 |
| S. marina | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 1 | 0 | 0 | 0 | 5 |
| A. aestivalis | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | '0 | 0 | 0 | 0 |  |
| S. urmer , | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 |  |
| Men. saxarilis' | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 |  |
| C. regalis | 0 |  | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 |  |
| C. ocellatus - | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 |  |
| P. gunnellus' | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 |  |
| Prionotus sp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 |  |
| $\underline{L}$ xanhuruis | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 |
| total | 39,426 | 17,700 | 1,359 | 1,204 | 6.482 | 2.043 | 3.059 | 10.562 | 2.609 | 1,404 | 3,544 | 4,674 | 6,198 | 3,388 | 7,935 | 7,209 | 4,894 | 2.638 | 4,725 | 5,984 | 6,414 | 8,592 | 6,108 | 6,013 | 5,535 | 169.919 |

${ }^{2}$ Fish identified to the lowest practical taxon.

| Year | 76.71 | 77.78 | 78.79 | 79-80 | 80-81 | 81-82 | 82-83 | 83.84 | 84-85 | 85.86 | 86-87 | 87.88 | 88-89 | 89-90 | 90-91 | 91.92 | 92-93 | 93-94 | 94-95 | 95-96 | 96-97 | 97-98 | 98-99 | 99-00 | 00-01 | Tatal |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Number of samples | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 1.878 |
| Taxon ${ }^{\text {a }}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| P. annericanus | 924 | 1.338 | 1,165 | 2.078 | 2.143 | 1,908 | 2.332 | 2.182 | 1,452 | 1.807 | 1,499 | 1,844 | 2,410 | 1,259 | 1,531 | 1,008 | 1,621 | 1,532 | 2.160 | 1,078 | 995 | 827 | 836 | 801 | 44 | 37,174 |
| S. chinsops | 693 | 1,465 | 388 | 425 | 903 | 800 | 1,436 | 1,843 | 1,264 | 391 | 1,528 | 1,315 | 996 | 747 | 1,369 | 7,903 | 3,749 | 277 | 1.848 | 282 | 67 | 514 | 2,042 | 749 | 1,628 | 34,622 |
| S. aquosus | 203 | 294 | 107 | 176 | 150 | 161 | 216 | 294 | 199 | 312 | 322 | 761 | 591 | 682 | 357 | 308 | 616 | 1,000 | 845 | 558 | 486 | 230 | 407 | 416 | 264 | 9,953 |
| Raja spp. | 97 | 90 | 44 | 91 | 144 | 112 | 191 | 358 | 136 | 312 | 344 | 385 | 473 | 448 | 469 | 543 | 299 | 484 | 401 | 556 | 343 | 312 | 312 | 405 | 238 | 7,587 |
| Menidia spp. | 287 | 718 | 805 | 728 | 190 | 146 | 75 | 115 | 80 | 144 | 128 | 33 | 60 | 26 | 24 | 261 | 1.915 | 107 | 112 | 82 | 107 | 57 | 705 | 193 | 145 | 7,263 |
| T, edspersus | 632 | 666 | 227 | 1,022 | 596 | 342 | 207 | 76 | 68 | 27 | 9 | 9 | 12 | 35 | 31 | 17 | 90 | 8 | 31 | 21 | 8 | 24 | 46 | 17 | 49 | 4,270 |
| Anchoa spp. | 165 | 58 | 806 | 0 | 44 | 354 | 1 | 20 | 13 | 1,799 | 95 | 41 | 11 | 1 | 3 | 16 | 2 | 2 | 3 | 12 | 0 | 5 | 183 | 1 | 7 | 3,642 |
| M. achurus | 45 | 87 | 56 | 72 | 162 | 176 | 208 | 242 | 76 | 59 | 126 | 85 | 111 | 84 | . 62 | 47 | 122 | 90 | 147 | 31 | 113 | 141 | 68 | 45 | 137 | 2,592 |
| Cadide | 18 | 69 | 63 | 62 | 423 | 315 | 140 | 194 | 94 | 135 | 35 | 373 | 21 | 20 | 17 | 24 | 36 | 23 | 26 | 50 | 23 | 26 | 25 | 19 | 7 | 2.238 |
| P. denutus | 73 | 40 | 16 | 9 | 24 | 49 | 37 | 53 | 80 | 39 | 107 | 121 | 61 | 10 | 63 | 75 | 113 | 108 | 94 | 69 | 150 | 112 | 123 | 195 | 195 | 2,020 |
| C. striata | 8 | 2 | 0 | 3 | 5 | 39 | 13 | 24 | 25 | 43 | 241 | 8 | 32 | 46 | 49 | 35 | 47 | 5 | 277 | 34 | 7 | 22 | 478 | 248 | 321 | 2.012 |
| P. triaccunitus | 5 | 2 | 12 | 3 | 4 | 9 | 4 | 7 | 0 | 3 | 10 | 5 | 929 | 10 | 328 | 60 | 24 | 16 | 1 | 7 | 0 | 0 | 6 | 1 | 0 | 1,446 |
| Prionotus spp. | 42 | 30 | 30 | 46 | 66 | 72 | 31 | 67 | 38 | 31 | 104 | 27 | 36 | 215 | 76 | 25 | 19 | 42 | 98 | 136 | 10 | 14 | 46 | 109 | 13 | 1.423 |
| E. microstromus | 6 | 0 | 0 | 0 | 1 | 17 | 4 | 15 | 14 | 34 | 107 | 39 | 59 | 12 | 85 | 82 | 86 | 31 | 96 | 36 | 33 | 77 | 178 | 227 | 111 | 1,350 |
| Urophycis spp. | 2 | 3 | 7 | 5 | 21 | 23 | 182 | 45 | 19 | 29 | 11 | 26 | 49 | 25 | 59 | 13 | 43 | 59 | 93 | 44 | 24 | 68 | 58 | 376 | 39 | 1,323 |
| M. Lilinearis | 101 | 15 | 2 | 36 | 109 | 48 | 38 | 52 | 26 | 38 | 4 | 4 | 23 | 51 | 47 | 47 | 73 | 5 | 31 | 23 | 7 | 16 | 69 | 34 | 35 | 974 |
| T. uniris | 63 | 70 | 86 | 68 | 47 | 27 | so | 41 | 46 | 47 | 23 | 17 | 42 | 18 | 16 | 14 | 9 | 24 | 17 | 28 | 19 | 24 | 15 | 26 | 55 | 892 |
| H. unericanus | 7 | 5 | 11 | 19 | 62 | 96 | 115 | 60 | 16 | 7 | 1 | 1 | 0 | 2 | 2 | 1 | 0 | 0 | 0 | 5 | 3 | 3 | 7 | , | 0 | 425 |
| O. mordax | 1 | 6 | 14 | 0 | 9 | 19 | 29 | 2 | 4 | 16 | 8 | 4 | 4 | 4 | 16 | 6 | 134 | 1 | 12 | 7 | 20 | 2 | 35 | 2 | 1 | 356 |
| S. juscus | 0 | 14 | 9 | 12 | 24 | 18 | 12 | 12 | 25 | 11 | 7 | 15 | 13 | 7 | 5 | 9 | 10 | 11 | 8 | 7 | 8 | 16 | 5 | 6 | 10 | 274 |
| A. pseudoharengus | 1 | 216 | 8 | 3 | 0 | 4 | 0 | 1 | 0 | 3 | 4 | 1 | 1 | 0 | 0 | 2 | 7 | 17 | 3 | 1 | 0 | 0 | 0 | 0 | 0 | 272 |
| C. Iumpus | 8 | 1 | 6 | 16 | 4 | 0 | 4 | 0 | 4 | 0 | 0 | 7 | 2 | 0 | 4 | 0 | 1 | 0 | 0 | 0 |  | 1 | 0 | 0 | 117 | 182 |
| S. maculatus | 2 | 1 | 0 | 0 | 1 | 0 | 0 | 2 | 0 | 0 | 1 | 0 | 4 | 3 | 8 | 22 | 16 | 1 | 7 | 5 | 4 | 1 | 9 | 19 | 15 | 121 |
| Clupeidec | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 110 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 2 | 113 |
| C. regalis | 1 | 8 | 0 | 1 | 0 | 3 | 1 | 0 | 0 | 1 | 4 | 2 | 10 | 0 | 1 | 1 | 1 | 0 | 1 | 2 | 1 | 0 | 8 | 3 | 0 | 49 |
| P. oblongus | 1 | 0 | 1 | 0 | 3 | 1 | 1 | 2 | 5 | 3 | 7 | 1 | 2 | 0 | 0 | 0 | 1 | 2 | 1 | 1 | 0 | 4 | 5 | 2 | 2 | 45 |
| B. grannus | 1 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 | 24 | 0 | 3 | 0 | 2 | 0 | 2 | 5 | 0 | 44 |
| M. americana | 2 | 7 | 1 | 0 | 6 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 2 | 0 | 2 | 0 | 1 | 1 | 1 | 1 | 0 | 0 | 1 | 8 | 35 |
| H. erectus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 1 | 0 | 1 | 0 | 2 | 6 | 0 | 0 | 0 | 4 | 0 | 4 | 8 | 4 | 0 | 32 |
| Lipuris spp. | 1 | 3 | 4 | 2 | 2 | 2 | 5 | 1 | 3 | 0 | 0 | 0 | 4 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 1 | 0 | 0 | 30 |
| A. supidissima | 15 | 2 | 0 | 0 | 4 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 1 | 25 |
| Alosa spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 7 | 1 | 0 | 0 | 0 | 3 | 5 | 0 | 2 | 0 | 20 |
| M. hispidus | 2 | 0 | 2 | 0 | 0 | 0 | 1 | 0 | 3 | 1 | 0 | 0 | 1 | 2 | 2 | 0 | 0 | 3 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 19 |
| G. aculeatus | 1 | 0 | 1 | 1 | 0 | 2 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 2 | 1 | 0 | 1 | 3 | 1 | 0 | 1 | 0 | 18 |
| D. solitans | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 4 | 1 | 2 | 3 | 0 | 1 | 0 | 0 | 2 | 0 | 0 | 0 | 1 | 0 | 17 |
| O. 1 tu | 1 | 0 | 0 | 1 | 4 | 1 | 1 | 0 | 2 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 3 | 17 |
| M. octodecemspinusus | 0 | 0 | 2 | 2 | 1 | 6 | 3 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 16 |
| A. asstivalis | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 9 | 0 | 0 | 2 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 16 |
| M. saxatilis | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 5 | 1 | 2 | 1 | 1 | 3 | 0 | 0 | 16 |
| C. harengus | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 2 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 7 | 0 | 2 | 0 | 15 |
| F. tabacaria | 1 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 3 | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 3 | 0 | 13 |
| M. acglefinus | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 1 | 0 | 3 | 0 | 0 | 0 | 0 | 5 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 11 |
| A. scioeps | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 2 | 1 | 0 | 1 | 0 | 0 | 1 | 0 | 1 | 0 | 1 | 0 | 10 |
| A. americumes | 0 | 2 | 0 | 1 | 1 | 0 | 2 | 0 | 0 | 2 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 9 |


| Year | 76-77 | 77.78 | 78.79 | 79-80 | $80-81$ | 81.82 | 82.83 | 83-84 | 84.85 | 85.86 | 86-87 | 87.88 | 88.89 | 89.90 | 90-91 | 91.92 | 92-93 | 93-94 | 94.95 | 95-98 | 96-97 | 97-98 | 98.99 | 99-60 |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Number of samples | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 1,956 |
| Taxon ${ }^{2}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| A. rostrara | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 2 | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 9 |
| P. crientaus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 0 | 0 | 2 | 2 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 8 |
| $L$ xamilurus | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 2 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 6 |
| S. setapinnis | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 6 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 6 |
| C. hippos | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 6 |
| P. arenarus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 |
| M. canis | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 |
| A. quadracus | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 3 |
| C. ocellarus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 3 |
| Gobiidae | 0 | 0 | 0 | 0 | - 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 3 |
| P.alsa | 0 | 0 | 0 | -0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 3 |
| S.foetens | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 3 |
| O. marginarum | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 3 |
| C.corsas | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| E. cimbrius | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | $\cdots 2$ |
| L.americanus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| M. auromus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| S. urmer | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 2 |
| U. subbifircata | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 2 |
| Lactrophys spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| P. soluarix | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 1 |
| P. pungitius | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| S. scombrus | 0 | 1 | 0 | 0 | 0 | 0 | . 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| T. Jathami | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | . 1 |
| T. macularus | 0 | 0 | 0 | 0 | 0 | 0 | . 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | ... 1 |
| C. schorpf | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | $\cdot 1$ |
| C. oceanicus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 |
| total | 3,468 | 5,246 | 3,889 | -4,890 | 5,190 | 4,779 | 5,382 | 3,752 | 3.727 | 5,427 | 4,796 | 3, 169 | 6,002 | 3,742 | 4,647 | 10,55 | 9.080 | 3,876 | 6,350 | 3,102 | 2,478 | 2.549 | 5,699 | 3,939 | 3.909 | 123,639 |

[^9]| Year | 76-77 | 77.78 | 78-79 | 79-80 | 80-81 | 81-82 | 82-83 | 83-84 | 84.85 | 85-86 | 86-87 | 87.88 | 88-89 | 89.90 | 90-91 | 91.92 | 92-93 | 93-94 | 94.95 | 95-96 | $96-97$ | 97.98 | 98-4) | 99-00 | 00-01 | Total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Number of samples | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 77 | 78 | 81 | 78 | 78 | 78 | 77 | 78 | 78 | 1.954 |
| Taxon ${ }^{\text {a }}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| P. umericanus | 1.422 | 991 | 1,262 | 1,039 | 885 | 385 | 1,665 | 1,302 | 795 | 1,064 | 781 | 617 | 660 | 648 | 551 | 540 | 640 | 656 | 598 | 564 | 432 | 362 | 387 | 432 | 381 | 19,059 |
| S. cinysops | 441 | 250 | 162 | 214 | 65 | 12 | 713 | 374 | 26 | 332 | 426 | 196 | 77 | 205 | 608 | 4,301 | 628 | 17 | 384 | 132 | 11 | 323 | 445 | 598 | 1,205 | 12,145 |
| Menidia spp. | 479 | 183 | 201 | 446 | 202 | 22 | 73 | 251 | 152 | 161 | 1,491 | 232 | 45 | so | 23 | 238 | 220 | 91 | 35 | 96 | 14 | 30 | 86 | 191 | 763 | 5,835 |
| T. adspersus | 97 | 78 | 90 | 232 | 191 | 263 | 209 | 120 | 73 | 23 | 28 | 15 | 148 | 34 | 55 | 88 | 162 | 50 | 61 | 46 | 6 | 37 | 854 | 512 | 394 | 3,866 |
| P. guniellus | 20 | 35 | 45 | 37 | 171 | 189 | 137 | 55 | 35 | 116 | 78 | 107 | 278 | 40 | 42 | 78 | 48 | 104 | 220 | 174 | 65 | 160 | 515 | 406 | 410 | 3,603 |
| Anclioa spp. | 285 | 9 | 32 | 4 | 6 | 6 | 4 | 24 | 25 | 326 | 283 | 164 | 256 | 492 | 15 | 1,407 | 4 | 1 | 1 | 16 | 0 | 1 | 62 | 0 | 29 | 3,452 |
| Raja spp. | 63 | 64 | 42 | 19 | 45 | 25 | 66 | 239 | 41 | 109 | 84 | 111 | 130 | 108 | 334 | 191 | 196 | 238 | 102 | 209 | 118 | 157 | 104 | 109 | 49 | 2,953 |
| Gadidac | 38 | 116 | 68 | 75 | 420 | 341 | 171 | 131 | 134 | 186 | 97 | 88 | 16 | 17 | 61 | 58 | 67 | 331 | 52 | 69 | 27 | 36 | 39. | 92 | 82 | 2,812 |
| S. aquosus | 135 | 110 | 87 | 108 | 65 | 30 | 93 | 199 | 108 | 155 | 107 | 65 | 74 | 98 | 82 | 64 | 119 | 148 | 77 | 137 | 145 | 78 | 53 | 41 | 23 | 2,403 |
| M. aenaeus | 33 | 115 | 9 | 46 | 70 | 113 | 98 | 133 | 82 | 76 | 57 | 97 | 159 | 26 | 27 | 45 | 102 | 111 | 137 | 59 | 35 | 90 | 147 | 66 | 153 | 2,193 |
| P. dentarus | 80 | 35 | 19 | 16 | 9 | 9 | 65 | 57 | 149 | 53 | 157 | 85 | 35 | 10 | 68 | 75 | 110 | 33 | 33 | 40 | 104 | 58 | 51 | 103 | 124 | 1,578 |
| O. mordax | 47 | 164 | 31 | 0 | 72 | 11 | 2 | 21 | 216 | 274 | 224 | 227 | 89 | 3 | 8 | 3 | 7 | 1 | 0 | 6 | 1 | 3 | 1 | 8 | 0 | 1,441 |
| T. onitis | 71 | 106 | 59 | 57 | 22 | 20 | 37 | 18 | 15 | 31 | 57 | 30 | 36 | 20 | 40 | 35 | 91 | 50 | 20 | 74 | 46 | 30 | 79 | 133 | 262 | 1,439 |
| S. juscus | 7 | 13 | 15 | 37 | 27 | 39 | 65 | 84 | 124 | So | 57 | 72 | 136 | 12 | 18 | 21 | 67 | 77 | 27 | 92 | 28 | 24 | 58 | 76 | 65 | 1.291 |
| Uroplycis spp. | 8 | 22 | 19 | 17 | 9 | 21 | 81 | 45 | 42 | 37 | 5 | 14 | 23 | 18 | 108 | 15 | 53 | 37 | 87 | 36 | 17 | 19 | 19 | 199 | 35 | 979 |
| C. striata | 3 | 0 | 0 | 0 | 0 | 6 | 1 | 6 | 1 | 4 | 45 | 0 | 1 | 3 | 5 | 10 | 9 | 2 | 64 | 4 | 1 | 16 | 443 | 135 | 120 | 879 |
| B. ņranuus | 0 | 0 | 0 | 0 | 0 | .0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 520 | 3 | 3 | 5 | 4 | 0 | 0 | 2 | 0 | 4 | 3 | 4 | 549 |
| E. micrustronus | 11 | 0 | 0 | 1 | 1 | 3 | 5 | 8 | 8 | 22 | 35 | 14 | 4 | 4 | 54 | 23 | 54 | 25 | 29 | 41 | 8 | 32 | 17 | 60 | 16 | 475 |
| H. unericanus | 2 | 8 | 3 | 36 | 49 | 82 | 145 | 80 | 28 | 8 | 1 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 446 |
| A. quadracus | 1 | 3 | 1 | 0 | 3 | 1 | 0 | 5 | 21 | 13 | 85 | 23 | 6 | 0 | 0 | 0 | 19 | 87 | 1 | 2 | 1 | 0 | 0 | 0 | 1 | 273 |
| M. bilineuris | 65 | 2 | 1 | 4 | 41 | 4 | 9 | 4 | 4 | 5 | 3 | 0 | 0 | 4 | 3 | 9 | 41 | 1 | 3 | 3 | 0 | 5 | 1 | 1 | 10 | 223 |
| C. Iunpus | 11 | 10 | 16 | 39 | 4 | 0 | 7 | 0 | 18 | 1 | 1 | 31 | 4 | 1 | 2 | 1 | 15 | 7 | 1 | 2 | 39 | 4 | 1 | 0 | 7 | 222 |
| Primutus spp. | 23 | 7 | 5 | 5 | 4 | 3 | 7 | 1 | $y$ | 3 | 15 | 0 | 4 | 7 | 5 | 8 | 12 | 13 | 2 | 3 | 2 | 4 | 7 | 9 | 0 | 158 |
| H. crecius | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 1 | 8 | 8 | 3 | 1 | 6 | 20 | 3 | 0 | 0 | 5 | 0 | 1 | 1 | 8 | 17 | 85 |
| P. incaumathus | 1 | 1 | 2 | 0 | 4 | 0 | 0 | 11 | 0 | 0 | 5 | 1 | 26 | 4 | 14 | 3 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 3 | 77 |
| C. harengus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 63 | 0 | 2 | 0 | 0 | 1 | 3 | 0 | 1 | 0 | 0 | 0 | 0 | 2 | 0 | 2 | 74 |
| F. rubucaria | 1 | 1 | 0 | 0 | 2 | 0 | 1 | 0 | 8 | 1 | 2 | 0 | 0 | 1 | 11 | 5 | 5 | 2 | 1 | 0 | 0 | 0 | 0 | 8 | 19 | 68 |
| A. rosirata | 1 | 3 | 0 | 1 | 1 | 8 | 9 | 3 | 5 | 4 | 0 | 1 | 1 | 1 | 0 | 5 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 44 |
| C. regalis | 1 | 11 | 1 | 0 | 0 | 5 | 0 | 0 | 0 | 0 | 2 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 7 | 11 | 41 |
| S. mactulams | 3 | 2 | 1 | 0 | 0 | 0 | 0 | 1 | 3 | 1 | 0 | 0 | 0 | 2 | 3 | 3 | 0 | 0 | 0 | 5 | 1 | 2 | 2 | 1 | 9 | 39 |
| A. unericunus | 3 | 2 | 6 | 0 | 6 | , | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 1 | 0 | 14 | 0 | 36 |
| Liparis spp. | 1 | 1 | 1 | 2 | 1 | 3 | 3 | 1 | 5 | 0 | 1 | 1 | 2 | 4 | 0 | 0 | 0 | 0 | 1 | 0 | 4 | 1 | 1 | 0 | 0 | 33 |
| M. lispidus | 0 | 1 | 2 | 0 | 0 | 0 | 6 | 0 | 2 | 4 | 1 | 0 | 0 | 1 | 2 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 22 |
| S. setapiuvis | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 16 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 17 |
| O. marginatum | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 10 | 16 |
| A. pseudoharengus | 1 | 3 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 11 |
| A. scluepfi | 0 | 1 | 0 | 1 | 1 | 0 | 0 | 1 | 0 | 1 | 1 | 0 | 2 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 10 |
| M. americana | 1 | 4 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 10 |
| M. canus | 2 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 4 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 10 |
| P. purgizius | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 | 0 | 5 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 10 |
| O. 1 au | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 1 | 2 | 0 | 2 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 10 |
| Lactuphins spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 1 | 3 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 9 |
| A. sapidissima | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 6 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 8 |


|  | APPENDIX IV. (continued). |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Year | 76-77 | 77.78 | 78.79 | 79.80 | 80.81 | 81.82 | 82-83 | 83-84 | 84.85 | 85-86 | 86.87 | 87.88 | 88.89 | 89.90 | 90.91 | 91.92 | 92-93 | 93.94 | 94.95 | 95-9 |  | 97-98 | 98-99 | 99-00 | 00-01 | Toxal |
|  | Number of samples | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 77 | 78 | 81 | 78 | 78 | 78 | 77 | 78 | 78 | 1.954 |
|  | Taxon ${ }^{\text {a }}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | Alosa spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 2 | 4 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 8 |
|  | C. wieationdi | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 1 | 0 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 8 |
|  | S. borealis | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 2 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 6 |
|  | P. saltaitx | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 6 |
|  | T. maculams | 3 | i | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 |
|  | Gobildae | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 3 |
|  | $L$ xantiurus | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 |
|  | T. latiami | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 |
|  | U. subbifurata | 0 | 1 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 4 |
|  | M. saxatifis | 0 | 0 | . 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 1 | 0 | 4 |
|  | M. octodecemspinosus | 0 | 0 | 2 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 |
|  | P. arenarus | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 |
|  | D. molions | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 8 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
|  | Ecimbrius | 0 | 0 | 0 | - 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 0 | 0 | 2 |
|  | Gastcrosteidae | 0 | 0 | . 0 | 2 | 0 | 0 | . 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | $\therefore 0$ | 0 | 0 | - 2 |
|  | P. cruentarus | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | . 0 | 0 | 2 |
|  | S. tromer | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |  | 0 | 2 |
|  | A. aestivalis | . 0 | 0 | 0 | 0 | , | 0 | . 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 1 |
|  | A. mediacris | 1 |  | 0 |  |  |  | 0 |  |  | 0 |  |  |  | 0 | 0 | 0 |  | 0 | 0 | 0 | 0 | 0 | - 0 |  | 0 | 1 |
|  | A. maculanus | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 0 | 0 | 0 | 0 | 0 | 0 | 0 | . 1 |
|  | C.oceanicus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | D. centroura | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | . 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 00 | 1 |
|  | D. mactrellus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | Fundulue stp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | Lamericanus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | M. aeglefinus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | Monacanthus spp. | 0 | 0 | 0 | 0 | . 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | . M. auratus | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | S. marina | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | T. myops | , 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | U. panius | 0 | 0 | . 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
|  | total | 3.376 | 2,375 | 2.351 | 2,493 | 2.394 | 1,609 | 3.679 | 3,400 | 3.098 | 3,160 | 4,496 | 2,592 | 2.277 | 2.341 | 2,158 | 7,298 | 2,707 | 2,202 | 1,941 | 1.819 | 1,132 | 1,483 | 3,387 | 3,223 | 4.217 | 71,208 |
| $\underline{\underline{n}}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |

. Fish identified to the lowest pracical taxon.

| Year | 76-77 | 77.78 | 78.79 | 79-80 | 80.81 | 81.82 | 82-83 | 83-84 | 84.85 | 85-86 | 86-87 | 87.88 | 88-89 | 89-90 | 90-91 | 91.92 | 92-93 | 93-94 | 94-95 | 95-96 | 96-97 | 97.98 | 98-99 | 99-00 | 00-01 | Toal |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Number of samples | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 1.936 |
| Taxon ${ }^{2}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| P. umericanus | 2.775 | 1.191 | 1.145 | 2,714 | 3,863 | 3,823 | 7,569 | 3.543 | 4,890 | 2,472 | 4,108 | 3,463 | 6,763 | 3.621 | 4.504 | 4,916 | 5,214 | 1,400 | 2,185 | 869 | 1,467 | 585 | 1,030 | 1,252 | 1,360 | 76,722 |
| Menidia spp. | 1,203 | 315 | 58 | 121 | 73 | 159 | 315 | 49 | 15 | 93 | 1,549 | 318 | 639 | 220 | 317 | 431 | 481 | 75 | 97 | 34 | 352 | 124 | 349 | 359 | 924 | 8,670 |
| M. ucrucus | 57 | 30 | 58 | 161 | 201 | 349 | 387 | 232 | 274 | 174 | 450 | 205 | 631 | 374 | 492 | 273 | 451 | 142 | 299 | 167 | 442 | 375 | 278 | 96 | 122 | 6.961 |
| S. uquosus | 76 | 40 | 32 | 134 | 158 | 165 | 230 | 225 | 244 | 136 | 302 | 236 | 173 | 342 | 363 | 234 | 225 | 292 | 94 | 97 | 77 | 110 | 64 | 54 | 39 | 4.142 |
| P. dentutus | 41 | 29 | 16 | 14 | 21 | 100 | 82 | 63 | 163 | 75 | 226 | 214 | 191 | 20 | 149 | 173 | 248 | 105 | 165 | 68 | 75 | 228 | 75 | 180 | 235 | 2.956 |
| S. fuscus | 20 | 11 | 13 | 31 | 61 | 178 | 132 | 80 | 70 | 117 | 114 | 177 | 128 | 35 | 112 | 90 | 69 | 182 | 87 | 88 | 64 | 29 | 49 | 51 | 20 | 2.028 |
| Prionorus spp. | 97 | 21 | 2 | 6 | 4 | 63 | 200 | 2 | 2 | 3 | 44 | 1 | 19 | 147 | 98 | 267 | 37 | 12 | 262 | 53 | 6 | 40 | 20 | 465 | 70 | 1,941 |
| T. onits | 39 | 16 | 30 | 45 | 25 | 129 | \% | 16 | 11 | 22 | 110 | 15 | 57 | 28 | 105 | 51 | 24 | 13 | 14 | 128 | 41 | 40 | 126 | 240 | 324 | 1,739 |
| G. aculeanus | 16 | 12 | 44 | 22 | 186 | ys | 38 | 7 | 162 | 86 | 8 | 19 | 63 | 8 | 14 | 410 | 135 | 20 | 60 | 3 | 115 | 54 | 79 | 4 | 6 | 1,690 |
| A. quadracus | 8 | 3 | 22 | 27 | 190 | 764 | 76 | 6 | 91 | 115 | 22 | 29 | 25 | 11 | 18 | 100 | 50 | 3 | 4 | 4 | 13 | 1 | 46 | 8 | 7 | 1.643 |
| Anchioa spp. | 10 | 193 | 0 | 2 | 0 | 0 | 11 | 2 | 5 | 16 | 8 | 11 | 168 | 734 | 2 | 12 | 1 | 0 | 12 | 2 | 14 | 3 | 21 | 139 | 0 | 1,368 |
| B. grannus | 0 | 13 | 1 | 0 | 1 | 1 | 0 | 1 | 0 | 31 | 10 | 3 | 0 | 788 | 2 | 182 | 23 | 17 | 41 | 0 | 3 | 0 | 2 | 106 | 102 | 1,327 |
| Gadude | 5 | 9 | 11 | 8 | 152 | 185 | 71 | 43 | 14 | 209 | 12 | 5 | 32 | 29 | 29 | 12 | 8 | 16 | 20 | 44 | 86 | 66 | 124 | 15 | 11 | 1,216 |
| S.ctrysops | 58 | 10 | 11 | 5 | 6 | 38 | 27 | 57 | 2 | 3 | 22 | 15 | 3 | 5 | 10 | 175 | 119 | 0 | 83 | 38 | 2 | 12 | 4 | 94 | 263 | 1,062 |
| P. gunnellus | 0 | 1 | 10 | 4 | 14 | 29 | 69 | 32 | 21 | 10 | 18 | 41 | 47 | 58 | 83 | 30 | 48 | 44 | 26 | 26 | 54 | 33 | 142 | 29 | 20 | 889 |
| O.tau | 96 | 21 | 7 | 17 | 27 | 34 | 24 | 21 | 22 | 31 | 53 | 50 | 56 | 29 | Ss | 17 | 5 | 9 | 18 | 11 | 6 | 8 | 7 | 5 | 8 | 637 |
| T. adspersus | 14 | 4 | 1 | 11 | 7 | 91 | 58 | 60 | 16 | 15 | 38 | 14 | 9 | 15 | 9 | 8 | 10 | 9 | 13 | 8 | 3 | 20 | 113 | 27 | 23 | 602 |
| C. striata | 18 | 0 | 0 | 1 | 2 | 3 | 2 | 0 | 1 | 13 | 107 | 0 | 3 | 2 | 20 | 39 | 1 | 0 | 9 | 6 | 1 | 2 | 13 | 39 | 58 | 340 |
| O. mordax | 55 | 86 | 2 | 0 | 6 | 13 | 30 | 1 | 3 | 37 | 4 | 5 | 34 | 4 | 7 | 3 | 10 | 2 | 1 | 2 | 5 | 0 | 3 | 0 | 0 | 313 |
| A rerrrata | 16 | 11 | 7 | 4 | 8 | 24 | 14 | 21 | 15 | 27 | 26 | 20 | 14 | 4 | 15 | 2 | 2 | 4 | 10 | 7 | 8 | 2 | 5 | 7 | 3 | 276 |
| Clupeidae | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 258 | 0 | 0 | 0 | 0 | 0 | 259 |
| S. maculatus | 11 | 5 | 0 | 0 | 7 | 11 | 12 | 7 | 2 | 1 | 1 | 0 | 2 | 4 | 13 | 22 | 3 | 2 | 17 | 15 | 1 | 3 | 1 | 28 | 8 | 176 |
| Urophycis spp. | 0 | 0 | 0 | 0 | 1 | 2 | 7 | 4 | 3 | 10 | 1 | 22 | 11 | 6 | 9 | 4 | 2 | 5 | 18 | 16 | 2 | 9 | 17 | 0 | 22 | 171 |
| Alusa spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 3 | 14 | 11 | 2 | 16 | 111 | 0 | 0 | 0 | 3 | 3 | 0 | 164 |
| Gubiidae | 3 | 0 | 0 | 0 | 4 | 0 | 0 | 2 | 9 | 6 | 2 | 5 | 10 | 2 | 19 | 15 | 1 | 23 | 1 | 3 | 1 | 4 | 3 | 0 | 3 | 118 |
| H. erecrus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 1 | 10 | 3 | 2 | 2 | 8 | 34 | 1 | 0 | 0 | 0 | 0 | 0 | 6 | 15 | 22 | 106 |
| A. americamus | 1 | 0 | 75 | 1 | 0 | 2 | 14 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 96 |
| H. umericanus | 0 | 0 | 0 | 3 | 7 | 6 | 20 | 35 | 9 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 1 | 0 | 1 | 0 | 0 | 86 |
| Raja spp. | 0 | 0 | 0 | 0 | 2 | 1 | 1 | 3 | 0 | 2 | 0 | 0 | 4 | 2 | 4 | 0 | 20 | 2 | 6 | 2 | 9 | 18 | 2 | 4 | 1 | 83 |
| A. pseudoharengus | 0 | 47 | 0 | 2 | 2 | 2 | 3 | 1 | 2 | 1 | 3 | 0 | 1 | 0 | 3 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 69 |
| E. microstromus | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 3 | 0 | 1 | 4 | 3 | 4 | 1 | 3 | 7 | 8 | 1 | 5 | 1 | 1 | 6 | 3 | 0 | 10 | 62 |
| A. sapidissima | 2 | 0 | 0 | 4 | 1 | 1 | 0 | 9 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 40 | 0 | 0 | 1 | 0 | 58 |
| A. arstivalis | 0 | 7 | 1 | 9 | 0 | 0 | 0 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 9 | 0 | 0 | 1 | 7 | 0 | 0 | 0 | 0 | 0 | 37 |
| M.americuna | 3 | 0 | 1 | 5 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 9 | 0 | 0 | , | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 26 |
| D. volitans | 2 | 0 | 0 | 0 | 0 |  | 3 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 3 | 2 | 2 | 1 | 1 | 4 | 1 | 0 | 0 | 3 | 25 |
| F. rabacaria | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 7 | 3 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 3 | 4 | 21 |
| Morune saxatilis | 0 | 0 | 2 | 1 | 0 | 1 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 4 | 1 | 0 | 0 | 2 | 2 | 0 | 3 | 1 | 0 | 1 | 0 | 20 |
| P. rriacuuthus | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 8 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 6 | 20 |
| M. bilinearis | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 4 | 0 | 0 | 4 | 0 | 1 | 0 | 2 | 0 | 17 |
| Liparis spp. | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 9 | 0 | 1 | 0 | 1 | 1 | 0 | 0 | 1 | 0 | 0 | 1 | 15 |
| C. lumpus | 0 | 0 | 1 | 1 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 2 | 0 | 0 | 0 | 2 | 1 | 1 | 0 | 0 | 1 | 1 | 0 | 0 | 2 | 14 |
| Fundulus spp. | 0 | 0 | 0 | 0 | 0 | 3 | 2 | 0 | 0 | 1 | 1 | 0 | 1 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 12 |


| Year | 76-77 | 77.78 | 78.79 | 79.80 | 80-81 | 81.82 | 82.83 | 83.84 | 84.85 | 85.86 | 86-87 | 87.88 | 88.89 | 89.90 | 90-91 | 91.92 | 92.93 | 93.94 | 94.95 | 95.96 | 96-97 | 97.98 | 98-99 | 99-00 | 00-01 | Toral |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Number of samples | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 81 | 78 | 78 | 78 | 78 | 78 | 78 | 78 | 1.956 |
| Taxon ${ }^{2}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| P. oblongus | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 7 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 12 |
| Gasterosteldae | 0 | 0 | 0 | 11 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 11 |
| O. marginatum | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 | 1 | 11 |
| P. satatrix | 0 | 1 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 6 |
| S. borealis | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 6 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 6 |
| Ment, saxarills | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 1 | 0 | 0 | 6 |
| C. harengus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 5 |
| C. oceanicus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 |
| S. foerens | 0 | 1 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 4 |
| M. hlspidus | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 3 |
| P. pungitius | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 |
| G. wheatlendi | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | $\bigcirc$ | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 3 |
| M. cephalus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 3 |
| C. ocellatus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| Lactophirys spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| P. ferruginew | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| C. regalis | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 2 |
| T. maculans | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| B. ocellatus | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| C. variegatus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| Monocanthus spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| Af. canis | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| Myoxocephalus spp. | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | - 1 |
| P.alta | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| P. arenatus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 |
| S. rrutra | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| S. retifer | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| S. setapinis | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| total | 4.708 | 2,159 | 1.637 | 3,445 | 5,111 | 6.360 | 9,797 | 4.609 | 6.133 | 3,794 | 7.375 | 4.962 | 9,184 | 6.633 | 6.572 | 7.619 | 7,288 | 2.486 | 3.747 | 2.044 | 2.976 | 1,855 | 2.666 | 3,309 | 3.761 | 120.230 |

[^10]
## Winter Flounder Studies

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## Winter Flounder Studies

## Introduction

Since 1973, the winter flounder (Pseudopleuronectes americanus; classification of Pleuronectidae revised by Cooper and Chapleau 1998) has been the subject of environmental impact studies at Millstone Power Station (MPS). Results of these studies have been provided to the Connecticut Department of Environmental Protection (DEP) in a series of annual reports and special submissions. This species is an important sport and commercial fish in Connecticut (Smith et al. 1989) and a dominant member of the local demersal fish community. Its abundance tends to be cyclical and population sizes fluctuate considerably (Jeffries and Johnson 1974; Jeffries and Terceiro 1985). Of particular concern has been the effects of the station on the local Niantic River spawning population.
The winter flounder occurs from Labrador to Georgia, but is most numerous in the central part of its range (Scott and Scott 1988), which includes Long Island Sound (LIS). Seasonal movements and reproductive activities of winter flounder are welldocumented (e.g., Klein-MacPhee 1978). Details of winter flounder, abundance, distribution, and size information in LIS were provided by Gottschall et al. (2000) and spawning information by Howell and Molnar (1996, 1998). Most adult fish enter inshore waters in late fall and early winter and spawn nocturnally in upper portions of estuaries during late winter and early spring at temperatures between 1 and $10^{\circ} \mathrm{C}$ (peaking at $2-5^{\circ} \mathrm{C}$ ) and salinities of 10 to $35 \%$ (Bigelow and Schroeder 1953; Pearcy 1962; Scarlett and Allen 1992; Stoner et al. 1999). Females require 3 years for oocyte maturation and one batch of eggs matures each year (Dunn and Tyler 1969; Dunn 1970; Burton and Idler 1984). In eastern LIS, females begin to mature at age-3 and most are mature at age-4 with most males maturing at age-2 (NUSCO 1987), although Johnson et al. (1998) reported'a small percentage of both age- 2 females and age-1 males in LIS as mature. The current estimated mean fecundity of Niantic River females is approximately 583,775 eggs per fish. Spawning behavior in a large experimental laboratory aquarium was described by Stoner et al. (1999). Crawford (1990) reported that winter flounder deposit the demersal and adhesive eggs on gravel bars, algal mats, eelgrass (Zostera marina) beds, and near freshwater springs. Viable hatch of winter flounder eggs is greatest at $3^{\circ} \mathrm{C}$ in
salinities of 15 to $35 \%$ and decreases with increasing temperature (Rogers 1976). Egg incubation ranges from 5 to 31 days (Rogers 1976) with Keller and Klein-MacPhee (2000) reporting hatching on 20 days at $4.1^{\circ} \mathrm{C}$ and 30 days at $1.6^{\circ} \mathrm{C}$. Larval development through metamorphosis takes about 6 to 8 weeks (Laurence et al. 1979) and is also temperaturedependent. Small larvae are planktonic and although many remain near the estuarine spawning grounds, others are carried into coastal waters by tidal currents (Smith et al. 1975; NUSCO 1989; Crawford 1990). Some of the displaced larvae are returned to the estuary on subsequent incoming tides, but many of them are swept away from the area into coastal waters in which their survival may be reduced.
Older, larger larvae maintain some control over their position by vertical movements and may spend considerable time on the bottom. Following metamorphosis, most young-of-the-year winter flounder settle or move into shallow inshore waters. Yearlings (age-1 fish) become photonegative and are typically found in deeper waters (Pearcy 1962; McCracken 1963; Casterlin and Reynolds 1982). Some adult fish remain in estuaries following spawning while others disperse offshore. By summer, most adults leave warmer shallow waters as their preferred temperature range is $12-15^{\circ} \mathrm{C}$ (McCracken 1963), although a few remain in estuaries, apparently avoiding temperatures above $22.5^{\circ} \mathrm{C}$ by burying themselves in cooler bottom sediments (Oila et al. 1969). Other aspects of winter flounder life history were summarized by Klein-MacPhee (1978) and Pereira et al. (1999). Because the early life history of the European plaice (Pleuronectes platessus) has many similarities to that of the winter flounder, relevant literature for this species was also reviewed for this report to gain further insights into winter flounder population dynamics.
MPS operation can result in the impingement of juvenile and adult winter flounder on the traveling screens of the cooling-water intakes and the entrainment of larvae through condenser cooling-water systems. The impact of impingement at MPS was largely mitigated by the installation and operation of fish return sluiceways at all units, with winter flounder survival exceeding $90 \%$ at Unit 3 (NUSCO 1986c, 1988a, 1994b). A fish return at Unit 2 began operation in May 2000 and its effectiveness during the first year of operation was evaluated in DNC (2001d). Although only 16 winter flounder were
collected in this latest study, all survived impingement and a subsequent $72-\mathrm{h}$ holding period. Also in regards to overall station impact, note that although Unit 1 was officially retired in July 1998, the unit was effectively shut down in November 1995. Thus, numbers of fish impinged or entrained at MPS relative to previous three-unit operation (1986-95) have been reduced accordingly as Unit 1 utilized approximately $23 \%$ of former total water volume entrained at the station.
Unlike many other marine fishes with large coastwide populations, the mortality of entrained winter flounder larvae potentially has greater significance as larvae are products of local spawning from geographically isolated stocks associated with specific estuaries or coastal areas (Lobell 1939; Perlmutter 1947; Saila 1961). In particular, the population of winter flounder spawning in the nearby Niantic River has been studied in detail to assess the long-term effect of larval entrainment through the MPS cooling-water system. Presently, a combination of sampling programs and analytical methods are used to examine current abundance and trends of the Niantic River population. This report updates results given most recently in DNC (2001a) and includes abundance estimates of various life history stages of winter flounder and inferences that can be drawn from the relatively long time-series of available information. Due to the large amount of material presented on MPS winter flounder studies, except for some general information on methodology immediately following this section, detailed materials and methods (e.g., field sampling, data analyses) are presented within appropriate subsections of Results and Discussion.
A mass-balance model, discussed below in the section entitled MPS Impact Assessment, provides an annual estimate of the fraction of the Niantic River winter flounder reproductive output removed by larval entrainment at MPS. This model was reevaluated in DNC (2001c) and its sensitivity to input parameters was analyzed by both DNC and two independent reviewers. Studies of more direct methods (i.e., larval stock identification) to assign entrained winter flounder larvae to source populations were undertaken in 2000 and 2001, including the use of genetic and chemical composition (microelements) techniques (Crivello 2002; Moran 2002), reports of which were submitted to DEP. The goal of these studies was to provide direct quantitative estimates of entrainment impacts to the Niantic River and other nearby winter flounder populations. The fractions of entrained winter flounder larvae attributed to the Niantic River from this
work are compared to results of the mass-balance model later in this report.
A computer simulation model, the winter flounder stochastic population dynamics model (SPDM), first described in NUSCO (1990), has been used to examine long-term effects of MPS operation. The SPDM simulates the long-term effects of historical and projected rates of fishing mortality and simultaneous plant operation. Plant operation results in annual losses from impingement of juveniles and adults and, using estimates of production loss to the Niantic River stock provided by the mass-balance model, the entrainment of larvae through the MPS cooling-water system (Lorda et al. 2000). Within the series of annual reports, results of SPDM simulations were last presented in NUSCO (2000). This model was also modified for use in evaluating various cooling-water system alternatives in DNC (2001c). This work was an extensive evaluation of the MPS cooling-water system and included investigations of the feasibility of reducing either once-through condenser cooling water flow or larval fish entrainment. The-effectiveness of various altematives in reducing entrainment mortality on winter flounder and other fishes was determined. In both DNC (2001c) and in the present application, an assumption was made that the period of MPS operation would be extended beyond the existing U.S. Nuclear Regulatory Commission (NRC) license period for both Units 2 and 3. Note that although DNC is currently investigating the possibility of a 20 year license renewal for both units, which would extend their operational lifetimes to 2035 and 2045, respectively, no formal submission has been made to the NRC.

## General Materials and Methods

Data needed to assess MPS impact on the winter flounder come from several biological sampling programs (Fig. 1). Specific timing and seasonal duration of sampling was designed to reflect the annual life cycle of Niantic River winter flounder. Some programs (e.g., Niantic River adult and larval surveys, age-0 survey) investigate specific life history stages of winter flounder, whereas information on juvenile (age-0 and age-1) winter flounder was obtained from three sources: a special sampling program specifically targeted post-larval young-of-the-year; catches of age-0 juveniles from specific stations and months of the trawl monitoring program (TMP); and the Niantic River adult spawning abundance surveys during which age-1


1. February-April sampling (spawning season) of adults and juveniles throughout the Niantic River.
2. February-June larval sampling at three stations in the Niantic River and one in Niantic Bay.
3. Year-round monitoring of all ichthyoplankton at the MNPS discharges.
4. Late May-September sampling of age-0 juveniles at two stations in the Niantic River.
5. .Year-round monitoring of all benthic fishes at six (1976-95) or three (1996 and later) stations near. MPS. Juvenile catch data come from two stations in November (all years), four (1976-95) or three (1996 and later) stations in December, and six (1976-95) or three (1996 and later) stations in January and February that allow the year-0 cohort to be followed into a second calendar year.

Fig. 1. Current sampling programs contributing data for computation of winter flounder abundance indices. Darkened areas show months from which data were used in this report for the two year-round sampling programs.
juveniles are incidentally collected. Thus, data on: juvenile fish abundance were available from about May of their birth year into April of the following year. Other programs sample an entire local fish community year-round. These include the TMP, in which catches of winter flounder constitute a major component, and the entrainment ichthyoplankton monitoring program at MPS. Additional information used in various assessments was presented in NUSCO (1987), which summarized various life history studies of the winter flounder prior to the operation of Unit 3.
Data from various field sampling programs were used in the calculation of annual and seasonal indices of relative abundance, often with a $95 \%$ confidence interval (CI) or $\pm 2$ standard errors (SE) indicating . precision of the estimate. Indices of abundance were computed for various life-stages of winter flounder, such as various developmental stages of larvae, age-0 and 1 juveniles, and adults, which include spawning females and estimates of egg production. For some indices a long-term mean was calculated with variability described by the coefficient of variation (CV $=100 \times[S D /$ mean], where SD $=$ standard . deviation). Spearman's rank-order correlation (Snedecor and Cochran 1967; Hollander and Wolfe 1973) was used to examine the correlation among various
time-series of abundance indices. In some instances, a nonparametric, distribution-free Mann-Kendall test (Hollander and Wolfe 1973) was used to determine the direction of change if an annual abundance timeseries represented a significant ( $p \leq 0.05$ ) trend. Most data analyses were carried out using the Statistical Analysis System (SAS) computer programs (SAS Institute Inc. 1990a-d). The type of abundance index selected in each case depended upon the particular stage of life, sampling effort, and suitability of the data. A detailed description of each index is given below in various subsections of the Results and Discussion.

## Results and Discussion

## Seawater Temperature

Water temperature is important because it affects many biological processes such as adult distribution and spawning; larval growth, development, and mortality; and the settlement, distribution, growth, and mortality of demersal young. Information on water temperature was obtained from : continuous temperature recorders at the intakes of MPS. Daily mean temperatures ( ${ }^{\circ} \mathrm{C}$ ) were determined from
available records of $15-\mathrm{min}$ average temperatures, from which monthly, seasonal, or annual means were calculated. During annual Niantic River adult winter flounder surveys, representative water temperature measurements were taken at the surface and bottom with a YSI Model 30 Salinity/Temperature/ Conductivity meter.

Monthly mean seawater temperatures recorded at the MPS intakes set records for warmest temperatures or were at least well above average in both 1999 and 2000. This trend continued to a lesser extent in 2001 (Table 1). The January and April mean water temperatures approximated long-term, 26-year averages, but the February mean temperature of $4.03^{\circ} \mathrm{C}$ was more than $1^{\circ} \mathrm{C}$ above average. Means for March and May through December were also above average, although none were record-setting, except for December $\left(10.37^{\circ} \mathrm{C}\right)$, which was $1.22^{\circ} \mathrm{C}$ warmer than any previous monthly mean for that month.
The seasonal mean winter temperature of $4.26^{\circ} \mathrm{C}$ in 2001 was above average, but not exceptionally warm (Table 2). Although the mean of $11.54^{\circ} \mathrm{C}$ during spring was the fifth highest recorded, it was less than
means found during 1999 and 2000 and approximated the value for 1998. The mean summer temperature of $20.45^{\circ} \mathrm{C}$ tied 2000 as the second warmest in 25 years, trailing only the 1999 record of $20.85^{\circ} \mathrm{C}$. Thus, although individual monthly means in spring and summer were not exceptionally high, in combination they indicated relatively warm seasons, with less variation found between months than during many other years. The fall water temperature of $13.55^{\circ} \mathrm{C}$ also established a new record high. Because of the generally warm water temperatures in 2001, the annual mean was $12.50^{\circ} \mathrm{C}$, the fourth highest of the series (Table 1). Annual mean temperatures exceeding $12^{\circ} \mathrm{C}$ only occurred in $1990,1991,1995$, and 1998-2001.

Monthly mean temperatures were most variable during January through March (monthly CV $=25$ $38 \%$; Table 1), when winter flounder spawning and early larval development occur. Temperatures were most stable (CV $=3-7 \%$ ) from May through October, a period during which collections of winter flounder were dominated by juveniles and when most energy is allocated to growth rather than to reproduction.

TABLE 1. Monthly and annual mean seawater temperature $\left({ }^{\circ} \mathrm{C}\right)$ from January 1976 through December 2001 as calculated from mean daily water temperatures recorded continuously at the intakes of MPS Units 1 and 2 through July 2001 and at Units 2 and 3 thereafter.

| Year | Jan | Feb | Mar | Apr | May | Jun | Jul | Aug | Sep | Oct | Nov | Dec | Annual mean |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1976 | 3.65 | 3.31 | 4.81 | 7.55 | 10.75 | 15.11 | 18.29 | 19.60 | 18.93 | 15.04 | 9.28 | 4.73 | 10.90 |
| 1977 | 0.67 | 0.36 | 2.85 | 5.66 | 10.72 | 14.92 | 19.08 | 20.33 | 19.41 | 15.58 | 12.18 | 6.72 | 10.84 |
| 1978 | 3.01 | 1.09 | 1.67 | 4.85 | 9.10 | 14.24 | 17.68 | 19.82 | 19.24 | 16.14 | 12.47 | 7.74 | 10.64 |
| 1979 | 4.53 | 1.48 | 3.35 | 5.93 | 10.50 | 15.57 | 18.84 | 20.91 | 20.05 | 15.99 | 12.41 | 8.60 | 11.57 |
| 1980 | 5.16 | 2.38 | 2.80 | 6.38 | 10.44 | 14.76 | 18.44 | 20.23 | 20.16 | 16.07 | 10.25 | 5.73 | 11.10 |
| 1981 | 1.06 | 2.63 | 3.36 | 6.40 | 10.19 | 15.48 | 19.51 | 20.86 | 19.94 | 14.75 | 11.07 | 6.29 | 11.01 |
| 1982 | 2.20 | 1.56 | 3.04 | 5.41 | 10.06 | 14.16 | 17.98 | 21.10 | 20.01 | 15.95 | 12.47 | 8.97 | 11.13 |
| 1983 | 5.58 | 3.74 | 4.55 | 7.07 | 10.50 | 15.05 | 19.10 | 19.17 | 20.57 | 17.37 | 12.57 | 7.90 | 11.98 |
| 1984 | 4.84 | 4.02 | 3.98 | 6.58 | 10.84 | 15.53 | 18.90 | 20.60 | 19.52 | 16.41 | 13.04 | 9.07 | 11.97 |
| 1985 | 4.36 | 2.36 | 4.17 | 7.02 | 10.95 | 14.99 | 18.98 | 21.24 | 20.44 | 17.46 | 13.14 | 7.95 | 11.98 |
| 1986 | 4.62 | 3.38 | 4.11 | 7.25 | 11.32 | 15.99 | 18.83 | 20.62 | 18.80 | 16.53 | 12.43 | 8.19 | 11.89 |
| 1987 | 5.28 | 3.27 | 4.53 | 7.51 | 11.26 | 15.91 | 19.19 | 20.47 | 19.30 | 15.70 | 11.10 | 7.16 | 11.78 |
| 1988 | 2.65 | 2.67 | 4.49 | 7.01 | 10.67 | 14.69 | 18.30 | 20.31 | 18.86 | 14.91 | 11.41 | 7.20 | 11.12 |
| 1989 | 4.49 | 3.24 | 3.67 | 6.21 | 10.59 | 15.25 | 18.95 | 20.31 | 19.92 | 15.83 | 12.25 | 4.87 | 11.34 |
| 1990 | 3.60 | 4.28 | 4.96 | 6.84 | 10.73 | 14.93 | 18.65 | 20.80 | 20.23 | 17.74 | 12.47 | 9.12 | 12.08 |
| 1991 | 5.72 | 4.76 | 5.61 | 8.11 | 12.26 | 16.61 | 19.53 | 20.48 | 19.99 | 17.11 | 12.00 | 8.17 | 12.59 |
| 1992 | 5.20 | 3.68 | 4.42 | 6.80 | 10.72 | 15.42 | 18.43 | 19.62 | 19.20 | 15.17 | 11.12 | 7.28 | 11.45 |
| 1993 | 5.09 | 3.10 | 3.12 | 6.09 | 11.37 | 15.64 | 18.96 | 20.88 | 19.88 | 15.35 | 11.73 | 8.47 | 11.69 |
| 1994 | 3.15 | 1.59 | 2.81 | 6.62 | 9.96 | 15.37 | 20.30 | 20.78 | 19.27 | 16.27 | 13.21 | 9.15 | 11.60 |
| 1995 | 6.60 | 4.11 | 5.14 | 7.82 | 10.98 | 15.28 | 19.30 | 21.06 | 20.43 | 18.33 | 13.41 | 6.89 | 12.51 |
| 1996 | 3.55 | 2.12 | 2.87 | 5.38 | 9.36 | 14.40 | 17.93 | 19.44 | 18.86 | 15.35 | 10.34 | 6.99 | 10.58 |
| 1997 | 3.72 | 3.20 | 4.16 | 6.50 | 9.38 | 13.66 | 17.94 | 19.47 | 19.06 | 15.83 | 10.99 | 6.31 | 10.90 |
| 1998 | 4.89 | 4.09 | 4.94 | 7.58 | 11.85 | 15.16 | 18.79 | 20.85 | 20.34 | 16.40 | 11.51 | 8.68 | 12.14 |
| 1999 | 4.97 | 4.59 | 5.30 | 8.27 | 12.01 | 17.38 | 20.05 | 21.46 | 21.07 | 17.37 | 13.08 | 9.00 | 12.92 |
| 2000 | 4.62 | 2.73 | 5.12 | 7.48 | 11.43 | 16.40 | 19.81 | 21.05 | 20.47 | 17.06 | 12.72 | 7.38 | 12.22 |
| 2001 | 4.08 | 4.03 | 4.64 | 6.74 | 11.35 | 16.52 | 19.53 | 20.91 | 20.93 | 17.08 | 13.17 | 10.37 | 12.50 |
| Overall mean | 4.13 | 2.99 | 4.02 | 6.75 | 10.74 | 15.33 | 18.90 | 20.47 | 19.78 | 16.26 | 11.98 | 7.64 | 11.63 |
| CV (\%) | 34 | 38 | 25 | 12 | 7 | 5 | 4 | 3 | 3 | 6 | 9 | 18 | 6 |

TABLE 2. Seasonala mean seawater temperature ( ${ }^{\circ} \mathrm{C}$ ) from 1976 through 2001 as calculated from mean daily water temperatures recorded continuously at the intakes of MPS Units I and 2 through July 2001 and at Units 2 and 3 thereafter.


- Winter defined as January-March, spring as April-June, summer as July-September, and fall as October-December.

The mean water temperatures given above reflected conditions in Niantic Bay, where the MPS intakes are located. Water temperature in the Niantic River usually has a larger annual range with colder temperatures found during winter and warmer ones in summer. This most likely occurs because the relatively smaller water volume of the river responds more quickly than LIS to changes in air temperature and solar radiation. During March, when considerable spawning, egg incubation, and larval development take place, mean water temperature in the Niantic River was determined from spot readings taken on several days each week during the annual adult winter flounder surveys. From 1976 through 2001, these means differed from those recorded at MPS by $0.6^{\circ} \mathrm{C}$ or less. In about half of the years, the water temperature in March was slightly warmer in the river than in Niantic Bay and in the remainder, the river was slightly cooler.' Data were insufficient to calculate comparative means during three surveys. In 2001, water temperature in the Niantic River' was about $0.6^{\circ} \mathrm{C}$ cooler than at the MPS intakes in Niantic

Bay, mostly because of colder temperatures occurring early in the month. After a winter storm, water temperature decreased rather abruptly by about $0.7^{\circ} \mathrm{C}$ from late February (ca. $3.2^{\circ} \mathrm{C}$ ) to early March $\left(2.5^{\circ} \mathrm{C}\right)$, with temperatures also cooler in the upper than the lower river. However, temperatures increased relatively rapidly to $4^{\circ} \mathrm{C}$ by mid-March and further to about $4.6^{\circ} \mathrm{C}$ in late March, with the upper river then having warmer temperatures than found in the lower river.


Sampling methodology of the adult winter flounder spawning surveys in the ,Niantic River has been consistent since 1983 (Danila 2000). Surveys usually begin between mid-February and mid-March after most ice cover disappears from the river and continue into April. Sampling ceases when the proportion of
reproductively active females decreased to less than $10 \%$ of all females examined for 2 consecutive weeks, indicating that most spawning is completed. In these surveys, the Niantic River was divided into a number of sampling areas, which are referred to as stations (Fig. 2). No samples were taken outside of the navigational channel in the lower portion of the river during 1979-2000 because of an agreement made with the East Lyme-Waterford Shellfish Commission to protect bay scallop (Argopecten irradians) habitat. With the present lack of both bay scallops and extensive eelgrass beds in the river, permission was obtained to sample shallow areas this year. A feiv tows were randomly taken in the area that is found immediately south of station 51, west of station 6 , and east of the navigational channel, extending as far south as the juvenile winter flounder station WA shown on Figure 2.
Winter flounder were sampled on at least 2 days of each survey week using two survey vessels. A $9.1-\mathrm{m}$ otter trawl with a $6.4-\mathrm{mm}$ bar mesh codend liner was used to capture winter flounder, which were held in water-filled containers aboard the boat before processing. Since 1983, all fish larger than 20 cm were measured to the nearest mm in total length and had gender ascertained. Before 1983, at least 200 randomly selected winter flounder were measured during each week of sampling. Fish not measured were classified into various length and gender groupings and, at minimum, all winter flounder examined were classified as either smaller or larger than 15 cm . Gender and reproductive condition of larger winter flounder were determined by either observing eggs or milt, or as suggested in Smigielski (1975), by noting the presence (males) or absence (females) of ctenii on left-side caudal peduncle scales. Before release, healthy fish larger than 15 cm (1977-82) or 20 cm (1983 and after) were marked in a specific location with a number or letter made by a brass brand cooled in liquid nitrogen.
Since 1983, the brand mark and location have been varied in a unique manner such that the year of marking was apparent for recaptured fish. During 1999-2001, a mark (termed photonic tag) was also applied by the injection of fluorescent pigmented particles into the dorsal, anal, or caudal fin of most freeze-branded winter flounder using a special injector (NEW WEST Technologies BIOMETRIX System 1000 MICRO-Ject $^{T \mathrm{M}}$ or SuperMICRO-Ject ${ }^{T \mathrm{M}}$ portable injectors). Because of problems with the MICRO-Ject ${ }^{\text {TM }}$ applicator in 1999, not all freezebranded fish received a photonic tag that year, but this was corrected prior to the 2000 work. Some fish were also not marked this year, again because of


Fig. 2. Location of stations sampled in the Niantic River during 2001 for adult winter flounder from February 26 through April 11 (numbers) and age-0 winter flounder from May 24 through September 24 (letters).
injector problems. The efficacy of this system versus freeze branding was examined by marking winter flounder using each technique and comparing marks observed on recaptured fish, including fish marked during the 1999-2001 spawning survey and in the 2000 special post-spawning sampling survey (for details of the latter work, see the section entitled Post-Spawning Survey of Adult Winter Flounder in DNC 2001a). This sampling was also expected to provide additional information on the utility of the annual CPUE index as a measure of spawning winter flounder abundance because a comparison could be made between the proportions of fish recaptured in 2001 marked in either the regular and post-spawning surveys.

## Relative Annual Abundance

Data Analyses. Trawl catch-per-unit-effort (CPUE) is used to describe the relative annual abundance of winter flounder $\sum 15 \mathrm{~cm}$ taken in the Niantic River during the late February-early April spawning season. Following standardization of collection methods and catch data, a $\Delta$-mean CPUE was established as the annual relative abundance index of Niantic River winter flounder spawners (NUSCO 1988c). Components of standardization for CPUE calculation included tow length, tow duration, weekly effort, minimum fish length, and, in some instances, gender (see Spawning Stock Size and Egg Production, below). Tow distance was initially measured using radar or LORAN, but more recently with differential Global Positioning System. Tow distance was fixed in 1983 (with exceptions noted below) because using the same tow length at all stations was expected to reduce this component of variability. Previously, tows of variable length had been taken at all stations and catch was standardized by time of tow. A distance of 0.55 km was selected as the standard because it represented the maximum length of a tow that was formerly possible at station 1. However, in 1990, tow distance at station 1 was reduced to 0.46 km because of a new bridge that was constructed at the river mouth. Particularly during 1987, 1989, 1990, and 2000, tows one-half to two-thirds of the standard length were commonly' taken, mostly in the upper river at stations 6 and 51, to avoid overloading the trawl with macroalgae and detritus. Because catch data from station 2 were also used in the TMP, tows there were made over 0.69 km , the standard for that particular sampling program, although shorter tows were often made at this location as well due to high detrital loads.

Because a standard tow distance was not set prior to 1983, catches of winter flounder made during each spawning survey were standardized to a $15-\mathrm{min}$ tow duration at stations 1 and 2 and the shallow flats area and to a $12-\mathrm{min}$ tow time at the other locations. . In general, tow duration was usually greater in the lower river than in the upper river due to differences in tidal currents and amounts of material collected in the trawl, even though distance was similar. To lessen error in the calculation of CPUE, data from either exceptionally long or brief tows, most of which occurred prior to 1983, were excluded from the analyses. The minimum fish length of 15 cm used for CPUE calculation was smaller than the 20 cm used for mark and recapture estimates described below because of data limitations from the 1977-82
surveys. Finally, effort was standardized within each year by replicating as necessary the median CPUE value for a given week such that the number of tows used in calculating CPUE was the same for each week sampled that year.
The 2001 'Spawning Survey. ${ }^{\text {A }}$ Although monthly mean water temperatures in Niantic Bay during December 2000 and January 2001 were not unusually cold (Table 1), ice covered much of the upper half of the Niantic River: With the ice clearing out in late "February, the adult winter flounder spawning survey began on February 26. Sampling in 2001 took place over a 7 -week period and the survey was completed on April 11 (Table 3). Effort was relatively consistent, with 44 to 53 trawl tows taken weekly.
'The rate of spawning by females was determined by observing weekly changes in the percentage of gravid fish larger than 26 cm , the size at which about half of all observed females were mature (NUSCO 1988b). Even though most spawning was completed

TABLE 3. Time period and duration of annual surveys in the Niantic River during the winter flounder spawning season from 1976 through 2001.

|  | Year | Dates sampled | Number of weeks sampled |  |
| :---: | :---: | :---: | :---: | :---: |
|  | 1976 | March I-April 13 | 7 |  |
|  | 1977 | March 7 - April 12 | 6 |  |
|  | 1978 | March 6-April 25 | 8 |  |
|  | 1979 | March 12 - April 17 | 6 |  |
|  | 1980 | March 17 - April 15 | -. 5 |  |
|  | 1981 | March 2 - April 14 | 7 |  |
|  | 1982 | February 22 - April 6 | 7 |  |
|  | 1983 | February 21 - April 6 | 7 |  |
|  | 1984 | February 14-April 4 | 8 |  |
|  | . 1985 | February 27 - April 10 | 7 |  |
|  | 1986 .: | February 24 - April 8 | 7 |  |
|  | 1987 | March 9-April 9 | 5 |  |
|  | 1988 | March 1-April 5 | 6 |  |
|  | 1989 | $\therefore$ February 21 - April 5 | $7{ }^{1}$ |  |
|  | 1990 | February 20-April 4 | 7 |  |
|  | 1991 | February 13-March 20 | 6 | . |
|  | 1992 | February 18 - March 31 | -. 7 |  |
|  | 1993 | February 16-April 7. | . $8^{\text {b }}$ |  |
|  | 1994 | March 22 - April 13 | 4 |  |
|  | 1995 | February 28 - April 6 | 6 |  |
|  | 1996 | February 27 - April 3 | 6 |  |
|  | 1997 | February 18 - March 25 | 6 |  |
|  | 1998 | February 9 - April 1 | 8 |  |
|  | 1999 | February 23 - April 6 | 7 | - |
|  | 2000 | - March 1-April 5 | 6 |  |
|  | 2001 | February 26 - April 11 | 7 |  |

[^11]by late March, as illustrated by the proportion of gravid females observed (Fig. 3), sampling was extended into April to keep effort relatively consistent with that of other recent years and to increase the number of marked fish released this year. Because of the comparatively later start of the 2001 survey and the cooler winter water temperatures found within the river, about two-thirds of observed females were gravid at the start of the survey, unlike the previous 2 years when which spawning occurred much earlier in the season. However, even in 2001 the weekly fractions of gravid females decreased relatively rapidly through late March, similar to declines observed in previous years. Unlike 2000 and 2001, when the number of gravid females decreased sharply, increases in the proportion of gravid females occurred during mid-March in both 1998 and 1999. These increases followed periods of cold weather, which may have reduced the rate of spawning. Because a majority of females were spent at the start of surveys in most years, much of the spawning apparently occurs earlier in winter. Exceptions include extremely cold years (e.g., 1978, 1996, 2001), when many gravid females were present even after relatively late starting dates of the surveys because of heavy ice cover (Fig. 4). In contrast, the proportions of gravid females in warmer years (e.g., 1989-91) were not only lower at the start of sampling, spawning was also completed earlier, even though the surveys commenced earlier in the season.
The large concentrations of macroalgae that hindered sampling in many areas of the river during the late 1980s and early 1990s were mostly absent during the past few years, with some notable exceptions. Similar to the past few years, less material was taken in most areas of the northwestern river arm and the trawl often contained mud, leaves, and molluse shells, particularly at station 53 .


Fig. 3. Weekly percentage of gravid Niantic River female winter flounder larger than 26 cm taken during the 19982001 adult population abundance surveys.


Fig. 4. Weekly percentage of gravid Niantic River female winter flounder larger than 26 cm taken during examples of warm (1989-91) and cold (1978, 1996, 2001) years of adult population abundance surveys. (Note that the vertical scales differ between the graphs).

However, beginning in mid-March, a filamentous alga, Ectocarpus siliculosus, became common and by clogging trawl mesh openings reduced sampling efficiency in stations 6 and 51 and in the shallow area south of station 51 . Also, for the first time in the 26 years of these surveys, large masses of the blue mussel (Mytilus edulis) were found covering the bottom throughout most of station 1 in the lower river navigational channel. The extensive mussel beds likely considerably reduced the demersal habitat favored by winter flounder in this area of the Niantic River.
As winter flounder abundance decreased through the 1990s, adults appeared to concentrate into relatively small areas of the Niantic River, such as the upper river arm (stations $52-54$ ) and particularly at station 54 . Few fish were taken at stations 1,2 , or within large portions of 51 . Only two winter flounder larger than 20 cm were taken at station 1 in 2001, likely because of the mussel cover noted above. Along the eastern shoreline of the river (stations 6 and adjacent 51), ripe males were more abundant than females, indicating that these areas were likely
spawning sites. Because large adults were also common in the shallow area of northernmost station 54, winter flounder probably complete most spawning in shallower areas of the river.
Relative Abundance Index. Because of an increasing frequency of zero catches in the mid1990s, the relative abundance index was changed from a median to a $\triangle$-mean CPUE (described in NUSCO 1988c), which is the best estimator of the population mean when the data come from a distribution that contains numerous zero values and the distribution of the non-zero values is approximately lognormal (Pennington 1983, 1986). Prior to 1993, 97-100\% of tows taken each year had at least one winter flounder $\geq 15 \mathrm{~cm}$, but the fraction each year since then varied from as high as $95 \%$ to a low of $69 \%$, with the latter value found in 2001. As a result of the sparse catches of winter flounder in 2001, a $\Delta$-mean CPUE of 1.4 fish per standardized tow was determined (Table 4; Fig. 5). This was the


Fig. 5.: Ȧnnual $\triangle$-mean CPUE and 95\% confidence interval of Niantic River winter flounder larger than 15 cm . from 1976 through 2001.
lowest CPUE found in 26 years, although based on broadly overlapping $95 \% \mathrm{Cls}$, this value was not significantly different from the $\Delta$-mean of 1.6 found in 1996. In the DEP trawl survey, which takes place

TABLE 4. Annual $9.1-\mathrm{m}$ otter trawl adjusted $\triangle$-mean CPUE of winter flounder larger than $15 \mathrm{~cm}^{\mathrm{b}}$ taken throughout the Niantic River during the 1976 through 2001 adult population abundance surveys.'

| Survey year | Weeks used for CPUE computation ${ }^{\text {c }}$ | Tows acceptable for CPUE ${ }^{d}$ | Adjusted number of tows used ${ }^{\text {c }}$ | Non-zero observations | $\Delta$-mean CPUE estimate | Standard crror | $95 \%$ confidence interval for $\Delta$-mean CPUE |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1976 | 7 | 169 | 224 | 222 | 48.4 | 2.9 | 42.7-54.1 |
| 1977 | 6 : | 223 | 264 | 261 | 27.5 | - 1.7 | 24.3-30.8 |
| 1978 | 6 | 150 | 174 | $\bigcirc 174$ | 31.2 | 2.3 | 26.6-35.7 |
| 1979 | 5 | 127 | 140 | 140 | 41.0 | 4.0 | 33.0-48.9 |
| 1980 | 5 | 117 | 150 | 149 | $41.5{ }^{\circ}$ | 2.9 | 35.9-47.1 |
| 1981 | 7 | 181 | 232 | 232 | 50.8 | 2.5 | 45.9-55.7 |
| 1982 | . 5 | 118 | 149 | -149... . | 47.8 | 3.5 | 41.0-54.6 |
| 1983 | 7 | 232 | 238 | 237 | 31.3 | 1.3 | 28.8-33.9 |
| 1984 | 7 | 246 | 287 | 286 | 18.4 | 0.7 | 17.1-19.7 |
| 1985 | 7 | 268 | 280 | ' 277 | 17.1 | 0.7 | 15.7-18.5 |
| 1986 | 7 | 313 | 343 | 341 | 12.2 | 0.5 | 11.2-13.3 |
| 1987 | 5 | 234 | 270 | 267 | 16.9 | 0.9 | 15.2-18.6 |
| 1988 | 6 | 292 | 312 | 310 | 17.9 | 0.7 | . 16.5-19.3 |
| 1989 | 6 | - 272 | 306 | 302 | 13.9 | 0.7 | 12.6-15.1 |
| 1990 | 7 | 307 | 343 | $342{ }^{\text { }}$ | 11.2 | 0.5 | 10.3-12.2 |
| 1991 | 6 | 301 | $\because 330$ | - 324 - | 16.7 | 0.9 | - 14.9-18.5 |
| 1992 | 7 | 380 | 406 | 395 | 7.7 | 0.3 | '7.0-8.3 |
| 1993 | 7 | 288 | 392 | 344 | 3.4 | 0.2 | 3.0-3.7 |
| 1994 | 4 | 185 | 212 | 201 | 6.4 | 0.5 | 5.5-7.3 |
| -1995 | 6 | 320 | 342 | 283 | 2.6 | 0.1 | $\therefore$ 2.4-2.9 |
| 1996 | 6 | 310 | 342 | 242 , | 1.6 | 0.1 | 1.4-1.8 |
| 1997 | 6 | 302 | 348 | 288 | 2.4 | 0.1 | . 2.2-2.6 |
| 1998 | 7 | 363 | 385 | 306 | 2.1 | 0.1 | 1.9-2.3 |
| 1999 | 7 | 336 | 364 | 297 | 2.7 | 0.2 | 2.4-3.1 |
| 2000 | 6 | 305 | 324 | 248 ': | 3.0 | 0.2 | 2.5-3.4 |
| 2001 | 7 | - '357 | $\cdots 371$. | '255: . | - 1.4 | 0.1 | 1.3-1.6 |

[^12]throughout LIS, Shake (2001) reported that the 1999 and 2000 abundance indices for winter flounder (fish of all ages taken in April and May) were the two lowest of the time-series and about one-half of the long-term (1984-2000) mean. Age-2 and 3 fish had particularly low abundances. The age-4+ index (adults) in 2000 was the sixth lowest of the series and just under the long-term average. In the same DEP annual report, MacLeod (2001) noted that the recreational fishing total catch and harvest of winter flounder in 1999 decreased considerably from 1998 and was the lowest ever recorded. A conclusion of this DEP report was that winter flounder did not appear to be responding to management measures to increase abundance as have some other recreational species. Preliminary indices from the 2001 DEP trawl survey (D. Molnar, CT DEP, Old Lyme, CT, pers. comm.) indicated an increase of about $20 \%$ in the winter flounder abundance index for fish of all ages and a $15 \%$ increase for fish age- $4+$, although the former remained as the third lowest of the timeseries.
In comparison to 2000 , the fraction of female winter flounder in the 29 - to $39-\mathrm{cm}$ size-classes decreased sharply this year (Fig. 6). Relatively similar abundances of both smaller and larger fish, however, were found during the past several years. Various peaks in abundance that occurred between 30 and 36 cm (probably ages-4 through 6; NUSCO 1987) likely represented variation in year-class strength of spawning females. The decline in female winter flounder abundance over the entire period of sampling was evident when catches from 1981 (largest CPUE since 1976; Table 4), 1985, 1990, and 2001 were compared (Fig. 7). The high abundance of females from 23 to 34 cm in 1981 were likely fish from he strong year-classes produced during the mid to late 1970s. Conversely, the low abundance of mid-sized spawners in 2001 was likely related to relatively low production of juveniles in the mid1990s (see Juvenile Winter Flounder, below).
Since the early 1980 s , large decreases in abundance have occurred for all size-classes of female winter flounder, with the exception of the very largest ( $\geq 40$ cm ) females, which were never very numerous. Larger fish have made up a relatively greater proportion of adult female winter flounder in the Niantic River since the mid-1990s. Shake (2001) also noted that large winter flounder were more common in the DEP LIS spring trawl survey in 1999 and 2000 than in previous years. However, despite relatively high abundance of age-0 fish produced in 1988, 1992 and 1994, female winter flounder from

20 to 30 cm in length did not become particularly abundant in subsequent years. Similar trends were noted by Simpson et al. (1996) for the 1988 and 1992 year-classes of winter flounder in LIS and by Desfosse et al. (1998) in Massachusetts. They reported that apparently strong year-classes observed at ages- 1 and 2 did not result in expected higher abundances at age- 3 and older. Among the explanations suggested for this phenomenon were variable discard mortality of juveniles in the commercial fishery, high rates of fishing mortality once the winter flounder recruited to the fisheries, and nonrandom fishing effort that occurred in overfished stocks (Simpson et al. 1996; Desfosse et al. 1998). Factors critical to the formation of winter flounder year-class strength are discussed in a section appearing later in this report.
Another factor affecting winter flounder spawner abundance in the Niantic River during recent years has been seal predation. Particularly in years when abundance is low, this source of mortality may be of importance. Harbor seals (Phoca vitulina concolor) were first noted in the Niantic River during the 1993 winter flounder spawning season. This year, individual seals were occasionally seen in waters inside and outside the mouth of the river and three seals were frequently observed as they hauled out at the Niantic River boat launching ramp during mid-March (M. Male, Old Lyme, CT, pers. comm.). At least ten dead or injured winter flounder were captured in 2001 that had wounds consistent with seal attacks; half of these fish were taken during the first 2 weeks of sampling when most spawning took place. This total represents the largest number of deaths or injuries observed during these surveys to date. Since passage of the Marine Mammal Protection Act in 1972, harbor seals have increased in numbers (Blaylock et al. 1995) and have expanded their range southward from northern New England (Payne and Schneider 1984). Although some studies show harbor seals prefer to feed on small, schooling fishes, they will also opportunistically prey on seasonally dominant species and have the capacity to shift prey selection rapidly in response to shifts in availability (Payne and Selzer 1989). Sampling also indicated that suitable alternative prey species were not common in the Niantic River during the winter flounder spawning season.
Recaptures from the 2000 Regular and Special Post-spawning Surveys. In 2000, a $\Delta$-mean of 11.8 winter flounder larger than 20 cm was recorded during the special post-spawning survey, a value larger than all annual spawning survey CPUE


Fig. 6. Comparison of annual standardized catch by length of female winter flounder 20 cm and larger taken in the Niantic River during the spawning season from 1999 through 2001.


Fig. 7. Comparison of annual standardized catch by length of female winter flounder 20 cm and larger taken in the Niantic River during the spawning season in 1981, 1985, 1990, and 2001.
estimates made since 1991 (DNC 2001a). A total of 784 fish was uniquely marked and released during this mid-April through mid-May sampling, considerably more than the 593 marked during the March-early April spawning survey (both surveys were 6 weeks in duration). Although the postspawning survey CPUE was considered somewhat biased because effort was not allocated to all areas of the river and was concentrated in the relatively deeper waters of the upper river basin (sta. 51; Fig. 2 ), the relatively high value nevertheless indicated that many more-winter flounder were available for capture in a large portion of the Niantic River than were found in that area during the regular spawning survey. In addition, larger-sized fish and females were caught in significantly greater proportions in the post-spawning than in the regular spawning survey. However, percentages of fish marked during the 1998 and 1999 spawning surveys (combined totals) and recaptured were nearly the same in each of the 2000 surveys ( $4.4,5.4 \%$ ). These findings suggested that winter flounder were perhaps more common in the Niantic River than indicated by the lower CPUE of 3.0 determined in the regular spawning survey. It was also possible that some fish from other spawning stocks could have entered the river to feed as part of normal post-spawning movements in spring. However, recaptures of the fish marked in both 2000 surveys during 2001 provided additional insights regarding the status of these winter flounder. The fractions of fish that were marked in the 2000 surveys and subsequently recaptured in 2001 were practically identical: $2.9 \%$ for the spawning survey fish and $2.8 \%$ for the post-spawning survey specimens. The similar recapture rates of previously marked fish found in both 2000 and 2001 indicated that the winter flounder found in the Niantic River from winter into spring likely represented the same stock, with numbers not appreciably increased by immigration from other populations.
These results further illustrated that the magnitude of annual CPUE can be affected by when and where effort is expended. In particular, surveys from 1976 through 1982, when annual winter flounder CPUE values were generally high, were less structured in terms of areal effort allocation. Sampling effort often occurred where fish were most common, and the surveys also went later into spring when winter flounder were apparently more available. As spring progresses, winter flounder withdraw into the upper river basin from other areas and this alone probably accounted for the appreciable increase in CPUE seen during 2000. Increases in weekly catches seen during the final weeks of sampling in 2001, as well
as in most other years, likely reflected a similar movement of winter flounder into deeper areas of the river from shallow flats not sampled as water temperatures increased. However, the relatively few tows taken on the shallow flat south of station 51 this year did not produce any more winter flounder than were found in deeper portions of the upper river. Therefore, additional winter flounder present in the river must have been located in other shallow-water areas not surveyed, including Smith Cove, which cannot be sampled because of numerous moorings and soft sediment.

## Absolute Abundance Estimates

Data Analyses. Absolute abundance estimates of winter flounder spawning in the Niantic River were obtained using mark-and-recapture methodology and the Jolly (1965) stochastic model. This model is appropriate for open populations as long as basic assumptions are approximately met (see Cormack 1968; Southwood 1978; Begon 1979; Pollock et al. 1990 for details). Annual absolute abundance estimates for Niantic River winter flounder larger than 20 cm were calculated by pooling together all fish marked and released during each annual survey and by observing the recaptures made in subsequent years. Absolute abundance estimates could not be generated for years prior to 1984 because of uncertainty in data records and ambiguity caused by re-use of brands from earlier surveys. Estimates of annual population size $(N)$ and other model parameters, including survival ( $\phi$ ), recruitment ( $B$ ), and sampling intensity ( $p$ ), were made using the computer program 'JOLLY' (Pollock et al. 1990).
Comparison of Marking Methodology. For the past 3 years, most winter flounder $\geq 20 \mathrm{~cm}$ in length were tagged with both a freeze brand and a photonic pigment mark. Observations of recaptured fish were expected to provide information on the efficacy of both marking methodologies (i.e., whether marks were successfully retained for 1 or more years). The rationale for this work was that each year a few winter flounder have been caught with an apparent freeze brand, but the exact year of marking could not be ascertained because the mark was indistinct. This loss of mark and recapture information warranted an examination of another tagging methodology.
Observations of both within-year and followingyear recaptures showed that no winter flounder were caught that had a photonic brand, but lacked a freeze brand. Thus, initially the freeze brand appeared to be the more stable mark. Only $82 \%$ of 55 within-year
recaptures of freeze-banded fish in 1999 had an observable photonic mark, although this result was confounded because about one-third of the fish were not tagged or were poorly marked due to equipment problems with this marking system. In addition, even though the photonic mark fluoresces under UV light, difficulties were encountered in the field while using these lights and the light boxes constructed for this purpose. Subsequently, this practice was abandoned. As a result of increased expertise in using the photonic marking equipment, $100 \%$ and $96 \%$ of within-year freeze-brand recaptures seen during the 2000 regular (total of 69 observed) and special postspawning ( 178 ) surveys, respectively, also had a photonic mark. In 2001, 93\% (41) of fish had both marks, indicating that these brands were relatively comparable in the short term, as long as there were no photonic injector problems. Only $31 \%$ (36) of the 1999 photonic marks were observed on freeze-band recaptures in 2000 (both surveys combined) and 20\% (10) in 2001 , percentages which were expected to be low because not all fish were released with a photonic mark. Using an estimate of fish not having a photonic mark but released in 1999 resulted in corrected recapture rates of $56 \%$ and $36 \%$, respectively, for these 2 years. Nearly all fish received a photonic mark in 2000 , but only $72 \%$ (18) and $85 \%$ (26) of fish recaptured in 2001 during the regular and special post-spawning surveys, respectively, had both brand types. The photonic marks were more difficult to apply, particularly on larger winter flounder, which have a tough integument resistant to retention of the pigment particles. Observations of recaptures were also more difficult, with the pigment mark often very small or faint. Because of occasional equipment problems along with less than complete photonic mark retention in the long-term, the freeze brand remains as the tagging method of choice for the Niantic River winter flounder surveys. Photonic marks will therefore no longer be applied to winter flounder as of the 2002 survey.

The 2001 Spawning Survey. During 2001, the number of winter flounder $\geq 20 \mathrm{~cm}$ in length that were caught and freeze-branded in the Niantic River varied from 25 in the second week of sampling to 69 . during the final week. Numbers marked (62-69) were relatively consistent during the last 4 weeks of the survey. . As noted previously, environmental effects (especially water temperature) and sampling efficiency due to macroalgal loading likely influenced the susceptibility of fish to capture as did the aforementioned availability of fish. Because of low abundance of Niantic River winter flounder population since 1995 , only 370 winter flounder were
marked with a freeze brand and released during 2001, a few fish less than in 1996 (376), which was the previous low, and more than several hundred fewer than marked during the past 4 years (Table 5). Twenty-nine previously-marked fish were recaptured in 2001, similar to the numbers taken in 1999 and 2000, which were the lowest totals to date. The proportion of recaptured fish to the total observed within a year was particularly low (4.1-6.0\%) during 1997-2000 in comparison to 1992-96 (7.1-14.8\%), and the recapture rate of $7.8 \%$ in 2001 represents a modest increase from the past few years. The fate of marked adults between spawning seasons is unknown. A previous summary of movements and exploitation of Niantic River winter flounder given in NUSCO (1986a, 1987) indicated that nearly all fish moved from the river to offshore waters by late spring following spawning. Most fish moved to the east, with many fish taken by the sport and commercial fisheries in Connecticut and New York waters of LIS, still others off Rhode Island and southern Massachusetts, and finally, some individuals moved as far as Nantucket Shoals or were taken off southern Cape Cod. These fish return to Connecticut and Niantic Bay waters in fall and early winter prior to entering the river for spawning and are subjected to a local trawl fishery during that period.
Most ( $17=59 \%$ ) of the fish recaptured in 2001 were marked in 2000, with others branded in 1999 (8), 1998 (3), and 1996 (1). Excluding 1984 and 1985, years at the start of the mark-recapture data series, and 1994 (36\%) and 1997 (33\%), fish marked during the preceding year accounted for about onehalf to two-thirds of all recaptures. During both 1994 and 1997, however, fewer recaptured fish were marked during the immediately preceding year in comparison to other years. It appears that in recent years fewer fish which had been marked more than 3 years previous to the survey were recaptured in comparison to previous years, but whether this was due to fewer fish being branded and available for recapture, higher rates of mortality following release, or a combination of the two is unknown.
Absolute Abundance Estimates. The markrecapture data from 2001 provided an initial abundance estimate of 9,300 winter flounder larger than 20 cm and present in the Niantic River during the 2000 spawning season (Table 6). 'Because standard errors of $N$ appear to be correlated with $N$ due to'the particular form of Jolly's variance formula, the $95 \%$ Cls computed are generally considered unreliable as a measure of sampling error except at very high sampling intensities (Manly 1971; Roff

TABLE 5. Mark and recapture data from 1983 through 2001 used to estimate abundance of winter tlounder larger than 20 cm in the Niantic River during the spawning season.

|  | Survey year |  | Number observed |  |  |  | Number notpreviously marked |  |  |  | Number marked and released |  |  |  | Number recaptured |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 1983 |  |  |  | 5,615 |  |  |  |  |  |  |  |  |  |  |  | ) |  |
|  | 1984 |  |  |  | 4,103 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1985 |  |  |  | 3,501 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1986 |  |  |  | 3,031 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1987 |  |  |  | 2,578 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1988 |  |  |  | 4,333 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1989 |  |  |  | 2,821 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1990 |  |  |  | 2,297 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1991 |  |  |  | 4,333 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1992 |  |  |  | 2,346 |  |  |  |  |  |  |  |  |  |  |  |  |  |
|  | 1993 |  |  |  | 984 |  |  |  | 38 |  |  |  | 2 |  |  |  |  |  |
|  | 1994 |  |  |  | 1,035 |  |  |  | 1 |  |  |  |  |  |  |  |  |  |
|  | 1995 |  |  |  | 682 |  |  |  | 8 |  |  |  | 1 |  |  |  |  |  |
|  | 1996 |  |  |  | 379 |  |  |  | 42 |  |  |  | 6 |  |  |  |  |  |
|  | 1997 |  |  |  | 642 |  |  |  | 05 |  |  |  | 2 |  |  |  |  |  |
|  | 1998 |  |  |  | 637 |  |  |  | 00 |  |  |  | 2 |  |  |  |  |  |
|  | 1999 |  |  |  | 617 |  |  |  | 92 |  |  |  | 4 |  |  |  |  |  |
|  | 2000 |  |  |  | 594 |  |  |  | 88 |  |  |  | 3 |  |  |  |  |  |
|  | 2001 |  |  |  | 371 |  |  |  | 42 |  |  |  | 0 |  |  |  |  |  |
| Number of fish marked in a given year that were recaptured during subsequent annual surveys |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Survey year | y 83 | 84 | 85 | 86 | 87 | 88 | 89 | 90 | 91 | 92 | 93 | 94 | 95 | 96 | 97 | 98 | 99 | 00 |
| 1984 | 130 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 1985 | 47 | 94 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 1986 | 23 | 45 | 76 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 1987 | 2 | 13 | 27 | 73 |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 1988 | 7 | 22 | 31 | 62 | 104 |  |  |  |  |  |  |  |  |  |  |  |  |  |
| 1989 | 2 | 11 | 9 | 33 | 32 | 145 |  |  |  |  |  |  |  |  |  |  |  |  |
| 1990 | 1 | 7 | 4 | 15 | 13 | 37 | 78 |  |  |  |  |  |  |  |  |  |  |  |
| 1991 | 1 | 4 | 4 | 12 | 25 | 30 | 50 | 126 |  |  |  |  |  |  |  |  |  |  |
| 1992 | 0 | 0 | 1 | 1 | 3 | 21 | 20 | 50 | 114 |  |  |  |  |  |  |  |  |  |
| 1993 | 0 | 0 | 0 | 1 | 0 | 4 | 3 | 15 | 20 | 103 |  |  |  |  |  |  |  |  |
| 1994 | 0 | 0 | 0 | 1 | 0 | 0 | 4 | 5 | 12 | 25 | 27 |  |  |  |  |  |  |  |
| 1995 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 2 | 8 | 8 | 17 | 41 |  |  |  |  |  |  |
| 1996 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 2 | 4 | 5 | 4 | 20 |  |  |  |  |  |
| 1997 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 1 | 6 | 3 | 5 | 7 | 13 |  |  |  |  |
| 1998 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 2 | 0 | 4 | 5 | 2 | 23 |  |  |  |
| 1999 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 3 | 4 | 17 |  |  |
| 2000 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 1 | 5 | 18 |  |
| 2001 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 3 | 8 | 17 |

1973; Pollock et al. 1990). Abundance estimates for the most recent years are subject to change as additional recaptures are made during future surveys. For example, because of the recaptures made in 2001, the initial abundance estimate given in DNC (2001a) of 6,749 fish for 1999 increased by 342 to 7,091 . The 1997 estimate also increased ( +376 ), but those for $1998(-178)$ and $1996(-386)$ decreased relative to results presented last year.
Sampling intensity ( $p$ ), or the probability that a fish will be captured, was estimated as 0.062 for 2000 ,
which was below the series average of 0.071 for this parameter. Some of the highest estimates occurred from 1997 through 1999, which may have reflected relatively intense sampling effort on fish concentrated in a few small areas of the river. Sampling intensities of at least 0.10 are recommended to obtain reliable and precise estimates of population size and survival rates with the Jolly model (Bishop and Sheppard 1973; Nichols et al. 1981), although Hightower and Gilbert (1984) found that low sampling effort may give acceptable estimates if

TABLE 6. Estimated abundance ${ }^{2}$ of winter flounder larger than 20 cm taken during the spawning season in the Niantic River from 1984 through 2000 as determined by the Jolly (1965) mark and recapture model.


- Annual estimates of these parameters may vary from those reported in DNC (2001a) because of mark and recapture data added from the 2001 adult winter flounder population survey (Table 5).
population size is relatively large and the number of marked animals is also relatively high. However, Gilbert (1973) and Carothers (1973) reported that $N$ was underestimated and had low accuracy when sampling intensities were low (0.05-0.09), regardless of population size or number of fish marked. Estimates of $p$ only approximated or exceeded 0.10 in 1992 (0.138) and 1998 (0.112), so $N$ has likely been underestimated. Loss of information because
brands were missed or from mortality of fish handled also 'requires increased sampling effort. Other sampling errors, model assumptions, and biases inherent in the Jolly'model that could have affected these estimates were discussed in NUSCO (1989) and Pollock et al. (1990).
Although the Jolly estimates are subject to some error, annual $\Delta$-mean CPUE and Jolly abundance estimates were significantly correlated (Spearman's
rank-order correlation; $r=0.942 ; p=0.0001 ; n=17$; Fig. 8). Thus, based on a $\Delta$-mean CPUE of 1.4 for winter flounder 20 cm and larger in 2001, absolute abundance of winter flounder this year likely was less than 10 thousand fish. By extrapolation from the CPUE in 1981, total abundance for that year of extreme abundance in the river may have been between 150 to 200 thousand winter flounder larger than 20 cm .

Estimates of survival ( $\Phi$ ) have varied considerably from year to year (0.189-0.908; Table 6). The initial value of 0.470 for 1999 was similar to the long-term average of 0.461 , after several years of lower than average estimates, which reflected low rates of recapture in those years. Except for 1993 ( 12,184 fish), estimated recruitment ( $B$ ) values were relatively low following 1990. Estimates of both $\Phi$ and $B$ are considered to be much less reliable than those of abundance when using the Jolly model (Manly 1971; Cormack 1972; Bishop and Sheppard 1973; Buckland 1980, 1982; Arnason and Mills 1981; Hightower and Gilbert 1984; NUSCO 1986b) and have not been used as quantitative estimators of survival or recruitment in the Niantic River winter flounder studies. Estimates of $B$ are also relatively imprecise, having $95 \%$ Cls with wide ranges. As for other parameter estimates based on only I year of recapture information, estimates of $\Phi$ and $B$ based on only 1 or 2 years of recapture data may change considerably with the addition of data from subsequent annual surveys.


Fig. 8. Comparison between estimates of absolute abundance of winter flounder larger than 20 cm in the Niantic River during the spawning season and the corresponding $\Delta$-mean CPUE from 1984 through 2000.

## Spawning Stock Size and Egg Production

A second index of relative abundance was based on the size distribution of female fish from adult spawning survey catches standardized by weekly and
yearly effort (i.e., number of tows). Catches were adjusted by effort to insure that each size group of fish was given equal weight within each week of sampling, among weeks in each survey year, and to adjust for varying effort among years (see NUSCO 1989 for more details). To avoid confusion with the CPUE index, this adjusted catch is referred to as "annual standardized catch" throughout the remainder of this report. The annual standardized catch of Niantic River female winter flounder spawning stock was the basis for the calculation of annual recruitment and egg production.

The annual standardized catch of spawning females and the production of eggs by this stock were determined from available data on abundance, sex ratios, sexual maturity, and size-frequencies. The sex ratio of winter flounder larger than 20 cm during the 2001 spawning season in the Niantic River was 2.39 females to each male (Table 7; Fig. 9). In most years more females than males were taken and this sex ratio was larger than the long-term average of 1.56 for the 25 -year time-series. In general, more highly

- skewed ( $\geq 1.70$ ) sex ratios have been found since

TABLE 7. Female to male sex ratios of winter flounder taken during the spawning season in the Niantic River from 1977 through 2001.

| Year | All fish captured | Measured <br> fish $>20 \mathrm{~cm}$ |
| :---: | :---: | :---: |
| 1977 | 1.03 | 1.26 |
| 1978 | 2.23 | 1.95 |
| 1979 | 1.37 | 1.21 |
| 1980 | 2.66 | 2.03 |
| 1981 | 1.42 | 1.61 |
| 1982 | 1.16 | 1.50 |
| 1983 | 1.52 | 1.52 |
| 1984 | 1.07 | 1.07 |
| 1985 | 1.37 | 1.37 |
| 1986 | 0.92 | 0.92 |
| 1987 | 0.78 | 0.78 |
| 1988 | 1.50 | 1.50 |
| 1989 | 1.32 | 1.32 |
| 1990 | 1.24 | 1.24 |
| 1991 | 1.22 | 1.22 |
| 1992 | 1.26 | 1.26 |
| 1993 | 1.47 | 1.47 |
| 1994 | 1.70 | 1.70 |
| 1995 | 2.70 | 2.70 |
| 1996 | 1.78 | 1.78 |
| 1997 | 2.40 | 2.40 |
| 1998 | 2.18 | 2.18 |
| 1999 | 2.36 | 2.36 |
| 2000 | 2.15 | 2.15 |
| 2001 | 2.39 | 2.39 |
|  | 1.56 | 1.56 |
| Geometric mean |  |  |



Fig. 9. Female to male sex ratio of Niantic River winter flounder larger than 20 cm from 1976 through 2000.

1994 as winter flounder abundance decreased. Since 1986,' a highly significant increasing trend in sex ratio has occurred (Mann-Kendall test, $Z=3.42 ; \mathrm{n}=$ $16 ; \mathrm{p}<0.001$ ). The reasons for the preponderance in females are unknown. Mature females are larger than males, so greater exploitation of females might be expected, which would tend to disproportionately decrease their numbers in comparison to males. However, this seems to be contrary to the above findings and may indicate susceptibilities that differ by gender for fishing (e.g., availability), natural mortality (e.g., predation, disease, or longevity), or their availability to trawl sampling during the winter flounder survey (e.g., males remaining to spawn in shallow areas less frequently sampled). Ratios of 1.50 to 2.33 also in favor of females were reported by Stoner et al. (1999) for the Navesink River, New Jersey, and by Saila (1962a, 1962b) and Howe and Coates (1975) for other winter flounder populations in southern New England. Witherell and Burnett (1993) found greater proportions of female winter flounder in Massachusetts waters, particularly in older age-classes." Based on evidence of greater senescent mortality reported for males by Burton and Idler (1984), they believed that males likely have a higher natural mortality rate, which would tend to skew sex ratios in favor of females.
The gender of winter flounder larger than 20 cm during the Niantic River surveys was ascertained by external examination of collected specimens. Correct classification to gender was readily obvious in fish that were ripe or gravid, as it was for recently spawned females having distinctly thin, flaccid ovaries. However, fish not in spawning condition were sexed by palpation of their scales'for ctenii according to the method of Smigielski (1975). Bejda and Phelan (1998) noted that the latter methodology may be subject to error, but, as discussed in NUSCO (1999), a systematic misclassification of fish by
gender was unlikely during the spawning survey. Although errors in sexing fish can and probably do occur during external examinations, the skewed sex ratio found for Niantic River winter flounder during the spawning season appears to be real and likely occurs for the reasons given above.
Stoner et al. (1999) observed that female winter flounder held in a large laboratory aquarium spawned on average 40 times during the season. "Most individuals spawned over a duration of about 1 week. Similarly, in laboratory spawning of winter flounder by NUSCO in the early 1980s, some females were observed to spawn more than once over a period of several days. However, throughout the many years of study in the Niantic River only a few females were observed in a condition intermediate between gravid and spent. These observations suggest that perhaps some additional spawning by individuals may occur or that a spawning event, which takes place at night (Stoner et al. 1999), may have been interrupted. However, most females examined appeared to have been either fully gravid, completely spent after recent spawning, or in a post-spawning (i.e., feeding) state. Because Burton and Idler (1984) reported that winter* flounder only produce one batch of mature eggs each year, fecundity estimates are likely unaffected even if eggs are released during more than one spawning event, unlike a serially spawning species, such as the tautog (Tautoga onitis), which continuously produces batches of eggs over an extended period.
The proportion of mature female winter flounder in each $0.5-\mathrm{cm}$ length increment beginning at 20 cm was estimated from qualitative observations of reproductive condition (percent maturity by $0.5-\mathrm{cm}$ size-classes) made from 1981 through the present. About half of all observed females were observed to be mature at $26^{\circ} \mathrm{cm}$ (NUSCO 1988b), which is comparable to $\mathrm{L}_{50}$ estimates of size-at-maturity of 25.8 cm reported for LIS by Johnson et al. (1998) and 28.3 and 27.6 cm for Massachusetts waters by Witherell and Burnett (1993) and O'Brien et al. (1993), respectively. Pooled estimates were adjusted to give continuously increasing fractions of mature fish through 34 cm . All females 34 cm or larger were considered to be mature (Fig. 10). The fecundity (annual egg production per female) was estimated for each $0.5-\mathrm{cm}$ size-class by using the following relationship determined for Niantic River winter flounder (NUSCO 1987):

$$
\begin{equation*}
\text { fecundity }=0.0824 \text { (length) }{ }^{4.000} \text {. } \tag{1}
\end{equation*}
$$

The length-fecundity relationship was used with the annual standardized catch of mature females and


Fig. 10. Empirically derived proportion of mature Niantic River female winter flounder by length used in the calculation of stock and recruitment. All females smaller than 20 cm were immature and larger than 34 cm were mature.
their length composition to calculate egg production. Although some density-dependent regulatory mechanisms have been proposed for adult flatfishes, size-specific fecundity appears to be stable over a wide range of abundances and may decrease only at the very highest densities (Rijnsdorp 1994). Thus, no adjustments were considered for the Niantic River winter flounder length-fecundity relationship. Annual mean fecundity was determined by dividing the sum of all individual egg production estimates by the standardized catch of females spawning per year.
Since an excellent correlation ( $r=0.942$ ) exists between $\triangle$-mean trawl CPUE and Jolly abundance estimates, the annual standardized catches of all fish $\geq 20 \mathrm{~cm}$ for 1984-99 (the 2000 estimate was considered as preliminary and not used) were divided by the corresponding annual total abundance estimate from the Jolly model. Annual ratios ranged between 2.2 and $5.4 \%$ and had a geometric mean of $3.39 \%$. Thus, the relative numbers of females and eggs produced each year, as determined from the standardized catches, were conservatively assumed to represent about $3.39 \%$ of the absolute values and a multiplier of 29.5 was used to scale up standardized catch indices to absolute numbers of female winter flounder spawning in the Niantic River. In using this scaling factor, it was assumed that ratios of annual standardized catch to absolute abundance during 1977 through 1983 would have been similar to those of later years had estimates of absolute abundance been available for the earlier period. Annual estimates of the number of spawning females were also used to determine egg production and for the derivation of a relationship between stock and recruitment for Niantic River winter flounder. As this scaling factor differed from that used in DNC
(2001a) and previous reports, revised estimates of female stock size, recruitment, and egg production resulted.
Estimates of female stock size ranged from a maximum of 71,140 fish in 1982 to a minimum of 2,009 in 2001 (Table 8). Female abundance in 2001 was less than one-half of the 2000 estimate. Despite the preponderance of larger mature females in the Niantic River, the number of spawning females has been very low since 1995 because of low overall abundance of winter flounder. Mature females usually comprised approximately one-third to onehalf of all winter flounder 20 cm and larger. Highest fractions of mature females were most often found in recent years, including 1995 (63\%) and 1997 (60\%). The high fractions were related to highly skewed sex ratios and larger fish present during these years. Including only females larger than 20 cm , mature fish made up from 51 to $94 \%$ of annual female totals. Higher values indicated that, in many years, smaller and younger non-spawning females were not present in the Niantic River or that incoming year-classes were very weak. For some years, including 1999 ( $65 \%$ ) and 2001 ( $69 \%$ ), smaller fractions of mature females reflected the presence of larger numbers of immature winter flounder, individuals which were expected to recruit into the spawning population in subsequent years. However, this apparently has not resulted in increased stock size as shown by abundance estimates for the past 2 years.
Annual egg production estimates were determined using annual catch, age, maturity, and length information with the length-fecundity relationship for Niantic River winter flounder. Yearly egg production during 1977-2001 ranged between about 1.7 to 43.7 billion (Table 8). Differences in percent maturity resulting from variable annual lengthfrequency distributions affected mean fecundity, which was low during the late 1970 s when smaller fish were more abundant. However, mean fecundity has been relatively high since 1992 because of increasing proportions of older and larger fish. Total egg production was greatest from 1981 through 1983 because of peak population abundance and moderate mean annual fecundity estimates. Estimates were also relatively high in 1988, 1989, and 1991 as older and larger females dominated moderately-sized reproductive stocks. Total egg production decreased to relatively low values in recent years because of very low winter flounder abundance, with 2001 again marking the lowest estimate because of a small abundance estimate. Estimated long-term mean fecundity of Niantic River female winter flounder, determined by dividing the total number of eggs by

TABLE 8. Relative and absolute standardized catch of female winter flounder spawners and corresponding egg production in the Niantic River from 1977 through 2001.

| Survey year | Relative index of spawning females ${ }^{\text { }}$ | \% mature females (of all fish) ${ }^{\text {b }}$ | ```% mature females (of all females)c``` | Average fecundity ${ }^{d}$ |  | Relative index of total egg production ${ }^{\text {e }}$ | Total female stock size ${ }^{\text {r }}$ | Total egg production $\left(\times 10^{5}\right)^{\prime}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1977 | 1,069 | 36 | 61 | 450,470 |  | 481.4 | 31,526 | 14.202 |
| 1978 | 1,644 | 51 | 76 | 503,750 |  | 828.3 | 48,511 | 24.437 |
| 1979 | 1,197 | 37 | 64 | 472,405 |  | 565.7 | 35,328 | 16.689 |
| 1980 | 1,004 | 34 | 51 | 466,919 |  | 468.9 | 29,626 | 13.833 |
| 1981 | 2,411 | 44 | 70 | 519,461 |  | 1,252.5 | 71,140 | 36.955 |
| 1982 | 2,551 | 49 | 77 | 580,082 |  | 1,479.8 | 75,265 | 43.660 |
| 1983 | 1,705 | 47 | 78 | 577,885 |  | 985.5 | 50,313 | 29.075 |
| 1984 | 822 | 40 | 77 | 577,774 |  | 474.9 | 24,253 | 14.013 |
| 1985 | 846 | 43 | 75 | 608,374 |  | 514.9 | 24,969 | 15.190 |
| 1986 | 604 | 42 | 86 | 666,132 |  | 402.5 | -17,826 | 11.875 |
| 1987 | 780 | 39 | 85 | 622,995 |  | 485.9 | 23,010 | 14.335 |
| 1988 | 1,172 | 53 | 89 | 676,469 |  | 792.8 | 34,579 | 23.391 |
| 1989 | 891 | 52 | 90 | 726,652 |  | 647.1 | 26,274 | 19.092 |
| 1990 | 498 | 41 | 73 | 631,150 |  | 314.4 | 14,695 | 9.275 |
| 1991 | 963 | 47 | 85 | 600,832 |  | 578.7 | 28,418 | 17.075 |
| 1992 | 485 | 53 | 94 | 730,945 |  | 354.2 | 14,296 | 10.450 |
| 1993 | 247 | 54 | 93 | -815,957 |  | 201.6 | 7,288 | 5.947 |
| 1994 | 458 | 56 | 87 | 648,741 |  | 297.3 | 13,523 | 8.773 |
| 1995 | 192 | 63 | 87 | 772,187 |  | 148.1 | 5,659 | 4.370 |
| 1996 | 88 | 52 | 82 | 843,825 | : | 74.3 | 2,598 | 2.192 |
| 1997 | 166 | 60 | 83 | 796,337 |  | 131.8 | 4,884 | 3.889 |
| 1998 | 118 | 55 | 79 | 755,317 |  | 89.1 | 3,479 | 2.627 |
| 1999 | 107 | 45 | 65 | 698,084 |  | 75.0 | 3,168 | 2.211 |
| 2000 | 167 | 55 | 80 | 737,566 |  | 123.0 | 4,919 | 3.628 |
| 2001 | 68 | 49 | 69 | 823,565 |  | 56.1 | 2,009 | 1.655 |

- Based on proportion of the relative annual standardized catches of winter flounder that were mature females.
- As a propórtion of all winter flounder 20 cm or larger.
c As a proportion of all female winter flounder 20 cm or larger.
- Total egg production divided by the number of spawning females.
- A relative index for year-to-year comparisons and not an absolute estimate of production.
- Calculated on the assumption that the relative annual standardized catches were approximately $3.3 \%$ of absolute values.
the total number of mature females since 1977, was approximately 583,775 , a modest increase from the value of 582,000 given in DNC (2001a). Each total annual egg production estimate was divided by the present value of mean fecundity to produce an estimate of scaled parental stock size used in a StockRecruitment Relationship (SRR), which is discussed below in a following section.


## Comparisons Among Regional Abundance Indices

Comparisons were made among the abundance of Niantic River winter flounder and regional abundance indices, including the spring CPUE of winter flounder from the DEP LIS-wide stratified random trawl survey (Shake 2001; D. Molnar, CT DEP, Old Lyme, CT, pers. comm.), the mean CPUE from a spring and fall stratified random trawl survey.
conducted by the Rhode Island Department of Environmental Management, Division of Fish and Wildlife (RIFW) in Narragansett Bay, RI and nearby waters (Lynch 2000), an annual CPUE developed from weekly trawl sampling conducted by the University of Rhode Island (URI) near Fox Island in upper Narragansett Bay (sampling program described in Jeffries and Johnson 1974; Jeffries and Tereceiro 1985; Jeffries et al. 1989; data provided by D. Taylor, University of Rhode Island Graduate School of Oceanography, Narragansett, RI, pers. comm.), and a stratified random trawl survey conducted during spring by the Massachusetts Division of Marine Fisheries (MDMF) in state waters extending from the tip of Cape Cod to the Rhode Island border (Howe et al. 1999; T. Currier, MDMF, Pocasset, MA, pers. comm.).

Most regional abundance indices of winter flounder were : strongly correlated (Spearman's rank-order correlation) with one another (Table 9). These

TABLE 9. Matrix of Spearman's rank-order correlations among various regional winter flounder abundance indices.

| Abundance Index | RIFW - Narragansett Bay | URI- Fox Island | CTEEP-LIS | MDMF - Southern Mass. |
| :---: | :---: | :---: | :---: | :---: |
| Niantic River adult winter flounder survey ( 215 cm ) $\Delta$-mean CPUE | $\begin{gathered} 0.7629^{5} \\ 0.0001 * * \\ 22 \end{gathered}$ | $\begin{gathered} 0.7908 \\ 0.0001 * * \\ 26 \end{gathered}$ | $\begin{gathered} 0.5501 \\ 0.0180 * \\ 18 \end{gathered}$ | $\begin{gathered} 0.6693 \\ 0.0003 * * \\ 24 \end{gathered}$ |
| RIFW Narragansett Bay trawl survey spring and fall mean CPUE |  | $\begin{gathered} 0.8701 \\ 0.0001 * * \\ 22 \end{gathered}$ | $\begin{gathered} 0.3848 \\ 0.1272 \text { NS } \\ 17 \end{gathered}$ | $\begin{gathered} 0.6201 \\ 0.0021 \text { ** } \\ 22 \end{gathered}$ |
| URI Fox Island annual geometric mean CPUE |  |  | $\begin{gathered} 0.4613 \\ 0.0540 \mathrm{NS} \\ 18 \end{gathered}$ | $\begin{gathered} 0.7998 \\ 0.0001 \text { ** } \\ 24 \end{gathered}$ |
| CT DEP LIS spring geometric mean CPUE |  |  |  | $\begin{gathered} 0.1768 \\ 0.4827 \mathrm{NS} \\ 18 \end{gathered}$ |

* The three statistics shown for each correlation matrix element are: correlation coefficient $(r)$,
probability of a larger $r$ (NS - not significant $[p>0.05], *$ - significant at $p \leq 0.05$, ** - significant at $p \leq 0.01$ ), and number of annual observations (sample size).
correlations were positive, indicating no inverse trends in abundance among areas. The $\Delta$-mean CPUE of adult winter flounder spawning in the Niantic River was significantly correlated with two Rhode Island winter flounder CPUE indices, the RIFW Narragansett Bay trawl index (1979-2000; $\mathrm{r}=$ 0.7629 ) and the URI Fox Island series (1976-2001; 0.7908 ), and with the MDMF Southern Massachusetts trawl index having catches expressed as biomass per tow (1978-2001; 0.6693). A weaker, but still significant correlation was found with the DEP LISwide CPUE (1984-2001; 0.5501). This finding may be due to a shorter time-series for comparison and because the DEP trawl survey had limited sampling in eastern LIS (Gottschall et al. 2000).
Comparisons of normalized indices of abundance for these regional data sets showed good correspondence among them (Figs. 11 and 12). The two Rhode Island CPUE indices each had a peak in 1979, 2-3 years before the peak in abundance was observed in the Niantic River, perhaps because the Rhode Island catches had a larger component of younger fish than the mostly older evident by the mid-1980s in each of these winter flounder abundance indices. Gibson (1998) noted that winter flounder abundance in Narragansett Bay began declining in the late 1970s and remained low for more than a decade, not rebounding to levels seen previously. Although the Rhode Island catches increased from the early 1990s to a small peak in 1995, perhaps reflecting catch of fish : from the relatively strong 1992 year-class, CPUE subsequently decreased in years following to the present. In


Fig. 11. Comparison of regional winter flounder abundances from 1976 through 2001. Included are the $\Delta$-mean CPUE of lish larger than 15 cm taken in the Niantic River during the spawning season, the mean CPUE from Rhode Island Fish and Wildlife (RIFW) spring and fall trawl surveys in Narragansett Bay, and the annual geometric mean CPUE of winter llounder taken at the University of Rhode Island (URI) Fox Island trawl station in Narragansett Bay. Each CPUE series was normalized by dividing all values by the corresponding largest estimate and multiplying by 100 .
comparison, a larger increase in abundance was seen for the DEP LIS CPUE through 1996, but this index has also decreased subsequently to the lowest values of this series in 1999 and 2000, although a modest increase occurred in 2001. In contrast, the Niantic River population size remains small, with another downturn in seen in 2001 after 4 years of increasing abundance since 1996 (Fig. 12).
Coherence among abundance indices of winter flounder within a relatively small geographical


Fig. 12. Comparison of regional winter flounder abundances from 1984 through 2001. Included are the: $\triangle$-mean CPUE of fish larger than 15 cm taken in the Niantic River during the spawning season, the annual geometric mean CPUE of winter flounder taken at the University of Rhode Island (URI) Fox Island trawl station in Narragansett Bay, and the annual geomettic mean CPUE of winter flounder taken during April-June by Connecticut Department of Environmental Protection Marine Fisheries in Long Island Sound (DEP LIS). Each CPUE series was normalized by dividing all values by the corresponding largest estimate and multiplying by 100 .
region should be expected. Fox et al. (2000) reported synchrony in the recruitment of European plaice throughout the entire waters of the United Kingdom, although abundances in adjacent areas tended to be most similar. According to NEFSC (1999), which is a recent stock assessment of winter flounder on the northeastern coast of the United States, each winter flounder abundance survey discussed above sampled a distinct geographical area and was likely providing measures of different components of the same aggregated stock. Further, older winter flounder dominated catches in the Niantic River surveys, but younger fish were prevalent in the DEP LIS sampling. Based on catch-at-age data and lengthfrequency information presented in NEFSC (1999) and Lynch (2000), respectively, younger winter flounder made up a large component of the catches in these surveys as well. Thus, direct comparisons of these indices may need some qualification, even though temporal trends appeared to have some similarities. Also noted in NEFSC (1999) was that regional abundance surveys tended to present a continuum of optimistic to pessimistic trends in abundance and illustrated variable tracking of yearclass strength when they were compared among one
another. This was attributed to possible differences in the availability of winter flounder to sampling; this was also noted for the Niantic River studies. A further indication was given by the Atlantic States Marine Fisheries Commission (ASMFC) Winter Flounder Management Board, who recently reported that stock biomass of the Southern New England and Mid-Atlantic winter flounder remained at below target levels, even though $F$ was believed to be at 0.3-0.4 (ASMFC 2001). Some state surveys indicated that stock biomass has actually decreased. With improvement in stock status not observed in some member state waters, some ASMFC members also questioned whether winter flounder distribution patterns have changed over time.
Finally, in a recent study on long-term recruitment trends of New England groundfish, Brodziak et al. (2001) concluded that nine of eleven stocks examined had a significantly declining trend in spawning stock biomass. Among these stocks, the 1963-96 time-series of Southern New England winter flounder showed significant declining trends for both spawning stock biomass and recruits-per-spawner. This result adds to the mounting empirical evidence supporting the notion that the decline in winter flounder abundance during the past 20 years or so has taken place over a wide regional scale and well beyond any possible influence of MPS.

## Larval Winter Flounder

## Field Sampling, Laboratory Processing, and Data Analyses

Winter flounder larvae have been collected in Niantic Bay at station NB since 1979 and in the Niantic River at stations A, B, and C since 1983 (Fig. 13). To collect larvae, a $60-\mathrm{cm}$ bongo plankton sampler weighted with a $22.7-\mathrm{kg}$ oceanographic depressor and fitted with $3.3-\mathrm{m}$ long nets with mesh size of $202 \mu \mathrm{~m}$ was deployed during February and March. Net size was changed to $333 \mu \mathrm{~m}$ in April to increase catchability of larger larvae. Volume of water filtered was determined from a single General Oceanics flowmeter mounted in the center of each bongo opening. The sampler was towed at approximately 2 knots using a stepwise oblique tow pattern, with equal sampling time at surface, middepth, and near bottom. The length of tow line necessary to sample the mid-water and bottom strata was determined by water depth and tow-line angle measured with an inclinometer. Nets were towed for 6 minutes and each net typically filtered about 120


Fig. 13. Location of stations (denoted by letters) sampled for larval winter flounder during 2001.
$\mathrm{m}^{3}$ of water. Only one of the duplicate samples was retained for laboratory processing, typically the bongo that sampled the greatest volume. When present, jellyfish medusae at the three river stations were removed from the samples using a $1-\mathrm{cm}$ mesh sieve and their mass estimated volumetrically to the nearest 100 mL .
The larval winter flounder sampling schedule for Niantic River and Bay was based on knowledge gained during previous years and was designed to increase data collection efficiency (NUSCO 1987).
Larval sampling at the Niantic River stations usually begins in early to mid-February. In 2001, sampling in the river began on February. 13, but only at stations C and B due to heavy ice cover in the upper river. On February 20, all stations were clear of ice and sampling continued at each station until May 7, after which stations A and B were dropped due to lack of larvae in samples. Stations C and NB were last sampled on June.15.. Daytime tows were made within 1 hour of low slack tide through the end of March. During the remainder of the season until the disappearance of larvae at each station, tows were
made at night during the second half of a flood tide. From 1983 through 1990, collections were taken on 2 days each week. Starting in 1991, sampling was reduced to I day a week (NUSCO 1991a). Through 1992, station NB was sampled during day and night every two weeks during February and at least once a week from March through the end of the larval winter flounder season. Beginning in 1993, station NB was sampled weekly only during the day from the start of the larval season through March and at night from April through the remainder of the larval season. Water temperature and salinity measurements (surface, midwater, and bottom) were recorded at each sampling station using a YSI Model 30 Salinity/Temperature/Conductivity meter. All the ichthyoplankton samples, including entrainment collections described below, were preserved with $10 \%$ formalin.
Winter flounder larvae entrained through the MPS cooling-water system have been sampled at the plant discharges (station EN, Fig. 13) since 1976. These collections provide information not only for the estimation of entrainment impact, but also on the
abundance and temporal distribution of winter flounder larvae in Niantic Bay. In the earliest years of study, collections usually alternated between the discharges of Units 1 and 2, depending upon plant operation and the resulting water flow from the condenser cooling-water pumps. In more recent years, collections were also taken at the Unit 3 discharge. Collections were eliminated at the Unit 1 discharge in 1995 when this unit was retired from operation. In 2001, entrainment sampling was conducted at only Unit 2 because the Unit 3 sampling gantry was inoperable. Larvae were collected with a $1.0 \times 3.6-\mathrm{m}$ conical plankton net of $333-\mu \mathrm{m}$ mesh deployed from a gantry system into the thalweg of the discharge water from each unit. Four General Oceanic (GO) Model 2030 flowmeters were positioned in the net mouth to account for horizontal and vertical flow variation. Sample volume was determined by the average of four volume estimates from the flowmeters. Under normal operations, the net is usually deployed for 3 to 4 minutes (filtering about $200 \mathrm{~m}^{3}$ ), with variation in sampling time dependent upon the number of circulating water pumps in operation and tidal stage. Samples of longer duration ( $6-10 \mathrm{~min}$ ) were taken when fewer than the normal complement of circulating water pumps were in operation. Sampling frequencies and volume filtered have varied since 1976 (NUSCO 1987, 1994a). In recent years, sampling was conducted during both day and night once per week in February and 3 days and nights per week during March through June. Entrainment samples were processed similarly to field-collected larvae, which is described below.
During laboratory processing and in accordance with laboratory sorting protocols, preserved ichthyoplankton samples were split at minimum to one-half volume and to lesser volumes if larvae were relatively abundant. Sample material was viewed through a dissecting microscope and winter flounder larvae were removed and counted. Up to 50 randomly selected larvae were measured to the nearest 0.1 mm in standard length (snout tip to notochord tip). The developmental stage of each measured larva was recorded using the following identification criteria:

Stage'1. Yolk-sac present or eyes not pigmented (yolk-sac larvae);
Stage 2. Eyes pigmented, no yolk-sac present, no fin ray development, and no flexion of the notochord;
Stage 3. Fin rays present and flexion of the notochord begun, but left eye not migrated to the midline;

Stage 4. Left eye reached the midline, but juvenile characteristics not present;
Stage 5. Transformation to juvenile stage complete and intense pigmentation present near the caudal fin base.
Larval data analyses were based on standardized densities (number $500 \mathrm{~m}^{-3}$ of water sampled). A geometric mean of weekly densities was used in analyses because the data generally followed a lognormal distribution (McConnaughey and Conquest 1993) and weekly sampling frequencies varied among some stations and years. Because older larvae apparently remained near the bottom during the day and were not as susceptible to entrainment or the bongo sampler, data from daylight samples collected after March at stations EN and NB were excluded from abundance calculations, except for estimating entrainment at MPS.
The distribution of larval abundance data over time is usually skewed because densities increase rapidly to a maximum and then decline slowly. A cumulative density over time from this type of distribution results in a sigmoid-shaped curve, where the time of peak abundance coincides with the inflection point. The Gompertz function (Draper 'and Smith 1981; Gendron 1989) was used to describe this cumulative abundance distribution because the inflection point of this function is not constrained to the mid-point of the sigmoid curve. The form of the Gompertz function used was:

$$
\begin{equation*}
C_{t}=A \times \exp (-\exp [-k \times\{t-p\}]) \tag{2}
\end{equation*}
$$

where $C_{t}=$ cumulative density at time $t$
$t=$ time in days from February 15
$A=$ total or asymptotic cumulative density
$p=$ inflection point scaled in days since February 15
$k=$ shape parameter
The time of peak abundance was estimated by the parameter $p$. The origin of the time scale was set to February 15, which is the approximate date when winter flounder larvae first appear in the Niantic River. Least-squares estimates, standard errors, and asymptotic $95 \%$ confidence intervals (CI) for these parameters were obtained by fitting the above equation to the cumulative abundance data using nonlinear regression methods (SAS Institute Inc. 1990d). Cumulative data were obtained as the running sums of the weekly geometric means of the abundance data. The $A$ parameter of the cumulative curve was used as an index to compare annual abundances.

A "density" function was derived from the first derivative of the Gompertz function (Eq. 2) with respect to time. This density function, which directly describes the larval abundance over time (abundance curve), has the form:

$$
\begin{align*}
\mathrm{d}_{t}= & A^{\prime} \times k \times \exp (-\exp [-k \times\{t-p\}]-k \times \\
& {[t-p]) } \tag{3}
\end{align*}
$$

where $d_{t}=$ density at time $t$ and all the other parameters are as described for Equation 2, except for $A^{\prime}$, which was re-scaled by a factor of 7 (i.e., $A^{\circ}=7 A$ ) because the cumulative densities were based on weekly geometric means and, thus, accounted for a 7 -day period.
Larval mortality rates were estimated from data collected at the three Niantic River stations. Data from 1983 were excluded as smaller larvae were undersampled then because of net extrusion (NUSCO 1987). The abundance of 3 mm and smaller larvae was used to calculate an index of newly-hatched larvae because 3 mm was the approximate length at hatching. The decline in the frequency of larvae in progressively larger size-classes (in 1 -mm groups) was attributed to both natural mortality and as a result of tidal flushing from the river. Hess et al. (1975) estimated the loss of larvae from the entire river as $4 \%$ per tidal cycle and also determined that the loss from the lower portion of the river was about $28 \%$ per tidal cycle. Thus, the weekly abundance estimates of larvae 3 mm and smaller at station C in the lower portion of the river were re-scaled by a factor of 1.93 to compensate for the $28 \%$ decline per tidal cycle (two cycles per day). The abundance of larvae in the $7-\mathrm{mm}$ size-class was used to calculate an index of larval abundance just prior to metamorphosis. Because previous studies (NUSCO 1987, 1989) showed a net import of larger larvae into the Niantic River, the weekly abundance of larvae in the $7-\mathrm{mm}$ size-class at station C was not adjusted for tidal flushing. To calculate each annual rate of mortality, sums were made of weekly mean abundance indices (three stations combined) of newly-hatched larvae (after adjusting for tidal flushing) and larvae in the $7-\mathrm{mm}$ size-class. Survival rates from hatching through larval development were estimated as the ratio of the abundance index of the larger larvae ( $7-\mathrm{mm}$ size-class) to that of the smaller larvae ( $3-\mathrm{mm}$ and smaller size-classes).
The presence of density-dependent mortality was investigated by relating annual larval abundance in the $7-\mathrm{mm}$ and larger size-classes from station EN to the annual egg production estimate for the Niantic

River using the Ricker nonlinear relationship (Ricker 1975):

$$
\begin{equation*}
\mathrm{L}=a \times \mathrm{E} \times \exp (-b \times \mathrm{E}) \tag{4}
\end{equation*}
$$

where $\mathrm{L}=$ annual larval abundance of larvae $7-\mathrm{mm}$ and larger at EN as estimated by $A$ (see Equation 2)
$\mathrm{E}=$ annual estimate of egg production in the Niantic River
$a=$ slope at the origin of the larval recruitment curve
$b=$ instantaneous mortality dependent upon annual egg abundance
Density-dependent mortality may be assumed when the slope $(b)$ is significantly different from zero.
Regression analyses were used to examine possible relationships between variables and, at times, to make predictions. Ordinary least-squares linear regression was used when the independent variable was assumed to be measured without error (e.g., water temperature). The test of a relationship was based on the slope being significantly ( $\mathrm{p} \leq 0.05$ ) different from zero. Functional regression methods developed by Ricker $(1973,1984)$ were used in the cases where the independent variable was measured with error (e.g., abundance indices). For functional regressions, the probability that the correlation coefficient r was significantly ( $\mathrm{p} \leq 0.05$ ) different from zero was the criterion used to decide whether a valid relationship existed prior to determining the slope and its $95 \% \mathrm{CI}$.

## Abundance and Distribution

The $A$ parameter of the Gompertz function (Eq. 2) was used as an index for temporal (year to year) and spatial (Niantic River and Bay) abundances of winter flounder larvae. Based on the $A$ parameter estimates, larval abundance in aggregate during 2001 in Niantic Bay (stations EN and NB combined) was the highest observed during the 19 -year period (Table 10). In contrast, abundance in the Niantic River (stations A, $B$, and $C$ combined) was the fifth lowest of the series, although about double the abundance found in 2000, which was the lowest ever observed (Table 10). In general, annual abundances in the bay have varied less than in the river. In most years the abundance of larval winter flounder in the river was two to six times greater than the bay. However, in 1988 and 1989, abundances in the river were more than ten times greater than in the bay, but in 1983 and 1999, river abundance was less than two times as much as in the bay. The reason that larval abundance is

TABLE 10. Index of annual abundances and $95 \%$ confidence intervals based on the $A$ parameter from the Gompertz function for larval winter flounder taken in Niantic River and Bay from 1983 through 2001.


- Stage I larvac undersampled in 1983 because a $202-\mu \mathrm{m}$ mesh net was not used during the early portion of the larval season.
generally greater in the river than the bay is probably related to preferred winter flounder spawning areas, as discussed below in the comparison of spatial distribution of developmental stages. Biological and physical processes occurring during larval development are variable as no significant relationship was found between the indices of annual abundances (excluding 1983) for the two areas (Spearman's rank-order correlation; $\mathrm{r}=0.187 ; \mathrm{p}=0.458 ; \mathrm{n}=18$ ). This year (2001) was the second time since 1984 that abundance was higher in the bay than in the river, the first being 2000. Low abundance in the river during 1983 was attributed, in part, to undersampling because smaller larvae were likely extruded through the $0.333-\mu \mathrm{m}$ mesh net then used throughout the year (NUSCO 1987). However, this was rectified in 1984 when a net with smaller mesh ( $202 \mu \mathrm{~m}$ ) was deployed during the early portion of the larval season. The reason for the higher abundance in the bay in 2000 and 2001 is unknown, but may be related to relatively high rates of flushing early in the season and higher survival for larvae found in LIS relative to those remaining in the Niantic River.
Annual spatial abundances of the first four larval developmental stages were based on cumulative weekly geometric means at each of the stations sampled (Figs. 14 and 15). The abundance distribution of Stage 5 individuals (i.e., newly transformed to juveniles) was not examined because so few of
them were collected by ichthyoplankton sampling gear. Cumulative density data (the annual sum of the weekly geometric means) were used to compare abundances as a surrogate for the $A$ parameter from the Gompertz function (Eq. 2) because in some instances this function could not be fitted. This usually occurred when a developmental stage was rarely collected at a station (e.g., Stage 1 at stations EN and NB or Stage 4 at station A). Cumulative weekly geometric means and the corresponding $A$ parameters were found to be highly correlated (Spearman's rank-order correlation; $r=0.999 ; p<$ $0.001 ; \mathrm{n}=120$ ) in a previous comparison (NUSCO 1989), indicating that the cumulative means could be used as another index of larval abundance.
Stage 1 abundances by station during 2001 were considerably below average in both the Niantic River (Fig. 14) and Niantic Bay (Fig. 15). A comparison of annual Stage 1 abundance among years at the three river stations showed a similar relative ranking at the three stations, with 1988, 1989, and 1997 ranked the highest and 1983, 1986, 1993, and 2000 the lowest. Except for a greater abundance at station $A$ in some years, annual abundances at the three river stations have been similar in most years; significant ( $p \leq$ 0.05 ) positive correlations were found among Stage I annual abundances for all river stations (Table 11). This indicated a somewhat homogeneous distribution of Stage 1 larvae throughout the river. Because winter flounder eggs are demersal and adhesive and


Fig. 14. Index of cumulative density by developmental stage of larval winter flounder at Niantic River stations A, B, and C from 1983 through 2001. (Note that the vertical scales differ among the graphs).
the duration of Stage 1 is short (about 10 days), the homogenous distribution suggested either that spawning was not restricted to a specific area of the river or that the river is well-mixed.

On average, abundance of Stage I larvae was about 50 times less at the two Niantic Bay stations (Fig. 15) than in the river (Fig. 14), indicating that little, if any, spawning occurred in the bay. Abundance at station

NB was consistently greater than at EN, possibly because NB was located closer to the river mouth, the likely source of Stage 1 larvae, and because undersampling occurred at EN as a result of extrusion of Stage 1 larvae through the $333-\mu$ m mesh net used at the MPS discharges. At NB, ranks of annual abundance indices were similar to those of the river stations suggesting that most Stage 1 larvae


Fig. 15. Index of cumulative density by developmental stage of larval winter flounder at the Niantic Bay stations EN and NB from 1983 through 2001. (Note that the vertical scales differ among the graphs).
collected in the bay probably originated from the Niantic River. However, even though abundance at station A was correlated with EN and NB, there were less clear associations between $B$ and $C$ and the two Niantic Bay sites (Table 11).
Annual abundance of newly hatched winter flounder larvae should be related to adult spawner egg production and the fraction of eggs that hatch.

However, from 1995 through 1999 and in 2001, abundances of Stage 1 larvae at the three river stations were greater than expected from the low annual egg production estimates determined for the Niantic River spawning stock (Fig. 16). The index of Stage' 1 larval abundance was the $A$ parameter from the Gompertz function (Eq. 2) for the Niantic River (stations A, B, and C combined) and the previously

TABLE 11. Matrix of Spearman's rank-order correlations among stations for the indices of annual cumulative abundance of each developmental stage of larval winter flounder from 1983 through 2001.


[^13]discussed egg production estimates were given on Table 9. To examine egg survival, a relative annual survival rate index was determined by computing the ratio of the Stage 1 abundance index (in thousands) to annual egg production estimates (in billions), followed by grouping years as either 1984-87, 1990-

94 and 2000 or 1988-89, 1995-99, and 2001. The mean ratio (relative survival rate index) for the first yearly group was $0.28(\mathrm{SE}=0.06)$ and for the latter group was $1.27(\mathrm{SE}=0.18)$, indicating that average egg survival appeared to be greater by more than five times during most recent years in comparison to the


Fig. 16. Comparison between the index of annual Stage 1 abundance in the Niantic River ( $A$ parameter of the Gompertz function) and estimated annual egg production from 1984 through 2001. Ricker stock-recruitment functions were fitted to data from 1984 to 1994 and 2000 (dashed line): Stage $1=$ 0.0217 ( $\mathrm{Egg}^{1.424} \times e^{-122 \mathrm{Kd1} \times \mathrm{Ess})} ; \mathrm{p}=0.0031$, and 1995.99 and 2001 (solid line): Stage $1=2.0353\left(\mathrm{Egg}^{3 \times x: 31} \times \mathrm{e}^{0.0374} \cdot \mathrm{E}_{\mathrm{Es}}\right) ; \mathrm{p}=$ 0.0010 .
other period. In addition, there was a significant difference between ratios for the two yearly groups ( $p=0.001$; Wilcoxon two-sample test; Sokal and Rohlf 1969). These relationships were explored further using nonlinear regression techniques fitting the data to the Ricker stock-recruitment equation (Ricker 1975). Statistically significant relationships were found for both groups of years, demonstrating an apparently higher egg survival in 1988-89, 199599, and 2001 (Fig. 16). The stock-recruitment relationship in these years approached a theoretical asymptote at high levels of egg production, whereas the model for the years 1984-87, 1990-94, and 2000 became depressed at high egg production levels, indicating stronger effects of compensatory processes that increased egg mortality and limited abundance of Stage 1 larvae. The biological significance of these relationships has yet to be determined and warrants closer examination of mechanisms regulating egg survival, such as predation and egg quality.
Stage 2 abundances at the three river stations in 2001 were among the lowest observed (Fig. 14). On average (1983-2001), the abundance of Stage 2 larvae in the river was about $60 \%$ of Stage 1 , but ratios for each year have been variable. For example, in 1999 Stage 2 abundance was less than $20 \%$ of that of Stage 1, in 2000 abundances were similar, and in 2001 Stage 2 abundance was about $30 \%$ of that for Stage 1. In general, annual ranks of Stage 2 abundance at the three river stations have been similar to those of Stage 1 , which implied a similar annual rate of larval loss (mortality and flushing) during larval development from Stage 1 to 2 . In
most years, including 2001, Stage 2 larvae occurred predominantly in the river, but were more prevalent in the bay compared to Stage. 1. At the two bay stations, Stage 2 abundances in 2001 were the eighth highest (NB) or second highest (EN) recorded, indicating a possible high rate of flushing this year. Annual abundances were always greater at station NB than at EN (Fig. 15). Significant ( $\mathrm{p} \leqslant 0.05$ ) positive correlations of abundance were found among all river stations and between station NB and EN, but not among the river and bay stations (Table 11).
Abundance of Stage 3 larvae in 2001 was the lowest observed at station A, second lowest at station B, fourth lowest at station C, and in the upper third of all years at EN and NB (Figs. 14 and 15). Of the four larval developmental stages, Stage 3 annual abundances were the most similar among all the stations, although numbers tended to be lower at A than elsewhere. The pattern of annual abundances among stations showed positive and significant correlations ( $p \leq 0.05$ ) among all stations, except between stations A and NB and B and NB (Table 11).

Annual abundances of Stage 4 larvae were generally lowest in the upper river (station A) and greatest in the lower river (C), with intermediate abundances found at station B (Fig. 14). This suggests a gradual flushing to the lower river as development proceeds and perhaps an emigration of older, pre-settling larvae transported or emigrating from the bay into the river late in the season. During 2001, no Stage 4 larvae were taken at A, abundance was less than the median value at $B$, and was second highest at C. Annual abundance of Stage 4 larvae collected at station EN was the third highest of the 19-year period and was the highest ever observed at NB (Fig. 15). Including data from 2001, abundance of Stage 4 larvae at station $A$ was only significantly correlated with that at $B$ and $B$ was only similar to $C$ (Table 11). In Niantic Bay, abundances at NB and EN were significantly correlated.
Dates of peak abundance, estimated from the inflection point $p$ of the Gompertz function (Eq. 2), were used to compare the times of occurrence in the river (station $A, B$, and $C$ combined) and bay (EN and NB combined) for each developmental stage (Table 12). Dates of peak abundance of: Stage 1 larvae were not estimated for bay stations because during several years this larval stage was rarely collected outside of the Niantic River., In 2001, the dates of peak abundance for both Stage 1 and 3 larvae in the Niantic River were near the middle of the dates of the 19-year period, but was among the earliest for Stage 2 and the latest for Stage 4.

TABLE 12. Estimated annual dates of peak abundance of larval winter flounder by developmental stage in the Niantic River and Bay and the number of days corresponding to the $95 \%$ confidence interval from 1983 through 2001.

| Year | Stage 1 | Stage 2 | Stage 3 | Stage 4 |
| :---: | :---: | :---: | :---: | :---: |
| Niantic River |  |  |  |  |
| 1983 | March 5 (3) | March 15 (2) | April 18 (1) | May 2 (4) |
| 1984 | March 7 (5) | March 9 (5) | April 24 (5) | May 19 (10) |
| 1985 | March 11 (1) | March 16 (2) | April 25 (3) | Mayl6 (7) |
| 1986 | February 26 (1) | March 11 (5) | April 20 (3) | May 12 (10) |
| 1987 | March 10 (2) | March 17 (3) | April 20 (2) | May 9 (4) |
| 1988 | February 29 (1) | March 9 (1) | April 7 (4) | May 1 (5) |
| 1989 | March 8 (6) | March 12 (5) | April 14 (3) | May 11 (9) |
| 1990 | February 17 (3) | February 18 (5) | April 21 (2) | May 9 (14) |
| 1991 | February 27 (3) | March 14 (11) | April 13 (5) | April 29 (3) |
| 1992 | March 16 (4) | April 6 (3) | April 16 (2) | May 2 (2) |
| 1993 | March 9 (2) | March 14 (8) | April 11 (7) | $\rightarrow$ |
| 1994 | March 22 (4) | March 31 (5) | April 24 (1) | May 10 (3) |
| 1995 | March 2 (3) | March 7 (5) | April 20 (2) | May 4 (2) |
| 1996 | March 14 (6) | March 21 (4) | April 19 (8) | May 17 (5) |
| 1997 | March 4 (3) | March 10 (2) | April 2 (4) | April 18 (4) |
| 1998 | February 27 (2) | March 5 (2) | April 19 (22) | May 5 (5) |
| 1999 | February 28 (3) | March 28 (13) | April 10 (2) | April 24 (6) |
| 2000 | March 4 (1) | March 9 (4) | April 17 (5) | April 25 (5) |
| 2001 | March 7 (3) | March 4 (2) | April 9 (2) | May 24 (21) |
| Niantic Bay |  |  |  |  |
| 1983 | - | April 7 (2) | April 23 (1) | May 10 (4) |
| 1984 | - | April 8 (2) | May 4 (3) | May 25 (8) |
| 1985 | - | April 1 (4) | April 29 (6) | May 18 (3) |
| 1986 | - | April 5 (3) | April 28 (3) | May 11 (2) |
| 1987 | - | April 6 (6) | April 28 (2) | May 16 (4) |
| 1988 | - | March 24 (3) | April 22 (2) | May 9 (5) |
| 1989 | - | April 13 (1) | April 23 (2) | May 17 (3) |
| 1990 | - | April 3 (8) | April 23 (2) | May 7 (5) |
| 1991 | - | March 28 (5) | April II (3) | April 29 (4) |
| 1992 | - | April 15 (4) | April 30 (2) | May 7 (4) |
| 1993 | - | April 3 (44) | May 6 (8) | May 23 (11) |
| 1994 | - | April 14 (2) | May 2 (2) | May 20 (3) |
| 1995 | - | April 4 (5) | April 21 (4) | April 28 (3) |
| 1996 |  | April 7 (4) | April 30 (2) | May 24 (9) |
| 1997 | - | March 23 (7) | April 13 (1) | April 30 (9) |
| 1998 | - | March 25 (6) | April 23 (3) | May 14 (6) |
| 1999 | - | April 9 (3) | April 23 (6) | May 9 (16) |
| 2000 | - | April 1 (3) | May 3 (11) | May 22 (4) |
| 2001 | - | April 9 (3) | ${ }^{-1}$ | May 23 (3) |

- Due to low abundance during the 1993 and 2001 sampling, the Gompertz function could not be fitted to the data.

Similarly, the date for Stage 2 larvae in Niantic Bay was one of the earlier ones, and for Stage 4 was among the latest. No estimate was made for Stage 3 larvae in Niantic Bay because the Gompertz function could not be fitted to the data.
Water temperature in Niantic Bay was warmer than average in February $\left(4.03^{\circ} \mathrm{C}\right.$; Table 1) and peak winter flounder spawning probably occurred prior to mid-February (see Adult Winter Flounder, above). Rogers (1976) and Buckley (1982) reported egg incubation times for winter flounder and Keller and Klein-MacPhee (2000) recently noted a 20 -day
incubation period for eggs at $4.1^{\circ} \mathrm{C}$ and 30 days at $1.6^{\circ}$ C. Buckley et al. (1990) observed that egg developmental time was inversely related to water temperature during oocyte maturation and egg incubation. The combination of warm water temperatures in February followed by somewhat warmer than average March temperatures (Table 1) probably contributed to the average date of peak abundance for Stage 1 larvae in 2001. A comparison between February water temperatures and the annual dates of Stage 1 peak abundance in the river during 1983 through 2001 showed a significant negative
relationship (Spearman's rank-order correlation; $r=$ $-0.638 ; \mathrm{p}=0.003 ; \mathrm{n}=19$ ). Dates of peak abundance for later developmental stages were likely related to temperaturé-moderated'developmental rates and perhaps variable rates of flushing, which could accelerate the buildup of larval abundance sequentially at stations in the Niantic River and Bay. The relationship between water temperature and larval developmental rate is discussed in more detail in the following section.

## Development and Growth

The length-frequency distribution of each larval stage has remained relatively consistent since developmental stage determination began in 1983. Stage-specific length-frequency distributions by $0.5-\mathrm{mm}$ size-classes in 2001 showed some separation in predominant size-classes by developmental stage (Fig. 17). Nearly all (89\%) Stage 1 larvae were in the 2.5 to $3.5-\mathrm{mm}$ size-classes, $99 \%$ of Stage 2 were 3.0 to $5.0 \mathrm{~mm}, 97 \%$ of Stage 3 were 5.0 to 7.0 mm ; and $98 \%$ of Stage 4 were 6.0 to 8.0 mm . These predominant size-classes for each developmental stage were generally similar to those reported for previous years (DNC 2001a). Consistent stagespecific length-frequency distributions from year to year indicated that developmental stage and length of larval winter flounder were closely related. These data agreed with laboratory studies on larval winter flounder, which showed that there were positive correlations between growth and developmental rates (Chambers and Leggett 1987; Chambers et al. 1988; Bertram et al. 1996).

Length-frequency distributions of larvae (all stages combined) collected in the Niantic River (stations A, $B$, and $C$ combined) differed from those obtained for Niantic Bay (EN and NB combined) for all years of sampling, including 2001 (Fig. 18). Smaller ( $\leq 4.0$ mm ) size-classes usually predominated in the river due to the high abundance of Stage 1 larvae. The cumulative total of $69 \%$ of larvae $\leq 4.0 \mathrm{~mm}$ in 2001 was slightly less than the historical (1983-2000 combined) length-frequency distribution, which showed about $75 \%$ of larvae in these smaller sizeclasses. In addition, length-frequency distributions in the river for some previous years showed a slight increase or near-stable frequencies of larvae in the 7.0 to $8.0-\mathrm{mm}$ size-classes; this was also apparent during 2001. These findings suggested that some older larvae were imported into the , river. Importation of larger size-classes was also apparent in the length-frequency distribution at a station





Fig. 17. Combined length-frequency distribution of larval winter flounder by developmental stage at all stations sampled in the Niantic River and Bay during 2001.



Fig. 18. Length-frequency distribution of larval winter flounder in the Niantic River and Bay during 2001.
located at the river mouth, which was sampled in 1991-93 during maximum flood tidal currents (NUSCO 1994a).
The size-class distribution for Niantic Bay in 2001 was also consistent with most previous findings. Generally, a skewed unimodal distribution in lengthfrequency was found with the larger size-classes predominating, except for 1996 (NUSCO 1997b). Typically, the greatest size frequencies found in the bay were the $5.0-$ to $7.0-\mathrm{mm}$ size-class ( $1983-2000$ combined, $53 \%$ ), as they were in 2001 ( $70 \%$; Fig. 18). The sharp decrease of larvae 8.0 mm and larger occurs as larvae metamorphose to the demersal Stage 5 juvenile.
Length-frequency data from entrainment collections taken from 1976 through 2001 (station EN) were used to estimate larval winter flounder growth rates in Niantic Bay. Weekly mean lengths during a season formed a sigmoid-shaped curve (NUSCO 1988b). The linear portion of the sigmoid curve usually occurred in the middle of the larval season and growth rates were estimated by fitting a linear model to individual larval length measurements during this time period. This linear model
adequately described growth and all slopes (growth rate as $\mathrm{mm} \cdot \mathrm{day}^{-1}$ ) were significantly ( $\mathrm{p} \leq 0.001$ ) different from zero (Table 13). In addition, most intercepts of the linear regression were about 3 , the approximate size of winter flounder larvae at hatching. Annual growth rates for station EN were variable and ranged from 0.046 to $0.062 \mathrm{~mm} \cdot \mathrm{day}^{-1}$, with growth in 2001 among the lowest rates observed in the 19 -year monitoring program. To validate this estimation technique, growth rates were estimated from length data collected at station NB from 1979 through 1989 (NUSCO 1990); annual growth rates were highly correlated ( $r=0.89 ; p \leq 0.001 ; n=11$ ) with those from station EN:
In laboratory and mesocosm studies, water temperature affected the growth rate of winter flounder larvae (Laurence 1975; NUSCO 1988b; Casas 1998; Keller and Klein-MacPhee 2000). To examine the effect of temperature on estimated annual growth rates, mean water temperatures in Niantic Bay, determined from data collected by continuous recorders in the MPS intakes, were calculated for a 40 -day period starting at the beginning of the week when the first larval length measurements were used to estimate the annual growth rate (Table 13). The mean temperatures used were not necessarily indicative of the actual annual seasonal water temperatures because annual starting points varied from February 28 (1993) to April 3 (1977). A positive exponential relationship was found between growth rate and water temperature, with 1999 having the largest growth rate as well as the highest water temperature (Fig. 19). Growth rate in 2001 appeared to be lower than indicated by the $5.9^{\circ} \mathrm{C}$ mean water temperature. A similar exponential relationship between temperature and growth was reported for larval plaice by Hovenkamp and Witte (1991). Keller and Klein-MacPhee (2000) also reported larval winter flounder growth to be significantly greater in warmer than cooler experimental mesocosms. If temperature affects growth rate, then the length of a larva at a specific time during the season should be related to water temperatures to which it has been exposed. Therefore, the mean length of larvae collected at station EN during the period of April 1-15 for each year was compared to the mean March water temperatures (Fig. 20). There was a positive relationship, with larger mean lengths associated with warmer March temperatures (e.g., 1999). The mean length found for 2001 was slightly higher than the predicted value due to the warmer than average March mean water temperature (Table 1).

TABLE 13. Annual larval winter flounder growth rates in Niantic Bay as estimated from a linear regression fitted to length data collected at station EN from 1976 through 2001. The $95 \%$ confidence intervals and mean water temperatures during the first 40 days of the time period are also given.


- Time period of the larval lengths used to estimate growth rate.
- Mean during a 40-day period starting at the beginning of the week that the first weekly mean length was used in estimating growth rate.

As concluded previously from comparisons 'of annual length-frequency distribution and developmental stages, growth and larval development are closely related. If water temperature affects growth rates, then it should also affect larval developmental time. The timing of peak larval abundance should


Fig. 19. Exponential relationship between mean water temperature $\mathrm{T}\left({ }^{\circ} \mathrm{C}\right)$ and the estimated growth rate G ( mm per day) of winter flounder larvae at station EN from 1976 through 2001 ( $\left.G=0.0278 \times e^{018 \pi}\right)$.
therefore be related to the rates of recruitment and loss (including mortality and juvenile metamorphosis), which, in tün, would be affected by larval development. Annual dates of peak abundance of larval winter flounder collected at EN were negatively correlated with the mean water temperature in


Fig. 20. Relationship between annual mean March water temperature ( ${ }^{\circ} \mathrm{C}$ ) and the mean length of winter flounder larvae during April 1-15 at station EN for 1976 through 2001.

March and April; earlier dates of peak abundance were associated with warmer mean water temperatures (Fig. 21). This agreed with the results of Laurence (1975), who found that winter flounder larvae metamorphosed 31 days earlier at $8^{\circ} \mathrm{C}$ than at $5^{\circ} \mathrm{C}$. Annual dates of peak abundance varied by 41 days during the 25 -year period, possibly because of a $3.6^{\circ} \mathrm{C}$ difference in the March-April water temperature between one of the earliest (April 13, 1991) and the latest (May 23, 1978) dates of peak abundance. Despite the wide range in annual growth rates, a consistent relationship was found between lengthfrequency distribution and stage of development (Fig. 17). This was also consistent with laboratory observations of larval winter flounder, as Chambers et al. (1988) found that age at metamorphosis was more variable than length and larval age and length were independent of one another. Again, 2001 was somewhat inconsistent with respect to information on. growth as the relatively late date of peak abundance (May 21) was associated with a warmer than average water temperature of $5.6^{\circ}$ for the period.

Growth rates were also estimated for Niantic River larvae using length and temperature data from station $C$ with the methods given above. Station $C$ was selected for this analysis because all developmental stages were collected there in abundance (Fig. 14). A


Fig. 21. Relationship between March-April mean water temperature $\left({ }^{\circ} \mathrm{C}\right)$ and the annual date of peak abundance (estimated from the Gompertz function) of winter flounder larvae at station EN from 1976 through 2001.
linear model again provided a good fit and slopes (growth rates as $\mathrm{mm} \cdot \mathrm{day}^{-1}$ ) were significantly ( $\mathrm{p} \leq$ 0.001 ) different from zero. Estimated growth rates for larvae in the river were generally greater than for larvae from the bay. The 2001 growth rate of 0.106 $\mathrm{mm} \cdot \mathrm{day}^{-1}$ for the river was about average for the 19 year period, and it was associated with an average water temperature (Table 14). Growth of larvae in the river in 2001 was slightly higher than growth rates of laboratory reared fish, which ranged from

TABLE 14. Annual larval winter flounder growth rates in the Niantic River as estimated from a linear regression fit to length data collected at station C from 1983 through 2001. The $95 \%$ confidence intervals for the growth rate, a mean determined for water temperatures recorded at station $C$ during the first 6 weeks of the time period, and the annual abundance indices of Stage 2 larvae in the river are also given.

| Year | Time period included ${ }^{\text {a }}$ | Growth rate ( $\mathrm{mm} \cdot \mathrm{day}^{-1}$ ) | $95 \%$ contidence interval | Mean water temperature $\left({ }^{\circ} \mathrm{C}\right)^{\text {b }}$ | Stage 2 abundance index ${ }^{\text {c }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1983 | March 20-May 1 | 0.100 | 0.096-0.104 | 6.1 | 749 |
| 1984 | March 25 - May 6 | 0.100 | 0.094-0.105 | 6.4 | 1,501 |
| 1985 | March 31-May 26 | 0.084 | 0.080-0.088 | 7.7 | 4,676 |
| 1986 | March 23 - May 4 | 0.109 | 0.103-0.115 | 8.0 | 176 |
| 1987 | March 22 - May 10 | 0.099 | 0.095-0.103 | 7.2 | 829 |
| 1988 | March 20-May 21 | 0.099 | 0.094-0.104 | 6.8 | 4,469 |
| 1989 | March 26 - May 21 | 0.087 | 0.082-0.092 | 7.4 | 3,976 |
| 1990 | March 25 - May 13 | 0.106 | 0.099-0.113 | 7.5 | 365 |
| 1991 | March 10 - April 28 | 0.123 | 0.114-0.132 | 6.9 | 252 |
| 1992 | March 15-May 17 | 0.088 | 0.083-0.093 | 5.7 | 1,367 |
| 1993 | March 7 - May 16 | 0.070 | 0.065-0.075 | 4.1 | 133 |
| 1994 | March 20 - May 29 | 0.072 | 0.068-0.076 | 4.7 | 1,248 |
| 1995 | March 12 - April 30 | 0.130 | 0.121-0.140 | 6.8 | 2,023 |
| 1996 | March 24 - May 19 | 0.096 | 0.092-0.099 | 6.6 | 4,424 |
| 1997 | March 2 - April 20 | 0.136 | 0.127-0.144 | 5.2 | 8,449 |
| 1998 | March 22 - May 3 | 0.109 | 0.098-0.119 | 7.4 | 6,553 |
| 1999 | March 28 - April 25 | 0.150 | 0.129-0.171 | 8.9 | 288 |
| 2000 | March 26. April 30 | 0.103 | 0.082-0.124 | 8.1 | 402 |
| 2001 | March 25 - April 29 | 0.106 | 0.099-0.113 | 6.1 | 704 |

[^14]0.104 and $0.100 \mathrm{~mm} \cdot \mathrm{day}^{-1}$ at mean water temperatures of 6.9 and $7.5^{\circ} \mathrm{C}$, respectively (NUSCO 1988b).
Slight declines in growth rate caused by less than optimal food, unfavorable temperatures, disease, or pollution leads to longer developmental times, during which high rates of mortality can have a profound effect on recruitment (Houde 1987). Food availability and water temperature appeared to be the two most important factors controlling larval winter flounder growth (Buckley 1982). A laboratory study with larval winter flounder held at $8^{\circ} \mathrm{C}$ showed a decrease in growth as prey densities decreased, suggesting density-dependent growth due to food availability (Laurence 1977). Although Laurence (1975) demonstrated that the metabolic demands of larval winter flounder increased at higher temperatures, growth rate also increased if sufficient food resources were available, and other laboratory, studies showed that larval winter flounder growth rates depend upon prey availability (Laurence 1977; Buckley 1980). The effect of both larval density (i.e., density-dependency) and water temperature on larval growth rates were examined for data collected from the Niantic River by comparing annual growth rates to the abundance index for Stage 2 larvae, the developmental stage during which larvae begin to feed. This method assumed that prey availability was similar from year to year. The annual index of Stage 2 larval abundance was the $A$ parameter (Eq. 2) for all. three river stations combined (Table 14). A density-dependent relationship was previously reported (NUSCO 1990, 1991b, 1992a, 1993), but since 1993 , this relationship no longer has been significant. Because a strong relationship existed between growth and water temperature in the bay, both Stage 2 abundance and water temperature were used as independent variables in " a multiple' regression model to examine growth rates. The multiple regression, which included data from 1983 through 1996, was significant ( $p=0.035 ; \mathrm{R}^{2}=0.456$; $\mathrm{df}=11$, with a positive slope for temperature and negative slope for Stage 2 abundance. Although the Stage 2 slope was not always significant, this variable was needed for a significant multiple regression model. However, with the addition of 1997 and 1998 data, the relationship was no longer evident ( $p=$ $0.280 ; \mathrm{R}^{2}=0.178 ; \mathrm{df}=13$ ), probably because even though growth rates in those years were among the highest, Stage 2 abundances were also among the highest. With a further addition of data from 19992001, a positive relationship was found between growth rate and water temperature ( $p=0.008 ; r^{2}=$ $0.460 ; \mathrm{df}=16$ ), but there was no apparent density-
dependent relationship, even though Stage 2 abundances in these 3 years were among the lowest of the 19 -year period. The varying results from year to year suggested that factors affecting growth may be more complex than just water temperature and abundance, and may be related to factors such as prey abundance and diversity, for which no - information was collected. In summary, growth and development of winter flounder larvae in Niantic Bay appeared to be related to water temperature, with a similar, although less certain, relationship possible for larvae found within the Niantic River.

## Mortality

The 2001 estimate of instantaneous natural mortality rate (M) of 0.92 from hatching (Stage 1) to 7 mm (Stage 4) was the lowest observed in the timeseries. This instantaneous rate translated into a total mortality rate (A) of $59.8 \%$, which was also the lowest observed. From 1984 through 1999, total mortality from hatching to just prior to larval metamorphosis ranged from 82.4 to $97.9 \%$ with corresponding estimates of M of 1.73 to 3.88 (Table 15).

Based on length-frequency distributions in the river during previous years, most larval winter flounder mortality occurred between the 3.0 and $4.5-\mathrm{mm}$ sizeclasses, which included yolk-sac (Stage 1) and firstfeeding Stage 2 larvae. A decrease in abundance of these size-classes was also apparent in the river during 2001 (Fig. 18, upper). Constructing a catch curve from catch data of these smaller larvae resulted in an estimated M of 3.2. Based on a growth rate of $0.054 \mathrm{~mm} \cdot$ day $^{-1}$ determined in 2001 for Niantic River larvae, it would take approximately 27 days for a larva to grow from 3 to 4.5 mm (Table 13). Daily M was therefore 0.119 , in contrast to a higher long-term (1983-98) average of $0.174 \cdot$ day $^{-1}$ (NUSCO 2000). This apparently lower than average mortality during the transition to first-feeding larvae in 2001 was illustrated by a smaller than average decline in abundance from Stage 1 to Stage 2 larvae at the three river stations and because Stage 2 larvae were particularly abundant in Niantic Bay this year (Figs. 14 and 15). The decrease in abundance of older larvae was examined using larvae collected in Niantic Bay (Fig. 18, lower). Here the catch curve from 6 to 7.5 mm (Stages 3 to 4) resulted in an M of 0.76 . At a growth rate of $0.106 \mathrm{~mm} \cdot \mathrm{day}^{-4}$ in Niantic Bay (Table 14), a larva would take approximately 14 days to grow from 6 to 7.5 mm . Thus, an M of $0.054 \cdot \mathrm{day}^{-1}$ was indicated.

TABLE 15. Estimated larval winter flounder total instantaneous natural mortality rate ( $M$ ) from hatching to the $7-\mathrm{mm}$ size-class from 1984 through 2001.

|  | Abundance index <br> Newly <br> hatched | \%-mm <br> size-class | mortality <br> rate <br> $(A))^{2}$ | Instantaneous <br> mortality rate <br> $(M)^{2}$ |
| :---: | ---: | ---: | :---: | :---: |
| 1984 | 6,500 | 654 | 89.9 | 2.30 |
| 1985 | 13,773 | 452 | 96.7 | 3.42 |
| 1986 | 2,483 | 438 | 82.4 | 1.73 |
| 1987 | 6,480 | 474 | 92.7 | 2.62 |
| 1988 | 24,561 | 678 | 97.2 | 3.59 |
| 1989 | 19,192 | 394 | 97.9 | 3.88 |
| 1990 | 7,915 | 653 | 91.7 | 2.49 |
| 1991 | 3,992 | 560 | 86.5 | 2.00 |
| 1992 | 8,020 | 609 | 92.4 | 2.58 |
| 1993 | 1,874 | 88 | 95.3 | 3.06 |
| 1994 | 7,270 | 761 | 89.5 | 2.26 |
| 1995 | 13,088 | 1,536 | 88.3 | 2.14 |
| 1996 | 11,151 | 576 | 94.8 | 2.96 |
| 1997 | 14,894 | 1,645 | 89.0 | 2.20 |
| 1998 | 9,306 | 921 | 90.1 | 2.31 |
| 1999 | 4,658 | 791 | 83.0 | 1.77 |
| 2000 | 1,725 | 1,758 | .6 | $-b$ |
| 2001 | 4,252 | 1,708 | 59.8 | 0.92 |
|  |  |  |  |  |
|  |  |  |  | mean = 2.48 |

2 Survival $=1-A=e^{M}$.

- Could not be calculated as more larger than smaller larvae were found.

In other studies, Pearcy (1962) reported a greater mortality for young winter flounder larvae ( $20.7 \% \cdot$ day $^{-1}$ ) compared to older individuals ( $9.1 \%$ day ${ }^{-1}$ ) in the Mystic River, CT. Keller and Klein-MacPhee (2000) reported mortality rates of 5.6 to $8.9 \% \cdot \mathrm{day}^{-1}$ in their warm mesocosm tanks and 3.6 to $4.0 \% \cdot$ day $^{-1}$ in cold tanks, although predators in these experimental tanks were relatively low in comparison to natural systems. Larval mortality rates in the Niantic River appear to be similar.
Gelatinous zooplankton, including jellyfish and ctenophores, are potential predators of winter flounder larvae (see Formation of Winter Flounder Year-class Strength, below). During 7 of the 18 years (1983, 1984, 1986, 1989, 1990, 1994, and 1998) sampled, weekly mean larval abundance of larvae at station A was negatively correlated ( $p \leq$ 0.05 ; Spearman's rank-order correlation coefficients $r$ ranged between -0.736 and -0.927 ) with the weekly mean jellyfish volume during the period when both medusae and larvae were collected. Weekly jellyfish volumes in 2001 were generally lower than the long-term average (1983-2000) from early March through early May (Fig. 22) and were significantly negatively correlated with weekly mean larval abundance $(r=-0.829, p=0.0212)$. This year,


Fig. 22. Comparison of weekly mean volume and $95 \%$ confidence interval of gelatinous zooplankton (primarily Cyanea sp.) collected at station A in the Niantic River for the period of 1983 through 2000 with the weekly mean volume taken in 2001.
larval abundance at station A was relatively low throughout the sampling period compared to the long-term average, but larvae were consistently captured at the station through mid-April and abundance peaked in late March. Jellyfish abundance at Station A was considerably lower than the long-term average, reaching a peak volume of only $1.5 \mathrm{~L} \cdot 500 \mathrm{~m}^{-3}$ in mid-April (Fig. 22). In many years, including 2001, these gelatinous zooplankton generally appeared late in the larval season at station $A$ at a time when winter flounder larval abundance was declining in this area. Thus, the temporal distribution of jellyfish and winter flounder larvae may have overlapped very little. Although for some years there was an apparent negative relationship between larval abundance and gelatinous predators, it may just be coincidental, and due to a natural decline in larval abundance' by gradual flushing from the upper portion of the river. Thus, the role of jellyfish in cropping winter flounder larvae remains uncertain.
The possibility of density-dependent mortality of winter flounder larvae was examined using a nonlinear function (Eq. 4) provided by Ricker (1975) that requires estimates of annual spawning stock size and larval recruitment. The annual egg production estimate in the Niantic River (Table 9) was used as a measure of spawning stock size. The $A$ parameter from the Gompertz function fitted to the abundance of $7-\mathrm{mm}$ and larger larvae collected from 1976 through 2000 at station EN was selected as a measure of larval recruitment, even though many of these larvae may be progeny of other spawning stocks (see Mass-Balance Calculations, below). Abundance of larvae in the $7-\mathrm{mm}$ and larger size-classes was used as an index of early recruitment because these larvae would soon metamorphose into juveniles. This index of late larval abundance was plotted against egg
production estimates and the nonlinear slope ( $b$ in Eq. 4) was estimated by nonlinear regression (Fig. 23). A significant slope, $b=0.237$ ( $\mathrm{p}<0.0001 ; \mathrm{n}=$ 25 ), suggested the existence of compensatory mortality. As a measure of precision, the standard error for $b$ was only $16 \%$ of the estimated value. The 4 years (1995, 1997-99) with the highest abundance of $7-\mathrm{mm}$ and larger larvae were among the years with the lowest egg production estimates (Table 9) and apparently highest egg survival (Fig. 16). Because evidence exists that many of the winter flounder larvae collected at station EN did not originate from the Niantic River, the compensatory relationship suggested that annual egg production estimates for the Niantic River were consistent with regional trends in winter flounder egg production and similar early life history processes. Also, the apparent higher egg survival in the 1995-99 period may have resulted in better recruitment to larger larval size-classes. Although the production of yolk-sac larvae this year in the Niantic River appeared to be more in line with estimated egg production, the high abundance of older larvae appears to support the concept of compensation at low levels of abundance.
Another approach to detect the possible presence of density-dependent larval mortality of the Niantic River stock was a comparison of natural mortality rate (M) of larvae from the river (Table 15) to estimated egg production from 1984 through 1999 and 2001 (Table 9). In addition, the effect of water temperature on larval mortality was examined. A multiple regression model used egg production estimates with various combinations of seasonal water temperature recorded at MPS (Tables 1 and 2). The combinations included monthly means for


Fig. 23. Relationship between the annual winter flounder egg production in the Niantic. River and the annual abundance of 7 mm and larger larvae at station EN from 1977 through 2001. The curve represents predicted values from fitting Eq. 4.

March though May, combinations of March and April and April and May, and seasonal means for winter (January-March) and spring (April-June). The best model ( $\mathrm{R}^{2}=0.481 ; \mathrm{p}=0.010 ; \mathrm{df}=14$ ) indicated that larval mortality decreased as egg production ( $p=$ 0.033 ) decreased and April temperature ( $p=0.022$ ) increased (Fig. 24). This suggests that densitydependent larval mortality occurred in the Niantic River that was further :moderated by April water temperatures. The effect of temperature on mortality may be due to its positive relationship to rates of larval growth and development. For example, the low mortality rate in 1998 was likely related to one of lowest annual egg production estimates and the warmest April water temperature (Tables 1,9, and 14). Conversely, a relatively high mortality rate in 1988 was associated with high egg production and cool temperatures.


Fig. 24. Relationship between the instantaneous natural mortality rate (M) of winter flounder larvae and annual egg production in the Niantic River and April mean water temperature ( ${ }^{\circ} \mathrm{C}$ ) at the MPS intakes from 1984 through 1999 and 2001: $\mathrm{M}=5.297+0.078$ (egg production in billions) 0.482 (April water temperature).

> Juvenile Winter Flounder
> Age-0 Juveniles during Summer

- Field Sampling and Data Analyses. The Niantic River serves as a nursery ground for much of the first , year of life of post-larval age-0 winter flounder. Abundance of this life stage has been monitored from late May through the end of September since 1983 (LR) or late 1984 (WA) at two shallow (about $1-2 \mathrm{~m}$
in depth), nearshore stations in the Niantic River (Fig. 2). Through 1992, collections were made weekly, but in 1993 sampling frequency was reduced to biweekly, resulting in ten dates sampled during the 19-week season. Sampling occurred during daylight from about 2 hours before to 1 hour after high tide. A $1-\mathrm{m}$ beam trawl was used that had two tickler chains and nets of $0.8-, 1.6-, 3.2-$, and $6.4-\mathrm{mm}$ mesh. In 1983, triplicate tows were made at LR using nets of increasing larger mesh as the season progressed. Beginning in 1984, two frames with nets of successively larger mesh were used during each sampling trip; nets were deployed in a random order. A change to the next larger mesh of the four-net sequence was made when fish had grown large enough to become retained by it, as use of larger meshes reduced the amount of detritus, algae, and sevenspine bay shrimp (Crangon septemspinosa) collected. At each station, four replicate tows were made, two each with the two nets in use. Rarely, only three tows were taken at a station because of bad weather or net damage. Tow distance was estimated by releasing a measured line attached to a lead weight as the net was hauled at approximately 25 $\mathrm{m} \cdot \mathrm{min}^{-1}$. The length of each tow was increased from 40 to 100 m in $20-$ or $40-\mathrm{m}$ increments at a station as fish abundance decreased over time. However, in years when densities of young fish were high, maximum tow length may have been only 60 or 80 m . Surface water temperature was taken with a handheld mercury stem thermometer.
The catch of young-of-the-year winter flounder in each of the three or four replicated $1-\mathrm{m}$ beam trawl tows was standardized to a $100-\mathrm{m}$ tow distance before computing mean CPUE for each day and station; density was therefore expressed as the number per $100 \mathrm{~m}^{2}$ of bottom. A median CPUE abundance index was determined for each halfseason, with late May through July denoting the early season and August-September the late season. A $95 \% \mathrm{CI}$ was calculated for each median CPUE using a distribution-free method based on order statistics (Snedecor and Cochran 1967).
Nearly all of the age-0 winter flounder collected were measured fresh in either the field or laboratory to the nearest 0.5 mm in total length (TL). During the first few weeks of study, standard length (SL) was also measured to the nearest 0.5 mm because many of the smaller specimens had damaged caudal fin rays and total length could not be ascertained. A relationship between the two lengths determined by a functional regression was used to convert SL to TL whenever necessary:

$$
\begin{equation*}
T L=-0.2+1.212(\mathrm{SL}) \tag{5}
\end{equation*}
$$

Growth of age-0 winter flounder at each station was examined by following weekly mean lengths throughout the sampling season. Mean lengths of young taken at the Niantic River stations LR and WA from late July through September were compared using an analysis of variance. Significant differences among means were determined with Tukey's studentized range test (SAS Institute Inc. 1990c). The relationship between growth and abundance of young and water temperature was examined using multiple linear regression (SAS Institute Inc. 1990d) and functional regression methods described previously in the Larval Winter Flounder section.
To calculate M , all young were assumed to comprise a single cohort with a common birthdate. A catch curve was constructed such that the natural logarithm of density was plotted against age (time in weeks) and the slope of the descending portion of the curve provided an estimate of the weekly rate for $M$ (Ricker 1975). Once this rate was determined, the monthly mortality rate ( $\mathrm{M}_{\mathrm{mo}}$ ) was calculated as $\mathrm{M} \times$ ( 30.4 / 7) and monthly survival ( $\mathrm{S}_{\mathrm{mo}}$ ) as $e^{-\mathrm{Mmo}}$.
Abundance. Although beam trawls are much more efficient than small otter trawls for collecting juvenile flatfish (Kuipers et al. 1992), the densities of young winter flounder reported herein should be regarded as minimum estimates because of collection inefficiencies. For example, Berghahn (1986) caught more young European plaice at night using a beam trawl in comparison to samples taken during the day, while Rogers and Lockwood (1989) showed that replacing tickler chains normally used with even heavier, spiked chains nearly doubled catches. Efficiency of the NUSCO $1-\mathrm{m}$ beam trawl was discussed previously in NUSCO (1987, 1990). Although some Enteromorpha clathrata algal mats, which hampered sampling efficiency in the Niantic River during much of 1993 and occasionally in other years, was present at LR, no large mats developed during 2001 that affected the catches. Presence of extensive algal mats may also influence distribution of young winter flounder. Recently metamorphosed ( $11-15 \mathrm{~mm}$ ) European plaice avoided filamentous algal mats in comparison to bare sediment in laboratory experiments conducted by Wennhage and Pihl (1994). This preference could potentially affect year-class strength as well as measurements of abundance if large areas of nursery grounds are vegetated and young are crowded into uncovered patches of bottom.

During 2001, densities of Stage 4 larvae found in the Niantic River at station C and at station NB in

Niantic Bay ranked second and first, respectively, of the abundance time-series (Figs. 14 and 15). Even so, at the start of sampling in late May, young settling in the Niantic River were relatively scarce at WA ( $7.5 \cdot 100 \mathrm{~m}^{2}$ ), although they were moderately abundant (ca. $50 \cdot 100 \mathrm{~m}^{2}$ ) at LR (Fig. 25). However, densities increased rapidly at both stations, with numbers at WA appearing to lag behind those at LR by one biweekly sampling interval. Densities reached a peak of about $250 \cdot 100 \mathrm{~m}^{2}$ during late June through mid-July at LR and in mid-July at WA. Abundance at both stations steadily declined during the remainder of the summer, with a particularly sharp linear decrease seen at WA until mid-August, after which the rate of loss moderated. Increasing abundance during the early part of the season may have reflected a lengthier than usual period for recruitment of settling older larvae entering the river from Niantic Bay as this year the dates for peak abundance of Stage 4 larvae in both Niantic River and Bay were among the latest of the time-series (May 23-24; Table 12). Also, there may have been a time lag in the shoreward movement of previously settled juveniles from deeper areas of the river. The rate of loss of age-0 winter flounder is discussed below in a subsection discussing their mortality.


Fig. 25. Weekly mean CPUE ( $\pm 2$ standard errors) of age- 0 winter flounder taken at Niantic River stations LR and WA by $1-\mathrm{m}$ beam trawl during 2001.

As measured by early (May-July) and late (AugustSeptember) seasonal median CPUE, abundance of age-0 winter flounder during the first half of the season in 2001 was higher ( $180.100 \mathrm{~m}^{2}$ ) at LR than at WA (141.3; Table 16). In late summer, fish were more than twice as abundant at LR (50.0) than at WA (21.7) in contrast to 2000, when the opposite occurred. Following 4 years of relatively similar early summer abundance, the early summer median CPUE (computed using catches at both Niantic River stations combined) in 2001 was among the highest observed since 1984 (Fig. /26). Although this summer started with a potential for high production of young, abundance decreases occurring in mid to late summer resulted in the formation of a relatively strong, but not exceptional, 2001 year-class. Abundance of age- 0 winter flounder at the end of summer has steadily increased since 1996. Other relatively abundant year-classes included those of 1988, 1994, 1995, and 2000.
Beginning in 1988, DEP sampled at eight seine stations along the Connecticut coast during September (Howell 2001). As was found for the


Fig. 26. Early and late summer seasonal $1-m$ beam trawl median CPUE and $95 \%$ CI at Niantic River stations LR and WA combined from 1984 through 2001. (Note that the vertical scales differ between the graphs).

TABLE 16. Seasonal 1 -m beam trawl median CPUE (number• $100 \mathrm{~m}^{-2}$ ) of age- $0^{4}$ winter ilounder at two stations in the lower Niantic River (LR and WA) from 1983 through 2001.

| Survey year | Station | Season ${ }^{\text {b }}$ | Tows used for CPUE | $\begin{gathered} \text { Median } \\ \text { CPUE } \\ \text { estimate } \end{gathered}$ | 95\% confidence interval for median CPUE | Coctficient of skewness ${ }^{\text {c }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1983 | LR | Early | 30 | 32.7 | 20.0-50.7 | 2.29 |
|  | LR | Late | 27 | 10.0 | 8.0-13.3 | 0.49 |
| 1984 | LR | Early | 40 | 18.8 | 16.7-25.0 | 0.63 |
|  | LR | Late | 36 | 6.3 | 3.8-7.5 | 0.58 |
|  | WA | Late | 32 | 11.3 | 8.0-17.5 | 0.94 |
| 1985 | LR | Early | 40 | 13.3 | 10.0-16.3 | 0.91 |
|  | LR | Late | 32 | 7.0 | 6.0-8.0 | 0.97 |
|  | WA | Early | 40 | 15.0 | 10.0-20.0 | 0.81 |
|  | WA | Late | 32 | 9.0 | 8.0-10.0 | 0.70 |
| 1986 | LR | Early | 39 | 33.8 | 23.3-40.0 | 0.33 |
|  | LR | Late | 36 | 13.8 | 12.5-17.5 | 0.80 |
|  | WA | Early | 40 | 21.7 | 12.5-26.7 | 1.49 |
|  | WA | Late | 36 | 18.1 | 15.0-20.0 | 2.03 |
| 1987 | LR | Early | 40 | 59.2 | 53.3-73.3 | -0.12 |
|  | LR | Late | 36 | 17.9 | 12.5-26.7 | 0.70 |
|  | WA | Early | 40 | 28.3 | 21.7-38.3 | 0.27 |
|  | WA | Late | 36 | 10.6 | 6.0-13.8 | 0.83 |
| 1988 | LR | Early | 40 | 61.3 | 52.5-72.5 | 0.37 |
|  | LR | Late | 36 | 60.0 | 50.0-70.0 | 1.17 |
|  | WA | Early | 40 | 40.0 | 32.5-51.7 | 0.13 |
|  | WA | Late | 36 | 38.3 | 33.3-51.7 | 0.22 |
| 1989 | LR | Easly | 40 | 17.5 | 11.7-21.7 | 0.09 |
|  | LR | Late | 36 | 8.8 | 7.0-11.3 | 0.84 |
|  | WA | Early | 40 | 10.0 | 8.3-13.8 | 1.16 |
|  | WA | Late | 34 | 5.5 | 4.0-10.0 | 0.66 |
| 1990 | LR | Early | 40 | 156.3 | 137.5-187.5 | 1.05 |
|  | LR | Late | 36 | 20.0 | 15.0-52.5 | 1.10 |
|  | WA | Early | 40 | 68.8 | 50.0-95.0 | 0.62 |
|  | WA | Late | 36 | 13.5 | 10.0-19.0 | 1.20 |
| 1991 | LR | Early | 44 | 77.5 | 51.7-90.0 | 0.96 |
|  | LR | Late | 36 | 21.7 | 18.3-28.3 | 0.75 |
|  | WA | Early | 44 | 37.9 | 30.0-43.3 | 1.34 |
|  | WA | Late | 36 | 25.8 | 21.3-31.7 | 1.27 |
| 1992 | LR | Early | 40 | 90.0 | 57.5-122.5 | 1.16 |
|  | LR | Late | 36 | 28.1 | 23.8-33.3 | 0.51 |
|  | WA | Early | 40 | 74.6 | 56.7-82.5 | 1.35 |
|  | WA | Late | 36 | 30.0 | 27.5-32.5 | 0.23 |
| 1993 | LR | Early | 20 | 10.6 | 7.0-15.0 | 0.68 |
|  | LR | Late | 20 | 5.0 | 3.0-7.0 | 1.15 |
|  | WA | Early | 20 | 5.0 | 3.8-7.5 | 2.57 |
|  | WA | Late | 20 | 5.5 | 4.0-10.0 | 0.77 |
| 1994 | LR | Early | 20 | 128.8 | 125.5-172.5 | 0.38 |
|  | LR | Late | 20 | 62.9 | 38.3-75.0 | 0.26 |
|  | WA | Early | 20 | 126.3 | 92.5-192.5 | 0.31 |
|  | WA | Late | 20 | 49.2 | 35.0-55.0 | -0.79 |

TABLE 16. (continued).

|  | Survey year | Station | Season ${ }^{\text {b }}$ | Tows used for CPUE | Median CPUE estimate | $95 \%$ conlidence interval for median CPUE | - |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1995 |  | LR | Early | 20 | 87.5 | 52.5-140.0 |  | 1.82 |
|  |  | LR | Late | 20 | 15.8 | 12.0-26.7 |  | 1.96 |
|  |  | WA | Early | 20 | 116.3 | 85.0-137.5 |  | 2.31 |
|  |  | WA | Late | 20 | 55.0 | 28.3-70.0 |  | 0.59 |
|  | 1996 | LR | Early | 20 | 8.8 | 5.0-15.0 |  | 0.27 |
|  |  | LR | Late | 20 | 3.0 | 3.0-6.0 |  | 1.42 |
|  |  | WA | Early | 20 | 21.7 | 11.7-27.5 |  | 1.30 |
|  |  | WA | Late | 20 | 6.2 | 3.0-7.0 |  | 1.76 |
|  | 1997 | LR | Early | 20 | 19.2 | 16.7-25.0 |  | 1.03 |
|  |  | LR | Late | 20 | 7.0 | 2.0-10.0 |  | 0.29 |
|  |  | WA | Early | 20 | 53.8 | 35.0-80.0 |  | 0.59 |
|  |  | WA | Late | 20 | 9.5 | 6.3-20.0 |  | 0.92 |
| - | 1998 | LR | Early | 20 | 46.3 | 32.5-60.0 |  | 0.18 |
|  |  | LR | Late | 20 | 15.0 | 6.7-20.0 |  | 1.08 |
|  |  | WA | Early | 20 | 45.0 | 25.0-65.0 |  | 1.42 |
|  |  | WA | Late | 20 | 13.0 | 8.0-25.0 |  | 1.00 |
|  | 1999 | LR | Early | 20 | 47.5 | 25.0'-80.0 |  | 1.30 |
|  |  | LR | Late | 20 | 29.2 | 16.0-35.0 |  | 0.57 |
|  |  | WA | Early | 20 | 37.5 | 27.5-57.5 |  | 2.16 |
|  |  | WA | Late | 20 | . 17.1 | 11.7-18.8 |  | 1.13 |
|  | 2000 | LR | Early | 20 | 38.8 | 10.0-67.5 |  | 0.52 |
|  |  | LR | Late | 20 | 27.5 | 22.5-35.0 |  | 1.34 |
|  |  | WA | Early | 20 | 40.0 | 22.5-58.3 |  | 0.35 |
|  |  | WA | Late | 20 | 47.9 | 35.0-52.5. |  | 0.75 |
|  | 2001 | LR | Early | 20 | 180.0 | 107.5-247.5 |  | 0.27 ' |
|  |  | LR | Late | 20 | 50.0 | 36.7-82.5 |  | 1.73 |
|  |  | WA | Early | 20 | 141.3 | 45.0-227.5 |  | 0.10 |
|  |  | WA | Late | 20 | 21.7 | 10.0-32.5 |  | 1.71 |

- For age-0 fish, the year-class is the same as the survey year.
${ }^{\text {b }}$ Early season corresponds to late May through July and late to August through September.
c Zero for symmetrically distributed data.

Niantic River, catches throughout LIS showed relatively strong year-classes of winter flounder produced during 1988, 1992, 1994, and 1995. However, the 2001 index was the lowest since this sampling began in 1988, with time-series lows found at stations in Greenwich, Bridgeport, Milford, New Haven, and Old Lyme (D. Molnar, CT DEP; Old Lyme, CT, pers. comm.). One of the DEP stations was at a beach on the Waterford shoreline of the Niantic River adjacent to the beam trawl station WA. The geometric mean seine catch of 9.1 in 2001 was a conisiderable decrease from the mean of 41.6 at this location in 2000, which was the highest of the DEP seine series. The proximity of the two sampling sites allowed for comparison of the annual late summer median 1-m beam trawl CPUE index at WA and the

DEP seine CPUE using functional regression (Fig. 27). Despite relatively wide confidence intervals associated with each of these two abundance estimates; a significant positive relationship ( $r=$ $0.576 ; \mathrm{p}=0.031 ; \mathrm{df}=12$ ) was found between these two indices. The point for 2001 showed a somewhat larger deviation from the computed functional regression line than for many other years with the seine catch lower than expected from the long-term relationship. Some differences in relative abundance found by these two sampling methods appeared to have been explained by summer water temperatures affecting fish distribution (NUSCO 1999), but this effect was less certain when re-examined in NUSCO (2000). A temperature effect may also have reduced the seine catch in 2001 if young avoided shallow


Fig. 27. Relationship (functional regression) between the late summer (August-September) seasonal $1-m$ beam trawl median CPUE at station WA in the Niantic River and the geometric mean seine catch during September at a nearby shoreline station sampled by CT DEP from 1988 through 2001.
nearshore waters because of high water temperature; September 2001 water temperatures were the second highest in 26 years (Table 1). In general, however, both sampling methods allowed for the identification of both particularly strong and weak year-classes of winter flounder and serve to corroborate one another. Continued sampling of juvenile winter flounder following their movement out of shallow waters during late fall and early winter is discussed in the following section of this report and provides additional evidence of relative year-class strength during the first year of life.
Growth. Growth of young winter flounder is important because size of fish often has important implications for survival, which in turn affects recruitment. Growth of age-0 winter flounder was measured by increases in mean length over time. In some years, increases in weekly or biweekly mean length occurred steadily throughout the season, whereas in other years asymptotes appeared to be reached by mid-summer. During 2001, mean lengths showed small, but steady increases before leveling off in mid-August (Fig. 28). Based on overlapping error bars associated with each biweekly mean length, little difference in size was seen between the two stations. The relatively small standard errors also indicated little variability in size until September, although an occasional individual much larger than average was taken. Nearly all (99\%) fish collected in 2001 were 60 mm or smaller, but a few individuals ranged up to 90 mm .
Fast growth after settlement followed by a rapid decline in growth rate was reported for young winter flounder in New Jersey bays by Sogard and Able


Fig. 28. Weekly mean length ( $\pm 2$ standard errors) of age- 0 winter flounder taken at Niantic River stations LR and WA by 1 -m beam trawl during 2001.
(1992), who found nearly imperceptible growth rates by the time young winter flounder reached 50 mm in length. It is likely that growth compensation occurs in winter flounder where size-at-age, which may diverge in larval stages, converges during the early juvenile phase and progressive declines are seen in size-at-age differences (Bertram et al. 1993). The few larger individuals taken this year may have reached a size giving them a competitive advantage over other fish and perhaps were able to utilize a food resource not available to smaller winter flounder.
In most years since 1984, mean length of fish at LR during late summer (i.e., July through September) was significantly greater than that of fish at WA (Table 17). In some years, including 2000 ( 12 mm ), a relatively large ( $\geq 8 \mathrm{~mm}$ ) difference was found in mean length between the two stations. Differences were moderate ( $3-4 \mathrm{~mm}$ ) in a few years (e.g., 1986, 1992, 1998) and mean lengths were similar ( -2 to 1 mm ) in other years (e.g., 1991, 1994), including 2001. This year, mean lengths of young were only 29.6 mm at LR and 31.1 mm at WA ; both were the smallest means attained at each station since 1983. A trend in significantly smaller mean lengths at WA has been seen since 1994, whereas mean lengths at LR appeared to have varied more from year to year.
Growth of young winter flounder was probably affected by water temperature, with both positive and negative effects possible. Warmer temperatures result in increased growth rates, but after some point increasing respiratory and other metabolic demands inhibit growth and optimal growth temperatures for young winter flounder can be exceeded (Sogard and Able 1992; Meng et al. 2000). With some exceptions, smaller mean lengths tended to occur in years

TABLE 17. Comparison of the mean lengths (mm) of age-0 winter flounder taken at stations LR and WA in the Niantic River from midJuly through September of 1983 through 2001. Seasonal mean lengths are ranked from largest to smallest and significant differences in means were tested by one-way analysis of variance and Tukey's studentized range (HSD) test.


[^15]when summer water temperatures (Table 2) were warmest with larger means found during cooler years. In this regard, summer water temperatures from 1999 through 2001 were the highest in 26 years (Table 2) and likely influenced the small mean lengths observed for fish taken at WA during these summers. Some differences in mean lengths noted each year between LR and WA (Table 17) were also likely due to water temperature, as WA was generally warmer (ca. $0.5-1.0^{\circ} \mathrm{C}$ ) than LR. No numerical
relationships have been found between mean length and water temperature (see below) nor have the annual differences in mean length at the two stations sampled been consistent.
Another physicochemical factor potentially affecting growth, distribution, and survival is dissolved oxygen (DO). Growth and survival of young winter flounder were depressed in habitats where DO concentrations were low (ca. $<2 \mathrm{mg} \cdot \mathrm{L}^{-1}$ ) during extended periods (Bejda et al. 1992; Phelan et
al. 2000; Meng et al. 2001). Data provided in Goldberg et al. (2000) showed that at a mid-Niantic River station ( 2 m in depth at low tide) in August 1997, DO levels in association with a period of warm water temperatures averaged $3.42 \mathrm{mg} \cdot \mathrm{L}^{-1}$ and periodically decreased to below $2 \mathrm{mg} \cdot \mathrm{L}^{-1}$. DO levels at the shallower shoreline stations sampled for age-0 winter flounder are unknown, but low DO in combination with warm water temperatures in late summer could depress growth rates of young winter flounder.
The occurrence of relatively large mean lengths associated with low age-0 winter flounder abundance in some years and small mean lengths in years when fish were abundant suggested that growth was density-dependent. DeLong et al. (2001) reported that age- 0 winter flounder growth rate was negatively related to their density. However, numerous accounts of age- 0 European plaice were inconclusive with respect to effects of density on growth. Bergman et al. (1988) and Zijlstra et al. (1982) reexamined reports of density-dependent growth in British waters by Steele and Edwards (1970), Lockwood (1972), and Rauck and Zijlstra (1978). They concluded that increases in length corresponded to maximum growth expected from prevailing water temperatures and that growth was not densitydependent. Nash et al. (1994) suspected a complex interaction between density of age- 0 European plaice and water temperature upon growth. Bergman et al. (1988) and Van der Veer et al. (1990) noted that growth of young European plaice in northwestern Europe was not food-limited, but was related to prevailing water temperatures and the length of the growing season in different nursery areas. Pihl and Van der Veer (1992) also believed that growth of young European plaice in Swedish bays was affected by ambient water temperatures and was not foodlimited, although Modin and Pihl (1994) later found evidence for density-dependent growth in a year of extremely high densities. In contrast, Berghahn (1987) and Karakiri et al. (1989) suggested that food limitation and not water temperature may have been responsible for differences in growth of European plaice observed among years in the German Wadden Sea. Also, within the Wadden Sea of The Netherlands, growth of European plaice was slower in an area of tidal sand flats than in an area of mixed sediments with a richer benthos (Berghahn et al. 1995). Benthic food production and its availability also may differ among areas within the Niantic River and likely changes from year to year with variable effects on growth. Juvenile winter flounder growth also appears to be independent of the presence or
absence of eelgrass or macroalgae (Meng et al. 2000; Phelan et al. 2000), although Sogard (1992) reported faster growth in unvegetated in comparison to vegetated areas.
The effects of density (median CPUE during July and August) and water temperature (both cumulative degree-days and running average temperature during the periods of both February I and May 15 through September 30) on growth (mean lengths achieved during late summer) at each station were examined using a multiple linear regression model. Despite the findings of others regarding the growth of young flatfishes, water temperature generally did not appear to consistently affect age-0 winter flounder growth in the Niantic River, at least using these simple models. Using functional regression, the mean length of young at LR during late summer was significantly negatively correlated ( $r=-0.739 ; p=0.0003 ; \mathrm{df}=$ 18) with density (Fig. 29). Although several mean


Fig. 29. Relationship (functional regression) between the annual density (inedian catch per $100 \mathrm{~m}^{2}$ ) and mean length of age-0 winter flounder during late summer (AugustSeptember) at stations LR and WA in the Niantic River from 1983 (LR) or 1984 (WA) through 2001. For WA, data points designated by an ' $x$ ' for 1988 and 1991 were not included in the regression. (Note that the vertical and horizontal scales differ between the graphs).
lengths at high densities appeared to be greater than would have been expected, the value for 2001 was somewhat less than predicted by this relationship. Also, mean lengths were smaller than expected in several years when densities were moderately low. The relationship between abundance and late summer mean length at station WA was also significant and negative ( $r=-0.590 ; p=0.016 ; d f=15$ ), but only if data outliers from 1988 and 1991 were excluded. Mean lengths were much larger than expected from the relatively high densities found during these 2 years. However, the mean length achieved in 2001 (smallest of the series) was less than expected from the calculated relationship.
Mortality. Estimates of monthly instantaneous mortality rate ( $\mathrm{M}_{\mathrm{mo}}$ ) at each station were obtained annually using catch curves constructed from weekly or biweekly abundance data. This method assumed that young comprised a single-age cohort throughout the season and also did not take into account any offstation movements, which would have positively biased estimates of mortality. With some exceptions, the catch curves generally fit the data well as indicated by relatively high $\mathrm{r}^{2}$ values (Table 18 ).
Poor fits (e.g., $r^{2}=0.47$ for LR in 1996) were generally indicative of relatively high variation in weekly abundance estimates. Also, no estimates could be made for WA in 1986, 1993, and 1996 because of considerable variation in weekly abundance. No mortality estimates were determined for LR and WA during the high abundance year of 1988 as slopes of these catch curves were not significantly different from zero. This also occurred at WA in 2000, as abundance steadily increased into late summer and remained relatively stable thereafter (DNC 2001a). High ( $\geq 0.90$ ) $r^{2}$ values were determined for the 2001 catch curves. The $\mathrm{M}_{\text {mo }}$ estimate for station LR in 2001 was 0.869 (equivalent to a survival rate $\mathrm{S}_{\mathrm{mo}}$ of $42.0 \%$ ), which was among the highest of age-0 mortality rates since 1983. Similarly, the $M_{\mathrm{mo}}$ of $1.197\left(\mathrm{~S}_{\mathrm{mo}}=30.2 \%\right)$ at WA was the highest mortality rate found at that station since 1985. The relationship between fish density and annual mortality rates is discussed below in the section on Formation of Winter Flounder Year-class Strength.

Long-term mean monthly mortality estimates were similar (0.62) at both Niantic River stations. This mortality rate was greater than the equivalent value of 0.371 reported by Pearcy (1962) for the Mystic River, CT' estuary and 0.336 for post-settlement European plaice in The Netherlands (Beverton and Iles 1992a; Jager et al. 1995), but was similar to various estimates (0.563-0.693) made for young

European plaice in British coastal embayments (Lockwood 1980; Poxton et al. 1982; Poxton and Nasir 1985; Al-Hossaini et al. 1989). At five different locations in Connecticut, $M_{m o}$ estimates were from 0.24 to 1.21 (Howell 1993; Meise et al. 1999) and in Narragansett Bay, RI values ranged between about 0.3 and 1.1 over an 11-year period (DeLong et al. 2001).
Niantic Bay was sampled for age-0 winter flounder in addition to the Niantic River during the summers of 1988 through 1992 (NUSCO 1994a). Mortality of young in the bay was much greater than in the river. Except for a station just outside the mouth of the Niantic River in 1988, no young were found in Niantic Bay from mid-summer through the end of September in each of these years. Even in 1988, densities at the bay station in late summer were only 10 to $15 \%$ of those in the river at the same time. High natural mortality of young winter flounder in Niantic Bay was the probable reason for declines in density following larval .. metamorphosis and settlement to the bottom, rather than from emigration inshere because none of these smaller, slower growing fish were found in subsequent collections in the river. Because of the apparent lack of young in Niantic Bay in comparison to the Niantic River, no further sampling was conducted in the bay after 1992. This work also suggested that nearly all winter flounder recruits in this area are likely produced in protected inshore embayments or estuaries rather than in more open areas of LIS. Despite the presence of many winter flounder larvae in coastal Atlantic waters (Smith et al. 1975), only one metamorphosed age-0 winter flounder specimen was captured by Steves et al. (1999) in sampling on the continental shelf of the New York Bight, indicating that survival of settled winter flounder is extremely low in deeper, offshore areas and that larvae found in these waters are likely surplus production.
Stock Identification. A winter flounder stock identification study performed by Crivello (2002) used a DNA-based genetic technique to classify winter flounder larvae entrained at MPS to their source of origin (see MPS Impact Assessment Larval Entrainment, below). As part of this study, age-0 juveniles collected at stations LR and WA in both early (late June or early July) and late (September) during 2000 and 2001 were also analyzed using the same techniques. A range of 18 to 78 juveniles 'were processed from 'this set " of collections by station and date. Fish were assigned to three source populations (Niantic River, Westbrook, Thames River) or as unknown, based on the collection and genetic analysis of yolk-sac larvae

TABLE 18. Monthly instantaneous natural mortality rate (M) estimates as determined from catch curves of age-0 winter flounder taken at two stations (LR and WA) in the Niantic River from 1984 through 2001

| Year | Station | $n^{*}$ | slope ${ }^{\text {b }}$ | Standard error | $r^{2}$ | Station | $\mathrm{n}^{2}$ | slope ${ }^{\text {b }}$ | Standard error | $\mathrm{r}^{2}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1984 | LR | 16 | -0.129** | 0.017 | 0.80 | WA | - | - | - | - |
| 1985 |  | 15 | -0.118** | 0.015 | 0.82 |  | 16 | -0.084** | 0.023 | 0.51 |
| 1986 |  | 15 | -0.127** | 0.012 | 0.89 |  | - | . | - | - |
| 1987 |  | 15 | -0.108 ** | 0.021 | 0.67 |  | 16 | -0.139 ** | 0.016 | 0.84 |
| 1988 |  | 19 | NS | - | - |  | 19 | NS | - | - |
| 1989 |  | 12 | -0.154** | 0.022 | 0.84 |  | 13 | -0.145** | 0.028 | 0.71 |
| 1990 |  | 13 | -0.322** | 0.028 | 0.92 |  | 15 | -0.235 ** | 0.028 | 0.84 |
| 1991 |  | 18 | -0.140** | 0.016 | 0.82 |  | 18 | -0.049** | 0.011 | 0.54 |
| 1992 |  | 18 | -0.129** | 0.019 | 0.7 .4 |  | 16 | -0.112*** | 0.009 | 0.91 |
| 1993 |  | 9 | -0.087 * | 0.028 | 0.57 |  | 10 | NS | - | - |
| 1994 |  | 9 | -0.110** | 0.008 | 0.96 |  | 9 | -0.124** | 0.020 | 0.84 |
| 1995 |  | 9 | -0.203** | 0.010 | 0.93 |  | 9 | -0.138** | 0.018 | 0.89 |
| 1996 |  | 9 | -0.116* | 0.046 | 0.47 |  | 8 | NS | - | - |
| 1997 |  | 8 | -0.185** | 0.025 | 0.90 |  | 9 | -0.187** | 0.020 | 0.93 |
| 1998 |  | 8 | -0.101** | 0.013 | 0.91 |  | 8 | -0.102 ** | 0.029 | 0.68 |
| 1999 |  | 9 | -0.111** | 0.024 | 0.74 |  | 9 | -0.128** | 0.029 | 0.73 |
| 2000 |  | 8 | -0.072 ** | 0.019 | 0.71 |  | 8 | NS | - | - |
| 2001 |  | 8 | -0.200** | 0.024 | 0.92 |  | 8 | -0.276 ** | 0.037 | 0.90 |
|  | Mortality ( $\mathrm{Manc}_{\text {m }}{ }^{\text {d }}$ |  |  | Survival ( $\mathrm{Smom}_{\text {mo }}$ ) |  | Mortality ( $\left.\mathrm{M}_{\text {noo }}\right)^{\text {d }}$ |  |  | Survival ( $\mathrm{S}_{\text {mo }}$ ) |  |
| 1984 | LR | 0.560 |  | 57.1\% |  | WA | - |  | - |  |
| 1985 |  | 0.512 |  | 59.9\% |  |  | 0.363 |  | 69.9\% |  |
| 1986 |  | 0.552 |  | 57.6\% |  |  | . ${ }^{\text {c }}$ |  | - |  |
| 1987 |  | 0.469 |  | 62.6\% |  |  | 0.604 |  | 54.7\% |  |
| 1988 |  | - |  | - |  |  | - |  | - |  |
| 1989 |  | 0.669 |  | 51.2\% |  |  | 0.630 |  | 53.3\% |  |
| 1990 |  | 1.398 |  | 24.7\% |  |  | 1.021 |  | 36.0\% |  |
| 1991 |  | 0.608 |  | 54.4\% |  |  | 0.213 |  | 80.8\% |  |
| 1992 |  | 0.560 |  | 57.1\% |  |  | 0.486 |  | 61.5\% |  |
| 1993 |  | 0.377 |  | 68.6\% |  |  | - |  | - |  |
| 1994 |  | 0.476 |  | 62.1\% |  |  | 0.538 |  | 58.4\% |  |
| 1995 |  | 0.883 |  | 41.4\% |  |  | 0.600 |  | 54.9\% |  |
| 1996 |  | 0.504 |  | 60.4\% |  |  | - |  | - |  |
| 1997 |  | 0.802 |  | 44.9\% |  |  | 0.811 |  | 44.4\% |  |
| 1998 |  | 0.437 |  | 64.6\% |  |  | 0.443 |  | 64.2\% |  |
| 1999 |  | 0.480 |  | 61.9\% |  |  | 0.556 |  | 57.4\% |  |
| 2000 |  | 0.312 |  | 73.2\% |  |  | - |  | - |  |
| 2001 |  | 0.869 |  | 42.0\% |  |  | 1.197 |  | 30.2\% |  |
|  | Mean | 0.616 |  | 54.0\% |  | Mean | 0.622 |  | 53.7\% |  |
|  | Standard error | 0.062 |  |  |  | Standard error | 0.079 |  |  |  |
|  | CV | 42\% |  |  |  | CV | 44\% |  |  |  |

- Weekly sampling during 1984-92 and biweekly sampling thereafter. WA was not sampled in 1984.
- Slope (M) deternined from a catch curve (natural logarithm of density plotted against time in weeks). The probability level that the slope of the catch curve differs from zero is shown:

NS - not significant $(p>0.05), *-$ significant at $p \leq 0.05, * *-$ significant at $p \leq 0.01$.
e Although having a significant slope, the catch curve for 1986 at station WA did not provide a reliable estimate of M because of considerable variation in weekly abundance.
d Monthly mortality rate $\left(\mathrm{M}_{1 \mathrm{ma}_{0}}\right)=(-\mathrm{M}) \times(30.4 / 7)$ and monthly survival $\left(\mathrm{S}_{\mathrm{ma}}\right)=\boldsymbol{e}^{-\mathrm{Mma}}$.
collected earlier in the year at each of the three study sites. He reported that most juveniles probably originated from the Niantic River or Westbrook, with fewer coming from the Thames River. Combining data for the two stations and months, about one-third
of the larvae each originated from the Niantic River and Westbrook, $15 \%$ from the Thames River, and the remainder could not be classified specifically to a source area or were unknown in 2000. During 2001, about $22 \%$ of the larvae apparently originated from
the Niantic River, 35\% from Westbrook, 15\% from the Thames River, with the rest not classified specifically to a source area or were unknown. The fraction of age- 0 fish assigned to the Niantic River was very similar to the $24 \%$ of entrained larvae examined by Crivello (2002) that were similarly classified to this source.

## Age-0 Juveniles during Late Fall and Early Winter

Field Sampling and Data Analyses. The collection of age- 0 fish during late fall and early winter occurs during a transitional period following the $1-\mathrm{m}$ beam trawl sampling of young in summer and preceding the catch of this cohort of fish as age-1 juveniles during the intensive winter-early spring adult winter flounder survey in the Niantic River and elsewhere in the MPS study area. Catches from the year-round trawl monitoring program (TMP; see the Fish Ecology section of this report for methods) were used to determine the abundance of age- 0 winter flounder during fall and *winter following the movement of these fish from shallow waters near the shoreline to deeper waters as water temperatures decrease. Young were first regularly captured by trawl in November at the two shallower inshore stations (NR and JC) adjacent to inshore nursery grounds and in December at the near-shore Niantic Bay stations (IN and NB). These fish next appeared during January at the deeper-water stations (TT and BR), formerly sampled in LIS as part of the TMP' (NUSCO 1996b). A $\Delta$-mean (NUSCO 1988c) index describing relative, year-class abundance was developed for age 0 fish using pooled TMP catch data, beginning with the months given above and continuing through the end of February. In January 1996, TMP stations BR, TT, and NB were deleted (NUSCO 1997a). Therefore, sample size was reduced from 42 during 1976-77 through 1994-95 to ${ }^{\text {t }}$ 28 in 1995-96 and 25 in 1996-97 and thereafter.

Abundance and Distribution. Based on the availability of data for this report, the most recent $\Delta$-mean CPUE is for the 2000 year-class. The $\Delta$-mean CPUE for $2000-01^{\circ}$ was 11.6, which is similar the value of 10.5 in 1998-99 (Table 19). However, relatively broad $95 \% \mathrm{CI}$ were computed for nearly all these seasonal $\Delta$-means and differences among years must be viewed with caution. In most years since 1984, when data were first available from Niantic River beam trawl sampling, the late fall-early winter abundance indices reflected densities of age-0 winter flounder found in the river during late summer
(Fig. 30). An exception was the 1996-97 $\Delta$-mean of 18.3 for the 1996 year-class, which appeared to be weak from sampling conducted during summer in the Niantic River (NUSCO 1997b). However, as noted above, the DEP seine catch of young winter flounder showed that the 1996 year-class of winter flounder was likely larger than indicated by the beam trawl catches in late summer. Similarly, the CPUE of 4.8 in 1995-96 and 11.6 in 2000-01 did not correspond to the relatively high abundance of age- 0 fish seen in the Niantic River during the preceding summer months. Greater concordance among abundance indices, however, was found in many other years, including 1994-95, as the $\Delta$-mean CPUE of 31.7 reflected the strong 1994 year-class. Also evident in recent years was good production of young in 1988 and 1992 as well as weaker year-classes in 1993 and 1997, based on $\Delta$-means of 29.6, 31.1 ; 7.4, and 2.3, respectively. The reason for the disparity found between the abundance age- 0 winter flounder in the summer of 2000 in the Niantic River and at the TMP stations during the subsequent fall and early winter is unknown.
Despite some discrepancies noted between the two age-0 abundance indices, they generally tracked one another and were significantly correlated (Spearman's rank-order correlation; $r=0.529 ; p=0.029 ; n$ = 17). However, the fall-winter TMP $\triangle$-mean CPUE and $\Delta$-mean CPUE of winter flounder smaller than 15 cm taken in the Niantic River during the subsequent (late February-early April) adult winter flounder survey (see Age-1 Juveniles during Late Winter, below) were not significantly related ( $r=-0.246 ; p=$ $0.235 ; n=25$; Fig. 31). The relationship between these abundance indices of temporally successive life-stages was unclear. More juvenile winter flounder from the 1984 and earlier year-classes were taken in the river than at the six TMP stations (five of which are outside of the Niantic River) during the preceding months. Since the 1988 year-class was produced, the catch index of young from the TMP in most years has been higher that that of age-1 fish in the river, although the indices were nearly the same magnitude in 1989, 1993, and 1997. The reason for this areal shift in magnitude of relative abundance that has occurred since the mid-1980s is unknown. The numbers of young taken during late fall and early winter by the TMP should be a predictor of age-1 fish abundance in the Niantic River during late winter and early spring. However, this assumes that the relative distribution of fish both inside and outside the river remains constant each year, which likely does not occur.

TABLE 19. The late fall-early winter seasonal ${ }^{4} \Delta$-mean CPUE $^{6}$ of age- $0^{c}$ winter flounder taken at six (1976-95) or three (1996-2001) trawl monitoring stations in the vicinity of MPS.

| Survey year | Number of samples | Non-zero observations | $\Delta$-mean ${ }^{\text {b }}$ | 95\% confidence interval |
| :---: | :---: | :---: | :---: | :---: |
| 1976-77 | 42 | 36 | 6.1 | 2.0-10.3 |
| 1977-78 | 42 | 38 | 5.1 | 2.3-7.9 |
| 1978-79 | 42 | 36 | 4.2 | 2.0-6.4 |
| 1979-80 | 42 | 38 | 4.2 | 2.2-6.2 |
| 1980-81 | 42 | 39 | 10.1 | 4.3-15.9 |
| 1981-82 | 42 | 39 | 7.7 | 2.9-12.5 |
| 1982-83 | 42 | 37 | 19.6 | 9.0-30.3 |
| 1983-84 | 42 | 39 | 6.6 | 3.2-10.0 |
| 1984-85 | 42 | 35 | 7.4 | 1.7-13.1 |
| 1985-86 | 42 | 39 | 8.1 | $4.4-11.7$ |
| 1986-87 | 42 | 39 | 11.7 | 3.4-19.9 |
| 1987-88 | 42 | 41 | 4.8 | 2.1-7.5 |
| 1988-89 | 42 | 41 | 29.6 | 11.8-47.3 |
| 1989-90 | 42 | 42 | 11.3 | 6.7 - 15.9 |
| 1990-91 | 42 | 40 | 21.7 | 6.7-36.8 |
| 1991-92 | 42 | 41 | 19.0 | 7.6-30.3 |
| 1992-93 | 42 | 39 | 31.1 | 7.4-54.8 |
| 1993-94 | 42 | 38 | 7.4 | 3.4-11.4 |
| 1994-95 | 42 | 41 | 31.7 | 7.3-56.1 |
| 1995-96 | 28 | 25 | 4.8 | 1.0-8.6 |
| 1996-97 | 25 | 2.4 | 18.3 | -4.1-40.7 |
| 1997-98 | 25 | 19 | 2.3 | 0.8-3.7 |
| 1998-99 | 25 | 22 | 10.5 | 2.1-18.9 |
| 1999-00 | 25 | 23 | 15.9 | -8.3-40.2 |
| 2000-01 | 25 | 21 | 11.6 | -2.7-25.8 |

* Data restricted during 1976-77 through 1994-95 to November-February for NR and JC, December-February for IN and NB, and JanuaryFebruary for TT and BR and during 1995-96 and thereafter to November-February for NR and JC and December-February for IN
b Catch per standardized tow of 0.69 km (see Materials and Methods of Fish Ecology section).
c For age-0 fish, the year-class is the same as the first year given.


Fig. 30. Comparison between the late fall-early winter seasonal $\Delta$-mean CPUE (solid line; $\pm 2$ standard errors) of age-0 winter flounder (all trawl monitoring program stations) and the late summer Niantic River (stations LR and WA combined) age-0 $1-\mathrm{m}$ beam trawl median CPUE (dashed line; $\pm 2$ standard errors) for the 1984-2000 yearclasses.

Relationships among abundance indices of juvenile winter flounder may have been obscured by differences in sampling gear used and variations in
fish behavior in response to environmental conditions. Major biases in abundance estimation can arise from size selectivity of the gear, spatial distribution of individuals in relation to the gear, and behavior of fish in the vicinity of the gear (Parrish 1963). From 1985 through 2000, annual mean lengths of age- 0 winter flounder taken by otter trawl from October through December were from 21 to 44 mm larger than those taken during August and September by $1-\mathrm{m}$ beam trawl (Fig. 32). The annual pattern of mean lengths for the two gear types appeared similar, although they were not significantly related (Spearman's rank-order correlation; $r=0.465 ; p=0.069 ; n=16)$. The differences in observed mean lengths could have been achieved entirely by growth of these individuals, although CPUE indices also may have been biased because smaller individuals were excluded from the TMP catch. Examination of length-frequencies of otter trawl-caught winter flounder from the TMP and the Niantic River adult winter flounder survey indicated that even though an occasional small fish was taken, small winter flounder did not begin to regularly appear in the


Fig. 31. Comparison between the late fall-early winter seasonal $\Delta$-mean CPUE (solid line; $\pm 2$ standard errors) of age-0 winter flounder (all trawl monitoring program stations) and the Niantic River (stations 1 and 2) spawning survey $\triangle$-mean CPUE (dashed line; $\pm 2$ standard errors) of winter flounder smaller than 15 cm for the 1976-2000 yearclasses. (Note that the vertical scales differ between the graphs).
catches until they were about $40-50 \mathrm{~mm}$ in length. The otter trawl was probably not fully efficient until fish were about $55-60 \mathrm{~mm}$ in length. Therefore, lengths achieved by age-0 winter flounder each year (Table 17) may have differentially biased otter trawl CPUE because smaller means occurred more frequently after the mid-1990s than in earlier years. Finally, the fixed locations of the otter trawl sampling stations in relation to the habitat available to juveniles also may have affected the catch comparisons as well as the deletion of three offshore TMP stations.

Movements of small juveniles were probably influenced by factors such as water temperature and tide. Moreover, their availability to sampling gear in fall and winter appeared to have varied from week to week and year to year. Relatively large $95 \%$ Cls around the $\Delta$-mean CPUE values were probably a consequence of this variation. In contrast, variation was usually less in data collected during summer by the relatively efficient $1-\mathrm{m}$ beam trawl, which is also


Fig. 32. Comparison between the mean length ( $\pm 2$ standard errors) of age -0 winter flounder taken during late summer (August-September) by $1-m$ beam trawl in the Niantic River and in fall (October-December) by otter trawl at three TMP stations.
when age-0 winter flounder remain relatively sedentary. Furthermore, sampling in summer occurred weekly or biweekly during the same tidal stage and in areas known to be preferred habitat of young winter flounder. Finally, a mixture of juveniles from a number of sources most likely was present throughout LIS during the winter, which would have influenced measures of abundance because of potential variable contributions from different stocks. These factors all contributed to weakening the strength of correlations among juvenile winter flounder abundance indices, which is discussed fürther below.

## Age-1 Juveniles during Late Winter

Field Sampling and Data Analyses. In addition to catches made during the TMP, juvenile winter flounder smaller than 15 cm in length (mostly age-1) were taken along with adults in the annual February-April Niantic River adult spawning surveys, the methods for which were described previously. These fish were processed similarly as the adults, although gender was usually not specified and these fish were not branded. In years when small winter flounder were abundant, a subsample of at least 200 fish was measured each survey week; otherwise, all specimens were measured. For some annual comparisons of abundance, data were restricted to stations 1 and 2 in the lower river navigational channel (Fig. 2) because the distribution of small winter flounder generally varied more than the adult fish, and also because no tows were made in the upper river from 1977 through 1980. Annual
$\Delta$-mean CPUE indices of age-1 juveniles were determined as described previously for fish larger than 15 cm . One CPUE was computed for fish found at stations 1 and 2 combined and, when sufficient data were available, others were calculated for all river stations combined or for the upper river basin and arm (stations 6 and 51-54). For comparative purposes, an annual $\Delta$-mean abundance index of juvenile fish of similar size was also determined using catch data from the five (or in 1996 and thereafter, two) TMP stations outside of the Niantic River during January through April, a period which overlapped the adult spawning surveys and also served to increase sample size. Annual effort for this abundance index included 45 trawl collections through 1995 and 18 in subsequent years.

Abundance and Distribution. The $\Delta$-mean CPUE for age-1 juveniles taken in the navigational channel of the lower Niantic River during 2001 was 0.3, which was the smallest abundance index found since

1976 (Table 20). Abundance of age-1 fish has been particularly low in the Niantic River during the past 6 years. The presence of a heavy blue mussel set at station 1, previously noted in Adult Winter Flounder - Relative Annual Abundance, most likely affected the presence of juveniles in this area as it did adults. Median CPUE of juveniles in the upper river was 2.2. This abundance index has shown a small, but steady increase since its lowest value in 1996 (Table 21).
Distribution of juvenile winter flounder during winter largely influences their availability to sampling and apparently differs from year to year, probably as a result of variable environmental conditions, including water temperature, winter storm events, and the aforementioned change in bottom type. The relative abundance of small winter flounder has also not been consistent between Niantic Bay and Niantic River from year to year as well as within areas of the river. A $\triangle$-mean CPUE computed for winter flounder smaller than 15 cm taken by the TMP

TABLE 20. Annual $9.1-m$ otter trawl adjusted $\triangle$-mean CPUE" of winter flounder smaller than $15 \mathrm{~cm}^{b}$ taken in the navigational channel of the lower Niantic River during the 1976 through 2001 adult population abundance surveys.

| Survey year | Weeks used for CPUE computation ${ }^{\text {c }}$ | Tows acceptable for CPUE | Adjusted number of tows used ${ }^{\text {c }}$ | Non-zero observations | $\Delta$-mean CPUE estimate | Standard error | $95 \%$ confidence interval for $\Delta$-mean CPUE |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1976 | 7 | 135 | 182 | 180 | 24.1 | 1.7 | 20.8-27.4 |
| 1977 | 6 | 206 | 264 | 256 | 25.6 | 2.5 | 20.7-30.6 |
| 1978 | 6 | 144 | 168 | 164 | 33.8 | 4.5 | 24.9-42.6 |
| 1979 | 5 | 118 | 135 | 133 | 65.1 | 9.7 | 46.1-84.1 |
| 1980 | 5 | 115 | 150 | 149 | 56.1 | 4.9 | 46.5-65.7 |
| 1981 | 7 | 97 | 154 | 154 | 86.2 | 7.9 | 72.8-99.5 |
| 1982 | 5 | 50 | 70 | 70 | 57.4 | 10.4 | 37.0-77.8 |
| 1983 | 7 | 77 | 77 | 76 | 52.5 | 6.4 | 39.9-65.0 |
| 1984 | 7 | 73 | 77 | 76 | 25.3 | 3.0 | 19.4-31.3 |
| 1985 | 7 | 83 | 84 | 84 | 33.9 | 3.7 | 26.6-41.2 |
| 1986 | 7 | 75 | 126 | 115 | 5.9 | 0.6 | 4.8-7.0 |
| 1987 | 5 | 41 | 50 | 44 | 6.6 | 0.9 | 4.9-8.3 |
| 1988 | 6 | 49 | 54 | 52 | 17.0 | 3.1 | 11.0-23.1 |
| 1989 | 6 | 50 | 56 | 50 | 10.6 | 1.9 | 6.9-14.3 |
| 1990 | 7 | 66 | 91 | 88 | 14.5 | 2.0 | 10.6-18.3 |
| 1991 | 6 | 45 | 60 | 56 | 7.4 | 1.2 | 5.0-9.8 |
| 1992 | 7 | 35 | 49 | 44 | 11.9 | 2.1 | 7.8-16.1 |
| 1993 | 7 | 36 | 49 | 45 | 6.6 | 1.0 | 4.6-8.5 |
| 1994 | 4 | 22 | 24 | 24 | 5.6 | 1.3 | 3.1-8.1 |
| 1995 | 6 | 42 | 54 | 50 | 6.3 | 1.1 | 4.2-8.4 |
| 1996 | 6 | 49 | 60 | 38 | 1.6 | 0.3 | 1.0-2.2 |
| 1997 | 6 | 43 | 48 | 41 | 3.2 | 0.5 | 2.2-4.2 |
| 1998 | 7 | 56 | 63 | 47 | 2.5 | 0.3 | 1.8-3.1 |
| 1999 | 7 | 50 | 63 | 39 | 1.0 | 0.2 | 0.7-1.3 |
| 2000 | 6 | 42 | 48 | 24 | 1.0 | 0.2 | 0.5-1.4 |
| 2001 | 7 | 41 | 63 | 16 | 0.3 | 0.1 | 0.1-0.4 |

[^16]TABLE 21. Comparison of annual $9.1-\mathrm{m}$ otter trawl adjusted $\triangle$-mean CPUE' of winter flounder smaller than $15 \mathrm{~cm}^{6}$ taken in the navigational channel of the lower Niantic River (sta. 1 and 2; Fig. 2) with those caught in the upper river arm and basin (sta. 6 and 51-54; Fig. 2) during the 1981 through 2001 adult population abundance surveys.


- Catch per standardized tow (see text for details).
- Mostly age-1 fish; predominant age-class was produced I year before the survey year.
- Effort equalized among weeks; during several years weeks with very low effort were not used for computing CPUE. Only tows of standard time or distance were considered.
from January through April at stations outside of the Niantic River was compared to the $\triangle$-mean CPUE for fish found in the navigational channel of the lower river during the spawning season. Generally, the catch of age- 1 winter flounder in winter and early spring fluctuated less outside than inside the Niantic River, although the latter had higher $95 \% \mathrm{CI}$ because of much smaller sample sizes (Fig. 33). As the number of age-1 winter flounder in the lower river declined in recent years, relative abundance of these fish in Niantic Bay increased. The CPUE of fish found in the bay during January through April was greater than that of fish taken in the lower river navigational channel in most years since 1986, including 2001. The catch outside the river during 1995 was the highest of this time-series, indicating that most fish from the relatively strong 1994 yearclass did not remain within or re-enter the Niantic River during the adult spawning season. ${ }^{1}$ Smaller peaks were seen in the bay catches during 1989, 1993, and 1997, which corresponded to relatively good year-classes produced the previous summer and fall.

Since the mid-1980s, annual effort (as measured by the adjusted number of tows used for calculation of CPUE) in the upper Niantic River basin and arm (stations 6,51-54) has increased relative to effort expended in the lower river navigational channel (stations I and 2), particularly since abundance in the latter area has decreased during the past decade. There has been some concern that the sharp decrease in abundance following the 1981 peak in $\Delta$-mean CPUE of age-1 winter flounder taken in the lower river navigational channel was an early indication of subsequent poor recruitment to the Niantic River adult spawning population. Although most of the decrease occurred prior to Unit 3 going online, a question also remained whether or not a plant entrainment effect contributed to this decrease. Since both sampling effort and abundance of age-1 juvenile winter flounder in this small area of the river has decereased from the earliest years of study, another abundance index was calculated for age-1 fish. A $\Delta$-mean CPUE was computed for winter flounder $<15 \mathrm{~cm}$ that were caught in the upper Niantic River basin and arm from 1981 through 2001. Insufficient effort was made in this area prior to 1981 for a


Fig. 33. Comparison between the annual January-April $\Delta$-mean CPUE (solid line; $\pm 2$ standard errors) at all trawl monitoring program stations except NR and the Niantic River (stations 1 and 2) spawning survey $\Delta$-mean CPUE (dashed line; $\pm 2$ standard errors) of winter flounder smalier than 15 cm from 1976 through 2001. (Note that the vertical scales differ between the graphs).
meaningful calculation. After 1989, a decrease was also observed in the upper river $\Delta$-mean CPUE timeseries, but the magnitude of the decline was considerably less than seen in the lower river (Fig. 34). Also, abundance, although remaining lower than in the 1980s, has increased steadily since the lowest value in 1996. As for the lower river CPUE, the upper river abundance may be compared with CPUE from the TMP stations outside of the Niantic River from January through April (same index as
shown on Fig. 33). Age-I winter flounder remained more abundant outside the Niantic River relative to the upper Niantic River for nearly all these years.
A small CPUE for age-1 winter flounder taken within the lower Niantic River may not reflect the abundance of the corresponding year-class, which may be more accurately described by age-0 indices discussed in the two previous sections. A relatively small increase in catch of age-1 fish in the much larger geographical area of Niantic Bay and nearby


Fig. 34. Comparison of $\Delta$-mean CPUE of age-1 winter flounder taken in the navigational channel (stations 1 and 2) of the lower Niantic River and in the upper river basin (stations 6,51-54) during the spawning survey (WFS) and the January-April $\Delta$-mean CPUE of age-1 winter flounder taken at all trawl monitoring program (TMP) stations except NR from 1976 through 2001.

## Comparisons among Life-Stages of Winter Flounder Year-Classes

LIS could have accounted for seemingly low abundance in the river as fish dispersed from a relatively limited and confined area into a much larger geographical space. Similarly, the upper Niantic River basin and arm offers considerably more spatial area for juvenile winter flounder, as do shallow areas within the river not sampled during the spawning surveys. Because of the differential distribution and abundance of age-1 juveniles, perhaps as a consequence of variable environmental conditions influencing their behavior and availability to sampling, the abundance indices determined from data ${ }^{-}$ taken during the :TMP and the adult spawning surveys remain generally unreliable predictors of future population size. Finally, further adjustments to year-class strength occur in the remaining several years of juvenile life. The dispersion of older juvenile winter flounder over large areas of LIS and adjacent estuarine systems that allows for considerable mixing of fish from many spawning stocks makes the task of following a specific cohort from a particular spawning population very difficult. The relationship between juvenile and adult abundance indices is explored further in the following section.

Abundance indices for various life-stages of the 1976 through 2001 year-classes of Niantic River winter flounder given throughout this report are summarized in Table 22. Coefficients of variation (CV) computed for most of these indices were used to compare annual variability in abundance among years (Table 23). Total adult female abundance (CV $=85 \%$ ) and egg production estimates ( $78 \%$ ) were among the least variable of the abundance indices.
Considering the first three adult female age-classes, variability decreased from age-3 (124\%) to ages-4 (103\%) and 5 (101\%). This likely reflected not only variation in recruitment of year-classes, but variable numbers of immature fish of age- 3 fish that were present in the river each year. This is also consistent with findings of Miller et al. (1991), who noted that interannual variability in the abundance indices of many flatfishes appeared to decrease with age.
Stage 1 larval abundance appeared to be the most stable ( $\mathrm{CV}=79 \%$ ) of the early life history stages and had nearly the same variation as found for egg production. Abundance was more variable for Stage 2 ( $109 \%$ ) and Stage 3 ( $121 \%$ ) larvae. Stage 4 larval abundance showed the largest annual variation (134\%), as relatively high densities of this stage were

TABLE 22. Comparison of abundance indices of various winter flounder life-stages from the 1976 through 2001 year-classes.

| $\begin{aligned} & \text { Year- } \\ & \text { class } \end{aligned}$ | Adult abundance indices |  |  | Larval abundance indices |  |  | Juvenile abundance indices |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Niantic female | River annual | Niantic River stations (Feb-Jun) |  |  |  | MPS | Age-0 <br> Niantic | Age-0 <br> Niantic | Age-0 <br> River/bay | Age-1 <br> $\Delta$-mean | Age-1 <br> $\Delta$-mean |
|  | spawners (Feb-Apr) | egg prod. | Stage 1 <br> ( 3 mm ) | Stage 2 <br> ( 3.5 mm ) | Stage 3 <br> ( 6 mm ) | Stage 4 <br> ( 7.5 mm ) | $\begin{gathered} \text { (EN) } \\ (\geq 7 \mathrm{~mm}) \end{gathered}$ | R. stas. (May-Jul) | R. stas. (Aug-Sep) | $\Delta$-mean (Nov-Feb) | lower rive (Feb-Apr) | (upper river) <br> (Feb-Apr) |
| 1976 | $\bullet$ | - | - | - | - | - | 854 | - | $\bullet$ | 6.1 | 25.6 | - |
| 1977 | 1,069 | 481.4 | - | - | - | - | 567 | - | - | 5.1 | 33.8 | - |
| 1978 | 1,644 | 828.3 | - | - | - | - | 754 | - | - | 4.2 | 65.1 | - |
| 1979 | 1,197 | 565.7 | - | - | - | - | 641 | - | - | 4.2 | 56.1 | - |
| 1980 | 1,004 | 468.9 | - | - | - | - | 845 | - | - | 10.1 | 86.2 | 7.9 |
| 1981 | 2,411 | 1,252.5 | - | - | - | - | 561 | - | - | 7.7 | 57.4 | 7.6 |
| 1982 | 2,551 | 1,479.8 | - | $\cdot$ | - | - | 610 | - | - | 19.6 | 52.5 | 9.9 |
| 1983 | 1,705 | 985.5 | - | 749 | 408 | 56 | 1,215 | - | - | 6.6 | 25.3 | 5.3 |
| 1984 | 822 | 474.9 | 2,601 | 1,501 | 573 | 67 | 917 | - | 7.5 | 7.4 | 33.9 | 10.3 |
| 1985 | 846 | 514.9 | 6,260 | 4,676 | 584 | 35 | 312 | 15.0 | 8.0 | 8.1 | 5.9 | 5.9 |
| 1986 | 604 | 402.5 | 1,279 | 176 | 301 | 24 | 510 | 26.7 | 15.6 | 11.7 | 6.6 | 3.4 |
| 1987 | 780 | 485.9 | 3,218 | 829 | 1,036 | 48 | 315 | 41.7 | 13.1 | 4.8 | 17.0 | 4.8 |
| 1988 | 1,172 | 792.8 | 14,491 | 4,469 | 1,531 | 210 | 419 | 52.5 | 51.7 | 29.6 | 10.6 | 8.9 |
| 1989 | 891 | 647.1 | 12,463 | 3,976 | 589 | 73 | 327 | 12.5 | 7.8 | 11.3 | 14.5 | 2.0 |
| 1990 | 498 | 314.4 | 4,728 | 355 | 258 | 57 | 508 | 117.5 | 18.0 | 21.7 | 7.4 | 1.8 |
| 1991 | 963 | 578.7 | 3,248 | 252 | 343 | 112 | 439 | 45.0 | 24.2 | 19.0 | 11.9 | 3.3 |
| 1992 | 485 | 354.2 | 5,476 | 1,367 | 2,339 | 195 | 1,003 | 77.1 | 29.4 | 31.1 | 6.6 | 2.9 |
| 1993 | 247 | 201.6 | 1,187 | 133 | 111 | $6{ }^{\circ}$ | 130 | 7.3 | 5.0 | 7.4 | 5.6 | 2.6 |
| 1994 | 458 | 297.3 | 3,692 | 1,248 | 429 | 90 | 834 | 126.3 | 52.1 | 31.7 | 6.3 | 1.3 |
| 1995 | 192 | 148.1 | 5,580 | 2,023 | 2,615 | 787 | 1,804 | 111.3 | 28.3 | 4.8 | 1.6 | 0.6 |
| 1996 | 88 | 74.3 | 4,376 ${ }^{\circ}$ | 4,424 | 152 | 31 | 462 | 14.2 | 4.5 | 18.3 | 3.2 | 1.1 |
| 1997 | 166 | 131.8 | 8,342 | 8,449 | 4,978 | 202 | 2,904 | 28.8 | 8.4 | 2.3 | 2.5 | 1.6 |
| 1998 | 118 | 89.1 | 3,138 | 6,553 | 1,550 | 393 | 1,509 | 46.3 | 14.4 | 10.5 | 1.0 | 1.9 |
| 1999 | 107 | 75.0 | 2,632 | 288 | 584 | 73 | 1,420 | 43.8 | 18.5 | 15.9 | 1.0 | 2.1 |
| 2000 | 167 | 123.0 | 408 | 403 | 444 | 66 | 1,134 | 38.8 | 35.0 | 11.6 | 0.3 | 2.2 |
| 2001 | 68 | 56.1 | 1,934 | 704 | 140 | 66 | 837 | 150.0 | 33.8 | - | - | - |

2 An approximation based on cumulative geometric weekly means because the Gompertz function could not be fit to the data as larvae were only collected during 2 weeks of sampling.
seen during several years and an extremely low value was found in 1993 (Table 22). Following metamorphosis, annual variability of newly settled juvenile
winter flounder abundance decreased to $79 \%$, a value similar to the CVs for egg production and yolk-sac larvae. Variability decreased slightly further to 70-

TABLE 23. Coefficients of variation (CV) of annual abundance indices' of various life stages of Niantic River winter flounder.

| Life stage | Abundance index used | Number of observations | CV |
| :---: | :---: | :---: | :---: |
| Female spawners | Annual standardized catch | 25 | 85\% |
| Age-3 females | Annual standardized catch | 23 | 124\% |
| Age-4 females | Annual standardized catch | 22 | 103\% |
| Age-5 females | Annual standardized catch | 21 | 101\% |
| Eggs | Egg production index | 25 | 78\% |
| Stage 1 larvae | A parameter of Gompertz function | 18 | 79\% |
| Stage 2 larvae | $A$ parameter of Gompertz function | 19 | 109\% |
| Stage 3 larvae | $A$ parameter of Gompertz function | 19 | 121\% |
| Stage 4 larvae | A parameter of Gompertz function | 19 | 134\% |
| Age-0 young | Median CPUE in Niantic River (May-July) | 17 | 79\% |
| Age-0 young | Median CPUE in Niantic River (August-Sept) | 18 | 71\% |
| Age-0 young | Fall-winter $\Delta$-mean at trawl stations | 25 | 70\% |
| Age-1 juveniles | $\Delta$-mean CPUE of fish $<15 \mathrm{~cm}$ in lower Niantic River | 25 | 112\% |
| Age-1 juveniles | $\Delta$-mean CPUE of fish $<15 \mathrm{~cm}$ in upper Niantic Rives | 21 | 74\% |

[^17]$71 \%$ in both late summer and in the upper river during the adult spawning surveys is considered, the CV is an approximately similar $74 \%$. However, a larger CV of $112 \%$ computed for age-1 juveniles taken in the navigational channel of the lower Niantic River was probably related to the previously discussed annual differences in distribution related to behavior as much as from actual variation in yearclass stréngth. Considerable variation occurs during the post-settlement juvenile life history stage. In some instances, compensation likely takes place in a density-dependent fashion, whereas other changes in abundance occur independently of density.
Rothschild and DiNardo (1987) reported a median CV of $70 \%$ for recruitment indices of various marine fishes, although various flatfishes had CV values mostly less than $75 \%$. This value is consistent with those found for Niantic River winter flounder. The CV of European flounder abundance decreased from $172 \%(n=9)$ in the larval stage to $99 \%(n=8)$ in newly settled young to $80 \%(n=8,12)$ in young during September and again at age-1 (Van der Veer
et al. 1991). As summarized by Van der Veer (1986), the highest CV for yearly abundance estimates of different life stages of European plaice in The Netherlands occurred during larval development in late winter ( $\mathrm{n}=4, \mathrm{CV}=95 \%$ ) and at first settlement of pelagic juveniles in spring following larval metamorphosis and settling ( $9,62 \%$ ). Smaller variation was found in post-larval young during midsummer ( $9,30 \%$ ) and age-2 recruits ( $9,35 \%$ ), which is less than the variability found for winter flounder. He attributed the decline in variation of abundance in older juveniles to a density-dependent regulatory mechanism that operated during and shortly after larval settlement.
Relationships among abundance indices of winter flounder for the same year-class are of interest for impact assessment (Tables 24 and 25). Knowledge of the earliest possible measure of relative year-class strength is desirable because it enables predictions of future recruitment to the adult stock, thus providing an early warning of decreases in stock abundance. If indices for all life-stages are assumed to be

TABLE 24. Matrix of Spearman's rank-order correlations among various winter flounder spawning stock and larval abundance indices. All indices refer to adults or larvae collected in the Niantic River, except for larvae 7 mm and larger taken at the kirS discharge.

| Index ${ }^{\text {a }}$ | Adult egg production | Stage 1 larvae | Stage 2 larvae | Stage 3 larvac | Stage 4 larvae | Larvac ( 27 mm ) at MPS discharge |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Female spawners | $\begin{gathered} 0.9662^{\mathrm{b}} \\ 0.0001^{* *} \\ 25 \end{gathered}$ | $\begin{gathered} 0.3787 \\ 0.1212 \mathrm{NS} \\ 18 \end{gathered}$ | $\begin{gathered} \hline-0.0263 \\ 0.9148 \mathrm{NS} \\ 19 \end{gathered}$ | $\begin{gathered} 0.0755 \\ 0.7588 \mathrm{NS} \\ 19 \end{gathered}$ | $\begin{gathered} -0.0562 \\ 0.8193 \text { NS } \\ 19 \end{gathered}$ | $\begin{gathered} -0.2931 \\ 0.1551 \text { NS } \\ 25 \end{gathered}$ |
| Adult egg production |  | $\begin{gathered} 0.4345 \\ 0.0716 \mathrm{NS} \\ 18 \end{gathered}$ | $\begin{gathered} 0.0211 \\ 0.9318 \mathrm{NS} \\ 19 \end{gathered}$ | $\begin{gathered} 0.1352 \\ 0.5812 \mathrm{NS} \\ 19 \end{gathered}$ | $\begin{gathered} -0.0421 \\ 0.8640 \mathrm{NS} \\ 19 \end{gathered}$ | $\begin{gathered} -0.3746 \\ 0.0650 \mathrm{NS} \\ 25 \end{gathered}$ |
| Stage 1 larvac |  |  | $\begin{gathered} 0.6760 \\ 0.0021^{* *} \\ 18 \end{gathered}$ | $\begin{gathered} 0.5741 \\ \therefore 0.0127 * \\ \therefore \quad 18 \end{gathered}$ | $\begin{gathered} 0.4762 \\ 0.0457^{*} \\ \hline \end{gathered}$ | $\begin{gathered} -0.0733 \\ 0.7726 \mathrm{NS} \\ 18 \end{gathered}$ |
| Stage 2 larvae |  |  |  | $\begin{gathered} 0.6626 \\ 0.0020^{* *} \\ \therefore 19 \end{gathered}$ | $\begin{gathered} 0.4697 \\ 0.0424 * \\ 19 \end{gathered}$ | $\begin{gathered} 0.2018 \\ 0.4075 \mathrm{NS} \\ 19 \end{gathered}$ |
| Stage 3 larvae |  | . |  | $\because$ | $\begin{gathered} 0.7260 \\ 0.0004 * * \\ 19 \end{gathered}$ | $\begin{gathered} 0.4353 \\ 0.0625 \mathrm{NS} \\ .19 \end{gathered}$ |
| Stage 4 larvae |  | . | - | $\cdots$ |  | $\begin{gathered} 0.5654 \\ 0.0116 * \\ 19 \end{gathered}$ |

[^18]TABLE 25. Matrix of Spearman's rank-order correlations among various larval and juvenile winter flounder abundance indices.

| Index ${ }^{\text {a }}$ | Niantic River Stage 4 larvae | Niantic River early summer age-0 juveniles | Niantic River late summer age0 juveniles | Fallearly winter river-bay age-0 juveniles | Niantic River winter-spring age-1 juveniles (lower river) | Niantic River winter-spring age-1 juveniles (upper river) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Larvae ( } 27 \mathrm{mlm} \text { ) } \\ & \text { at MPS } \\ & \text { disclarge } \end{aligned}$ | $\begin{gathered} 0.5654^{6} \\ 0.0116^{*} \\ 19 \end{gathered}$ | $\begin{gathered} 0.4412 \\ 0.0763 \mathrm{NS} \end{gathered}$ $17$ | $\begin{gathered} 0.3333 \\ 0.1765 \mathrm{NS} \\ 18 \end{gathered}$ | $\begin{gathered} -0.2105 \\ 0.3125 \mathrm{NS} \\ 25 \end{gathered}$ | $\begin{gathered} -0.2220 \\ 0.2862 \mathrm{NS} \\ 25 \end{gathered}$ | $\begin{gathered} -0.2740 \\ 0.2294 \mathrm{NS} \\ 21 \end{gathered}$ |
| Niantic River Stage 4 larvae |  | $\begin{gathered} 0.5313 \\ 0.0282^{*} \\ 17 \end{gathered}$ | $\begin{gathered} 0.4762 \\ 0.0457= \\ 18 \end{gathered}$ | $\begin{gathered} 0.1075 \\ 0.6712 \mathrm{NS} \\ 18 \end{gathered}$ | $\begin{gathered} -0.1907 \\ 0.4485 \mathrm{NS} \\ 18 \end{gathered}$ | $\begin{gathered} -0.2767 \\ 0.2663 \mathrm{NS} \\ 18 \end{gathered}$ |
| Niantic River early summer age-0 juveniles |  |  | $\begin{gathered} 0.7941 \\ 0.0001 \text { ** } \\ 17 \end{gathered}$ | $\begin{gathered} 0.4768 \\ 0.0618 \mathrm{NS} \\ 16 \end{gathered}$ | $\begin{gathered} 0.0324 \\ 0.9052 \mathrm{NS} \\ 16 \end{gathered}$ | $\begin{gathered} -0.2118 \\ 0.4311 \mathrm{NS} \\ 16 \end{gathered}$ |
| Niantic River late summer age0 juveniles |  |  |  | $\begin{gathered} 0.5288 \\ 0.0291 \text { * } \\ \hline 17 \end{gathered}$ | $\begin{gathered} -0.1448 \\ 0.5793 \text { NS } \\ 17 \end{gathered}$ | $\begin{gathered} -0.0564 \\ 0.8298 \mathrm{NS} \\ 17 \end{gathered}$ |
| Fall-early winter river-bay age-0 juveniles |  |  |  |  | $\begin{gathered} -0.2464 \\ 0.235 \mathrm{NS} \\ 25 \end{gathered}$ | $\begin{gathered} -0.0520 \\ 0.8229 \text { NS } \\ 21 \end{gathered}$ |
| Niantic River winter-spring age-1 juveniles (lower river) | - |  |  |  |  | $\begin{gathered} 0.7161 \\ 0.0003 \text { ** } \\ 21 \end{gathered}$ |

2- Indices used correspond to those given on Tables 22 and 23.

- The three statistics shown in each correlation matrix element are: correlation coefticient $(r)$, probability of a larger $r$ (NS - not significant $[p>0.05], *$ significant at $p \leq 0.05, * *$ significant at $p \leq 0.01$ ), and number of annual observations (sample size).
accurately and precisely measured each year, they should be correlated (after applying appropriate time lags), except when processes such as densitydependent mortality or size-selective fishing result in a lack of colinearity between two consecutive lifestages. Indices of spawning females and egg production were highly correlated, which was expected because calculation of the latter included adult female abundance as part of the methodology of estimation. Significant correlations were also found between the abundances of successive larval stages as well as between Niantic River Stage 4 larval abundance and age-0 juveniles collected in the Niantic River during both early and late summer, although considerable scatter was seen in these relationships (Fig. 35). Early and late summer age-0 juvenile abundances were correlated as were the late summer index and the late fall-early winter abundance. Annual density of larger ( $\geq 7 \mathrm{~mm}$ ) larvae collected in entrainment samples at MPS was significantly correlated with Stage 4 larval abundance, although with considerable scatter (Fig.
36), but was not correlated with other larval, juvenile, or adult abundance indices (Tables 24 and 25). As discussed previously, the abundance of age-1 winter flounder taken during the Niantic River spawning survey in the navigational channel of the lower river was not significantly correlated with that of young taken during the previous fall and early winter in the TMP (Fig. 31). This was probably mostly related to changes in distribution rather than an indication of any compensatory mortality.
Most Niantic River female winter flounder are not fully recruited to the spawning population until about age-4. Thus, abundance of age- 3 spawning females in particular probably was not indicative of year-class strength because only a fraction of these fish were present on the spawning grounds each year. Furthermore, the presence of immature fish in the river may vary from year to year because of environmental conditions. Females of ages- 3 through 5 were significantly positively correlated with age-1 juveniles taken in both the lower and upper river during the spawning season, but age- 4 and 5 fish


Fig. 35. Comparison between the abundance indices of successive winter flounder early life history stages in the Niantic River (varies among year-classes from 1983 through 2001, depending upon data availability), including larval abundance ( $A$ parameter of Gompertz function), age-0 juveniles in the Niantic River (early and late seasonal $1-\mathrm{m}$ beam trawl median CPUE at stations LR and WA combined); and the late fall-early winter seasonal $\Delta$-mean CPUE at all trawl monitoring program (TMP) stations. (Note that the vertical and horizontal scales differ among the graphs).
were negatively correlated with age-0 fish taken during fall and early winter (Table 26). However, the form of these relationships is unclear (Fig. 37; only lower river shown) and because of the scatter seen in the data plots, these relationships may not be very meaningful. Persistence of some negative correlations among abundance indices perhaps results from unknown processes operating after winter flounder become age-1 that produce fewer adult recruits from
more abundant year-classes of juveniles. This was also noted by Simpson et al. (1996) for the 1988 and 1992 year-classes of winter flounder in LIS and by Desfosse et :al. (1998): during recent : years in Massachusetts as abundant numbers of age-1 and 2 fish did not appear as such at age-3 and older. Suggested explanations include variable discard mortality of juveniles in the commercial fishery; high rates of fishing at recruitment to the fisheries


Fig. 36. Comparison between the abundance of Stage 4 winter flounder larvae taken in the Niantic River ( $A$ parameter of Gompertz function; 1983-2001) and larvae $\geq 7$ mm taken at station EN (MPS entrainment sampling). (Note that the horizontal scales differ between the graphs).
(Simpson et al. 1996; Desfosse et al. 1998); and nonrandom fishing effort, which may occur in overfished stocks. Meanwhile, until explanatory factors are identified, none of these life-stage indices can presently be used as a reliable measure of future year-class strength, a discussion of which is given further below in a section discussing the formation of winter flounder year-class strength.

Stock-Recruitment Relationship (SRR)
An SRR described by Ricker $(1954,1975)$ provides the basis for calculating survival from egg to immature fish during the first year of the life-cycle, which has been used with a population dynamics simulation model of Niantic River winter flounder, last presented in NUSCO (2000). The stock and recruitment data for determining the SRR form were derived from the catch-at-age of female winter flounder taken during the Niantic River spawning survey. Because the spawning stock is made up of many year-classes, the true recruitment consists of the total reproductive contribution over the life of each individual in a given year-class (Garrod and Jones 1974; Cushing and Horwood 1977). Therefore, the index of annual parental stock size was based on derived egg production and the index of recruits or year-class size was based on calculated egg production accumulated over the life-time of the recruits. This method accounted for variations in year-class strength and in fecundity by size and age. The assumptions and methods used to age Niantic River winter flounder and to calculate a recruitment index expressed as equivalent numbers of spawning females were described in detail in NUSCO (1989, 1990) and are summarized below.

TABLE 26. Matrix of Spearman's rank-order correlations among various winter flounder larval and Niantic River female spawner abundance indices.

| Index ${ }^{\text {a }}$ | Larvae ( 27 mm ) at MPS discharge | Niantic River early summer age-0 juveniles | Niantic River late summer age0 juveniles | Fall-early winter river-bay age-0 juveniles | Niantic River winter-spring age-1 juveniles (lower river) | Niantic River winter-spring age-1 juveniles (upper river) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Age-3 female spawners | $\begin{gathered} -0.0464^{c} \\ 0.8333 \mathrm{NS} \\ 23 \end{gathered}$ | $\begin{gathered} -0.2220 \\ 0.4456 \mathrm{NS} \\ 14 \end{gathered}$ | $\begin{gathered} -0.0571 \\ 0.8397 \mathrm{NS} \\ 15 \end{gathered}$ | $\begin{gathered} -0.3451 \\ 0.1068 \mathrm{NS} \\ 23 \end{gathered}$ | $\begin{gathered} 0.8031 \\ 0.0001 * * \\ 23 \end{gathered}$ | $\begin{gathered} 0.8193 \\ 0.0001 * * \\ 19 \end{gathered}$ |
| Age-4 <br> female spawners ${ }^{\text {b }}$ | $\begin{gathered} -0.0616 \\ 0.7856 \mathrm{NS} \\ 22 \end{gathered}$ | $\begin{gathered} -0.1429 \\ 0.6415 \mathrm{NS} \\ 13 \end{gathered}$ | $\begin{gathered} -0.2308 \\ 0.4273 \mathrm{NS} \\ 14 \end{gathered}$ | $\begin{gathered} -0.4436 \\ 0.0386^{*} \\ 22 \end{gathered}$ | $\begin{gathered} 0.8037 \\ 0.0001 \text { ** } \\ 22 \end{gathered}$ | $\begin{gathered} 0.7668 \\ 0.0002 * * \\ 18 \end{gathered}$ |
| Age-5 <br> female spawners ${ }^{\text {b }}$ | $\begin{gathered} 0.3429 \\ 0.1281 \mathrm{NS} \\ 21 \end{gathered}$ | $\begin{gathered} -0.1539 \\ 0.6331 \mathrm{NS} \\ 12 \end{gathered}$ | $\begin{gathered} -0.1429 \\ 0.6415 \mathrm{NS} \\ 13 \end{gathered}$ | $\begin{gathered} -0.5726 \\ 0.0067 * * \\ 21 \end{gathered}$ | $\begin{gathered} 0.6976 \\ 0.0004^{* *} \\ 21 \end{gathered}$ | $\begin{gathered} 0.6838 \\ 0.0025 * * \\ 17 \end{gathered}$ |

[^19]

Fig. 37. Comparisons between the late fall-early winter seasonal $\triangle$-mean CPUE of age-0 winter flounder taken at trawl monitoring program stations (TMP) and the $\Delta$-mean CPUE of age-1 winter flounder taken in the lower Niantic River (stations 1 and 2) during the adult winter flounder surveys (WFS), and the relative annual abundance of age-3, (1976-98 year-classes), age-4 (1976-97 year-classes), and age-5 (1976-96 year-classes) Niantic River female winter flounder taken during the spawning season. (Note that the vertical scales differ among the graphs).

Stock and Recruitment Indices. Methods used to calculate the annual standardized catch index and total egg production of the parental stock were given previously (see Adult Spawning Stock Size and Egg Production). A recruitment index was determined by applying an age-length key described in NUSCO (2000) to the annual standardized catches of females partitioned into length categories. Aging females allowed for the determination of abundance by yearclass present in each age-class during successive spawning seasons. A common age-length key was
used over all years because Witherell and Burnett (1993) reported that no trends were observed in mean leng'th-at-age during '1983-91' for". Massachusetts winter flounder despite a $50 \%$ reduction in biomass during that period.
From observations made of abundance and age over the years, a large fraction of age- 3 females and some age- 4 fish were apparently immature and not present in the Niantic River during the spawning season; all females age- 5 and older were assumed to be mature. The total number of females was reduced
to spawning females using length-specific proportions of mature fish estimated from annual catches in the Niantic River (Fig. 10). Because the estimates of age-3 fish present in the Niantic River were thought to be unreliable as a large fraction of them were immature and not present on the spawning grounds, this estimation process was only carried through the 1997 year-class (i.e., age-4 females taken in 2001). The adjusted numbers of mature fish provided an index of the fully recruited year-class expressed as the aggregated number of adult females passing through each age-class. An implied assumption was that catches in the Niantic River were representative of the population, with the exception of immature fish, most of which did not enter the river until fully recruited to the spawning stock. Although this recruitment index could be used together with the annual number of adult females to derive the SRR, this would ignore size composition differences that affected annual egg production. Therefore, the above index was adjusted for differences in fecundity among fish using the length-fecundity relationship of Niantic River winter flounder (Eq. 1). Also, since the recruitment index was based on total lifetime contribution of a year-class, estimates from more recent year-classes needed to be projected into the future. Therefore, an annual survival rate of 0.39 , which is equivalent to a total instantaneous total mortality rate ( Z ) of 0.94 (the current estimated instantaneous fishing mortality rate, F , for LIS winter flounder of 0.74 plus 0.2 for M ), was applied to generate estimates of year-class egg production through 2009 (i.e., age-12 fish from the 1997 yearclass). Finally, annual egg production was summed up over the lifetime of each year-class to determine a recruitment index as eggs, which was then converted to equivalent spawning females at the rate of one female for each 583,775 eggs, which is the calculated value of current mean fecundity for the Niantic River stock.
Stock and Recruitment Parameters. The Ricker SRR appeared best suited for use with the Niantic River winter flounder stock because the relationship between recruitment and spawning stock indices was a dome-shaped curve with substantial decline in recruitment when the stock was larger than average (NUSCO 1989). This particular form of a SRR has also been applied to other New England flounder stocks (Gibson 1989, 1993). Furthermore, Iles (1994) stated that a dome-shaped stock-recruitment relationship appeared to be generally consistent for a number of flatfish stocks and Brodziak et al. (2001) reported that Southern New England winter flounder
demonstrated a density-dependent form of a SRR. The mathematical form of the Ricker SRR is:

$$
\begin{equation*}
R_{t}=\alpha \times P_{t} \times \exp \left(-\beta \times P_{t}\right) \tag{6}
\end{equation*}
$$

where $R_{t}$ is the recruitment index for the progeny of the spawning stock $P_{t}$ in year $t$ and $\alpha$ and $\beta$ are parameters estimated from the data. The $\alpha$ parameter describes the growth potential of the stock and $\log _{e}(\alpha)$, the slope of the $\operatorname{SRR}$ at the origin, is equivalent to the intrinsic natural rate of increase (Roughgarden 1979) when the stock is not exploited. The $\beta$ parameter is the instantaneous rate at which recruitment declines at large stock sizes due to some form of density-dependent mortality. The natural logarithm of winter flounder recruitment was found to be correlated with mean water temperature during February at the intakes of MPS, which is when most spawning and early larval development occurs (NUSCO 1988b, 1989). Temperature and climatic effects are also discussed in the following section on the formation of winter flounder year-class strength. Using nonlinear regression methods (SAS Institute Inc. 1990 d ), the parameters $\alpha$ and $\beta$ were estimated initially by fitting Equation 6 to the data and then re-estimated under the assumption that there was a significant temperature effect. Following Lorda and Crecco (1987), Gibson (1987, 1993), Iles and Beverton (1998), Quinn and Deriso (1999: p. 91), and Planque and Frèdou (1999), an annual mean water temperature (here, mean February temperature, the rationale for which is discussed in the following section) was used as an explanatory variable added to Equation 6 to adjust the two-parameter SRR for temperature effects. This served to reduce recruitment variability and to obtain more reliable parameter estimates for the SRR. The tempera-ture-dependent SRR had the form:

$$
\begin{equation*}
R_{t}=\alpha \times P_{t} \times \exp \left(-\beta \times P_{t}\right) \times \exp \left(\phi \times T_{F c b}\right) \tag{7}
\end{equation*}
$$

where the second exponential describes the effect of February water temperature on recruitment and the added parameter $\phi$ represents the strength of that effect. This effect either decreases or increases the number of recruits-per-spawner produced each year because temperature was defined as the deviation ( $\mathrm{T}_{\mathrm{Fb}}$ ) of each particular mean February temperature from a long-term (1977-97) average of February water temperatures. When the February mean water temperature is equal to the long-term average, the deviation ( $\mathrm{T}_{\mathrm{Feb}}$ ) in Equation 7 becomes zero and the exponential term equals unity (i.e., no temperature
effect). Thus, Equation 7 reduces to its initial form (Eq. 6) under average temperature conditions.
Additionally, a depensatory form of SRR was investigated. Coincidentally, Walters and Kitchell (2001) recently emphasized that the risk of depensatory effects should be a goal of recruitment research. When depensation processes occur, the per capita recruitment rate decreases with decreasing parental stock size, which may lead to a rapid stock collapse. The two-parameter Ricker SRR (Eq. 6) was modified with the addition of another parameter, $\Omega$, as suggested by Saila and Lorda (1982). The depensatory version of the Ricker SRR has the form:

$$
\begin{equation*}
\mathrm{R}_{t}=\alpha \times\left(\mathrm{P}_{\ell}\right)^{\Omega} \times \exp \left(-\beta \times \mathrm{P}_{\ell}\right) \tag{8}
\end{equation*}
$$

where the parameter $\Omega$ must be greater than zero. When $\Omega>1$, the SRR can describe both depensation below some threshold stock size and compensation for larger stock sizes, as in the two-parameter SRR (Eq. 6 ). When $\Omega=1$, the above equation reverts to the standard two-parameter model.
Nonlinear regression methods (SAS Institute Inc. 1990d) were used for estimating the parameters in the above equations. The value of $\alpha$ as estimated from these stock and recruitment data was not used in the SPDM, which uses a value, termed $\alpha_{0}$, that was calculated from life history parameters, the derivation of which will be discussed below.
Biological Reference Points. The fishing mortality rate ( F ) is an important factor affecting the growth potential of the stock (Goodyear 1977) and, thus, is relevant for assessing other impacts. Because fishing and natural mortality ( $M=0.2$; total instantaneous mortality rate $Z=F+M$ ) of winter flounder take place concurrently through the year, the actual fraction of the stock removed by the fishery each year (i.e., the exploitation rate $u$ ) is obtained as:

$$
\begin{equation*}
u=(F / Z) \times(1-\exp [-Z]) \tag{9}
\end{equation*}
$$

Stock-recruitment theory and the interpretation of several biological reference points derived from Ricker's SRR model were discussed in detail in NUSCO (1989). The equilibrium or sustainable stock size of an exploited stock (i.e.; when $F>0$ ) is given by:

$$
\begin{equation*}
P_{E(F)}=\left(\log _{e}[\alpha]-F\right) / \beta \tag{10}
\end{equation*}
$$

Rearranging terms and solving for the rate of fishing that would achieve a given equilibrium stock size results in:

$$
\begin{equation*}
F=\log _{e}(\alpha)-\left(\beta \times P_{E(f)}\right) \tag{11}
\end{equation*}
$$

When $\mathrm{F}=0$, Equation 10 becomes the equilibrium or replacement level of the unfished stock:

$$
\begin{equation*}
P_{\text {rep }}=\left(\log _{e}[\alpha]\right) / \beta \tag{12}
\end{equation*}
$$

The fishing rate for "recruitment overfishing" was recently defined for winter flounder stocks as the rate of fishing that reduces the spawning stock biomass to less than $25 \%$ of the stock for maximum spawning potential (Howell et al. 1992). This concept is discussed in more detail in the section entitled Stochastic Simulation of the Niantic River Winter Flounder Stock.
Although Equations 10 through 12 can be used to calculate equilibrium stock sizes and fishing rates for the winter flounder, the results are only deterministic approximations that ignore age-structure effects. Therefore, these equations are primarily useful to calculate initial values of the corresponding biological reference points. These are better estimated through simulations using the SPDM or other similar population or production models that include age structure and both $M$ and $F$.
Sampling-based Estimates. Egg production estimated from annual spawning surveys was the basis for determining recruitment, because the abundance of other 'early life-stages have not been reliably correlated with adult winter flounder. Both parental spawning stock size (previously described in Absolute Abundance Estimates) and recruitment indices were scaled to absolute population size using annual estimates of abundance; size frequency, maturity, egg production, and overall population mean fecundity. . These scaled annual values were used with the Ricker SRR model as estimates of adult female spawning stock and : potential female recruitment (Table:27). The addition of new catch data from the 2001 adult winter flounder survey resulted in some differences among current estimates of spawning females and recruits and those previously reported since an SRR was first described in the 1989 annual report. Annual female recruitment has decreased substantially from the 1970-80s and, with a few exceptions, February water temperatures have been warmer than the long-term average.: 'The long-term trend in February mean water temperature :was significantly positive from 1977 through 2001 (Mann-Kendall test, $Z=2.76$; $n$ $=25 ; p=0.006$ ):

A two-parameter SRR model (Eq. 6) was initially fitted to the data. . The stock growth potential parameter $\alpha$ (scaled as numbers of fish). for this

TABLE 27. Annual Niantic River winter flounder stock-recruitment data based on indices of egg production in the 1977 through 1997 yearclasses with mean February water temperature and deviations ( $\mathrm{T}_{\mathrm{Fe}}$ ) from the mean.


2 Scaled number of female spawners and recruits from expected egg production; scaling factors used were 583,775 eggs per female (mean fecundity) and a multiplier of 29.5 to convert relative abundance to an absolute population size. Indices of female spawners and recruits differ from those reported in DNC (2001a) because of data added from the 2001 adult winter flounder population survey.
model was estimated as 1.056 with a standard error of 0.334 ( $32 \%$ of the parameter value). This estimate of $\alpha$ and the estimate of $\beta$ (the second model parameter) were used as initial values for fitting the three-parameter SRR model with temperature effects (Eq. 7) and the depensatory model (Eq. 8).
Fitting the depensatory model to the same data resulted in a parameter estimate for $\Omega$ that was not significantly different from 1 . Therefore, a depensatory form of the SRR was not indicated for the Niantic River winter flounder population. Myers et al. (1995) examined spawner-recruit data for 128 fish stocks and reported that evidence for depensation was only indicated in three populations. They concluded that most observed fish population collapses could not be reasonably attributed to depensatory fish population dynamics. In another meta-analysis, Liermann and Hilborn (1997) found that no depensation was likely for fishes of four broad taxonomic groups, including pleuronectid flounders, but cautioned that both depensation and hypercompensation were possible for some stocks, given the broad range of their data distribution. In a simulation analysis, Frank and Brickman (2000)
found that a SRR can appear to be compensatory even if the stock reproductive dynamics did not exhibit this behavior. They further noted, however, that species exhibiting social behavior, such as group mating or schooling were at most risk for depensation. Winter flounder do not exhibit these traits, which are mostly found in pelagic fishes. Also, spawning in inshore estuaries may protect winter flounder from fisheries directed specifically at spawning aggregations. However, fish moving to or from the spawning grounds are targeted by trawlers in fall and winter and spring, including within Niantic Bay. Furthermore, the Niantic River has had a unique night spear fishery for winter flounder that occurs during the spawning and immediate postspawning periods, which can be considered as a source of increased fishing mortality specific to this stock. In recent years, fishing mortality has apparently remained relatively high on LIS winter flounder (Johnson et al. 2000).
For the three-parameter SRR model with temperature effects, all parameter estimates were significantly different from 0 and the model explained $72 \%$ of the variability associated with the
recruitment index. Relationships resulting from fitting both the two- and three-parameter Ricker models separately to stock and recruitment data scaled to absolute population sizes are shown as the curved lines in the central portion of Figure 38 as follows: the unadjusted SRR (two-parameter model; Eq. 6) is shown as the more widely-spaced dashed line and the three-parameter model (SRR adjusted for $T_{\text {feb }}$; Eq. 7) is represented by the solid line. The outermost two more finely-spaced dashed lines illustrate low recruitment in the warmest year (Table 27; 1991, $\mathrm{T}_{\mathrm{Feb}}=+1.95$ ) and high recruitment in the coldest year (1977, $\mathrm{T}_{\mathrm{Feb}}=-2.45$ ). A similar plot, showing the effect of a range in water temperatures on the SRR of Irish Sea Atlantic cod (Gadus morhua), was shown by Planque and Frèdou (1999).

Using the three-parameter model, the current estimate of $\alpha$ was determined as 1.095 and had a standard error of 0.241 , which is $22 \%$ of the parameter value (Table 28). Differences among annual values of $\alpha$ seen in the time-series of estimates were likely caused by increased fishing mortality on winter flounder in addition to the inherent instability of parameter estimates fitted to small data sets. In particular, the apparent influence of the 1988-97 data points on the estimate of $\alpha$ were illustrative of higher recent exploitation and poor
recruitment. The Niantic River winter flounder population apparently now has a greatly diminished compensatory reserve in response to rates of fishing that increased from 0.71 in 1985 to $1.10-1.33$ in the mid-1990s. A recent fishing mortality rate as high as 1.02 was still reported for LIS winter flounder by Johnson et al. (2000). Relatively high abundance of juvenile winter' flounder from the 1988 year-class was expected to result in increased numbers of adult fish during 1992-94 that rwould dominate the spawning population. Unfortunately, winter flounder from these large year-classes wëre removed quickly by fishing (Simpson et al. 1996). Also, the apparent lack :of adult fish in Connecticut waters in more recent years, even though juvenile fish of the 1992 year-class appeared to be abundant, was noted by Simpson et al. (1996) as well as in the Niantic River studies. Increases in adult abundance resulting from relatively numerous age- 0 juveniles seen in the past several years were expected as these fish become mature, but this has not materialized. Other factors may have substantially reduced their numbers prior to adult recruitment, as was seen for the 1988 and 1992 year-classes.
The estimate of Ricker's $\beta$ parameter, which describes the annual rate of compensatory mortality as a function of the stock size, is an important factor


Fig. 38. Ricker SRRs of Niantic River winter flounder (see text for explanation of the four curves plotted). Calculated recruitment indices (see Table 32) of the 1977 through the 1997 year-classes are shown.

TABLE 28. Annual estimates of the modified Ricker stock-recruitment function (Eq. 7) determined for the Niantic River winter flounder population from 1989 through 2001.

| Year of estimation | Year-classes included ${ }^{\text {A }}$ | $\mathbf{a}^{\text {b }}$ | Standard error | $\begin{gathered} \beta \\ \left(\times 10^{-5}\right) \end{gathered}$ | $\begin{gathered} \hline \text { Standard } \\ \text { error } \\ \left(\times 10^{-5}\right) \end{gathered}$ | ¢ | Standard error |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1989 | 1977.85 | 2.646 | 0.599 | 2.228 | 0.456 | -0.259 | 0.095 |
| 1990 | 1977-86 | 2.502 | 0.399 | 2.466 | 0.372 | -0.264 | 0.064 |
| 1991 | 1977 -87 | 2.226 | 0.518 | 2.140 | 0.461 | -0.329 | 0.098 |
| 1992 | 1977 -88 | 2.149 | 0.543 | 2.466 | 0.567 | -0.357 | 0.010 |
| 1993 | 1977 -89 | 1.977 | 0.566 | 2.523 | 0.642 | -0.412 | 0.108 |
| 1994 | 1977-90 | 2.071 | 0.428 | 2.498 | 0.478 | -0.379 | 0.077 |
| 1995 | 1977-91 | 1.710 | 0.380 | 2.583 | 0.516 | -0.415 | 0.078 |
| 1996 | 1977-92 | 1.473 | 0.306 | 2.450 | 0.488 | -0.418 | 0.075 |
| 1997 | 1977.93 | 1.442 | 0.283 | 2.399 | 0.463 | -0.417 | 0.072 |
| 1998 | 1977-94 | 1.186 | 0.265 | 1.961 | 0.513 | -0.381 | 0.082 |
| 1999 | 1977-95 | 1.082 | 0.265 | 2.071 | 0.570 | -0.408 | 0.088 |
| 2000 | 1977-96 | 1.125 | 0.252 | 1.886 | 0.478 | -0.400 | 0.082 |
| 2001 | 1977-97 | 1.095 | 0.241 | 1.877 | 0.472 | -0.404 | 0.081 |
|  | Mean | - |  | 2.273 |  | -0.373 |  |

- Age-4 considered to be minimum age of recruitment.
b The compensatory reserve for an unfished stock ( $\alpha_{0}$ ) used in the SPDM is 5.20 (see NUSCO 2000 for the derivation of this value).
in SPDM simulations. The present value for $\beta$ is $1.877 \times 10^{-5}$, the lowest estimate of the series (Table 28). Previous values ranged between 1.886 and $2.583 \times 10^{-5}$. The current long-term (1989-2001) mean estimate of $2.273 \times 10^{-5}$ for $\beta$ was less than the average found during the mid-1990s because of four consecutive low estimates, suggesting a possible weakened capability to compensate. However, the present calculation does not include the contribution of year-classes after 1997. These year-classes had increasing numbers of juveniles found in the Niantic River (see Juvenile Winter, which should result in better recruitment if mortality later in the juvenile stage (ages-1 through 3) is not excessive. However, the contribution of these juveniles to future adult stock abundance remains to be seen.
The parameter $\phi$, which reflects the effect of February temperature deviations ( $\mathrm{T}_{\mathrm{Fet}}$ ) from the 1977-97 mean of $2.81^{\circ} \mathrm{C}$ (Table 27), has also been used as an SPDM input parameter and was estimated as -0.404 in 2001, a value nearly the same as found in both 1999 and 2000 (Table 28). The long-term mean for $\phi$ was -0.373 (range of -0.418 to -0.259 ). As noted above, February water temperatures have been generally warmer in recent years and estimates for the $\phi$ parameter have increased in magnitude. The effect of February temperatures on winter flounder recruitment is discussed in the following section.
Ricker's stock-at-replacement ( $\mathrm{P}_{\text {rep }}$; Eq. 12), a derived biological reference point, was estimated using the SRR parameter estimates as 72,532 female
spawners. This is the unfished equilibrium spawning stock size, the biomass for which is often referred to as maximum spawning potential (MSP). This reference point was calculated as $119,678 \mathrm{lbs}$ for Niantic River winter flounder (i.e., 1.65 lbs per spawner; see Table 30, below). Stocks with biomass less than the critical size of $25 \%$ of MSP (in this case, $29,919 \mathrm{lbs}$ ) are considered to be overfished (Howell et al. 1992).
Estimation of $\alpha_{0}$ for SPDM Simulations. The stock-recruitment-based estimates of $\alpha$ for the Niantic River winter flounder discussed above underestimated the true slope at the origin of this stock. The method of calculating annual recruitment included the effects of fishing on winter flounder age- 2 and older as well as the entrainment of larvae at MPS. Therefore, these direct estimates of $\alpha$ correspond to a compensatory reserve diminished by existing larval entrainment and exploitation rates. The concept of a compensatory reserve in fishing stocks and the effect of exploitation on the shape of the reproduction curve when the recruitment index is based on the exploited stock was discussed by Goodyear (1977: Fig. 1). Thus, if larval entrainment and fishing rates increase, the field estimates of recruitment will be smaller and so will the estimates of $\alpha$ (i.e., the remaining compensatory reserve). To assess impacts appropriately, the inherent potential of a stock to increase in the absence of fishing and plant effects must be determined. Crecco and Howell (1990) investigated the possibility of using indirect methods to estimate the true $\alpha$ parameter (i.e., $\alpha_{0}$ for
the unfished stock when $F=0$ ). They used four indirect methods (Cushing 1971; Cushing änd Harris 1973; Longhurst 1983; Hoenig et al. 1987; Boudreau and Dickie 1989) based on different life history parameters (Table 29). Because these methods did not depend upon direct estimates of recruitment, biases caused by changing fishing rates are avoided and independent means of validating SRR-based estimates are provided. The geometric mean of $\alpha_{0}=$ 5.20 calculated from these estimates was used in the SPDM: This parameter describes the inherent potential of a stock to increase because the natural logarithm of $\alpha$ is the slope of the SRR at the origin for the unfished stock (Ricker 1954) and that slope, in turn, corresponds to the intrinsic rate of natural increase of the population (Roughgarden 1979). Consequently, the large difference between the derived value of $\alpha_{0}(5.20)$ and regression estimates of $\alpha$ based on field data reflects the difference in potential growth between unfished and highly exploited stocks of winter flounder. Use of an unfished stock as a starting point for a population dynamics simulation has a number of advantages, depending upon the particular scenario selected. The data-based estimates of the other two SRR parameters ( $\beta_{\text {, }}$ and $\phi$ in Table 28) used in the population simulations, however, do not depend upon fishing and entrainment rates, and since they are not directly related to current winter flounder abundance, their estimates were obtained as long-term averages of their series.
Finally, both the mean weight and fecundity of a Niantic River female winter flounder was calculated for a theoretical population of ages 1 through 15 at
equilibrium for which only $M$ was assumed (i.e., the unfished population). These calculations used population data previously reported (NUSCO 1990, 2000) and an estimated $M$ of 0.2.'. The equilibrium calculation for this theoretical unfished stock showed a mean weight of 1.65 lbs per female and a mean fecundity of $1,322,994$ eggs per spawner (Table 30).


## Formation of Winter Flounder Year-Class Strength

Year-class strengths are likely determined during larval and early juvenile life stages of marine fishes, yet these phases of life history are least understood (Sissenwine 1984; Bailey and Houde 1989; Bradford 1992). Accordingly, formation of winter flounder year-class strength begins at egg deposition, or even before, given some known maternal influences on egg quality. For example, a study by Buckley et al. (1991) noted that female size and time of spawning affected various winter flounder reproductive parameters, including egg size, fecundity, and viability. They observed that eggs deposited earlier in the season by larger females appeared to have better survival than eggs produced by smaller fish late in the season. Biological consequences of the selective removal of large fish by fishing can be more profound than just reducing egg production because of the aspects of egg size and time of spawning relating to reproductive success (Trippel 1995; Conover 2000).
: Compared to the , winter flounder larval developmental period, less is known about the egg

TABLE 29. Methods of estimating the compensatory reserve parameter $\alpha_{0}$ of the untished Niantic River winter flounder based on several life history models (modified from Table 4 in Creceo and Howell 1990).

$2 \cdot r_{m}=$ annual intrinsic rate of population increase; weight is weight in Kcal at which $50 \%$ of female winter flounder first spawn; $t_{m}=$ mean generation time in days; $K$ and $L_{\infty}$ are parameters of the von Bertalanffy growth equation and $L_{m}$ is mean length; and FEC $=$ mean fecundity.
b $\alpha_{n}=\exp \left(r_{m} \times t_{m}\right)$, where $t_{m}=4.5$ years (mean time to maturation for females).

TABLE 30. Biomass calculations of the Niantic River female winter flounder spawning stock at equilibrium, based on an instantaneous natural mortality rate of $M=0.2$ and an instantaneous fishing mortality rate of $\mathrm{F}=0$ (i.e., an unfished stock).

| Age | $\begin{gathered} \text { Female } \\ \text { population } \\ \text { size } \end{gathered}$ | Fraction mature | Number of mature females | Mean length (cm) | Mean weight of mature females (lbs) | Eggs per mature female | Spawning stock biomass (lbs) | Egg production (millions) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2 | 1,000.00 | 0.00 | 0.00 | 18.0 | - | - |  | 0.000 |
| 3 | 818.73 | 0.25 | 204.68 | 27.0 | 0.497 | 232,088 | 101.77 | 47.504 |
| 4 | 670.32 | 0.80 | 536.26 | 31.0 | 0.776 | 432,517 | 416.36 | 231.940 |
| 5 | 548.81 | 1.00 | 548.81 | 34.5 | 1.096 | 700,390 | 601.73 | 384.381 |
| 6 | 449.33 | 1.00 | 449.33 | 37.5 | 1.435 | 1,019,793 | 644.71 | 458.224 |
| 7 | 367.88 | 1.00 | 367.88 | 39.0 | 1.628 | 1,216,926 | 599.04 | 447.683 |
| 8 | 301.19 | 1.00 | 301.19 | 40.5 | 1.839 | 1,442,512 | 553.95 | 434.470 |
| 9 | 246.60 | 1.00 | 246.60 | 42.0 | 2.068 | 1,699,372 | 510.01 | 419.065 |
| 10 | 201.90 | 1.00 | 201.90 | 43.5 | 2.316 | 1,990,489 | 467.61 | 401.880 |
| 11 | 165.30 | 1.00 | 165.30 | 45.0 | 2.584 | 2,319,011 | 427.09 | 383.332 |
| 12 | 135.34 | 1.00 | 135.34 | 46.5 | 2.872 | 2,688,253 | 388.70 | 363.828 |
| 13 | 110.80 | 1.00 | 110.80 | 48.0 | 3.182 | 3,101,703 | 352.54 | 343.669 |
| 14 | 90.72 | 1.00 | 90.72 | 49.5 | 3.514 | 3,563,020 | 318.78 | 323.237 |
| 15 | 74.27 | 1.00 | 74.27 | 51.0 | 3.869 | 4,076,040 | 287.36 | 302.727 |
| Total | 5,181.19 |  | 3,433.08 |  |  |  | 5,669.66 | 4,541.941 |
| Mean weight per mature female fish Mean fecundity (unfished stock) |  |  | $\begin{aligned} & (5,670 \mathrm{lbs}+3,433 \text { mature females }) \\ & 1,322,994 \text { eggs per female spawner } \end{aligned}=1.65 \mathrm{lbs}(-39.2 \mathrm{cr}$ |  |  |  |  |  |

stage. Environmental factors have important implications on egg survival and hatching (Bunn et al. 2000). Keller and Klein-MacPhee (2000) reported that in a mesocosm study completed in Rhode Island, winter flounder egg survival, percent hatch, time to hatch, and initial size were greater in cooler than warmer systems. Morrison et al. (1991) reported high mortality of demersal Atlantic herring (Clupea harengus) eggs in the Firth of Clyde, Scotland because of heavy deposition of organic matter resulting from a bloom of a diatom, Skeletonema costatum. The decomposing material caused a depletion of oxygen and egg death due to anoxia. This diatom was one of the most abundant of the phytoplankton collected at MPS during entrainment sampling from 1977 through 1980 (NUSCO 1981). However, highest densities occurred in summer, after the winter flounder egg incubation period. Skeletonema costatum was also a dominant form in the warm (about $1^{\circ} \mathrm{C}$ above current ambient mean water temperature) treatment in the mesocosm study of Keller and Klein-MacPhee (2000). Macroalgal and detrital mass varied considerably among years of study in the Niantic River since the mid-1970s and the amount of organic material on the river bottom may have had some influence on the survival of demersal winter flounder eggs.
Based on a comparison of estimates of egg production and abundance of Stage 1 larvae (see Larval Winter Flounder, Abundance and Distribution: Fig. 16), egg mortality from unknown causes
may be considerable in the Niantic River, but may also vary in a density-dependent fashion in some years. This apparent compensatory effect resulted in egg survival that could have been as much as four times higher during a period including most recent years (1995-99 and 2001), when adult abundance was particularly low, than in most previous years or in 2000. The mechanisms responsible for greater winter flounder egg survival were not studied, but hypotheses concerning this effect include less predation pressure because of fewer cues for predators under low egg abundance or better egg quality because of greater food resources shared among fewer adult females. Also, Buckley et al. (1991) suggested greater relative fecundity and viability of eggs produced by a spawning stock dominated by larger individuals. As noted previously in Adult Winter Flounder - Relative Annual Abundance, larger females have made up an increasing proportion of spawners in recent years.
Regarding predation on eggs, DeBlois and Leggett (1991) and Frank and Leggett (1984) found that both an amphipod (Calliopius laeviusculus) and winter flounder both preyed heavily upon demersal capelin (Mallotus villosus) eggs, removing up to $39 \%$ and $5 \%$ of the production, respectively. They suggested that invertebrate predation on demersal fish eggs may be an important regulatory mechanism for population size in marine fishes having demersal eggs. Because demersal fish eggs are immobile and found in relatively dense patches, they may attract many
predators (Bunn et al. 2000). In particular, the sevenspine bay shrimp has been observed feeding on winter flounder egg clusters in laboratory tanks (D. Taylor, University of Rhode Island, Graduate School of Oceanography, Narragansett, RI, pers. comm.)."
Larval mortality can vary considerably from year to year and can have profound effects on the number of young that metamorphose and settle. Larval winter flounder mortality was found to be related to both Jarval density and April water temperatures (see Larval Winter Flounder, Mortality: Fig. 24). Thus, when egg abundance was particularly low, as during 1995-99 and 2001, enhanced egg survival "or hatchability was apparent. Relatively high abundance of older larvae the past 2 years suggested that larval survival was also particularly good, perhaps reflecting a compensatory response related to initially low larval densities.
Intrinsic (e.g., growth) and extrinsic (e.g., predation) factors both affect survival. Several workers (Laurence 1977; Chambers et al. 1988) found that much of larval winter flounder mortality was concentrated in early life and at first feeding, a few weeks after hatching. Hjorleifsson (1992) showed that the ratio between RNA and DNA, an index of condition and growth rate, was lowest at the time of first feeding of winter flounder (about 4 mm ) and that these ratios were affected by food availability. Therefore, the strength of a year-class could be determined by the availability of sufficient food after completion of yolk absorption. However, Keller and Klein-MacPhee (2000) noted in a mesocosm study that food availability was not the most important factor related to larval mortality 'as mortality was highest in their warm treatment, which also had the highest food availability. Daily mortality rates increased directly with the abundance of active predators in the mesocosms, but this may have been an effect of the relatively low numbers of predators in their experimental system. They further noted that daily growth and mortality rates were significantly and inversely related.
Predation is often one of the most important causes of larval fish mortality (Bailey and Houde 1989) and this has relevance to winter flounder larval populations. The relationship between winter flounder larvae and their predators was studied by Williams and Brown (1992), who found that escape response increased with increasing larval size, but that it remained slower than that of other larval fishes examined. Larval winter flounder are likely vulnerable to both fish and invertebrate predators. Although susceptible to attacks by planktivorous fishes, the temporal occurrence and abundance of
fishes that could potentially prey on larval winter flounder are low, particularly during the early portion of the larval winter flounder season.
Most predation on winter flounder larvae is probably by invertebrate contact predators, such as anemones (Keller and Klein-MacPhee 2000), other cnidarians, and ctenophores, many of which have been previously identified as feeding on flatish larvae. Evidence of a causal predator-prey relationship on larvae of European plaice and European flounder by a scyphomedusañ (Aurelia aurita) and a ctenophore (Pleurobrachia pileus) was reported by Van der Veer (1985). However, most predation by these species occurred at the end of the larval European plaice season and likely did not ultimately affect year-class strength (Van der Veer 1985; Van der Veer et al. 1990). Laboratory studies showed that successful capture of European plaice larvae increased as medusal size of $A$. aurita increased (Bailey and Batty 1984). Pearcy (1962) stated that Sarsia tubulosa medusae were important predators of larval winter flounder in the Mystic River and had greatest impact on younger, less mobile larvae.
Crawford and Carey (1985) reported large numbers of the moon jelly (A. aurita) in Point Judith Pond, RI and believed that they were a significant predator of larval winter flounder. Potential predators of winter flounder larvae in the Niantic River were medusae of the lion's mane jellyfish (Cyanea sp.), which can be abundant in the upper river (Marshall and Hicks 1962), including station A. A laboratory study showed that winter flounder larvae contacting the tentacles of the lion's mane jellyfish were stunned and ultimately died, even if not consumed by the medusa (NUSCO 1988b). Also, in 1997 ctenophores, another larval fish predator, were present during the larval winter flounder season (NUSCO 1998). However, the relationship between larval winter flounder and jellyfish biomass did not support a cause and effect relationship and the role that these and other planktonic predators have each year on ultimate year-class strength is yet unknown.
' Physical, hydrodynamic processes occurring during the larval stage and settlement of juveniles may also affect the success of a year-class (Werner et al. 1997). This can occur by determining their rates of éxposure to predation and food resources and location of settlement. Many winter flounder larvae are flushed from natal estuaries and transported by water currents within LIS. Variability may also be imposed on the transport of larvae by winds and work on other flatfishes is illustrative of these effects. Densities of settled juveniles of the brown sole (Pleuronectes herzensteini) in Japan were related to
the frequency of onshore wind events (Nakata et al. 2000). Variable rates of transport in LIS can affect not only the availability of winter flounder larvae for entrainment at MPS, but likely also where larvae are able to settle as juveniles. Winter flounder, European plaice, and stone flounder (Platichthys bicoloratus) larvae each use selective tidal transport to enter preferred coastal nursery areas (Rijnsdorp et al. 1985; Yamashita et al. 1996a; Chant et al. 2000). As illustrated by a secondary peak in abundance of older larvae during the later stages of development, winter flounder larvae also appear to be entering the Niantic River. Nearly all pre-recruits are produced in the river and similar nursery habitats. The survival of metamorphosed juveniles settling in Niantic Bay or other deeper areas outside of inshore nursery grounds is poor, probably because of high rates of predation. Similarly, Yamashita et al. (2000) found that small estuaries produced disproportionately more stone flounder juveniles than geographically much larger nursery areas found in shallow, more exposed areas of Sendai Bay, Japan.
For newly settled young of flatfishes, Van der Veer et al. (1990) speculated that, in general, predation by crustaceans may be a common regulatory process, although this may not be a strong effect in all areas (Nash and Geffen 2000). Predation by various caridean shrimps has been suggested as the cause of high mortality after metamorphosis for several flatfishes, including the winter flounder (Witting and Able 1993, 1995), European plaice (Lockwood 1980; Van der Veer and Bergman 1987; Pihl 1990; Van der Veer et al. 1991; Pihl and Van der Veer 1992; Gibson et al. 1995; Van der Veer et al. 2000a), Japanese flounder (Paralichthys olivaceous; Seikai et al. 1993), and stone flounder (Yamashita et al. 1996b). Smaller ( $10-20 \mathrm{~mm}$ ) European plaice were preferentially preyed upon by the brown shrimp (Crangon crangon; Van der Veer and Bergman 1987). Similarly, Witting and Able (1993, 1995) found that the size of age-0 winter flounder significantly affected their probability of predation by sevenspine bay shrimp, with predation greatest at settlement for the smallest fish. Mortality decreased with size and young apparently outgrew predation by shrimp when they reached 17 to 20 mm in length, which meant that fish would have to double in length after settlement before attaining a size refuge from shrimp attacks. Predation was also related to shrimp density and steadily increased until reaching an asymptote at shrimp densities greater than $10.6 \cdot \mathrm{~m}^{-2}$ (Witting and Able 1995).
Other predators of juvenile winter flounder include the green crab (Carcinus maenus; Fairchild and

Howell 2000), larger fishes (e.g., grubby; Myoxocephalus aenaeus) and marine birds (e.g., double-crested cormorants; Phalacrocorax auritus). Predation likely continues from settlement throughout summer in shallow nursery habitats preferred by young winter flounder. Pearcy (1962) suggested that several of these species preyed upon juvenile winter flounder in the Mystic River, CT estuary. Winter flounder and other flatfishes occurred frequently in the diets of cormorants in northern New England and Canada (Pilon et al. 1983; Blackwell et al. 1995; Rail and Chapdelaine 1998) and cormorant predation on winter flounder has been frequently observed in the Niantic River. Winter flounder otoliths have been identified in regurgitated pellets of cormorants nesting in areas near MPS and over 100 active nests alone were found on Waterford Island, just outside the mouth of the Niantic River (M. Male, Old Lyme CT, pers. comm.). Birt et al. (1987) found that close to cormorant nesting colonies, predation depleted fish populations, including those of winter flounder, with higher fish densities observed in areas beyond the foraging range of these birds. Breeding pairs of cormorants have increased steadily since they were first counted as nesting in Connecticut in 1982 with populations increasing by $15 \%$ annually between 1986 and 1998 and by $14 \%$ in coastal New York (Victoria 2001). In Rhode Island, the annual increase from 1981 to 1990 was even greater (63\%). Manderson et al. (2000) reported that summer flounder (Paralichthys dentatus) preyed upon age-0 winter flounder in the Navesink River, NJ with selection increasing as juvenile winter flounder grew from 20 to 90 mm . Striped searobin (Prionotus evolans) prey heavily upon young winter flounder when they co-occur (Manderson et al. 1999). Green crab abundance in the Niantic River increased substantially in recent years, with a non-parametric Mann-Kendall test showing a significant ( $p<0.0001$ ) positive trend in the slope of the annual catch at the trawl monitoring program station in the river since 1976. However, green crab abundance dropped off sharply in 2001 (Fig. 39). Manderson et al. (2000) found that the presence of vegetation (eelgrass, macroalgae) decreased the vulnerability of young winter flounder to predation. Eelgrass beds in the Niantic River have decreased considerably in recent years (DNC 2001b), perhaps increasing the vulnerability of winter flounder to predation.
Another cause of mortality during the first year of life of winter flounder is from infection by the microsporidian parasite Glugea stephani, the severity of which can vary by fish size, density, and


Fig. 39. Total number of green crabs taken each year at TMP station NR from January 1976 through December 2001.
environmental factors such as water temperature (Takvorian and Cali 1981, 1984; Cali et al. 1986; Cali and Takvorian 1991; MacLean 1993). Irregular rates of infection in age- 0 winter flounder can also introduce variation in winter flounder recruitment.
As with larvae, the time duration that settled juveniles spend in a vulnerable size range, which is related to growth rate, affects the vulnerability of young to predators such as the sevenspine bay shrimp or green crab. Variation in growth, which depends upon specific location of settling, specific habitat within a location, temperature, food, or physicochemical parameters (Sogard 1990; Sogard and Able 1992; Bejda et al. 1992; Gibson 1994; Meng et al. 2000, 2001; Phelan et al. 2000; De Long et al. 2001), may have significant implications for young winter flounder survival after metamorphosis. Al-Hossaini et al. (1989) reported greater growth for cohorts of European plaice that settled relatively early in Wales, but these fish also had higher mortality. Conversely, growth was slower in late-settling cohorts, but survival was higher. In contrast to shrimp predation, larger-sized age-0 flatfish may be preferentially selected by birds and certain fishes (Van der Veer et al. 1997). In addition, environmental effects, such as water temperature, may greatly influence the ability of predators to prey on young winter flounder. In particular, the effects of February water temperature on the recruitment of winter flounder is discussed below.
Temperature also acts indirectly across : several early life history stages by affecting growth, development, and mortality, Van de Veer et al. (2000b) proposed that meristic elements, such as the number of vertebrae and fin rays, had effects seen later in the juvenile stage of young European plaice. Temperature-dependent characters, particularly the
numbers of vertebrae (established during the egg stage) and fin rays (during the larvae phase), appear to be related to growth and mortality experienced later on by juveniles. This was likely related to variable performance in locomotion and predator avoidánce responses. Thus, a non-genetic phenotypic plasticity found during very early life history that was influenced by environmental factors, in combination with events occurring later in life during the settled juvenile stage, could affect resulting yearclass strength.
Variation in annual year-class strength of European plaice was suggested to occur in either the pelagic larval phase (Zijlstra and Witte 1985; Van der Veer 1986) or after settlement during the juvenile stage on nursery grounds (Nash and Geffen 2000). Van der Veer (1986), Van der Veer and Bergman (1987), and Bergman et al. (1988) noted that recruitment variability in European plaice found in The Netherlands was stabilized between years as a result of a density-dependent regulatory process, predation on newly metamorphosed fish by caridean shrimp. In contrast, year-class strength of European plaice in Swedish bays varied to a greater degree (CVs $=67-$ 118\%). The latter was thought related to the effects of temperature variation during the larval stage and more variable crustacean predation on newly metamorphosed young European plaice found in móre northerly waters (Pihl 1990; Pihl and Van der Veer 1992). However, variable hydrographical effects on settlement of young Euröpean plaice may also have occurred in Sweden (Modin and Pihl 1994; Pihl et al. 2000). Thus, population regulation in flatfishes may be coarsely determined during the earliest life history stages by variable survival of eggs and larvae and then fine-tuned by mortality of newly metamorphosed "juveniles, which can be densitydependent beyond certain threshold levels of abundance (Van der Veer and Bergman 1987; Iles and Beverton 1991; Beverton and Iles 1992a, 1992b; Rose et al. 1996; Van der Veer et al. 2000a). .
High recruitment of winter flounder is associated with cold winters and a significant effect of February water temperature has reduced variability in the Niantic River winter flounder stock and recruitment relationship. The exact mechanism of how February temperatures, in particular, affect winter flounder recruitment remains unknown. Cold February water temperature was also suggested as an environmental influence that increased European plaice recruitment in both The Netherlands (Zijlstra and Witte 1985; Van der Veer 1986; Iles' 1994; Van der Veer and Witte 1999; Van der Veer et al. 2000a) and in a small bay on the Irish Sea (Nash et al. 1994). Similarly;
recruitment of European plaice throughout much of the waters around the United Kingdom were negatively correlated with water temperatures during February-June, the period encompassing the drift of their eggs and larvae and settlement (Fox et al. 2000). February coincides with most winter flounder spawning, egg incubation, and hatching. These processes and larval growth are all temperaturedependent. Buckley et al. (1990) noted that the winter flounder reproductive process appears optimized for cold winter temperatures that are followed by a gradual spring warming. Keller and Klein-MacPhee (2000) reported winter flounder egg survival, percent hatch, and initial size were significantly greater and mortality rates lower in cool as opposed to warm experimental mesocosms. Adult acclimation temperatures and egg and larval incubation temperature affected larval size and biochemical composition. Cold winters and warm springs produced large larvae that were in the best condition at first feeding, which favored high survival and partly explained the observed correlation between cold years and strong yearclasses of winter flounder. Townsend and Cammen (1988) noted that the metabolic rates of pelagic consumers are more sensitive to lower temperature than rates of photosynthesis by phytoplankton, which bloom more in response to the amount of solar radiation received, which is generally consistent over time each year. Therefore, a bloom in a cold year has the possibility of lasting longer before being grazed down by zooplankton. This allows for a greater contribution of organic matter to the benthos than in other years, benefiting juvenile demersal fishes that metamorphose just after the spring bloom of phytoplankton and have to outgrow various predators. Warmer winter water temperatures in Rhode Island have also been correlated with smaller winterspring phytoplankton blooms (Keller et al. 1999), which may have had consequences to marine food webs. The effect of temperature on potential prey or predators of larvae and newly metamorphosed juveniles, such as the sevenspine bay shrimp, may be an additional means for control of population abundance. The association of strong year-classes of European plaice with cold winters likely occurred because predatory brown shrimp suffered high mortality during low water temperatures or migrated out of the nursery areas (Zijlstra and Witte 1985; Van der Veer 1986; Pihl 1990; Pihl and Van der Veer 1992; Van der Veer et al. 2000a). Keller and KleinMacPhee (2000) also observed that the sevenspine bay shrimp remained inactive and buried within the
sediments of cool experimental mesocosms, whereas they were active in the warm systems.
Potential effects of temperature and winter climate on winter flounder are further illustrated in Figure 40 by comparing a long-term (1959-2001) CPUE index (annual sum of monthly means) of annual winter flounder (all sizes) developed from weekly trawl sampling conducted by the University of Rhode Island (URI) near Fox Island in upper Narragansett Bay (previously described in Adult Winter Flounder, Comparisons among Regional Abundance Indices) with the mean Providence, RI winter (DecemberFebruary) air temperature (Anonymous 2002a) and the North Atlantic Oscillation (NAO) winter (December-March) index (Anonymous 2002b). The NAO index (Hurrell 1995) is based on the difference of normalized sea level pressures between Lisbon, Portugal (representing the Azores high pressure system) and Stykkisholmur or Reykjavik, Iceland (Icelandic low). Negative phases of the NAO index result in colder, wetter, and windier winters in the northeastern United States and positive phases are related to warmer, milder winters (Greene and Pershing 2000). The NAO is considered to be a driving force in climatic systems in the northern hemisphere (Hurrell 1995; Ottersen et al. 2001). In recent years, considerable interest has developed in examining relationships between climate and biological processes. Climatic oscillations reflected by the NAO may be viewed as a proxy for various regulating forces in marine or other ecosystems, including changes to timing of reproduction, population dynamics, distribution, and in interspecific relationships, including competition and predator-prey (Ottersen et al. 2001). Using data from the past 47 years, the NAO and Providence winter air temperature indices were positively correlated (Spearman's rank-order correlation; $\mathrm{r}=0.457, \mathrm{p}=$ 0.0012 ). A Mann-Kendall test also showed that both of these climatic indices had significantly ( $p<0.001$ ) increasing trends from 1959 through 2000, with the majority of the positive values occurring since the mid-1980s. Conversely, the URI annual trawl CPUE index for winter flounder had a significantly ( $p<$ 0.001 ) declining trend during the same period. When the air temperature and NAO indices were advanced 3 years with respect to the URI trawl index, significant negative correlations (Spearman's rankorder correlation; Providence winter air temperature: $r=-0.493, p=0.0008, n=43$; NAO winter index: $r=-0.478, p=0.0012, n=43$ ) were found. This occurred even though the trawl catches were comprised of many age-classes of winter flounder


Fig. 40. Comparison from 1959 through 2001 between the annual URI winter flounder trawl CPUE abundance index at Fox Island in Narragansett Bay and the Providence, RI mean winter (December-February) air temperature index (top) and the North Atlantic Oscillation (NAO) winter (January-March) winter index (bottom). Both climatic indices were advanced 3 years relative to winter flounder abundance.
and each year's winter weather should have affected the reproductive success of only one year-class. There was likely a tendency for relatively large or weak year-classes to dominate trawl catches. This result showed that winter flounder abundance was highest during periods of relatively severe winters, such as occurring in the mid-1960s and late 1970s, when dominant year-classes were produced. Abundance was lowest when winters were milder, years during which fewer fish were produced. The trend of generally warm winters found from the mid-1980s through the present, most likely in combination with relatively high fishing mortality rates, has likely kept winter flounder abundance depressed. The climatic trend may continue as models have predicted that the current positive phase of the NAO may persist through the present and the following decade (Paeth et al. 1999).
Recruitment of many fishes may also be affected by density-dependent processes occurring during the first year of life following completion of the larval stage (Bannister et al. 1974; Cushing 1974; Sissenwine 1984; Anderson 1988; Houde 1989; Myers and Cadigan 1993a, 1993b; Bailey 1994). Bannister et al. (1974), Lockwood (1980), Van der Veer (1986), and Pihl et al. (2000) all reported density-dependent mortality for young European plaice. Examination of some of these findings, however, indicate that greatest rates of mortality occurred only when extremely large year-classes of European plaice were produced (i.e., three to more than five times larger than average). This was confirmed in analyses by Iles and Beverton (1991) and Beverton and Iles (1992a, 1992b), who reported that although density-dependent mortality was indicated for age-0 European plaice in the North Sea, below a specific density ( $1.8 \cdot \mathrm{~m}^{-2}$ ) mortality was likely density-independent. Pihl et al. (2000) noted that density-independent mortality was considerable, even at relatively high densities of European plaice. The high production of young Niantic River winter flounder that occurred in 1988 because of very low apparent mortality also showed no sharp peaks in abundance, with densities generally remaining below $1 \cdot \mathrm{~m}^{-2}$. However, high ( $>2 \cdot \mathrm{~m}^{-2}$ ) densities of young winter flounder at LR during some weeks in early summer of 1990, 1994, 1995, and 2001 were followed by the steepest declines in abundance. Mortality rate at WA was also high in 1990, even though only moderate densities of young were found there. However, mortality rates during 1994 and 1995 were about average, although abundances were among the highest ever observed there. In contrast, relatively low densities were found at both stations
during 1997, but the apparent mortality rate was high. Thus, the relationship between density and mortality rate for young winter flounder in the Niantic River is subject to considerable variability, with the population regulatory mechanism either not well-established or consistent from year to year. However, in a study examining 11 years of data from Narragansett Bay, DeLong et al. (2001) reported that the monthly mortality rate of age-0 winter flounder was density-dependent and also positively related to water temperature. They noted that mortality rates were highest in years with temperatures above $18^{\circ} \mathrm{C}$. However, the mean summer water temperature at the MPS intakes has exceeded this value for each of the past 26 years (Table 2) and water temperature is even somewhat higher at the shallow Niantic River nursery sites. The large difference found in values of $\mathrm{M}_{\mathrm{mo}}$ between 2000 and 2001 (Table 18) despite the same mean summer water temperature of $20.45^{\circ} \mathrm{C}$ suggests a less certain temperature-dependent effect on mortality in the Niantic River.
A comparison of early and late season median catch-per-unit-effort (CPUE) of age-0 winter flounder in the Niantic River showed that initially large numbers of young present during late spring and early summer in some years did not necessarily result in high densities of fish at the end of summer (Fig. 41). Differences found between early and late summer were largely related to variation in mortality rates, which affected year-class abundance. Notably, little observed mortality in 1988 and 2000 meant that modest initial sets of young resulted in relatively strong year-classes, whereas high mortality occurring during early summer in 1990, 1995, and 2001 considerably reduced initially high densities by late summer. Above-average survival rates found during


Fig. 41. Comparison between the density (median catch per $100 \mathrm{~m}^{2}$ ) of age- 0 winter flounder during early and late summer at Niantic River stations LR and WA combined from 1984 through 2001.
both 1999 and 2000 should help these winter flounder year-classes remain abundant. However, success also depends upon mortality rates that affect these fish during the several years remaining before recruitment as seemingly abundant year-classes of young winter flounder found during the 1990s did not result in numerous adults (Simpson et al. 1996; Desfosse et al. 1998). Thus, neither the 3 years of extended shutdowns at MPS nor the appearance of relatively numerous metamorphosed juveniles in recent years have resulted in a sharp rebound in abundance of Niantic River winter flounder. Considerable influence of natural mortality during the first several years of life and fishing mortality reducing adult spawner biomass remain the most important factors in determining recruitment and subsequent adult abundance. These factors also appear to be acting on a larger regional scale than just in the vicinity of MPS, which was discussed previously in Adult Winter Flounder - Comparisons Among Regional Abundance Indices.

## MPS Impact Assessment

## Larval Entrainment

Data Analyses. Field sampling and laboratory processing of winter flounder larvae entrained through the "MPS cooling-water system was previously described (see Larval Winter Flounder ': Field Sampling, Laboratory Processing, and Data Analyses). The estimated number of larvae entrained in the MPS condenser cooling water system each year was determined using larval densities at station EN (Fig. 13) and the measured volume of cooling water used by the three (now two) MPS units in operation (Table 31). The Gompertz density function (Eq. 3) was fitted to larval abundance data and daily densities (number. $500 \mathrm{~m}^{-3}$ ) were calculated. Daily entrainment estimates were determined after adjusting for the daily condenser cooling-water volume and an annual estimate was calculated by summing all daily estimates during the larval season.
Estimates of Larval Entrainment at MPS. The number of winter flounder larvae entrained in the condenser cooling water of MPS is the most direct measure of potential impact on the Niantic River winter flounder stock. Due to the importance of this measure, a review and evaluation of methods to estimate annual entrainment of larval winter flounder by MPS was conducted and provided in NUSCO (1991c). This topic was also summarized in NUSCO (2000).

Nearly all winter flounder larvae collected at station EN have been taken from February through June, with most ( $>90 \%$ ) during April and May. The entrainment estimate for 2001 of 376.2 million was the second largest annual estimate of the 26 -year period (Table 32). This estimate resulted from the second largest larval abundance ( $A$ parameter) found at EN and a relatively high (74.4\% of total; considering the rated capacity of all three MPS units) volume of seawater used for condenser cooling (Table 31). During the 2001 larval season, Unit 2 and Unit 3 were in full operation and Unit I did not operate due to decommissioning. If all three MPS units had been operating throughout this season, an estimated 507.8 million larvae would have been entrained, an estimate 1.35 times as large as the actual entrainment total. The fraction of entrainment foregone was the same as the proportion of cooling water not used by MPS in 2001 .
The percentage of available total permitted flow used by MPS from March through May during the three-unit (1986-2001) period ranged between $13.6 \%$ in 1997 to $96.5 \%$ in 1993. During the larval winter flounder seasons of 1996-98, all three units were shut down, resulting in the smallest circulating water flows since Unit 2 went online in 1976 (see the Introduction section of this report for a summary of plant operational history). However, variable larval abundance also influenced the entrainment estimates. During 1993, the estimate was the second lowest in 24 years because of larval abundance at EN that year was the lowest recorded. In contrast, the 1997 entrainment estimate was higher than expected because larval abundance that year was quite high. Similarly, relatively high abundance of winter flounder larvae in both 1992 and 2001 in combination with moderately high water use resulted in the two largest entrainment estimates.
As in previous years, larvae in Stage 3 of development predominated (58.8\%) in 2001 entrainment collections. This year, the fractions of total entrainment by developmental stage were $0.3 \%$ for Stage 1, $26.2 \%$ for Stage 2,58.8\% for Stage 3, 14.6\% for Stage 4, and $0.1 \%$ for Stage 5. For the period of '1983-2000, estimated composition' was $3.5 \%$ of the combined total for Stage 1, $19.0 \%$ for Stage 2, 63.3\% for Stage 3, $13.7 \%$ for Stage 4, and $0.5 \%$ for Stage 5 . In 2001, fewer Stage 1 and 3 larvae and more Stage i 2 larvae were entrained in comparison to the longterm averages.

- Entrainment Survival. For assessment purposes at MPS, all winter flounder larvae are presumed to die during passage through the condenser cooling-water system. However, results from early entrainment

TABLE 31. Annual average cooling-water flow and percent of nominal maximum flow at MPS Units 1 through 3 and station total during the April I-June 15 larval winter flounder season from 1971 through 2001.

| Nominal tlow at $100 \%$ capacity: | Unit 1" |  | Unit 2 |  | Unit 3 |  | MPS three-unit total |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $28.0 \mathrm{~m}^{3} \cdot \mathrm{sec}^{-1}$ |  | $36.1 \mathrm{~m}^{3} \cdot \mathrm{sec}^{-1}$ |  | $59.5 \mathrm{~m}^{3} \cdot \mathrm{sec}^{-1}$ |  | $123.6 \mathrm{~m}^{1} \cdot \mathrm{sec}^{-1}$ |  |
| Fraction of total MPS flow: | 0.227 |  | 0.292 |  | 0.481 |  | 1.000 |  |
|  | Average flow | $\%$ of maximum | Average flow | $\%$ of maximum | Average flow | \% of maximun | Average flow | $\%$ of maximum |
| 1971 | - | 68.0 | - | - | - | - | - | 15.4 |
| 1972 | - | 99.4 | - | - | - | - | - | 22.6 |
| 1973 | - | 23.7 | - | - | - | - | - | 5.4 |
| 1974 | - | 97.0 | - | - | - | - | - | 22.0 |
| 1975 | - | 84.4 | - | - | - | - | - | 19.2 |
| 1976 | 25.8 | 92.1 | 29.9 | 82.9 | - | - | 55.7 | 45.1 |
| 1977 | 26.4 | 94.1 | 15.0 | 41.5 | - | - | 41.3 | 33.4 |
| 1978 | 21.3 | 76.0 | 27.1 | 75.2 | - | - | 48.4 | 39.2 |
| 1979 | 10.0 | 35.7 | 21.8 | 60.4 | - | - | 31.8 | 25.7 |
| 1980 | 22.7 | 81.1 | 26.4 | 73.3 | - | - | 49.2 | 39.8 |
| 1981 | 1.8 | 6.5 | 32.2 | 89.2 | - | - | 34.0 | 27.5 |
| 1982 | 26.6 | 94.9 | 34.0 | 94.3 | - | - | 60.6 | 49.1 |
| 1983 | 26.2 | 93.7 | 28.5 | 78.9 | - | - | 54.7 | 44.3 |
| 1984 | 5.3 | 19.0 | 34.4 | 95.4 | - | - | 39.8 | 32.2 |
| 1985 | 26.7 | 95.4 | 15.3 | 42.3 | - | - | 42.0 | 34.0 |
| 1986 | 25.7 | 91.9 | 35.2 | 97.4 | 52.5 | 88.2 | 113.4 | 91.7 |
| 1987 | 24.9 | 88.9 | 36.0 | 99.6 | 50.9 | 85.7 | 111.8 | 90.5 |
| 1988 | 27.9 | 99.8 | 29.3 | 81.1 | 55.0 | 92.4 | 112.2 | 90.8 |
| 1989 | 9.4 | 33.6 | 29.4 | 81.6 | 41.7 | 70.2 | 80.6 | 65.2 |
| 1990 | 26.2 | 93.5 | 28.6 | 79.4 | 47.1 | 79.2 | 101.9 | 82.5 |
| 1991 | 2.8 | 10.1 | 27.7 | 76.7 | 52.3 | 87.9 | 82.8 | 67.0 |
| 1992 | 26.3 | 93.8 | 25.7 | 71.1 | 48.3 | 81.2 | 99.9 | 80.9 |
| 1993 | 26.5 | 94.5 | 35.6 | 98.5 | 57.3 | 96.3 | 119.3 | 96.5 |
| 1994 | 10.5 | 37.4 | 26.9 | 74.6 | 55.7 | 93.7 | 93.1 | 75.4 |
| 1995 | 27.7 | 98.9 | 18.5 | 51.2 | 31.3 | 52.7 | 77.5 | 62.7 |
| 1996 | 0.7 | 2.5 | 9.1 | 25.2 | 15.0 | 25.1 | 24.7 | 20.0 |
| 1997 | 0.0 | 0.0 | 8.3 | 22.9 | 8.5 | 14.3 | 16.8 | 13.6 |
| 1998 | 0.7 | 2.6 | 4.3 | 11.8 | 30.8 | 51.7 | 35.8 | 28.9 |
| 1999 | 0.7 | 2.7 | 25.0 | 69.3 | 32.7 | 54.9 | 58.4 | 47.3 |
| 2000 | 0.8 | 2.9 | 22.9 | 63.5 | 59.0 | 99.2 | 82.7 | 66.9 |
| 2001 | 0 | 0 | 33.0 | 91.4 | 59.0 | 99.2 | 92.0 | 74.4 |

* Unit I shut down on November 4, 1995 and its retirement was announced in July 17, 1998. Cooling-water tlow at Unit 1 included only the service water system during 1996-2000 and no cooling water was used as of 2001.
- No records of cooling-water flow were available for 1971-75; net electrical generation records were used to estimate flow.
survival studies at MPS (NUSCO 1975, 1984, 1987) and at other power plants (Itzkowitz and Schubel 1983; EA 1986) indicated variable survival rates of entrained winter flounder larvae. Laboratory thermal tolerance studies were conducted during 1973 and 1974 to estimate the effect of increased temperature on larvae during entrainment (NUSCO 1975, 1987). Larvae were grouped as pre-metamorphosed ( $<5 \mathrm{~mm}$ ) and metamorphosing ( $>5 \mathrm{~mm}$ ) and were exposed to water at about $70^{\circ} \mathrm{F}$ after acclimation at $46.5^{\circ} \mathrm{F}$ (i.e., a $\Delta \mathrm{T}$ of $23.5^{\circ} \mathrm{F}$ ). Pre-metamorphosed larvae died as exposure time increased. At 1 hour, $29 \%$ died, $48 \%$ died at 2 hours, $53 \%$ at 3 hours, and $89 \%$ at 6 hours. Metamorphosing larvae exposed to elevated tempera-
ture for up to 9 hours suffered no mortality. Because most entrained larvae are 5 mm or larger and during two- or three-unit operation would be exposed to elevated temperatures for considerably less than 9 hours, most larvae would be expected to survive the thermal increase associated with entrainment. A preliminary field study of entrainment survival was conducted in 1983 (NUSCO 1984). Winter flounder larvae $(\mathrm{n}=135)$ were collected in the quarry following passage through the condenser cooling-water system when water temperatures were about 70 to $72.5^{\circ} \mathrm{F} ; \Delta \mathrm{T}$ ranged between 14.5 to $20.7^{\circ} \mathrm{F}$. Larvae were collected by slowly hauling a $0.5-\mathrm{m}$ diameter plankton net from near the bottom to the surface (ca. 50 feet) by hand.

TABLE 32. Annual abundance index (A parameter of the Gompertz function) with $95 \%$ confidence interval of winter flounder larvae in entrainment samples and total annual entrainment estimates and the volume of seawater entrained at MPS each year from 1976 through 2001 during the larval period of occurrence.

${ }^{3}$ Values may differ from those reported in DNC (2001a) due to adjustments made in the larval period of occurrence.

Specimens were held at elevated temperatures for 2 to 4 hours (approximating estimated retention time in the quarry for three- and two-unit operation, respectively). Following this period, they were held at ambient water temperature without feeding to observe latent mortality for a total of 96 hours. All Stage 4 larvae collected were initially alive and $79 \%$ survived the holding period. Although some of both Stage 2 and 3 larvae were alive at time of collection ( 33 and $79 \%$, respectively), all of these larvae died during the 96 -hour holding period. Because of the relatively small (12\%) proportion of entrained larvae that are in Stage 4 and 5 of development, the overall survival of entrained larvae may be less than $10 \%$, if these estimates are indicative of actual entrainment survival at MPS. Besides the thermal component of entrainment, mortality may also be induced by mechanical damage (e.g., cavitation, pressure changes).

Further evidence of entrainment survival is important to evaluate entrainment impact to winter flounder. A more comprehensive entrainment survival study was completed at MPS in spring of both 2000 and 2001 using a specially designed pump sampler. The pump sampler was employed at the Unit 3 cooling water
discharge to evaluate the survival of entrained winter flounder larvae during spring in 2000 and 2001. The sampler consisted of a $1.0 \times 1.0 \mathrm{~m}$ conical screen of $333 \mu \mathrm{~m}$ mesh fitted with a 2-L codend bucket to hold larvae. The net was suspended inside a chamber connected to an intake hose situated in the chamber wall above the net and an outlet hose at the bottom of the chamber. The intake hose entered the chamber below the water line so that water entered the chamber by gravitational force. The unattached end of the intake hose on the sampler was placed in the discharge plume immediately below the' Unit 3 discharge structure. Water was pumped out of the bottom of the chamber using a gasoline powered pump, thereby filtering water through the net. Volume of water sampled was measured using an analog flow meter attached to the outlet line after the pump. Flow was adjusted with a valve at the terminus of the outlet line. The entire sampling apparatus was situated on a $3 \mathrm{~m}^{2}$ mobile floating platform.
The pump was typically run for 1 hour and filtered about $50 \mathrm{~m}^{3}$ of water. At the end of the hour sampling time, the net was slowly raised from the
chamber and gently rinsed to wash all organisms and debris into the codend bucket. The bucket was then removed, sealed, and placed in a bucket of discharge water (or intake area water depending on the site) to maintain temperature during transport to the laboratory. The sample was examined at the laboratory using a magnifying lens to remove larvae. Larvae were identified and assessed for signs of life. Transport and processing typically took between 20 and 30 minutes. Live winter flounder larvae were placed in flow-through beakers of sea water of similar temperature to the discharge water. Dead larvae were measured ( 1.0 mm ) and the developmental stage recorded. Beakers with live larvae were subjected to a 1 -hour acclimation period raising the temperature to ambient LIS water temperature to simulate the movement of water from the discharge into LIS then placed in a tank having flow-through sea water. Larvae were fed live rotifers to a satiated level. Numbers of larvae surviving entrainment were determined at $0,24,48$, and 96 hours after collection. Larvae experiencing latent mortality were measured and the developmental stage recorded.
In 2000, 18 Stage 3 and 67 Stage 4 winter flounder larvae and 3 Stage 5 juveniles were collected from the Unit 3 discharge water between May 18 and June 7. Volume of discharge water sampled in 2000 was approximately $465.5 \mathrm{~m}^{3}$ during 10 hours of sampling effort. Of the fish collected, 5 Stage 3, 44 Stage 4, and 3 Stage 5 fish were alive during the initial observation. No Stage 3 larvae survived while $21 \%$ of the Stage 4 larvae and $100 \%$ of the Stage 5 juveniles survived through 96 hours (Table 33). About 332 million winter flounder larvae were entrained by MPS in $2000^{\circ}$ (DNC 2001a). Placing the survival estimates into perspective with the numbers of winter flounder larvae entrained in 2000,
approximately 14.4 million Stage 4 larvae and 2.7 million Stage 5 juveniles would have survived entrainment, assuming that survival of entrained fish was similar across both units.
In 2001, 9 Stage 2, 292 Stage 3, and 115 Stage 4 winter flounder larvae and 1 Stage 5 juvenile were collected during 32 hours of sampling effort that filtered approximately $1,551.0 \mathrm{~m}^{3}$ of discharge water. While $44 \%$ of Stage 2 larvae were alive at time of initial examination, none survived the holding period. Stage 3 larvae showed $18 \%$ survival at the time of the initial observation, but only $1 \%$ survived through 96 hours. Stage 4 larvae showed $70 \%$ initial survival and $38 \%$ latent survival through 96 hours. The only Stage 5 individual collected in 2001 survived through 96 hours (Table 33). Applying these survival rates to the 2001 entrainment estimates, approximately 2.4 million entrained Stage 3 larvae, 19.6 million Stage 4 larvae, and 1.8 million Stage 5 juveniles would have survived entrainment. Cumulatively, these figures indicate that about $6 \%$ of the total number of winter flounder larvae entrained in 2001 survived.
Despite the small sample sizes of early developmental stage winter flounder larvae, entrainment survival appeared to increase with developmental stage. Survival of Stage 2 and 3 larvae was negligible, while Stage 4 survival was between $21 \%$ and $38 \%$ during this study. Survival of Stage 5 fish (transformed juveniles) was good at $100 \%$, although based on only 4 fish captured during the 2 -year study. Although this work was limited in scope and there were problems in estimating control mortality, the sampling indicated that some Stage 4 larvae and likely all of the relatively few Stage 5 juvenile winter flounder entrained could survive through-plant passage. Conservatively, as many as $6 \%$ of entrained larvae likely survive through-plant passage. This is

TABLE 33. Summary of winter tlounder larval entrainment survival at the MPS discharge during spring of 2000 and 2001. Volume is total volume sampled with the pump in $\mathrm{m}^{3}$; stage is developmental stage of winter tlounder larvae; total is the total number of larvae collected; number alive at time 0 is the number of larvae that were alive at the time of collection; and $\%$ alive at 96 h is the percent of winter flounder larvae alive at time 0 that survived through 96 hours.

particularly important as older and larger winter flounder larvae, which have greater potential for survival, suffer less entrainment mortality. Thus, based on preliminary studies done at MPS and other power plants, there is good likelihood that a portion of winter flounder larvae survive entrainment, even though impact assessments made to date (and including this report) assumed $100 \%$ mortality.
Effect of Entrainment on a Year-class. As noted previously in this section, both flow and larival densities vary over the larval season and among years (Fig. 42). The time-series of annual entrainment abundance (an index described by the $A$ parameter of the Gompertz function) and the annual volume of seawater used at MPS have relatively similar CVs ( 53 and $45 \%$, respectively). The entrainment estimates, however, have a CV of $72 \%$ because both larval densities and plant .operation affect this estimate and they vary independently from year to year. A high larval abundance for a particular year is not functionally associated with high flow and vice versa. In fact, these two measures are not significantly correlated (Spearman's rank-order correlation; $r=-0.123 ; p=0.548 ; n=26$ ). Thus, cooling-water flow alone cannot be used as a measure of plant effect, as the fraction of Niantic River larval production available to entrainment and its relation to total larval entrainment changes from year to year. Variable annual rates of entrainment


Fig. 42. Comparison between the annual estimates of larval winter flounder entrainment in millions, larval abundance at EN given as the annual mean density of larvae (A parameter of the Gompertz distribution), and seawater volume entrained in tens of millions of $\mathrm{m}^{3}$ at MPS from 1976 through 2001. The annual mean larval density estimates were divided by 10 to fit the same scale as the other two measures.
appear to be real and may be related to physical processes (e.g., precipitation, winds) that vary both intra- and interannually during the larval developmental period, when the abundance, distribution, natural mortality, and behavior of larvae are also variable.
The annual entrainment abundance index ( $A$ parameter of the Gompertz function) divided by the total volume of seawater entrained each year results in an index of the annual rate of entrainment, which is expressed as the number of larvae per unit volume of cooling water (Fig. 43). Based on the nonparametric Mann-Kendall test, the entrainment rate since 1976 has varied with no significant ( $p=0.877$ ) trend in slope. The entrainment rate time-series has a CV of $98 \%$, which is greater than the CVs for density, flow, and numerical entrainment estimates. Larval production and availability in Niantic Bay remained stable, despite increases in MPS coolingwater use during 1986-95, the effective period of full three-unit operation. Also, reduced recruitment has not been the result of reduced larval abundance, because the latter has exhibited no trend after many years of MPS operation.
'Relationships between larval entrainment estimates and various indices of juvenile abundance were examined to determine the effect of entrainment on a year-class. Annual entrainment estimates were significantly positively correlated with three abundance indices of age- 0 juvenile winter flounder, including the median CPUE of age-0 fish taken in both early and late summer in the Niantic River and with age-0 juveniles taken at TMP stations in late fall-early winter (Table 34). Entrainment estimates


Fig. 43. Annual rate of entrainment of winter flounder larvae at MPS, defined as the annual mean density of larvae ( $A$ parameter of the Gompertz distribution) divided by the volume of seawater entrained each year from 1976 through 2001. The data were scaled to result in the numerical rate as shown.

TABLE 34. Spearman's rank-order correlations between the annual estimates of larval winter flounder entrainment at MPS and the abundance indices of several post-entrainment early life history stages and a calculated apparent larval survival rate.

| Index ${ }^{2}$ | Niantic River early summer age-0 juveniles | Niantic River late summer age-0 juveniles | Fall-early winter river-bay <br> age-0 juveniles | Niantic River winter-spring age-1 juveniles (lower river) | Niantic River winter-spring age-1 juveniles (upper river) | Apparent survival rate |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Annual estimate of entrainment at MPS | $\begin{gathered} 0.6348^{6} \\ 0.0062 * * \\ 17 \end{gathered}$ | $\begin{gathered} 0.7977 \\ 0.0001{ }^{* *} \\ 18 \end{gathered}$ | $\begin{gathered} 0.4237 \\ 0.0348{ }^{*} \\ 25 \end{gathered}$ | $\begin{gathered} -0.1793 \\ 0.3911 \mathrm{NS} \\ 25 \end{gathered}$ | $\begin{gathered} 0.0000 \\ 1.0000 \mathrm{NS} \\ 21 \end{gathered}$ | $\begin{gathered} -0.3469 \\ 0.0893 \mathrm{NS} \\ 25 \end{gathered}$ |

- Indices used correspond to those given on Tables 22, 23, and 32, except for the apparent survival rate, which is the $\Delta$-mean CPUE of age-1 winter flounder taken in the lower Niantic River during adult population surveys divided by the $A$ abundance index of 7 mm and larger larvae at the MPS discharge (entrainment station EN).
- The three statistics shown in each correlation matrix element are: correlation coefficient ( $r$ ),
probability of a larger $r$ (NS - not significant $[p>0.05],{ }^{*}$ - significant at $p \leq 0.05, * *$ significant at $p \leq 0.01$ ), and number of annual observations (sample size).
were not significantly correlated with age-1 fish collected in either the lower or upper Niantic River during the adult winter flounder surveys, nor with a calculated apparent larval survival rate, which was defined as the $\triangle$-mean CPUE of age-1 winter flounder taken in the lower Niantic River during the adult population surveys divided by the $A$ abundance index of winter flounder larvae 7 mm and larger taken at the MPS discharge for the same year-class. Although statistically significant, the form of the relationships between the entrainment estimates and several age-0 abundance indices was not obvious, however (Fig. 44).

Densities of winter flounder larvae 7 mm and larger taken at EN (an index of abundance rather than one of estimated impact) were significantly correlated with abundance of Stage 4 larvae in the Niantic River (Table 25; Fig. 36). The significant correlation coefficients found between entrainment estimates and age-0 abundances were positive, implying no apparent entrainment effect and indicating that the more larvae available for entrainment, the more larvae that settled as demersal young in the Niantic River. However, even negative correlations between annual entrainment and abundance of early life history stages do not necessarily imply an entrainment impact unless positive correlations can be found between those early life history stages and mature female fish, which are lacking (Table 26).

Based on beam trawl catches in the Niantic River since 1983, most fish were produced in 1988, 1992, 1994, 1995, 2000, and 2001, with particularly weak year-classes formed during the mid-1980s, 1989, 1993, 1996 (although found to be numerous in fall and winter), and 1997. The highest larval winter flounder entrainment estimates at MPS during the
three-unit period were in 1992 and 1995 and lowest estimates were found in 1993 and 1996-98, years following the retirement of Unit 1 and during the extended shutdowns of Units 2 and 3. Subsequently, the second and third highest entrainment estimates occurred in 2000 and 2001. This suggests that in some years, year-class strength was strongly influenced by events during early life history stages resulting in abundant larvae and early settling juveniles. Thus, entrainment estimates only reflected emerging year-class strength rather than being an important factor affecting numerical abundance. The examples of 1988 and 2000 (low mortality) and 1990 and 2001 (high mortality) also illustrated that mortality during the first summer on the nursery ground, which is totally unaffected by MPS operation, profoundly influences year-class strength and, ultimately, recruitment to adult stocks.

Mass-balance Calculations. The magnitude of the impact of entrainment on the Niantic River winter flounder stock depends upon how many of the entrained larvae originated from this stock as well as larval densities in Niantic Bay. Hydrodynamic modeling (NUSCO 1976) and tidal-current drogue studies (NUSCO 1992b) showed that much of the condenser cooling-water used by MPS enters Niantic Bay from LIS. Other stocks are known to spawn both to the east and west of the bay (Howell and Molnar 1996, 1998) and results from tidal studies also indicated that a large number of winter flounder larvae entered Niantic Bay from LIS (NUSCO 1992a, 1992b). Mass-balance calculations were used to investigate whether the number of winter flounder larvae entering Niantic Bay from the Niantic River could sustain the number of larvae observed in the bay during the winter flounder larval season each


Fig. 44. Comparison between the $1-\mathrm{m}$ beam trawí median CPUE of age-0 winter flounder taken at Niantic River stations LR and WA combined during both early (1985: 2001) and late summer (1984-2001) and the late fall-early winter seasonal $\Delta$-mean CPUE of age- 0 winter flounder at trawl monitoring program stations (1976-2000) with annual entrainment estimates of winter flounder larvae at MPS. (Note that the vertical scales differ among the graphs).
year from 1984 through the present; 10 of these years (1986-95) occurred during full three-unit operation and 4 years (1996-99) with limited plant operation. The mass-balance model was not completed during 2000 because a reliable estimate of larval winter flounder mortality was not obtained (DNC 2001a). Recent assessments of the mass-balance model,
including sensitivity analyses, were conducted by staff and two independent consultants for DNC. Output of this model, at worse, should have biases well within $\pm 50 \%$ (DNC 2001c) and was far more sensitive to errors in larval density estimates than to errors in larval mortality rates.
Three potential inputs of larvae to Niantic Bay include eggs hatching in the bay, larvae flushed from the Niantic River, and larvae entering the bay from LIS across the boundary between Millstone Point and Black Point (Fig. 13). The few yolk-sac larvae collected annually in Niantic 'Bay suggested that minimal spawning and hatching occurred in the bay, which was therefore considered a negligible source of larvae. Larvae were known to be flushed from the river into the bay and this input to the bay was estimated from available data. The number of larvae entering Niantic Bay from LIS was unknown.' Four ways in which larvae may leave Niantic Bay include natural mortality, advection into the Niantic River during a flood tide, entrainment at MPS, and flushing from the bay into LIS. Estimates could be made for the number of larvae lost through natural mortality, advected into the Niantic River, and entrained at MPS, but little was known about the number of larvae flushed into LIS. The numbers of larvae flushed to and from LIS were combined as an unknown termed Source or Sink in the mass-balance calculations. Thus, the form of the mass-balance equation was:

$$
\begin{align*}
N B_{t}+5= & N B_{l}-\text { NumEnt }- \text { Mort }+ \text { FromNR }- \\
& T o N R \pm(\text { Source or Sink }) \tag{13}
\end{align*}
$$

where $t=$ time in days
$N B_{t+5}=$ number of larvae in Niantic Bay 5 days after day $t$ (instantaneous daily estimate)
$N B_{f}=$ initial number of larvae in Niantic Bay on day $t$ (instantaneous daily estimate)
NumEnt = number of larvae lost from Niantic Bay by entrainment in the condenser : cooling-water system (over a 5 -day period)
Mort $=$ number of larvae lost from Niantic Bay due to natural mortality (over a 5 -day period)
From $N R=$ number of larvae flushed from the
$\therefore \quad$ Niantic River (over a 5 -day period)
ToNR $=$ number of larvae entering the Niantic River (over a 5 -day period)
Source or Sink = unknown number of larvae in' Niantic Bay that flush out to LIS or enter the bay from LIS (over a 5 -day period)

Solving for the unknown Source or Sink term, the equation was rearranged as:

$$
\begin{align*}
\text { Source or Sink }= & N B_{t}+5-N B_{t}+\text { NumEnt }+ \\
& \text { Mort }- \text { FromNR }+ \text { ToNR } \tag{14}
\end{align*}
$$

Because these mass-balance calculations were based on the change in the number of larvae in Niantic Bay over a 5 -day period:

$$
\begin{equation*}
5 \text {-day change }=N B_{t}+5-N B_{t} \tag{15}
\end{equation*}
$$

Thus:
$\begin{aligned} \text { Source or Sink }= & 5-\text { day change }+ \text { NumEnt }+ \\ & \text { Mort }- \text { FromNR }+ \text { ToNR }\end{aligned}$
Daily abundance estimates were derived from the Gompertz density equation (Eq. 3) and the daily densities for Niantic Bay at two points in time ( $N B_{t}$ and $N B_{t}+5$ ) for each 5 -day period were calculated from data collected at stations NB and EN combined. These densities, adjusted for the volume of Niantic Bay (about $50 \times 10^{6} \mathrm{~m}^{3}$; E. Adams, Massachusetts Institute of Technology, Cambridge, MA., pers. comm.), provided an estimate of the instantaneous daily standing stock. The difference between these two estimates $\left(N B_{t}\right.$ and $\left.N B_{t}+5\right)$ was the term 5-day change in Equation 15. The selection of 5 days as the period of change was arbitrary and a cursory examination of results based on 10 -day periods showed that the same conclusions were reached with either 5 - or 10 -day periods.

Daily entrainment estimates were based on data collected at station EN and the actual daily volume of condenser cooling water used at MPS. The daily entrainment estimates were summed over each 5-day period (NumEnt). Annual stage-specific mortality rates for 1984-89 were determined by Crecco and Howell (1990), for 1990 by V. Crecco (CT DEP, Old Lyme, CT, pers. comm.), and for 1991 and thereafter by MPS Environmental Laboratory staff. Mortality was partitioned among developmental stages by comparing the rates of decline of predominant sizeclasses of each stage. Each developmental stage was assigned a portion of the total annual larval mortality rate $(Z)$; similar mortality rates were assumed for Stages 3 and 4. Although estimating stage-specific mortality in this manner was imprecise, sensitivity analysis on the mass-balance calculations (NUSCO 1991b) indicated that larval mortality was the least sensitive parameter in Equation 16 above. These annual rates were modified to daily stage-specific mortality rates by assuming a duration of 10 days each for Stages 1, 3, and 4 larvae, and 20 days for

Stage 2 larvae. The proportion of each stage collected at station EN during each 5 -day period was applied to the daily standing stock for Niantic Bay $\left(N B_{t}\right)$ to estimate the number of larvae in each developmental stage for stage-specific mortality calculations. The daily loss due to natural mortality (Mort) was summed for each 5-day period.

The 5-day input of larvae to Niantic Bay from the river (FromNR) was based on daily density estimates for station C in the lower river after adjusting for the rate of flushing between that station and the river mouth. To determine the relationship between the estimated daily density at station $C$ and the average density of larvae leaving the river on an ebb tide, the geometric mean density of samples collected during an ebb tide for ten import-export studies conducted at the mouth of the Niantic River during 1984, 1985, and 1988 (NUSCO 1985, 1986a, 1989) was compared to the estimated daily densities at station $\mathbf{C}$. The average density of larvae flushed from the Niantic River was estimated from the functional regression equation:

$$
\begin{align*}
\text { From } N R= & 9.751+0.473 \text { (daily density at } \\
& \text { station } \mathrm{C}) \tag{17}
\end{align*}
$$

The $95 \% \mathrm{Cl}$ for the slope ( $\mathrm{r}=0.969 ; \mathrm{p}=0.001$; $\mathrm{df}=$ 8) was $0.387-0.579$. The estimated average density, the average tidal prism of $2.7 \times 10^{6} \mathrm{~m}^{3}$ (Kollmeyer 1972), and about 1.9 tidal prisms per day were used to estimate the daily flushing of larvae from the river into Niantic Bay. This daily input to the bay was summed for each 5 -day period to calculate the term From $N R$ in the mass-balance equation.

Stepwise oblique tows were collected during 1991 in the channel south of the Niantic River railroad bridge (station RM) during a flood tide to estimate an average density to compute ToNR (NUSCO 1992a). In 1992 and 1993, sampling was conducted again at RM during a flood tide, but the collections were made by mooring the research vessel to the railroad bridge and taking continuous oblique tows (NUSCO 1994a). Comparison of densities from the paired stations of NB and RM showed a poor relationship. Therefore, daily densities at the two stations were estimated using the Gompertz density curve (Eq. 3). For station RM in 1992, the equation could only be adequately fit by smoothing the data using a 3 -week running average prior to calculating a weekly cumulative density. The Gompertz function could not be fit to data collected at station NB during 1993. Therefore, catches from stations NB and EN were combined to calculate the weekly geometric means prior to fitting the Gompertz function and estimating
daily densities for Niantic Bay. Daily density estimates for 1991-93 were combined and functional regression :was used to determine the relationship between abundance at stations NB and RM. . The average density of larvae flushed from Niantic Bay into the river was estimated by the functional regression equation:

$$
\begin{equation*}
T o N R=128.149+2.073\left(N B_{l}\right) \tag{18}
\end{equation*}
$$

The $95 \% \mathrm{Cl}$ for the slope ( $\mathrm{r}^{2}=0.705 ; \mathrm{p}=0.001$; $\mathrm{df}=$ 406) was $1.827-2.351$. After being adjusted for the average tidal prism and the number of tidal prisms per day, these daily estimates of the number of larvae entering the river during a flood tide were summed over each 5 -day period to calculate the term ToNR in the mass-balance equation. Because of the large intercept in the above regression line when no larvae were present in Niantic Bay $\left(N B_{l}=0\right)$, the term ToNR was conservatively set to zero. The term Source or Sink in Equation 16 represents the 5-day
net loss or gain of larvae to Niantic Bay from LIS required to balance the calculation. For a net loss of larvae (flushed to LIS), the Source or Sink term would be negative and for a net gain of larvae (imported from LIS), the Source or Sink term would be positive. Results from mass-balance calculations by developmental stage were used to estimate the number of larvae entrained at MPS each year from the Niantic River. If FromNR can support the number of larvae entrained by MPS, then the Source or Sink term is negative (i.e., no import) to balance the equation. These larval losses were then used to calculate conditional mortality rates for Niantic River larvae for under both actual operating conditions and projected full MPS three-unit operation, the results of which will be provided below. The results for each 5 -day period of the 2001 larval winter flounder season are . provided, as an example of these calculations (Table 35). Computations for other years were given in NUSCO (1993, 1994a, 1995, 1996a, 1997b, 1998, 1999, 2000).

TABLE 35. Results of mass-balance calculations during each 5-day period in 2001.

| Start of 5-day period, | 5-day change (X $10^{\prime \prime}$ ) | Number entrained (NumEnt) (X $10^{\prime \prime}$ ) | $\because$ $\therefore$ | Loss due to mortality (Mort) (X $10^{6}$ ) |  | $\begin{aligned} & \text { Number from the } \\ & \text { Niantic River } \\ & \text { (FromNR) } \\ & \left(\times 10^{\prime \prime}\right) \end{aligned}$ | Number to the Niantic River $($ ToNR $\left(\times 10^{6}\right)$ | Source or Sink (X 10") |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2.15 | $0.0^{\text {a }}$ | 0.0 |  | 0.0 | * | 5.6 . | 6.7 | 1.2 |
| 2-20 | 0.1 | 0.0 |  | 0.0 |  | 6.6 | 6.8 | 0.3 |
| 2-25 | 0.1 | 0.0 |  | 0.0 | . . | . . 7.6 | 6.9 | -0.5 |
| 3-02 | 0.3 | 0.0 |  | 0.0 |  | $\therefore 8.6$ | . 7.1 | -1.1 |
| 3-07 | 0.6 | 0.2 |  | 0.1 |  | 9.6 | 7.5 | -1.3 |
| 3-12 | 0.9 | 0.4 |  | 0.1 |  | 10.5 | 8.3 | -0.8 |
| 3-17 | 1.4 | 1.0 |  | 0.2 |  | 11.3 | 9.5 | 0.8 |
| 3-22 | 2.0 | 2.0 |  | 0.3 |  | 12.0 ! | 11.2 | 3.6 |
| 3-27. | 2.6 | 4.3 |  | 0.4 |  | 12.6 | 13.7 | 8.4 |
| 4-1 | 3.2 | 9.4 |  | 0.6 |  | $13.1{ }^{\circ}$ | 16.8 | 17.0 |
| 4.06 | 3.8 | 13.0 |  | 1.1 |  | 13.4 | 20.5 | : 24.9 |
| 4.11 | 4.2 | 16.0 |  | 1.4 |  | 13.6 | 24.8 | 32.8 |
| 4-16 | 4.3 | 20.0 |  | 2.2 |  | 13.7 | 29.4 | 42.2 |
| 4-21 | 4.3 | 22.7 |  | 2.5 |  | 13.7 | 34.1 | 49.9 |
| 4-26 | 4.1 | 24.9 |  | 3.0 |  | 13.6 | 38.7 | . 57.1 |
| 5-01 | 3.7 | 28.0 |  | 3.6 |  | 13.4 | 43.0 | . 64.9 |
| 5-06 | 3.2 | 27.0 |  | 3.8 |  | $13.1{ }^{\circ}$ | 46.8 | 67.8 |
| 5-11 | 2.5 | 28.7 |  | 4.0 |  | 12.7 . | 49.9 | . 72.4 |
| 5-16 | 1.8 | 27.1 |  | 4.0 |  | 12.3 . | 52.3 | - 72.9 |
| 5.21 | 1.1 | 25.7 |  | 4.6 |  | $11.8{ }^{\circ}$ | 54.0 | 73.6 |
| 5.26 | $\therefore 0.4$ | 24.3 |  | 5.0 |  | 11.4 | 54.9 | 73.2 |
| 5-31 | -0.2 | 22.3 |  | 5.0 |  | 10.8 | 55.1 | 71.4 |
| 6-05 | -0.7 | 20.1 |  | 5.0 |  | 10.3 | 54.6 | 68.7 |
| 6-10 | -1.2 | 17.9 | , | 4.9 |  | 9.8 | 53.6 | 65.5 |
| 6-15 | -1.6 | 15.8 | : | 4.8 |  | - 9.3 | 52.2 | 62.0 |
| 6-20 | -1.8 | 13.9 |  | 4.6 |  | 8.7 | 50.4 | - 58.2 |
| 6-25 | -2.1 | 11.6 | . | 4.4 |  | 8.2 | 48.3 | 54.0 |

[^20]The sign of the term 5 -day change (i.e., the difference in the number of larvae in Niantic Bay at the beginning and end of each 5 -day period) shifted in the 2001 larval season from positive to negative when the abundance of larvae in the bay began to decline on May 31 (Table 35). A negative Source or Sink term indicated a net loss of larvae from Niantic Bay during the first part of the larval season, but beginning with the 5 -day period starting on March 17, the Source or Sink term became positive. This indicated that larvae from other sources in LIS were required to support the change in larval abundance and balance the equation. Timing of this change in the Source or Sink term was similar to previous years (NUSCO 1993, 1995, 1996a, 1997b, 1998, 1999, 2000), except for an earlier date of February 25 in 1993 (NUSCO 1994a). Considerably fewer larvae were entrained (NumEnt) than were imported from LIS (i.e., positive Source or Sink), starting in mid to late March. Also in 2001, the weekly estimates of NumEnt were considerably smaller than the number entering the river from the bay (ToNR).
The proportion of entrained larvae from the Niantic River during each 5 -day period of the season was estimated from the ratio of larvae entering the bay from the river (FromNR) to the total input from both sources (FromNR + Source or Sink). This proportion was applied to the total number entrained in the same

5 -day period to estimate the number entrained from the Niantic River. During any 5 -day period when there was a net loss (negative Source or Sink term) or when the proportion from the river was greater than one, all larvae entrained during that time were assumed to have originated from the Niantic River. This estimate was conservative because the results of a dye study and larval dispersal modeling (Dimou and Adams 1989) showed that only about $20 \%$ of the water discharged from the Niantic River passed through MPS during full three-unit operation. Estimates of annual total entrainment and the annual number entrained from the Niantic River were then determined by summing over all 5 -day periods. In 2001, an estimated 80.7 million larvae were entrained from the river, which represented $21.4 \%$ of the annual total entrained (Table 36).
The potential impact of larval entrainment on the winter flounder population depends upon the age of each larva at the time it is entrained, as older individuals have a greater probability to contribute to year-class strength than younger ones. Therefore, the estimated number of each developmental stage entrained during each 5 -day period was based on the proportion of each stage collected at station EN. By applying the proportion of entrainment attributed to the Niantic River (FromNR / [FromNR + Source or $\operatorname{Sink}]$ ), the number of larvae in each stage was

TABLE 36. Estimates of the total number of larval winter flounder entrained, number of larvae entrained from the Niantic River, and the percentage of total entrainment attributed to the Niantic River from 1984 through 2000.

|  | Total entrainment <br> $\left(X 10^{6}\right)^{2}$ | Niantic River <br> larval entrainment <br> $\left(X 10^{6}\right)$ | \% entrainment <br> attributed to <br> (he Niantic River |
| :---: | :---: | :---: | :---: |
| 1984 | 84.3 | 32.0 | 37.9 |
| 1985 | 79.5 | 27.8 | 34.8 |
| 1986 | 122.6 | 28.1 | 22.4 |
| 1987 | 165.0 | 41.7 | 25.3 |
| 1988 | 183.7 | 39.5 | 21.3 |
| 1989 | 167.3 | 33.5 | 20.0 |
| 1990 | 132.6 | 38.5 | 30.9 |
| 1991 | 116.4 | 35.2 | 16.3 |
| 1992 | 492.3 | 80.3 | 13.9 |
| 1993 | 41.9 | 6.0 | 28.8 |
| 1994 | 173.2 | 77.9 | 36.4 |
| 1995 | 213.6 | 30.4 | 58.8 |
| 1996 | 51.2 | 9.3 | 12.3 |
| 1997 | 75.6 | 25.9 | 30.9 |
| 1998 | 84.1 | 32.9 | 22.5 |
| 1999 | 145.9 | 87.6 | 26.3 |
| 2000 | 331.1 | 80.7 | 21.4 |
| 2001 | 376.2 |  |  |

[^21]allocated to each of the two sources (Niantic River or other) for every 5 -day period. The annual total of each larval stage entrained from either source was estimated by summing over all 5 -day periods (Table 37). In most years, including 2001, larvae in Stage 3 of development were dominated the Niantic River larvae entrained. . However, Stage 2 larvae in 1984 and 1985 and Stages 1 and 2 larvae in 1996 predominated as the larvae attributable to the Niantic River. Consistently, however, most entrained Stage 3 and 4 larvae were determined as originating from sources other than the Niantic River.
The interpretation of mass-balance calculation results has been substantiated by results from several specially designed studies. Some of the larger larvae from other areas enter the Niantic River during a flood tide, forming a stable or increased frequency of larger ( 26 mm ) size-classes during some years (Fig. 18). In special bay-wide sampling in April and May of 1991 (NUSCO 1992a), months during which about $75 \%$ of Stage 3 larvae are entrained, more larvae entered Niantic Bay from LIS east of Millstone Point and passed by the MPS intakes during a flood tide than were flushed out of the bay to LIS during an ebb tide. Therefore, greater densities of Stage 3 larvae were expected at station EN during a flood tide than during an ebb tide. To determine how consistently more Stage 3 larvae were entrained during flood tides than during ebb tides, Stage 3 larval abundances at EN were examined by tidal stage (combined data, 1983-99). Data were restricted to collections when Stage 3 larvae were present and were tested with the Wilcoxon twosample test. Stage 3 larval densities were significantly ( $p<0.001$ ) higher during a flood tide ( 570 samples) than an ebb tide ( 583 samples), indicating that most entrained Stage 3 larvae likely entered Niantic Bay from LIS.
Comparison of the Mass-balance Model and DNA-based Entrainment Fraction Estimates. Estimates of the weekly fraction of entrained winter flounder originating from the Niantic River were calculated using results from both the mass-balance model and the larval DNA analysis reported by Crivello (2002). He examined 536 mostly Stage 1 and 2 larvae collected from three source populations (Niantic River, 164; Thames River, 174; Westbrook, 198). Based on their genetic characteristics, he then assigned 1,067 Stage 2 through 4 entrained larvae to source population. The entrained larvae were collected based on a stratified random sampling model, with larger sample sizes taken during weeks of peak entrainment. Overall, $24 \%$ of the larvae examined were assigned to the Niantic River source population,
$21 \%$ were from the Thames River, $34 \%$ were from -Westbrook, and the remainder could not be classified specifically or were from unknown sources. As in the mass-balance model, Niantic River larvae were more predominant in March and April, but less so in May and June.
A more precise estimate of loss to the Niantic River population was possible by applying the fraction of larvae assigned to the river by Crivello (2002) for each sampling date to the estimated entrainment during a comparable time period. This also allowed a comparison with the mass-balance model result of total Niantic River entrainment. Estimated numbers of winter flounder larvae entrained during each 5 -day interval (Number entrained in Table 35) from March 7 through June 25 were multiplied by the calculated fraction of larvae originating from the Niantic River based on the DNA analysis of Crivello (2002) or the mass-balance model (Table 38). Because DNA sampling intervals did not match exactly with time intervals used to estimate total entrainment of larvae, DNA fraction estimates were interpolated to match 5 day entrainment estimates from Table 35. Also, the first and last estimates of Niantic River entrainment based on DNA analysis results are the summed totals for all dates prior to and after the first and last DNA sample dates, respectively. Multiplying the DNA fraction estimates by the entrainment estimates produced estimates of the 5 -day totals of larvae entrained that originated from the Niantic River. These totals were then summed to estimate the total number of Niantic River larvae entrained. The massbalance estimated fraction of entrained winter flounder larvae originating from the Niantic River was derived by calculating the fraction of entrained larvae originating from the Niantic River over each 5 -day period beginning on March 7, when larvae first appeared in samples, and extending through June 25. The 5 -day fractions were then multiplied by the number of larvae entrained during each 5 -day period. These products were then summed to produce a total estimate of entrained larvae attributable to the Niantic River stock. Correlation analysis was used to examine the strength of the relationship between DNA analysis estimates and mass-balance modeling results and a $t$-test for two population means (method for paired comparisons using weekly estimates; SAS Institute Inc. 1990b) was used to investigate the significance of the difference between the means of the two independent estimates. Both analyses indicated that weekly fractions of entrained winter flounder larvae originating from the Niantic River were higher early in the larval period and declined over the course of the spring (Fig. 45). Weekly

TABLE 37. Estimated number of winter flounder larvae entrained at MPS by developmental stage from the Niantic River and other sources, based on mass-balance calculations for 1984 through 2001. Values reported in this table differ from those reported in previous reports (e.g. NUSCO 2000) due to adjustments made in the larval period of occurrence.

| Year | Source | $\begin{aligned} & \text { Stage } 1 \\ & \left(\begin{array}{ll} \times 10^{6} \end{array}\right) \end{aligned}$ | $\begin{aligned} & \hline \text { Stage } 2 \\ & \left(\mathrm{X} 10^{\circ}\right) \end{aligned}$ | $\begin{aligned} & \text { Stage } 3 \\ & \left(\times 10^{0 \prime \prime}\right) \end{aligned}$ | $\begin{aligned} & \hline \text { Stage } 4 \\ & \left(\times 10^{6}\right) \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1984 | Niantic River | 0.2 | 14.9 | 13.9 | 3.1 |
|  | Other | 0.1 | 23.9 | 24.7 | 3.5 |
| 1985 | Niantic River | 3.4 | 17.4 | 6.9 | 0.4 |
|  | Other | 0.8 | 10.4 | 34.1 | 6.4 |
| 1986 | Niantic River | 0.7 | 7.3 | 15.2 | 4.3 |
|  | Other | 1.5 | 24.0 | 59.1 | 10.7 |
| 1987 | Niantic River | 0.8 | 15.2 | 23.9 | 1.9 |
|  | Other | 0.6 | 30.1 | 85.0 | 7.5 |
| 1988 | Niantic River | 3.9 | 9.4 | 24.4 | 1.5 |
|  | Other | 1.1 | 7.6 | 113.2 | 22.7 |
| 1989 | Niantic River | 2.8 | 11.2 | 19.0 | 0.5 |
|  | Other | 4.1 | 40.4 | 81.1 | 8.4 |
| 1990 | Niantic River | 0.9 | 6.2 | 27.6 | 3.7 |
|  | Other | 0.8 | 12.1 | 72.4 | 9.0 |
| 1991 | Niantic River | 0.2 | 3.6 | 26.7 | 4.7 |
|  | Other | 0.6 | 8.8 | 65.4 | 6.4 |
| 1992 | Niantic River | 5.6 | 10.1 | 55.7 | 8.8 |
|  | Other | 30.0 | 54.0 | 295.0 | 33.0 |
| 1993 | Niantic River | 0.3 | 1.2 | 3.8 | 0.5 |
|  | Other | 1.2 | 4.9 | 22.4 | 7.5 |
| 1994 | Niantic River | 2.8 | 12.3 | 28.7 | 6.1 |
|  | Other | 2.6 | 24.2 | 80.1 | 16.5 |
| 1995 | Niantic River | 0.6 | 6.8 | 55.6 | 14.8 |
|  | Other | 1.1 | 13.5 | 104.0 | 17.4 |
| 1996 | Niantic River | 13.2 | 12.6 | 4.1 | 0.3 |
|  | Other | 1.6 | 4.2 | 11.5 | 3.9 |
| 1997 | Niantic River | 0.7 | 1.6 | 6.1 | 0.9 |
|  | Other | 5.2 | 12.1 | 43.0 | 6.1 |
| 1998 | Niantic River | 0.9 | 5.0 | 16.3 | 3.8 |
|  | Other | 0.3 | 2.8 | 37.9 | 17.1 |
| 1999 | Niantic River | 2.8 | 9.0 | 17.2 | 3.9 |
|  | Other | 7.7 | 21.4 | 64.8 | 19.1 |
| 2000 | Niantic River | 3.5 | 20.8 | 46.3 | 16.7 |
|  | Other | 6.0 | 55.7 | 139.3 | 42.9 |
| 2001 | Niantic River | 11.1 | 25.7 | 33.7 | 11.4 |
|  | Other | 0.0 | 65.8 | 160.8 | 67.9 |

TABLE 38. Five-day estimates of the total number of winter flounder larvae entrained (see Table 35), fraction attributable to the Niantic River based on the genetic DNA analysis of Crivello (2002), the number of Niantic River larvae entrained based on this analysis, fraction attributable to the Niantic River based on mass-balance modeling results, and the number of Niantic River larvae entrained based on massbalance modeling results.

| 5-day period | 5-day entrainment estimate ( $10^{6}$ ) | DNA-based fraction ${ }^{n}$ | Total ( $10^{\prime \prime}$ ) attributed to the Niantic River from genetic study ${ }^{\text {b }}$ | Mass-balance model fraction | Total ( $10^{\prime \prime}$ ) attributed to the Niantic River from the massbalance model |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 7 March | 0.2 |  | , | 0.561 | 0.11. |
| 12 March | 0.4 | ' , |  | 0.556 | 0.22 |
| . 17 March : $\because$. | 1.0 | - . ${ }^{\text {a }}$ | . | 0.543 | 0.54 |
| ' 22 March | 2.0 |  | $\because \quad \therefore$ | - 0.517 ־ | 1.03 |
| 27 March | 4.3 | 0.462 | 3.65* | 0.479 | 2.06 |
| 1 April | 9.4 | 0.304 | 2.86 | 0.438 | 4.12 |
| - 6 April | - 13.0 | 0.146 | 1.90 | 0.395 | 5.14 |
| 11 April | 16.0 | 0.221 | 3.54 | 0.354 | 5.67 |
| - 16 April | 20.0 | 0.295 | 5.90 | 0.318 | 6.36 |
| 21 April . | 22.7 | 0.356 | 8.08 | 0.287 | 6.50 |
| 26 April | 24.9 | 0.314 | 7.82 | 0.260 | 6.47 |
| 1 May | 28.0 | 0.272 | 7.62 | 0.238 | 6.65 |
| 6 May | 27.0 | 0.162 | 4.37 | 0.219 | 5.90 |
| 11 May | 28.7 | 0.167 . | 4.79 | 0.203 | 5.82 |
| 16 May | 27.1 | 0.172 | 4.66 | 0.190 | 5.16 |
| 21 May | 25.7 | 0.217 | 5.58 | 0.179 | 4.61 |
| 26 May | 24.3 | $\because 0.188$ | 4.57 | 0.172 | 4.18 |
| 31 May | 22.3 | 0.129 : | 2.88 | 0.164 | 3.65 |
| 5 June | 20.1 | 0.069 | 1.39 | 0.159 | 3.19 |
| 10 June | 17.9 | 0.185 | 3.31 | 0.155 | 2.77 |
| 15 June | 15.8 | 0.231 | 9.54* | 0.151 | 2.39 |
| 20 June | 13.9 |  | -: • | 0.147 | 2.05 |
| 25 June | 11.6 |  | . | 0.145 | 1.68 |
| Mean fraction |  | 0.22 |  | 0.21 | . |
| Total |  |  | 82.45 |  | 80.65 |

- Bolded values are actual fraction determined from DNA analysis (Crivello 2002), while non-bold values are interpolated.
- indicates values are products of the summed tails of the entrainment estimates and the DNA fraction.
estimates of entrained winter flounder larvae originating from the Niantic River determined by the DNA analysis were significantly positively correlated with estimates from the mass-balance model ( $\mathrm{r}=$ $0.74 ; \mathrm{p}=0.003$ ). Further, the weekly estimates were not significantly different (paired $t$-test; $\mathrm{p}=0.33$ ). On average across the entire winter flounder larval season in 2001, the fraction of entrained larvae attributable to the Niantic River was 0.22 and 0.21 as estimated by the DNA analysis and mass-balance modeling, respectively. These proportions translated into 82.45 million Niantic River larvae entrained as determined by the DNA analysis and 80.65 million Niantic River larvae entrained estimated by the massbalance modeling approach.
In a separate study having the same objective of identifying the stock of origin for entrained larvae, Moran (2002), using a technique analyzing individual larvae for a number of microelements, estimated that from about 10 to $20 \%$ of entrained larvae in 2001 originated from the Niantic River. However,
due to the demands of this technique, fewer larvae were analyzed than in the Crivello (2002) study, a total of 105 larvae from the same source sites and 142 entrained larvae. Also, the entrained larvae were only collected during the period of peak entrainment (mid-April through mid-May)." For these same weeks, $26 \%$ of the entrained larvae examined by Crivello (2002) were assigned to the Niantic River population.
Estimated Production Loss from the Niantic River Stock. ". Estimates of Niantic River larvae entrained by developmental stage were compared to annual abundance estimates for each larval stage in the river. The latter were computed by applying specific mortality rates to each stage of early life history, beginning with total annual egg production estimates (Table 9), which have declined in recent years (see Spawning Stock Size and Egg Production, above). This allowed for the determination of percent production loss (i.e.,' conditional entrainment mortality, ENT) of larvae from the Niantic River



Fig. 45. Fraction (top) and numbers (bottom) of entrained winter flounder larvae attributable to the Niantic River as determined by DNA analysis (solid line) and mass-balance modeling (dashed line).
stock (Table 39). Estimates of Niantic River Stage I larvae entrained were calculated from daily abundance estimates (Eq. 3) at station C, following an evaluation presented in NUSCO (1993). This study indicated that entrainment sampling may underestimate Stage 1 larval abundance because of net extrusion and was discussed in NUSCO (1997b). As noted previously, $20 \%$ of the Niantic River discharge passes through MPS during full three-unit operation; with the retirement of Unit 1, this fraction should now be $15 \%$. For theoretical full three-unit operation, $20 \%$ of daily estimates of Stage 1 larval densities at station C were used to calculate the entrainment of these larvae from the Niantic River in the estimation of production loss, but during periods of reduced plant operation the estimates were proportionally decreased, based on daily water volume use. Entrainment estimates for Niantic River Stages 2, 3, and 4 larvae were from the results of mass-balance calculations, which used actual entrainment sampling densities. The equivalent percentage of the Niantic River winter flounder egg production entrained in 1999 and 2001 were 29.8\% and $36.1 \%$, respectively, which were the second and third largest conditional mortality rates (ENT) of the

18 -year time-series. Note that the value for 2000 is not considered reliable due to the lack of a larval mortality estimate for that year (DNC 2001a). The largest equivalent egg production losses calculated for most recent years may be inflated due to greater egg survival to hatching during since 1995 (Fig. 16). The relationship between egg production and Stage 1 larval abundance indicated that survival during recent years may have been as much as four times that prior to 1995. For annual production loss calculations, the same egg survival rate was used for the entire series, but if egg survival since 1995 was much greater than occurred previously (1984-94), this process would proportionally reduce the estimated conditional mortality (ENT). This makes estimates of ENT for the past 6 years conservatively high. An example is 1999 , for which a reduction of $75 \%$ in egg mortality would reduce ENT from nearly $33 \%$ to about $8 \%$.
The total entrainment estimate in 2001 was the highest since 1992 , representing a $13 \%$ increase over 2000 and was more than 2.5 times the estimate for 1999 (Table 36). Nevertheless, entrainment estimates from 1996 through 1998 were the lowest since Unit 3 went online, except for 1993, when larvae were very low in abundance. Since 1977, annual egg production estimates in the river were the lowest from 1995 through 2001 (Table 9), but near-average (1996 and 2001) or high (1995 and 1997-99) larval abundance in the bay (Table 10) suggested increasing survival during early life history, which probably resulted from several factors. Egg survival for these years was apparently better than usual, as many more Stage 1 larvae were found than were expected (Fig. 16). Larval mortality rates in the river during 1995, 1997, 1998, and 1999 were among the lowest estimates found, although the mortality rate in 1996 was greater than the long-term average (Table 15). The larval recruitment indices for 1995-99 and 2001 at station EN (Fig. 23) were the highest calculated. Larval development is directly related to growth rate and shorter larval periods likely result in better survival (Houde 1987). For example, during 1995 and 1997-99, larval growth and development were among the fastest in comparison to previous years, as indicated by greater annual growth rates in the river (Table 14), had the largest mean length during the first 2 weeks of April (Fig. 20), and some of the earliest dates of peak abundance (Fig. 21). Higher larval survival in these years, which appeared to be related to faster growth and development, apparently also resulted in more Stage 3 larvae available for entrainment during the same years.

TABLE 39. Estimated abundance of winter flounder larvae in the Niantic River and the number and percentage of the production entrained from the Niantic River (ENT) by developmental stage from 1984 through 2001. Numbers of larvae entrained from the Niantic River were based on mass-balance calculations (see text) and differ from those reported in previous reports (e.g. NUSCO 2000) due to adjustments made in larval period of occurrence.

| Year | Stage of development | Niantic River abundance (X $10^{6}$ ) | Entrainment from the Niantic River (X $10^{\prime \prime}$ ) | \% production entrained (ENT) |
| :---: | :---: | :---: | :---: | :---: |
| 1984 | Stage 1 | 2,801 | 9.8 | 0.3 |
|  | Stage 2 | 670 | 14.9 | 2.2 |
|  | Stage 3 | 330 | 13.9 | 4.2 |
|  | Stage 4 | 230 | 3.1 | 1.3 |
|  | Total |  | 41.7 | 8.0 |
| 1985 | Stage 1 | 3,036 | 15.1 | 0.5 |
|  | Stage 2 | 727 | 17.4 | 2.4 |
|  | Stage 3 | 357 | 6.9 | 1.9 |
| , | Stage 4 | 249 | 0.4 | 0.2 |
|  | Total | . | 39.8 | 5.0 |
| 1986 | Stage 1 | 2,374 | 10.9 | 0.5 |
|  | Stage 2 | 667 | 7.3 | 1.1 |
|  | Stage 3 | 348 | 15.2 | 4.4 |
|  | Stage 4 | 243 | 4.3 | 1.8 |
|  | Total |  | 37.7 | 7.8 |
| 1987 | Stage I | 2,865 | 33.0 | 1.2 |
|  | Stage 2 | 805 | 15.2 | 1.9 |
|  | Stage 3 | 420 | 23.9 | 5.7 |
|  | Stage 4 | 293 | 1.9 | 0.6 |
|  | Total |  | 74.0 | 9.4 |
| 1988 | Stage 1 | 4,676 | 79.6 | 1.7 |
|  | Stage 2 | 699 | 9.4 | 1.3 |
|  | Stage 3 | 252 | 24.4 | 9.7 |
|  | Stage 4 | 181 | 1.5 | 0.8 |
|  | Total |  | 114.9 | 13.5 |
| 1989 | Stage 1 | 3,816 | 63.6 | 1.7 |
|  | Stage 2 | 532 | 11.2 | 2.1 |
|  | Stage 3 | 175 | 19.0 | 10.9 |
|  | Stage 4 | 118 | 0.5 | 0.4 |
|  | Total | ; | 94.3 | 15.0 |
| 1990 | Stage I | 1,854 | 31.7 | 1.7 |
|  | Stage 2 | 761 | 6.2 | 0.8 |
|  | Stage 3 | 210 | 27.6 | 13.2 |
|  | Stage 4 | 180 | 3.7 | 2.0 |
|  | Total |  | 69.1 | 17.7 |
| 1991 | Stage I | 3,413 | 12.6 | 0.4 |
|  | Stage 2 | 2,381 | 3.6 | 0.2 |
|  | Stage 3 | 724 | 26.7 | 3.7 |
|  | Stage 4 | 587 | 4.7 | 0.8 |
|  | Total | ., | 47.6 | 5.0 |
| 1992 | Stage 1 | 2,089 | 22.0 | 1.1 |
|  | Stage 2 | 875 | 10.1 | 1.2 |
|  | Stage 3 | 322 | 55.7 | 17.3 |
|  | Stage 4 | 258 | 8.8 | 3.4 |
| - | Total |  | 96.6 | 22.9 |

TABLE 39. (cont.).

| Year | Stage of development | Niantic River abundance (X $10^{6}$ ) | Entrainment from the Niantic River ${ }^{\text {a }}$ (X $10^{6}$ ) | \% production entrained (ENT) |
| :---: | :---: | :---: | :---: | :---: |
| 1993 | Stage 1 | 1,189 | 10.9 | 0.9 |
|  | Stage 2 | 614 | 1.2 | 0.2 |
|  | Stage 3 | 111 | 3.8 | 3.4 |
|  | Stage 4 | 78 | 0.5 | 0.6 |
|  | Total |  | 16.4 | 5.1 |
| 1994 | Stage 1 | 1,754 | 26.1 | 1.5 |
|  | Stage 2 | 962 | 12.3 | 1.3 |
|  | Stage 3 | 483 | 28.7 | 6.0 |
|  | Stage 4 | 420 | 6.1 | 1.5 |
|  | Total |  | 73.2 | 10.2 |
| 1995 | Stage 1 | 874 | 37.9 | 4.3 |
|  | Stage 2 | 551 | 6.8 | 1.2 |
|  | Stage 3 | 229 | 55.6 | 24.3 |
|  | Stage 4 | 153 | 14.8 | 9.6 |
|  | Total |  | 115.1 | 39.5 |
| 1996 | Stage 1 | 438 | 23.1 | 5.3 |
|  | Stage 2 | 155 | 12.6 | 8.1 |
|  | Stage 3 | 48 | 4.1 | 8.5 |
|  | Stage 4 | 33 | 0.3 | 0.9 |
|  | Total |  | 40.0 | 22.8 |
| 1997 | Stage 1 | 777 | 5.6 | 0.7 |
|  | Stage 2 | 511 | 1.6 | 0.3 |
|  | Stage 3 | 267 | 6.1 | 2.3 |
|  | Stage 4 | 151 | 0.9 | 0.6 |
|  | Total |  | 14.2 | 3.9 |
| 1998 | Stage 1 | 525 | 5.4 | 1.0 |
|  | Stage 2 | 312 | 5.0 | 1.6 |
|  | Stage 3 | 146 | 16.3 | 11.1 |
|  | Stage 4 | 84 | 3.8 | 4.5 |
|  | Total |  | 30.5 | 18.3 |
| 1999 | Stage I | 442 | 16.8 | 3.8 |
|  | Stage 2 | 169 | 9.0 | 5.3 |
|  | Stage 3 | 106 | 17.2 | 16.3 |
|  | Stage 4 | 89 | 3.9 | 4.4 |
|  | Total |  | 46.9 | 29.8 |
| $2000^{6}$ | Stage ! | 725 | 5.8 | 0.8 |
|  | Stage 2 | 267 | 20.8 | 7.8 |
|  | Stage 3 | 146 | 46.3 | 31.6 |
|  | Stage 4 | 120 | 16.7 | 13.9 |
|  | Total |  | 89.6 | 54.1 |
| 2001 | Stage 1 | 333 | 11.4 | 3.4 |
|  | Stage 2 | 270 | 25.7 | 9.5 |
|  | Stage 3 | 207 | 33.7 | 16.3 |
|  | Stage 4 | 165 | 11.4 | 6.9 |
|  | Total |  | 82.2 | 36.1 |
| Geometric mean |  |  |  | 11.5 |

[^22]
## Stochastic Simulation of the Niantic River Winter Flounder Stock

Modeling Strategy and Background. The SPDM developed for the Niantic River winter flounder stock includes a life-cycle module designed to generate new year classes of flounder that is based on the three-parameter Ricker form of the stock-recruitment relationship (SRR) given as Equation 7. Although the SRR equation does not appear explicitly in the model formulation, the SPDM equation which computes mortality through the first year of winter flounder life was derived from the SRR and assumes a Ricker-type form of recruitment. Beyond that point (i.e., age-1) in the life-cycle simulation, the population model simply keeps track of the annual changes in abundance of each year-class resulting from natural mortality and fishing, together with growth in size and increased fecundity. Population updates take place at the beginning of each model time-step of length equal to 1 year. The projection of adult fish populations over time has been implemented in many models by means of Leslie matrix equations (e.g., Hess et al. 1975; Saila and Lorda 1977; Vaughan 1981; Spaulding et al. 1983; Goodyear and Christensen 1984; Reed et al. 1984): In the SPDM, winter flounder were projected over time by grouping fish into distinct age-classes and by carrying out the computations needed (mostly additions and multiplications) iteratively over the age index ( 1 through 15) and over the number of years specified for each simulation. This approach was algebraically identical to the Leslie matrix formulation, which helps to conceptualize the model, and simplifies the computer code when describing the fish population either as numbers of fish or as biomass (allowing for size variation within each ageclass). A similar implementation of an adult fish population dynamics simulation was used by Crecco and Savoy (1987) in their model of Connecticut River American shad (Alosa sapidissima). $\because$. The Niantic' River winter flounder: SPDM and its application to impact assessment work at MPS was described in Lorda et al. (2000).

Model Components. The solid-line boxes in Figure 46 describe the SPDM as presently used, while the box with dashed lines corresponds to the previously. described - mass-balance calculations dealing with spatial larval distribution and entrainment loss estimates, which are not an integral part of the SPDM. Model components are briefly described in Figure 46, with' the more important model components, such as the one labeled age- 0 cohort
(box 3) and the two random inputs (boxes 3 A and 3B), described in more detail below.
The most critical aspects in the formulation of a stock-recruitment based population model are the specific equation and parameter estimates used to calculate total mortality during the first year of life of the fish (i.e., from egg to age-1). The equation used for this purpose in the SPDM was derived from Ricker's equilibrium equation for $\mathrm{Z}_{0}$ (total instantaneous mortality from egg to maturation age). This involved the extension of stock-recruitment theory, which was developed for fish that spawn only once, to iteroparous fish with multi-age spawning stocks.
The form of the equation as used in the present model is:

$$
\begin{align*}
\mathrm{Z}_{0, t}= & \log _{e}(\mathrm{FEC})+\log _{e}(\mathrm{ASF})-\log _{e}(\alpha)+\mathrm{n}_{t}- \\
& (\phi \times \mathrm{WT})-\mathrm{Z}_{1.2}+\left(\beta \times \mathrm{P}_{t}\right) \tag{19}
\end{align*}
$$

where the subscript $t$ denotes the time-step (each time-step represents a year) and non-subscripted terms remain constant from year to year; $\alpha, \beta$, and $\phi$ are the parameters of the SR function (see Eq. 7), but with $\alpha=\alpha_{0}$, the theoretical rate of increase in the absence of fishing, estimated from winter flounder life history parameters independently of the Niantic River stock and recruitment data (the derivation of $\alpha_{0}$ was discussed in the Stock and Recruitment section); FEC is the mean fecundity of the stock expressed as the number of female eggs produced per female spawner; ASF is a scaling factor to adjust $\alpha$ for the effect of a multi-age spawning stock; $\mathrm{n}_{t}$ and $\mathrm{WT}_{l}$ are independent random variates from two specified normal distributions described below; $\mathbf{Z}_{\mathrm{t}, 2}$ is the instantaneous mortality rate through the immature age-classes; and the last term ( $\beta \times \mathrm{P}_{\ell}$ ) is a feed-back mechanism that simulates stock-dependent compensatory mortality, which varies according to the size of the annual spawning stock $\mathrm{P}_{\mathrm{t}}$. The complete derivation of the above equation was given in NUSCO (1990: appendix to the winter flounder section). The scaling factor ASF is a multiplier that converts age-3 female recruits into their spawning potential throughout their Jifetimes. This spawning potential is defined as the cumulative number of mature females from the same year-class that survive to spawn year after year during the lifetime of the fish. The algebraic form of this multiplier is identical to the numerator of Equation A-4 in Christensen and , Goodyear (1988).
When simulating plant effects such as entrainment, the parameter $\alpha$ in Equation 19 is reduced by a factor equivalent to the projected reduction in annual


Fig. 46. Diagram of the stochastic fish population dynamics simulation model (SPDM) used to assess long-term effects of larval winter flounder entrainment at MPS. The computer implementation of the SPDM is in Fortran77. Brief descriptions of the computer program components referenced in the diagram follow:

1. Subprogram that process the input data files (see Table 40 for a sample of the main input file). The subprogram also verifies data ranges and stores parameters and data used for each simulation in common memory.
2. Auxiliary programs (not an integral part of SPDM) that estimate the annual fraction of Niantic River flounder production lost to larval entrainment. These estimates are based annual larval production, cooling water flow at the plant intakes, tidal exchange rates between Niantic Bay and River, and mass-balance calculations of weekly larval densities near the plant intakes in Niantic Bay.
3. Subprogram that calculates the number of young fish surviving to the end of their first year of life. Natural and compensatory mortality are described using a Ricker-type recruitment equation incorporating a temperaturedependent term (Eq. 7). Survival reduction due to larval entrainment is explicitly described as additional mortality (Eq. 19).
3A. Subroutine that generates random normal deviates of mean water temperature with given mean and variance derived from water temperature data during critical larval development (input to Eq. 19 in box 3).
3B. Subroutine that generates random standard normal deviates to simulate random variability in the natural mortality rates of early life stages of winter flounder (input to Eq. 19 in box 3).
4. Subprogram that updates the numbers of adult fish in each age group at the end of each time-step or year. This process is implemented with a Leslie matrix which accepts random variation in selected parameters. Natural and fishing mortality rates in addition to fish losses caused by impingement are used in the calculations.
5. Subroutine that calculates annual egg production from fecundity-at-age and the annually updated age structure calculated in box 4. This annual egg production is the population feed-back that starts each new cycle in box 3.
6. Subroutine that summarizes the adult population numbers and annual catch as biomass by age-class.
7. Subprogram that conducts a probabilistic risk analysis when the population dynamics is simulated as an stochastic process. Reference biological points "at risk" of being exceeded are provided with the initial input.
production. A similar scaling of $\alpha$ was described by Myers et al. (1999) in dealing with the estimation of $\alpha$ with low abundance of spawners. Stochasticity in the winter flounder model (Fig: 46) has two annual components: :a random term : $\mathrm{n}_{t}$ ) that represents uncertainties associated with the estimate of Ricker's $\alpha_{10}$ parameter and environmental variability in the form of random deviations ( $\phi \times \mathrm{WT}_{t}$ ) from the longterm mean February water temperature. These two components of annual variability are incorporated into the calculation of each new year-class via the mortality from egg to age-1 (Eq. 19). The random noise term $n_{t}$ is simulated as independent random variates from a normal distribution with zero mean and variance, equal to $\sigma^{2}$. The value of $\sigma$ was estimated during the model calibration runs as the amount of variance required to generate short timeseries of projected spawning stocks with a similar CV as that observed in field data. Similarly, the term ( $\phi$ $\times \mathrm{WT}_{t}$ ) represents the effect of annual variability of February water temperatures on larval survival. This effect becomes random when the input February water temperatures are generated as independent random variates from a normal distribution with mean and variance equal to the mean and variance of February water temperatures at the MPS intakes from 1977 through the current year.
A stochastic simulation of winter flounder population dynamics also provides the proper framework for applying probabilistic risk assessment methodology (box 7 of Fig. 46). The assessment is based on Monte Carlo methods (Rubinstein 1981), where many independent random replicates of the timeseries are generated so that the mean of the series and its standard error can be estimated. This simulation is a relatively simple application of Monte Carlo methods because only one population parameter, the age-0 survival rate, is stochastic. For this application of the SPDM, 100 replicate simulations per scenario were used, which was judged to be sufficient, given the amount of variability found (NUSCO 1990). Also, as noted in DNC (2001c), increasing the number of replicates from 500 to 1,000 produced very little additional variation in the SPDM output.
Monte Carlo replications of the projected stock size for any given year can also be used to derive the empirical cumulative frequency distribution (cdf) for that year without assuming any particular statistical distribution. This methodology was used to assess the risk of postulated stock reductions resulting from entrainment and impingement at MPS. The probabilities of stock reductions below some specified reference size were empirically derived from the cdf of stock size replications in selected years. Addition-
ally, approximate $95 \% \mathrm{Cls}$ associated with point estimates of annual stock sizes could be calculated.
Model Assumptions and Limitations. Major assumptions of the SPDM relate to the underlying form of the SRR used and the reliability of the SRR parameter estimates. Because the SPDM incorporated the Ricker form of SRR;'. it was assumed that stock-dependent compensation and the postulated effect of water temperature on larval survival (Eqs. 7 and 19) applied reasonably well to the Niantic River winter flounder stock. A second assumption was that the $\beta$ and $\phi$ parameters of the SRR could be estimated from annual time-series of field data and that, in particular the value of $\alpha_{0}$, which was based on life history parameters only, was a reasonable estimate.
Although the population was not assumed to be at steady state, the average ;ecundity and annual survival rates for fish age-1 and older were assumed to remain fairly stable over the period corresponding to the time-series data used to estimate the SRR parameters. Although this last assumption can generally be met in the case of fecundity rates and adult natural mortality, fishing mortality rates for winter flounder have been much less stable. Changes in exploitation rates from year to year should not cause estimation problems as long as the changes are not systematic (i.e., change in the same direction year after year). Because these assumptions are seldom completely met, early applications of the model (NUSCO 1990) included calibration runs to validate predictions under both deterministic and stochastic modes by comparing model results to recent series of stock abundance data.
Another assumption is that no temperature trend or large-scale environmental changes (e.g., global warming) has or will occur during the years simulated in each population projection. However, this assumption may not be entirely accurate as there has been a consistent pattern of warmer than average water temperatures in LIS during late winter and early spring in recent years (Foertch 2000; Fig. 47). Because these periods coincide with winter flounder spawning," egg incubation, larval development, metamorphosis and settling, and early demersal life, temperature-dependent effects could affect the reproductive success of winter flounder and its ultimate population size (see Formation of Winter Flounder Year-class Strength for a discussion of temperature effects). Finally, no trends in fishing rate were assumed to occur beyond 1999 in these simulations. However, this last assumption is very unrealistic, given the recent 20 -year history of dramatic changes in fishing rates that is presented below.


Fig. 47. Annual mean water temperature ( ${ }^{\circ} \mathrm{F}$ ) during the first (January-March) and second (April-June) quarters of the year calculated from average daily mean temperature at the intakes of MPS Units 1 and 2 from 1976 through 2000 (points connected by solid line). A regression line (solid line) with $95 \%$ confidence interval (dashed lines) determined for each series of annual quarterly means is also shown.

Model Input Data. The dynamics of the Niantic River winter flounder stock were simulated using the SPDM under a real-time scenario running from 1960, a decade before the operation of Unit 1 , to 2060, which encompasses a recovery period 15 years after
the projected shutdown of Unit 3 in 2045. This time period assumes that a 20 -year license renewal for Units 2 and 3 will have been received from the NRC. The scenarios used power plant effects that were based on actual or projected operation in each year
concurrently with estimates of $F$ based on historic or projected rates of commercial exploitation and sport fishing for winter flounder in Connecticut. Parameters used in the SPDM include: F , with an additional instantaneous mortality of 0.01 that accounts for impingement (IMP) losses (NUSCO 1992a); larval entrainment conditional mortality rates (i.e., ENT, the fraction of the annual production of Niantic River winter flounder removed as a result of power plant operation; discussed further below) estimated from the mass-balance calculations given previously in Tables 36, 37, and 39; a schedule of changes when any of these rates was not assumed constant; and the length of the time-series in years. The combined mortality of $\mathrm{F}+\mathrm{IMP}$ was used only during the simulation period (1971-2045) that corresponded to MPS operation. Based on improvements to the fish return at Unit 3 and the installation of a fish return at Unit 2, even this small value may be an overestimate of current impingement impact. Other data, rates, and inputs to the SPDM are summarized in Table 40 and included the number of age-classes, age-specific rates of maturation, natural mortality, average weight and fecundity at age, the estimated value of $\alpha_{0}$, longterm average estimates of $\beta$ and $\phi$, February water temperature statistics, and specific factors set for each simulation.

Because the ability of a fish stock to withstand additional stress is reduced by fishing mortality (Goodyear 1980), the actual fishing exploitation rates on the spawning stock are critically important in simulations of long-term effects of larval entrainment, particularly as F has been 3 to 7 times as much as natural mortality ( 0.2 ) of adult winter flounder in recent decades and remains the dominant force driving the dynamics of the spawning stock. The annual schedule of nominal $F$ values was based on DEP-generated estimates (D. Simpson, CT DEP, Old Lyme, CT, pers. comm.), for which the time-series begins in 1984. This schedule of $F$ rates was also used in DNC (2001c), a report evaluating cooling water system alternatives at MPS. These exploitation rates take into account both fishing effort and the effects of length-limits and other regulations implemented by the DEP to reduce fishing mortality in Connecticut waters. Values of $F$ used in the simulations were stepped up from 0.40 in the $1960 \mathrm{~s}^{\circ}$ and 1970s, prior to MPS operation, and reached a peak of 1.517 in 1990 (Table 41; Fig. 48), reflecting an historical period of high exploitation of winter flounder in the 1980s and 1990s (NEFSC 1999). Subsequently, estimates of $F$ decreased during the late 1990s. Based on discussions held with DEP (D. Simpson, CT DEP, Old Lyme, CT, pers. comm.)
prior to preparing DNC (2001c), the most recent estimate of fishing mortality (0.74) was based on the mean of the last available annual point estimate of $F$ provided by DEP (1999-2000) and the last available "three-point moving average of $F$ determined for age4+ winter flounder in LIS. This value was assigned to 1999 and subsequent years in simulating winter flounder stock size projections into the future. As a result of protective regulations, age-classes 1 through 3 are not fully vulnerable to fishing mortality, particularly in more recent ..years, so correction factors (Table 42) were used to reduce the annual values of $F$ for these ages that are found on Table 41.
The potential impact of entrainment depends upon the age of each larva at the time it is entrained, because an older individual has a grater probability of surviving to reproductive age than a younger one. The number of individuals entrained from each developmental stage was estimated from the proportion of each stage collected at the MPS discharge. Since the proportion of entrainment attributed to the Niantic River was estimated from massbalance calculations, the number of larvae for each stage could be allocated to either the Niantic River or other sources (Table 36). Most of the Stage 3 larvae entrained (the predominant stage collected at the MPS discharges) apparently originated from sources other than the Niantic River. Conditional mortality rates (i.e., production loss estimates) for larval entrainment (ENT) from 1984 through 2001 used in SPDM simulations (Table 41) were estimated directly using the mass-balance calculations under actual MPS operating conditions during these years (Table 39). The " conservative nature of the mass-balance calculations was previously discussed as were the larval stock identification studies (Crivello 2002; Moran 2002), which independently verified that fractions of entrained larvae attributed to the Niantic River winter flounder stock in 2001 were similar among the two studies and the mass-balance calculation. Larval losses for early years of plant operation (1971-83) were simulated in NUSCO (2000) by modifying a randomly chosen value of ENT from 1984 : through 1995 proportional to condenser cooling-water flows at MPS actually used during, the annual April 1-June 15 larval winter flounder season in those years and these values are given in Table 41. For years going forward in the simulations, a geometric mean ENT of $14.17 \%$ was calculated for the time-series of production loss estimates using the annual values for 1986-95 (years of MPS three-unit operation), 1999, and 2001. Values for the extended shutdown years of 1996-98 were excluded, as was the estimate for 2000 , which

TABLE 40. Data, rates, and other inputs used with the Niantic River winter flounder population dynamics simulation model (SPDM).

| Model input | Value used or available |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Number of age-classes in population |  | 15 |  |  |  |
| Earliest age at which all females are mature |  | 5 |  |  |  |
| Fraction mature, mean wt (lbs), and mean fecundity by age: |  |  |  |  |  |
| Age-1 females | 0 | 0.011 | 0 |  |  |
| Age-2 females | 0 | 0.125 | 0 |  |  |
| Age-3 females | 0.25 | 0.497 | 232,088 |  |  |
| Age-4 females | 0.80 | 0.776 | 432,517 |  |  |
| Age-5 females | 1.00 | 1.096 | 700,390 |  |  |
| Age-6 females | 1.00 | 1.435 | 1,019,793 |  |  |
| Age-7 females | 1.00 | 1.628 | 1,216,926 |  |  |
| Age-8 females | 1.00 | 1.839 | 1,442,512 |  |  |
| Age-9 females | 1.00 | 2.068 | 1,699,372 |  |  |
| Age-10 females | 1.00 | 2.316 | 1,990,489 |  |  |
| Age-11 females | 1.00 | 2.584 | 2,319,011 | $\checkmark$ |  |
| Age-12 females | 1.00 | 2.872 | 2,688,253 |  |  |
| Age-13 females | 1.00 | 3.182 | 3,101,703 |  |  |
| Age-14 females | 1.00 | 3.514 | 3,563,020 |  |  |
| Age-15 females | 1.00 | 3.869 | 4,076,040 |  |  |
| Age after which total annual mortality is constant |  | 4 |  |  |  |
| Instantaneous mortality rates $M$ and $F$ at age-1 |  | $5^{6} \mathrm{~F} \times$ |  |  |  |
| Instantaneous mortality rates M and F at age-2 |  | F $\times$ |  |  |  |
| Instantancous mortality rates M and F at age-3 |  | F |  |  |  |
| Instantaneous mortality rates $M$ and $F$ at age-4+ |  | F $\times$ |  |  |  |
| Initial number of female spawners |  | 72,532 |  |  |  |
| Mean fecundity of the stock (eggs per female spawner) |  | 322,99 |  |  |  |
| $\alpha_{0}$ for the unfished ( $\mathrm{F}=0$ ) stock |  | $5.20{ }^{\text {r }}$ |  |  |  |
| $\beta$ from the three-parameter SRR (mean value from 1989-2001) ${ }^{\text {c }}$ |  | $273 \times 1$ |  |  |  |
| $\phi$ from the three-parameter SRR (mean value from 1989-2001) ${ }^{\text {d }}$ |  | -0.373 |  |  |  |
| Mean February (1977-2001) water temperature ( ${ }^{\circ} \mathrm{C}$ ) |  | 2.99 |  |  |  |
| standard deviation |  | 1.13 |  |  |  |
| minimum temperature |  | 0.36 |  |  |  |
| maximum temperature |  | 4.76 |  |  |  |

- See Table 30 for the derivation of weight and fecundity at age.
- Re-calculated from the value of 0.5 used in NUSCO (2000) and based on a simulation used to calibrate model population size to that measured by sampling in the annual Niantic River winter flounder population spawning surveys (see Figure 50 and text for further explanation).
- Values of $M$ remain constant during all spawning cycles or years simulated. Fish at ages-1-3 are partially recruited and the multipliers shown ( $\geq 1987$; see Table 42 for previous years) are used to reduce $F$ accordingly (see Table 41 for annual values of $F$ used in the SPDM).
- Corresponds to the unfished stock at equilibrium, $\mathrm{P}_{\text {rp }}$ (see Eq. 12).
- Calculated for the Niantic River winter flounder female spawning stock at equilibrium in the absence of fishing (see Table 30).
r Indirectly calculated from life history parameters (see Stock-recruitment Relationship and Table 29).
- See Table 28.

TABLE 41. Schedule of conditional entrainment (ENT values) and fishing mortality rates ( $\mathbf{F}$ ) adjusted for impingement (IMP), including values of $F$ for less than fully vulnerable age-classes, as implemented in the current SPDM simulations.


- See text for explanation of values used.
- See Table 42 for adjustment factors.
although given in Table 39, was computed using a different methodology' due to a lack of a larval mortality estimate that year. Because of this, the geometric mean value was also used for 2000 in the simulations. This mean value was further reduced (based on unit cooling water demand) to $8.53 \%$ in 2035 following the projected retirement of Unit 2 and to 0 in 2045 after the projected retirement of Unit 3. Because there is .still some uncertainty regarding annual estimates of ENT, separate model projections were also made using both "high" and "low" mean values of ENT for 2002-2060. These were derived
by scaling the corresponding mean value of ENT by a factor of 1.5 and $1 / 1.5$, respectively (Table 41).
These :ranges were somewhat greater than values within two standard errors of the mean ENT.
Model Output and Calibration. :- All stock projections were given in units of spawning biomass (lbs) because overfishing criteria often rely on assessments of biomass, which tend to be more conservative, than those based on fish numbers. Furthermore, larval entrainment effects result in long-term stock reductions which can be quite different depending on whether the stock is expressed


Fig. 48. Historic and projected annual instantaneous mortality rate due to fishing ( F ), determined in consultation with CT DEP, plus a small ( 0.01 ) component accounting for impingement mortality (IMP) at MPS as implemented in the SPDM simulations for this report (see Table 41). The vertical dashed lines illustrate periods of actual or projected one-, two-, and three-unit operation at MPS with an assumed 20-year license renewal for Units 2 and 3, which would extend their operation to 2035 and 2045 , respectively.

TABLE 42. Factors used to adjust fishing mortality for less than fully vulnerable age-classes (ages-1-3) as used in the Niantic River winter flounder SPDM.

| Period | Age-1 | Age-2 | Age-3 |
| :---: | :---: | :---: | :---: |
| $\leq 1981$ | 0.04 | 0.75 | 1.00 |
| $1982-86$ | 0.02 | 0.50 | 1.00 |
| $\geq 1987$ | 0.02 | 0.25 | 0.60 |

as fish numbers or as biomass. Population reproductive capacity is more accurately reflected by biomass, which takes into account the size of individual females (egg production is a function of length or weight), as well as the number of spawners. Annual abundance of females by $0.5-\mathrm{cm}$ size-class was converted to weight (lbs) using a length-weight relationship determined for Niantic River winter flounder:

$$
\begin{equation*}
\text { weight }=\left(1.20173 \times 10^{-5}\right) \times(\text { length })^{3.226} \tag{20}
\end{equation*}
$$

and an annual mean weight was computed from 1977 through 2001. An overall mean weight was determined as 1.08 lbs , so during this period both number and weight were nearly synonymous (Fig. 49).


Fig. 49. Annual mean weight (lbs) of Niantic River female winter flounder spawners from 1977 through 2001. The overall grand mean weight (dashed line) was 1.08 lbs .

Although annual mean weight from 1995 through 2001 averaged about 1.3 lbs , this value would most likely decrease towards 1 lb if population size increased, since mean weight was determined to be almost exactly 1 lb during 1977-85, when winter flounder were more abundant than at present.
A complete simulation of MPS impact consisted of three stochastic time-series of female spawning stock sizes generated during separate and independent

SPDM simulation runs completed under a specified set of population parameters and plant operation conditions, including random variability. There were three basic scenarios: 1) a theoretical unfished stock, whose size was dependent only upon natural mortality and the dynamics of winter flounder reproduction and environmental variability (i.e., with no fishing or plant operational effects); 2) an exploited stock with reduced biomass as a result of fishing, but still with no power plant effects (i.e., the baseline time-series without MPS effects); and 3) a stock with a biomass reduced by all three types of anthropogenic mortality ( $\mathrm{F}, \mathrm{IMP}$, and ENT) occurring along with natural variation (i.e., the impacted stock). The first time-series with no fishing or plant effects was the reference series against which the potential for recruitment failure was evaluated when the largest reductions of stock biomass occurred during any of the other simulations. The second time-series represented the most likely trajectory of the exploited stock without MPS operation. The third time-series provided'a basis for quantitatively assessing MPS impact on the Niantic River winter flounder population. As noted, three subsets of the impacted stock were determined using the mean, low, and high values of ENT for the projections.
Prior to carrying out SPDM simulations presented in DNC (2001c), several calibration runs were conducted to insure that reasonable agreement existed between the population projections and recent abundance estimates; of the Niantic River winter flounder female spawning stock. Population parameter estimates used by the model were reviewed and updated such that model output would match more closely the observed abundance time-series. : The instantaneous natural annual mortality rate (M) of 0.50 for age-1 flounder used in previous applications of the SPDM (e.g., NUSCO 2000) was perhaps too low since it amounts to a monthly attrition rate of only $4.1 \%$ (annual survival $S=e^{-M}$ ). Field sampling showed consistently that numbers of age-0 juvenile winter flounder decline rapidly during the 4 months following metamorphosis and settlement (see Age-0 Juveniles during Summer), with a 17 -year (19842000) mean survival rate of only $9.2 \%$ through this 4 -month period, or an average monthly attrition rate of $45 \%$. Specific information is lacking to calculate a mortality rate during late fall and winter and when this group of fish becomes age-1 in the following spring. Although $M$ most likely decreases as these fish grow older and larger, mortality of these older immature fish nevertheless remains higher than that of adult fish. Data from a study of Mystic River, CT winter flounder (Pearcy 1962) that were discussed by

Klein-MacPhee (1978) in a synopsis of biological data for winter flounder suggested a total survival rate of 0.41 for winter flounder between the ages 12.4 and 22.4 months. Since this survival rate is equivalent to an annual natural mortality rate of $M=$ 1.07, or twice the rate formerly used in SPDM applications, it was decided to calibrate the model output by progressively increasing the value of $\mathrm{M}=$ 0.50 for age-1 winter flounder. After a few trial runs of the SPDM using the schedule of fishing rates (Table 41) and the same annual production losses (ENT) as given in NUSCO (2000), the model output converged to current stock levels as shown in Figure 50.. At the point in the calibration process at which model output matched field data-based adult stock sizes during the last 4 years, the simulated M for age1 flounder was 0.8425 . Since this value appeared reasonable, although still less than the estimate of Pearcy (1962), it was adopted as a better estimate for this and future model applications. Rates of M for older winter flounder remained unchanged from previous SPDM applications (Table 40).

Simulation Results. Plots of the stochastic variability for both baseline and impacted (mean ENT rate) stocks show the $95 \%$ CI relatively close to the mean, but the minimum and maximum replicates, which were a measure of the random variation in the simulations, had a relatively wide range during the initial (approximately 25-42 thousand lbs) years of the scenarios (Fig. 51). The smallest differences found between the minimum and maximum stock sizes were about 7.4 thousand lbs for the baseline and 5.9 thousand lbs for the impacted stock, each occurring in 1994, when biomass were near or at the lowest points of each simulation.
Based on the age and size structure of an unfished female winter flounder stock at equilibrium (Table 30), the unfished stock size used initially in all simulations was $119,678 \mathrm{lbs}$ (value of $\mathrm{P}_{\text {rep }}$ ), which was equivalent to 72,532 female spawners (Table 40; see also Stock-Recruitment Relationship, above).
This initial stock size represented the maximum spawning potential (MSP) for the unfished Niantic River female spawning stock, a biological reference point discussed in greater detail below. The geometric mean estimate of MSP from the SPDM simulations was $119,972 \mathrm{lbs}$, which was remarkably similar to the "deterministic estimate of $P_{\text {rep }}$ used to initiate the model runs. A critical stock size ( $25 \%$ of the MSP) was determined as $29,993 \mathrm{lbs}$ for these simulations, which is shown as the horizontal line in Figure 51. Allowing for natural variation in the simulation, all replicates, even the largest values, were less than $25 \%$ of MSP for the baseline stock


Fig. 50. Niantic River female winter tlounder biomass as projected with the SPDM using DEP fishing mortality rates for the purpose of calibrating the modeled time-series to recently observed numbers and biomass as estimated from annual spawning surveys. A natural mortality rate of 0.8425 was used for age-1 winter flounder in this model projection.
beginning in 1987 for the baseline stock and in 1985 for the impacted stock, with nearly all replicates remaining less than this critical value for the duration of the simulated time-series.
For the simulated baseline projection (shown as the solid line in Fig. 52), the stochastic mean size of the exploited stock under the starting nominal fishing rate of $F=0.40$ was quickly reduced to about 30 thousand lbs by 1980. The baseline responded as expected to the high rates of fishing through the mid1990s and the stock steadily declined to its lowest point of $5,630 \mathrm{lbs}$ in 1994, only about $19 \%$ of the critical stock size. With some reduction in F beginning in 1999 and holding this value at 0.74 from 1999 until the end of the simulation time-series allowed the stock to recover to some extent, but mean stock sizes mostly remained between 12 and 13 thousand lbs through 2060.
To determine the effect of MPS on the Niantic River female spawning stock, the baseline time-series is compared to the impacted (in this case, the mean ENT rate) time-series, shown as the dashed line in Figure 52. The impacted series corresponds to projections of the baseline stock, but with additional annual losses due to MPS operation (i.e., ENT + IMP), the effects of which first begin to be seen in 1974. Similar to the baseline, the lowest projected
stock biomass levels (about 4.3 thousand lbs) were attained in the mid-1990s. Absolute differences between the baseline and impacted stocks were initially small ( $<1.1$ thousand lbs) through 1979, but, in general, steadily increased to about 3 thousand lbs in 2000. However, the effects of the shutdown of Unit 1 and decreased fishing rates that were simulated beginning in the late 1990s began to propagate through the spawning population projections thereafter and stock sizes increased accordingly. The increase was greater for the baseline than for the impacted stock, with differences again steadily increasing from about 4.1 thousand lbs in 2004 to about 6 thousand lbs in 2018. Using a fixed level of ENT, the impacted stock projection shown here exhibited less variation than in previous applications of the SPDM (e.g., NUSCO 2000), where randomly selected entrainment production loss estimates (ENT) and unit cooling-water flows resulted in fluctuating biomass decreases and increases.
As the simulated retirement of Units 2 and 3 occurred in 2035 and 2045, respectively, impacted stock sizes began to approach those of the baseline. Differences decreased to about 5.5 thousand lbs about 7 years after the retirement of Unit 2 and to around 4 thousand lbs some 7 years after the retirement of Unit 3. The impacted stock biomass


Fig. 51 . Stochastic variability associated with projected Niantic River female winter flounder stocks expressed as biomass in lbs for: A. The baseline stock with simulated fishing rates ( F ) shown on Table 41 and Figure 48 with F in future years set at 0.74 , but with no effects from MPS operation, and B. The impacted stock with both fishing effects (same as A) and MPS impact (ENT + IMP). The solid lines are the geometric means and $95 \%$ confidence interval ( 100 Monte Carlo replications) of each stock size trajectory and are equal to the baseline and impacted (mean ENT rate going forward) stocks illustrated in Figure 52. The symbols above and below each solid line correspond to the largest and smallest stocks among the 100 replicates generated each year. The horizontal dashed line represents the critical stock size (here, $29,993 \mathrm{lbs}$ ); defined in Howell et al. (1992) as stock biomass equal to $25 \%$ of the maximum spawning potential, which was determined as the geometric mean of the unfished stock shown on Figure 54.


Fig. 52. Results of the SPDM simulation showing the combined effects of larval entrainment rate and fishing (with impingement) given as the dashed line labeled "ENT (mean) + IMP" on the biomass in lbs of Niantic River female winter flounder spawning stock. Entrainment (ENT) and fishing (F) rates changed annually during years of MPS units in operation (see text and Table 41 for details). A geometric mean ENT rate and an F of 0.74 were used going forward. The solid line labeled "Baseline (no impact)" represents the baseline with variable fishing effects only and no plant impact. All stock sizes are averages of 100 Monte Carlo replicates. The horizontal dashed line represents the critical stock size (here, $29,993 \mathrm{lbs}$ ), defined in Howell et al. (1992) as stock biomass equal to $25 \%$ of the maximum spawning potential, which was determined as the geometric mean of the unfished stock shown on Figure 54. The vertical dashed lines indicate proposed retirement dates of 2035 for Unit 2 and 2045 for Unit 3.
remained less than the baseline, with a difference of about 2.5 thousand lbs in 2060 at the conclusion of the simulation timeline. However, at the rate of $F$ used in the simulation, the impacted line would likely become equivalent to the baseline within another decade.
To examine the extent that uncertainties in production loss estimates for Niantic River winter flounder might have in stock projections of female spawner biomass, as noted above, three levels of ENT were used in going forward: a mean rate, a high rate (mean $\times 1.5$ ), and a low rate (mean $\div 1.5$ ); $F$ remained fixed at 0.74 . These three ENT rates resulted in stock trajectories that were parallel to some extent, but differed in magnitude (Fig. 53). Biomass levels in these three projections first began to diverge beginning in 2005. From this point on, biomass under a continuously applied high ENT rate generally continued to decrease, reaching a minimum level of 3.9 thousand lbs in 2035, the year Unit 2 was projected to be shut down. Following the projected
retirement of Unit 3 in 2045, a relatively steep increase in biomass followed, attaining about 8.3 thousand lbs at the end of the simulation time-series in 2060: The biomass projections under the mean ENT rate showed a somewhat steady increase from about 4.5 to 6 thousand lbs, although some negative perturbations were seen as a result of model stochasticity. The biomass increase accelerated after the retirement dates for Units 2 and 3 and terminated at about 10.4 thousand lbs in 2060 . Under the low ENT rate, biomass immediately began to increase, although again with some annual variation observed. Female spawner biomass exceeded 10 thousand lbs by 2052 and eventually reach about 11.4 thousand lbs in 2060. Differences between the three projections were greatest in 2037-38, just after the projected retirement of Unit 2, when biomass under the mean ENT rate was $79 \%$ of that under the low rate and biomass under the high rate was about twothirds of the mean rate and half that under the high rate. At the conclusion of the simulated time-series,


Fig. 53. Results of SPDM simulations for the years 2000-2060 showing the combined effects of larval entrainment rate and fishing (with impingement) on the biomass in lbs of Niantic River female winter flounder spawning stock. A fishing mortality rate of 0.74 was used going forward. Three rates of ENT are compared going forward: the mean, a high rate (mean $\times 1.5$ ), and a low rate (mean $\div 1.5$ ). All stock sizes are averages of 100 Monte Carlo replicates.
biomass under the mean rate was within $10 \%$ of that for the low rate and biomass under the high ENT rate was within about 20 and $25 \%$ of the mean and low rates, respectively.
The factors used in creating the high and low.ENT rates modeled encompassed the variation seen in the historical production loss rates' (Table 41). Unless physical conditions (e.g., current patterns, Niantic River discharge) permanently change, it is likely that annual variation will continue to be seen in production loss estimates due to natural physical and biological factors. Thus, effects to Niantic River female winter flounder biomass would most likely be more variable than seen in each of these three population projections. However, the high and low rates should be indicative of the minimum and maximum biomass levels, given that other conditions ( F in particular) remain as modeled.
The rate of fishing on winter flounder stock size is of great importance and population projections are only realistic for the particular fishing rates simulated. Actual spawner abundance could depart consid-: erably from predictions if fishing rates or other simulated conditions are not matched by actual conditions. For example, if fishing rates were to
become high (e.g., $>0.8$ ) again at any point in the future, differences between the baseline and impacted stock series would become wider and recovery would take longer, assuming that fishing would eventually decrease once again. The different nature of stock reductions caused directly by fishing and impingement, which affect mostly, adult fish, and, those resulting from larval losses through entrainment at MPS is related to the age structure of the spawning stock. Fishing reduces biomass of the stock at a greater rate than it reduces the number of spawners because it tends to select for larger fish and, thus, reduces the average weight of the spawners remaining in the stock. However, the most important difference between fishing and larval entrainment is that the former process removes individuals from each year-class every year for as long as any fish remain, while the latter causes a reduction only once in the lifetime of each generation and, then, very early in winter flounder life history. The relative effects of stock reductions due to fishing and MPS impact can be assessed by comparing the unfished stock projection line to those for the fished stock with and without plant effects (Fig. 54). Most biomass reductions can clearly be attributed to fishing.


Fig. 54. Comparison of Niantic River winter flounder female stock biomass in Ibs after the effects of fishing (the baseline stock, shown as a the lower solid line) and MPS operation under calculated entrainment and impingement rates (shown as the lower dashed line) as shown on Table 41 and in Figures 51-53 with the theoretical (SRR-based estimate) unfished stock (shown as the upper solid line). The baseline stock shown here was determined using simulated fishing rates ( $F$ ) found on Table 41, with $F$ going forward set at 0.74 . All stock sizes are averages of 100 Monte Carlo replicates.

However, as fishing mortality was reduced and stock biomass increased, winter flounder population size of the impacted stock was increasingly smaller in comparison to the baseline until MPS units ceased operation, when the two stocks began to converge.
Probabilistic Assessment of MPS Effects. The stochastic variability associated with stock projections for the baseline and impacted stocks as shown in Figure 51 formed the basis for probabilistic analyses. These analyses took into account not only the mean stock biomass predicted for each year, but also the empirical cumulative distribution frequency (cd) of 100 replicate predictions for each year, including stock sizes both smaller and larger than the mean. Stock sizes projected for each simulation scenario at ten decadal years from 1970 through 2060 are given in Table 43. The theoretical unfished stock in each of the years shown varied little, fluctuating between about 125 and 130 thousand lbs. Prior to MPS operation in 1970, each of the baseline and the impacted stocks were identical (geometric mean of $46,777 \mathrm{lbs}$ ) and represented about $37 \%$ of the unfished stock. By 1990, winter flounder spawning stocks under full MPS three-unit operation declined
to about one-quarter of the 1970 size, which was mostly the result of increased fishing, as the impacted stock was only about 1.6 thousand lbs less than the baseline. As noted previously, smallest stock sizes were predicted for the mid-to late 1990s as a result of high rates of exploitation during the early 1990s. In 2000, the baseline and impacted stocks were only about $6 \%$ and $4 \%$ of the unfished stock, respectively. In following years, the baseline stock responded more rapidly to decreased fishing than the impacted stock, with the latter only finally approximating the former in 2060, 15 years after MPS ceased operation.
To assess effects of MPS operation, the probability that the Niantic River female winter flounder spawning stock would fall below three selected threshold sizes was determined directly from the edf of selected annual stock sizes for each of the nine selected years. The threshold sizes were percentages ( 25,30 , and $40 \%$ ) of the biomass of spawning females for the unfished stock (i.e., the MSP) as suggested in the Atlantic States Marine Fisheries Commission management plan for inshore stocks of winter flounder (Howell et al. 1992). A stock that has been reduced to less than $25 \%$ of the MSP is

TABLE 43. ${ }^{\prime}$ Expected biomass in pounds of female winter flounder spawners at ten selected points in time for the theoretical unfished², baseline ${ }^{6}$, and impacted ${ }^{c}$ (baseline plus MPS impact for the low, mean, and high time-series of ENT values; see Table 42 and Figures 53 and 54) stocks of the Niantic River population. The baseline and impacted stock projections were determined under one common historical set of fishing mortality rates (1960-99) and a projected (2000-60) future fishing rate of 0.74 during SPDM simulations. Expected mean stock sizes are geometric means of 100 Monte Carlo replicates and $95 \%$ confidence intervals (CI) are given.

| Type of population simulated |  | 1980 | 1990 | $2000$ | 2010 | $2020$ | 2030 | $2040$ | 2050 | $2060$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Theoretical unfished stock: | $\therefore$ |  |  |  |  |  |  |  |  |  |
| Geometric mean | 125,419 | 126,126 | , 126,066 | 127,171 | 127,221 | $: 128,859$ | 125,977 | 126,298 | 129,555 | 126,721 |
| 95\% lower Cl | 123,299 | 123,588 | 123,408 | 124,732 | 124,440 | 125,774 | 123,671 | 123,743 | 126,721 | 124,488 |
| 95\% upper Cl | 127,575 | 128,717 | 128,782 | 129,657 | 130,065 | .132,019 | 128,326 | 128,906 | 132,452 | 128,995 |
| Baseline stock: |  |  |  |  |  |  |  |  |  |  |
| Geometric mean | 46,777 | 30,009 | 11,945 | 7,713 | 9,869 | . 11,426 | 12,143 | 12,135 | 12,887 | 12,869 |
| 95\% lower Cl | 45,462 | 28,813 | 11,319 | 7,236 | 9,227 | $\therefore 10,743$ | 11,450 | 11,417 | 12,179 | 12,237 |
| 95\% upper Cl | 48,130 | 31,254 | 12,607 | 8,222 | 10,554 | 12,151 | 12,878 | 12,899 | 13,636 | 13,533 |
| Impacted stock - low ENT |  |  |  |  |  |  |  |  |  |  |
| Geometric mean | 46,777 | 28,300 | 10,319 | 4,682 | 5,259 | 6,493 | 7,416 | 8,004 | 9,815 | 11,366 |
| 95\% lower Cl | 45,462 | 27,180 | 9,770 | 4,377 | 4,853 | 6,010 | 6,861 | 7.404 | 9,183 | 10,736 |
| 95\% upper Cl | 48,130 | 29,467 | 10,899 | 5,008 | 5,698 | 7,015 | 8,017 | 8,654 | 10,490 | 12,032 |
| Impacted stock - mean ENT |  |  |  |  |  |  |  |  |  |  |
| Geometric mean | 46,777 | 28,300 | 10,319 | 4,682 | 4,884 | 5,520 | - 5,983 | 6,374 | 8,241 | 10,360 |
| 95\% lower Cl | 45,462 | 27,180 | 9,770 | 4,377 | 4,507 | 5,100 | 5,507 | 5,853 | 7,658 | 9,734 |
| 95\% upper Cl | 48,130 | 29.467 | 10,899 | 5,008 | -. 5,293 | -5,975 | 6,500 | 6,940 | 8,869 | 11,028 |
| Impacted stock - high ENT |  |  |  |  |  |  |  |  |  |  |
| Geometric mean | 46,777 | 28,300 | 10,319 | 4,682 | 4,342 | 4,224 | 4,123 | 4,189 | 5,791 | 8,363 |
| 95\% loiver Cl | 45,462 | 27,180 | 9,770 | 4,377 | 4,006 | 3,893 | 3,767 | 3,803 | 5,309 | 7,758 |
| 95\% upper CI | 48,130 | 29,467 | 10,899 | 5,008 | 4,706 | 4,584 | 4,512 | 4,614 | 6,317 | 9,017 |

- No tishing ( $F=0$ ) or MPS effects.
b Fishing effects ( $\mathrm{F}=0.74$ going forward), but no MPS impact of entrainment or impingement.
c Combined effects of entrainment and impingement (low, mean, or high ENT + IMP) at MPS in addition to fishing rates used for the baseline stock.
considered overfished and its continued maintenance is questionable; spawner abundance may decline to even lower levels. : Alternatively, conservative fishing rates that preserve $40 \%$ or more of MSP allow for sustainability of stocks and maximize yield to fisheries in the long term. According to the management plan for eastern LIS, values of F ranging from 0.37 to 0.68 would be necessary to achieve maximum yield, depending upon various combinations of length ( 10,11 , or 12 inches) and trawl codend mesh ( $3.5,4.5,5.0$, or 5.5 inches) restrictions imposed on the commercial fishery. The difference between the probabilities found for the baseline and impacted stocks was an estimate of risk for MPS effects.

Using the above criteria for years of actual or projected MPS operation (i.e., for the decadal years shown in Table 43), only in 1980 did biomass for both baseline and impacted (mean ENT) stocks have any probability of exceeding some of the threshold
sizes. In 1980, the baseline stock was in all likelihood ( $p=0.99$ ) smaller than $40 \%$ of MSP, had high probability ( $\mathrm{p}=0.82$ ) of being less than $30 \%$ of MSP, but had an even chance $(p=0.50)$ of being less than $25 \%$ 'of MSP. In 1980 , the impacted stock was certain ( $p=1.00$ ) to be less than $40 \%$ of MSP and was most likely $(p=0.91)$ less than $30 \%$ of MSP, and a somewhat greater than even $(p=0.61)$ chance of being less than $25 \%$ of MSP. For all other years shown in Table 43, the probability that the biomass of both the baseline and impacted stocks were less than all three levels of MSP was certain $(p=1.00)$, including in 2060,-15 years after the retirement of Unit 3 and at the end of the simulation time-series. The effect of a constant fishing mortality rate of 0.74 in all future years was to keep the Niantic River winter flounder stock moderately depressed to levels as shown on Figures 51 through 54.

In summary, SPDM output shows Niantic River female winter flounder spawner biomass stabilizing and slowly increasing following the shutdown of MPS Units 2 and 3 in 2035 and 2045, but at levels constrained by the prevailing fishing mortality of 0.74 used in the projections. For a winter flounder stock to reach a desirable size, which according to Howell et al. (1992) is greater than $40 \%$ of MSP, it appears that fishing mortality would have to be reduced to much lower levels. However, given the uncertainty of the fishing rate currently operating on the Niantic River winter flounder stock and the almost impossible task of keeping it stable over many years, projected biomass gains become less reliable with elapsed time. As shown in other SPDM simulations of the Niantic River winter flounder population, even modest reductions in $F$ result in immediate and greater increases in stock biomass than from much
larger fractional reductions in larval entrainment (DNC 2001c).
Biomass Calculations. To provide some perspective on the Niantic River winter flounder stock size estimates found above, from both actual sampling and in the SPDM projections, recreational and commercial landings of winter flounder in Connecticut waters were obtained (NMFS 2002) for 1981 through 2000 (Table 44). Using the $F$ rates used in the SPDM simulations (Table 41), annual exploitation rates ( $u$ ) were calculated as follows:

$$
\begin{equation*}
\mathbf{u}=(F /[F+M]) \times(1-\exp [-F-M]) \tag{21}
\end{equation*}
$$

where $M$ is the instantaneous natural mortality rate for adult winter flounder (0.2). The annual exploitation rates were then used to estimate the annual winter flounder stock sizes ( N ) of winter flounder in

TABLE 44. Annual Connecticut recreational and commercial harvest (lbs) of winter flounder, total landings (C), instantaneous fishing mortality rate (F), calculated exploitation rate (u), calculated stock size in lbs of Long Island Sound (LIS) winter flounder, estimated exploitable biomass (lbs) of the Niantic River winter flounder spawning stock, and the fraction that the Niantic River stock made up of the LIS winter flounder resource from 1981 through 2000.


[^23]Connecticut waters, which was presumed to be the aggregate of all stocks in LIS:

$$
\begin{equation*}
N=C / u \tag{22}
\end{equation*}
$$

where $C$ is total annual Connecticut landings of winter flounder in lbs. :Because of variable F rates, LIS biomass estimates also varied. Most stock sizes from 1981 through 1990 exceeded 2 million lbs and were as high as 4.4 million lbs in 1982 (Table 44). However, biomass decreased to less than 2 million lbs in 1991 and to less than 1 million lbs in 1999 and 2000. Data from annual winter flounder abundance surveys were used to calculate the size of the exploitable biomass of Niantic River winter flounder for the same years. The annual standardized catches of both male and female winter flounder present during the spawning season in the Niantic River were determined using the minimum legal size for retention (see Table 2 of NUSCO 2000), which was lower in some years for the recreational fishery than the commercial fishery. Since 1981, minimum size has increased from 20.3 to 30.5 cm . Weights were determined using Equation 20 with length and abundance information and summed over all sizeclasses. Annual totals represented exploitable biomass for the Niantic :River stock (Table 44). - : Exploitable stock size decreased from more than 120 thousand lbs in 1981-82 to 4-7 thousand lbs in 19962000 as abundance decreased and minimum size for exploitation increased. Through 1995, these values represented from 0.9 to $3.2 \%$ of the LIS aggregated stock biomass, but were only about $0.5-0.7 \%$ during the last 4 years. The geometric mean of the timeseries indicated that the Niantic River winter flounder population probably made up less than $2 \%$ of the exploitable stock biomass in LIS during the past two decades, which gives an indication of the size of this population relative to the total resource in Connecticut waters.

## Conclusions

Abundance of adult winter flounder spawners in the Niantic River peaked in the early 1980s as a result of extraordinarily large year-classes produced during abnormally cold winters occurring during the late 1970 s . Thereafter, abundance decreased because of stock and recruitment effects (i.e., decreasing per capita recruitment at high adult stock sizes for several years) coupled with increased exploitation rates as well as a warming winter temperature trend, which appears to have negatively affected winter flounder
reproductive success. Beginning in 1992, abundance was further depressed, primarily as a result of historically high fishing mortality rates that prevailed from the late 1980s through the mid-1990s (NEFSC 1998). Similar declines in abundance of winter flounder stocks found throughout Southern New England were also attributed to high rates of fishing (Desfosse et al. 1998; Brown and Gabriel 1998).. Brodziak et al. (2001) noted a long-term declining trend for Southern New England winter flounder biomass and recruitment and that higher recruitment was more likely at higher spawner abundance, even though it was likely that this stock was best represented by a density-dependent SRR. There also was no indication of depensatory stock dynamics in the Niantic River population.
Niantic River winter flounder spawning stock abundance was significantly and positively correlated with several abundance indices from Connecticut, Rhode Island, and Massachusetts. These region-wide trends indicated that similar factors were influencing winter flounder population dynamics across a broad geographical scale. At present, winter flounder numbers remain low despite the production of several relatively good year-classes and high abundance of juveniles since 1988. The Niantic River spawning stock most likely makes up less than $2 \%$ of the exploitable winter flounder biomass in LIS. Based on a previous tagging study, Niantic River winter flounder range widely throughout ${ }^{\text {S }}$ Southern New England during most of the year and are subject to a variety of fisheries.
Despite current low abundance of female winter flounder spawners and resulting low egg production, abundance of newly hatched larvae was greater than expected in Niantic River and Bay during 5 of the last 6 years. Relatively numerous yolk-sac larvae during these years indicated that egg survival within the Niantic River could have been as much as four times higher than in previous years. The cause of this apparent density-dependent effect is unknown, but increased egg survival could have been related to decreased predation on eggs or from possible effects of adult female size. For example, Buckley et al. (1991) demonstrated that a spawning stock dominated by larger individuals had greatest egg viability and hatching success.
Larval winter flounder abundance in Niantic Bay was much higher than expected given the numbers of larvae observed in the Niantic River. Abundance of Stage 2 larvae was among the lowest ever observed in the Niantic River, but above average in Niantic Bay, suggesting that a high rate of flushing may have
occurred that quickly removed larvae from the river. Further, abundance of Stage 3 larvae was low at all three river sampling stations, but in the upper third of the range in Niantic Bay, indicating an accumulation of larvae in the bay. Stage 4 larval abundance was well above average at the lower river site due to an influx of larvae from the bay during incoming tides as Stage 4 abundance in the bay was among the highest observed. The timing of peak abundance of Stage 4 larvae was later than average in both the river and the bay, suggesting that some of these larvae may have originated from sources other than the Niantic River. Results of the mass-balance study and the genetic stock identification analysis support this hypothesis, with the proportion of entrained larvae attributed to the Niantic River declining over the course of the spring. The high abundance of older larvae during the last 2 years also indicated that larval survival was especially enhanced, perhaps reflecting a compensatory response. Similarly, the 4 years with the greatest abundance of winter flounder larvae $\geq 7 \mathrm{~mm}$ occurred in 1995, 1997-99, and 2001, which coincided with some of the lowest egg production estimates.
Increased larval abundance was also apparent in entrainment densities (a measure of relative abundance) at MPS during recent years, when some of the highest mean larval densities and entrainment estimates were recorded. This suggested that better egg or larval survival occurred region-wide and not only for the Niantic River winter flounder spawning stock. Entrained larvae are a mixture from several sources, including the Niantic River, stocks associated with the Thames and Connecticut Rivers, and perhaps larvae from more distant locations. Because larval densities and entrainment estimates vary independently from year to year, high larval abundance was not functionally associated with high cooling-water flow and the fraction of Niantic River production entrained has changed each year. As annual entrainment estimates were positively correlated with several juvenile winter flounder abundance indices, no entrainment effect was implied. As such, the more larvae that were available for entrainment, the more that settled as juveniles in Niantic River or Bay. However, based on previous sampling, those fish settling in the bay probably do not contribute appreciably to year-class abundance, likely due to higher rates of mortality in areas outside of the inshore nursery grounds (e.g., Niantic River) known to produce most of the recruits.

The accumulation of winter flounder larvae in Niantic Bay and the high numbers of larvae entrained sharply contrast with the lower than average numbers
of larvae observed in the Niantic River in 2001. These observations accentuate the need to understand the dynamics of larval import from other spawning stocks. Results of the mass-balance model and the genetic stock identification stock analysis showed good agreement in estimating the proportion (21$22 \%$ ) of winter flounder larvae entrained at MPS during 2001 that originated in the Niantic River. Further, both analyses also showed that the fraction of entrained larvae originating in the Niantic River decreased over the course of the larval period. The high level of agreement between these approaches gives credence to the reliability of the results.
Adult recruitment is largely influenced by natural biological and physical processes occurring in early life during larval and demersal juvenile stages, from the first summer through the next several years of life. For example, a very small adult spawning stock in 2001 nevertheless produced large numbers of late stage larvae, likely as a result of very low larval mortality rate this year. The initially high densities of settled juveniles were greatly reduced throughout the summer, however, by a relatively high mortality rate for these fish. This resulted in a good, although not exceptionally large, year-class of juveniles. In addition, their growth was relatively poor and most fish entering fall and winter were small with unknown consequences to subsequent survival rate. As seen in these studies, long-term abundance indices indicated that apparent losses during later juvenile life stages appeared to have disproportionately reduced the good recruitment at age- 4 expected from larger year-classes of young winter flounder. These processes occurred independently of MPS operation as demersal age-0 winter flounder, in particular, are unaffected by plant operation.
Simulations conducted this year using the SPDM continued to clarify the issue of the different scales by which winter flounder spawning biomass responds to changes in mortality due to larval entrainment and to exploitation of adult fish. For commercially exploited and long-lived species like the winter flounder, the long-term effect of larval entrainment on adult fish biomass is relatively minor when compared to the effects of commercial exploitation. This occurs because larval entrainment impacts each year-class not only once, but early in life when natural mortality is high, while commercial fishing impacts the year-class after the fish become vulnerable to the fishery year after year for as long as the year-class persists. Additionally, annual exploitation rates can be as high as $65 \%$ or more, such as for winter flounder in the early 1990 s, while entrainment rates are generally much less. This was recently
demonstrated by O'Connor (2001), who conducted a generic Leslie matrix-based comparative analysis and concluded that "...fishing has a more severe effect on future populations than chronic impacts that decrease only first-year survival or fecundity." However, it is difficult to quantify the difference in how stock biomass changes under these two types of mortality because the response depends on complex factors, such as density-dependent processes early in the life of the fish and the age structure of the spawners, which changes with variable annual recruitment and fishing pressure. Population dynamics models like the SPDM, with explicit representation of the age structure and integration of compensatory effects, are well-suited to simulate and help quantify the stock response to losses from both entrainment and the fisheries.

A substantial recovery of the Niantic River and some other regional winter flounder stocks has not occurred despite regulatory efforts to reduce fishing mortality. Even though some year-classes demonstrated high juvenile winter flounder abundance throughout the region, their numbers declined rapidly when they reached a fishable size (Simpson et al. 1996; Desfosse et al. 1998). Fishing or high rates of predation or other natural mortality may also be removing juvenile winter flounder before they recruit to the 'adult spawning' population. Sinclair and Murawski (1997) noted that persistent overfishing of pre-recruitment fish was a major factor in the decline of a number of Northwest Atlantic groundfish stocks, although environmental conditions probably have also affected recruitment and will influence how rapidly a stock can rebuild (Brodziak et al: 2001). The trends in the population dynamics of winter flounder in Southern New England were similar to those of other exploited marine fish populations on the verge of collapse. Greater than'average predation on pre-recruit sizes of winter flounder may be occurring, which could account for the disappearance of relatively abundant young before they become spawning adults. In recent years, highly abundant populations of green crabs, cormorants, and striped bass may be preying on juvenile winter flounder and increasing numbers of harbor seals may be feeding on larger juveniles $\because$ and adult spawners. Unfortunately, little quantitative information is available to explain how either fishing or predation has affected juvenile winter flounder mortality.

Even though the Niantic River population of winter flounder 'remains at a low level of abundance, it appears to have remained viable as some relatively large year-classes of young fish were produced in recent years. This likely occurred as a result of
compensatory responses during early life history, which helped mitigate low abundance of spawning adults. The retirement of Unit 1 in July 1998 resulted in a $23 \%$ reduction in MPS cooling-water demand and entrainment. The installation of a fish return sluiceway at Unit 2 further reduced an already low impact of impingement due to high survival of winter flounder in these systems. The high numbers of Stage 4 larvae entrained in 2001 also emphasized the importance of the entrainment survival study; where these larvae had relatively good (approximately 30\%) survival compared to earlier stage larvae. In any case, no adjustment for any larval survival was made when performing impact analyses using the SPDM. The substantial and permanent reduction in station operation and the extended shutdowns of Units 2 and 3 during 1997-99; however, did not result. in markedly stronger year-classes or enhanced recruitment of adult winter flounder. Continued efforts to reduce fishing mortality could ensure a quicker recovery of the Niantic River winter flounder population. Similarly, variable environment effects (e.g., water temperature: Keller and Klein-MacPhee 2000; hydrodynamic processes: Werner et al. 1997) and interactions with other species, especially during larval and juvenile life history, appear also to be important factors in the recruitment processes of winter flounder.

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[^0]:    - The Entire Staff of Millstone Environmental Laboratory

[^1]:    * DNGL refers to the daily net generation log, EL to Millstone Environmental Laboratory records, EDAN to the environmental data acquisition network, and MOSR to the monthly nuclear plant operating status report.

[^2]:    * Data were seasonally restricted to June-October for scup, October-February for silversides, but unrestricted (Junc-May) for the remaining taxa.

[^3]:    2. Fish eggs not identified prior to 1979. Data were seasonally restricted to May-August for T. adspersus, May-August for T. onitis, and June-August for Anchoa spp.

    - Data seasonally restricted to July-September for Anchoa spp., March-June for P. americanus, December-May for A. americanus, JuneDecember for B. tyrannus, February-May for M. aenaeus, January-May for P. gunnellus, June-August for $T$. adspersus, June-August for T. onitis, March-May for Liparis spp., April-September for S. fuscus, April-June for U. subbifurcata, April-July for E. cimbrius, MayOctober for $S$. aquosus, June-September for $P$. triacanthus, and February-May for $C$. harengus.
    - Period of occurrence (December-May) extends across calendar year; year given for $\Delta$-means is that of the latter 5 months.

[^4]:    - Volume was determined from the condenser cooling water flow at MPS during the season of occurrence for each taxa.

[^5]:    - Data seasonally restricted to June-August for eggs and July-September for larvae.

[^6]:    - Data seasonally restricted to June-December.

[^7]:    - Data seasonally restricted to February-May.

[^8]:    ${ }^{2}$ Data seasonally restricted to December-June at IN, but yearround (June-May) at JC and NR.

[^9]:    Fish idenifilied to the lowest practical taxon.

[^10]:    * Fish identified to lowest practical taxon.

[^11]:    * Limited sampling during week 2 because of ice formation.
    - Almost no sampling during week 3 and limited sampling during
    $\therefore$ weeks 2 and 5 because of ice and weather conditions.
    - An additional 6 weeks of post-spawning sampling conducted from April 10 Jrough May 17.

[^12]:    - Catch per standardized tow (see text for details).
    - Mostly age- 2 and older fish.
    c Effort equalized among weeks; during several years weeks with very low effort were not used to compute CPUE.
    d Only tows of standard time or distance were considered.

[^13]:    * The two statistics shown in each correlation matrix element are:
    correlation coefficient ( $r$ ), and
    probability of a larger $r($ NS - not significant $[p>0.05]$, * significant at $p \leq 0.05, * *$ signilicant at $p \leq 0.01)$.
    The number of observations (sample size) was 19 for each paired comparison.

[^14]:    * Time period of the larval lengths used to estimate growth rate.
    - Mean during a 6 -week period starting the week of the first larval length used in estimating growth rate.
    c A parameter from the Gompertz function for Stage 2 larvae in the Niantic River (three stations combined).

[^15]:    Means with the same letter are not signiticantly ( $\mathrm{p} \leq 0.05$ ) different from each other.

[^16]:    - Catclı per standardized tow (see text for details).
    - Mostly age- 1 fish; predominant age-class was produced 1 year before the survey year.
    - Effort equalized among weeks; during several years weeks with very low effort were not used for computing CPUE.
    - Only tows of standard time or distance were considered.

[^17]:    - Indices used correspond to those given on Table 22, except for age-3 through age-5 females.

[^18]:    2 Indices used correspond to those given on Tables 22 and 23.

    - The three statistics shown in each correlation matrix element are:
    correlation coefficient $(r)$,
    probability of a larger $r(N S$ - not significant $[p>0.05]$. * - significant at $p \leq 0.05, * *-$ significant at $p \leq 0.01)$, and number of annual observations (sample size).

[^19]:    ${ }^{2}$ Early life history indices used correspond to those given on Tables 22 and 23.
    b Determined by applying an age-length key (see Figure 4) to the length distribution of annual standardized female abundances.
    c The three statistics shown in each correlation matrix element are: correlation coefficient $(r)$.
    probability of a larger $r(N S$ - not significant $[p>0.05], *-$ significant at $p \leq 0.05, * *-$ significant at $p \leq 0.01$ ), and number of annual observations (sample size).

[^20]:    - Due to rounding, any zero value represents less than 50,000 larvae.

[^21]:    ${ }^{\text {a }}$ Values differ from those reported in NUSCO (2000) due to adjustments made in the larval period of occurrence.

[^22]:    - Entrainment estimates attributed to the Niantic River are higher than those in Table 36 due to adjustments made for Stage 1 entrainment.
    - Values listed for 2000 are based on estimated stage-specific mortality rates due to lack of actual mortality estimates for that year.

[^23]:    - From NMFS (2002).
    - As given on Table 41.
    c. $\mathbf{u}=(\mathrm{F} /[\mathrm{F}+\mathrm{M}] \times[1-\exp (-\mathrm{F}-\mathrm{M}])$, where $\mathrm{M}=0.2$, the instantaneous natural mortality rate of adult winter flounder.
    ${ }^{\star} \mathrm{N}=\mathrm{C} / \mathrm{u}$.
    e To be conservative, based on the minimum legal size for retention by the recreational tishery, which in some years had lower size limits than the commercial fishery. Estimates include the calculated biomass of both males and females found in the Niantic River during the annual spawning survey.
    r No data available from NMFS (2002).

