# The Relationships Between Several Limnological Factors and Bluegill Growth in Michigan Lakes 

Charles H. Theiling

# The Relationships Between Several Limnological Factors and 

 Bluegill Growth in Michigan Lakes.
## by

Charles H . Theiling

A thesis submitted in partial fulfillment of the requirements for the degree of

Masters of Science
School of Natural Resources
The University of Michigan
1990

Committee members
Assistant professor Michael J. Wiley, Chairman
Associate professor James S. Diana
Adjunct professor W. Carl Latta
Ex-officio examiners
James C. Schneider
Dr. James Breck
Table of Contents
page
ACKNOWLEDGEMENTS ..... ii
LIST OF TABLES. ..... iii
LIST OF FIGURES. ..... v
LIST OF APPENDICES. ..... vi
ABSTACT ..... vi
INTRODUCTION. ..... 1
METHODS. ..... 6
RESULTS. ..... 18
DISCUSSION. ..... 35
LITERATURE CITED. ..... 43

## ACKNOWLEDGEMENTS

I would like to thank everyone who helped in the successfull completion of this project. I thank my advisor, Mike Wiley, for the opportunity to come to $U$ of $M$ as well as his support and encouragement throughout the project.

I must also thank my friends at the Institute for Fisheries Research. Jim (Gappy), Roger and AI helped to gather benthic samples during some chilly weather. Donna Francis, Mary Walsh and Amy Kay helped wash and sort samples. Roger Haro helped with benthic taxonomy and provided stimulating conversation. Jim Schneider, my mentor, Jim Breck and Carl Latta were critical reviewers of earlier drafts and supporters all along.

At the School of Natural Resources Jim Diana was instrumental in providing the backround I needed for this type of work. His time in and out of the classroom is greatly appreciated. All the other graduate students in the basement also helped me through this work.

A special thanks to the people who put-up with me during the last 3 years I love them all and will always remember their encouraging words.

This project was funded by the Michigan Department of Natural
Resources through a grant from the Dingell-Johnson act.

## LIST OF TABLES

Table ..... Page

1. Locations, growth index (deviation in inches from the Michigan average) and classification ("good" $=G$, "poor" = P) of the study lakes. ..... 7
2. Benthic invertebrates in food item and large food item categories, and the regression coefficients used to estimate dry weight ( mg ) from lengths (mm). All large food items were considered as food items as well. ..... 12
3. Mean nutrient concentrations for each study lake. Lakes are listed in rank by bluegill growth rate with the slowest growth first. ..... 28
4. Secchi disk transparency, chlorophyll a, dissolvedoxygen rank and average alkalinity for the study lakes.Lakes are in rank order as in Table 3.32
5. Macrophyte densities expressed as littoral density index and as percent surface area coverage. Lakes are in rank as in Table 3. ..... 33
6. Zooplankton sizes for three categories (all zooplankters, large zooplankters and small zooplankters) and total densities (no./m3). Large and small zooplankton represent the size exceeded by $10 \%$ and $90 \%$ of the population, respectively. Lakes are in rank as in Table 3. ..... 34
7. Benthic invertebrate biomass for food items from the study lakes. Lakes are in rank as in Table 3. (--) indicates no profundal stratum. Benthic invertebrate biomass ( $\mathrm{mg} / \mathrm{m}^{2}$ ).37
8. Benthic invertebrate biomass for large food items from the study lakes. Lakes are in rank as in Table 3.
(--) indicates no profundal stratum. Benthic invertebrate
biomass ( $\mathrm{mg} / \mathrm{m}^{2}$ ). ..... 38
9. Depth and area (as percent of the total) of the littoral ( $0-3 \mathrm{~m}$ ), sublittoral ( $3-9 \mathrm{~m}$ ) and profundal (> 9 m ) zones, as well as whole lake area and maximum depth for the study lakes. Lakes are in rank as in Table 3.39
10. Results of One-way ANOVAs for each variable measured and its relation to bluegill growth. The classification factor is bluegill growth performance (good or poor) alpha $=0.05$, d.f. $=1$.

## LIST OF FIGURES

Figure
Page

1. The relationship between plant precent coverage 21 (natural log) and growth index of bluegills for each study lake.
2. The relationship between large-size zooplankton 26 and growth index of bluegills for each study lake.

## LIST OF APPENDICES

Appendix Page

1. Benthic invertebrates found in the study lakes. ..... 48
2. Correlations between macrophyte density andother variables. ( $\mathrm{N}=30$, alpha $=0.05$ ). $\mathrm{NS}=$ notsignificant.50
3. Correlations between algal standing crop and other variables. ( $\mathrm{N}=30$, alpha $=0.05$ ) . $\mathrm{NS}=$ not significant. ..... 51
4. Correlations between zooplankton and other variables. ( $\mathrm{N}=30$, alpha $=0.05$ ) . $\mathrm{NS}=$ not significant. ..... 52
5. Correlations between food items and other variables. ( $\mathrm{N}=30$, alpha $=0.05$ ). $\mathrm{NS}=$ not significant. ..... 53
6. Correlations between large food items and other variables. ( $\mathrm{N}=30$, alpha $=0.05$ ). $\mathrm{NS}=$ not significant. ..... 54
7. Correlation between growth index and other variables. ( $\mathrm{N}=30$, alpha $=0.05$ ). $\mathrm{NS}=$ not significant. ..... 55

## ABSTRACT

Thirty lakes in southern Michigan were studied to determine if food availability, chemical and physical factors and habitat type influenced the growth rate of bluegills (Lepomis macrochirus). The lakes studied spanned a continuum of bluegill growth rates which allowed for comparisons of the relative importance of several limnological factors. The limnological factors considered were; benthic biomass from discrete lake zones, zooplankton size and density, macrophyte density, algal concentration, nutrients (nitrates, ammonia, orthophosphate and total phosphorus),secchi disk transparency, chlorophyll, dissolved oxygen, alkalinity and morphometric variables (lake area, area of discrete lake zones and maximum depth). Data on the growth rate of bluegill were available from the Michigan Department of Natural Resources (MDNR) and were ranked according to the growth index for Michigan fishes.

Few variables differed for lakes with good or poor growth.
Macrophyte density correlated negatively and zooplankton size correlated positively with bluegill growth rate. The multiple linear regression model developed in the final analysis used bluegill growth rate as the dependent
variable and macrophyte density, zooplankton size and profundal benthos as the independent variables $\left(r^{2}=0.598\right.$, alpha $\left.=0.05\right)$.

A number of relationships among the variables studied were noted. Macrophytes played a key role in the size distribution and abundance of zooplankton. Lake morphology, similarily, played a key role in the distribution of macrophytes.

## INTRODUCTION

The bluegill (Lepomis macrochirus) is one of the most common fish in southern Michigan. Some lakes consistently produce populations of fastgrowing bluegills while many others consistently produce slow-growing bluegills. This study, a cooperative effort between the Michigan Department of Natural Resources (MDNR) and the University of Michigan, School of Natural Resources, examines limnological factors such as; food availability, physical and chemical factors and habitat type, that may be affecting the growth of bluegills.

The growth rate of bluegills, young and adult, is typically density dependent (Grice 1957; Parker 1958; Gerking 1962; Cooper et al. 1971; Latta and Merna 1977; Beard 1982). Larval bluegills must initiate exogenous feeding about four days after birth. Beard (1982) states that the fate of the year-class is determined in the first four days. In addition, he states that larval survival appears to be a key to controlling the fate of fish populations. Both density dependent and density independent factors are important determinants of reproductive success. The time of spawning is temperature dependent and occurs when water temperatures reach $21^{\circ} \mathrm{C}$ (Beard 1982). At this time of year (May and June in southern Michigan),
food items of the appropriate size (copepod nauplii and immature cladocera) are usually abundant (Lemly and Demmick 1982). Thus spawning usually occurs when biotic and abiotic factors, barring extreme shifts in temperature, are optimal for the survival of young. Large year classes of bluegills and the abundance of other animals feeding on the same food items can, however, deplete food resources (Engel 1985). This densityrelated reduction in food can lead to low rates of larval survival and reduced growth rates of the surviving individuals (Gerking 1962; Cooper et al. 1971; Latta and Merna 1977; Beard 1982; Weiner and Hanneman 1982). Latta and Merna (1977) identified an optimal density of larval bluegills at $41 \mathrm{fry} / \mathrm{m}^{3}$ of water in research ponds.

Adult bluegill growth can also be density dependent. Early investigators conducted removal experiments, and growth rates of bluegills improved after fish removal (Grice 1957; Parker 1958; Gerking 1962). Later investigators manipulated fish densities, as well as macrophyte and predator densities, in ponds. Fish from low density populations under similar environmental conditions showed faster growth than high density populations (Werner and Hall 1977; Crowder and Cooper 1979,1982). Wiener and Hanneman (1982) found density dependent factors to be so critical to growth that it masked pH -related factors affecting
growth.

Habitat use by bluegills is determined by physiological needs, feeding habits and predator avoidance behavior. They are found in shallow water during the spawning season and afterwards are widely distributed in all but anoxic areas of lakes (Werner et al. 1977; Werner 1979). Their feeding habits are generalized. They can be found feeding on zooplankton, benthos, terrestrial insects and plants (Laarman and Schneider 1972; Beard 1982; Engel 1985) as environmental conditions may dictate (Werner and Hall 1977, 1979; Mittelbach 1981; Werner and Mittelbach 1981). This generalized feeding accounts for their wide distribution in lakes (Werner 1977). Many times, however, their foraging is restricted by their predator avoidance behavior (Werner et al. 1983; Savino and Stein 1979, 1982, 1989). Savino and Stein $(1979,1982,1989)$ found that small bluegills can avoid predation in dense vegetation. Restricted movement and successful predation avoidance can lead to overcrowding of the littoral zone and subsequent resource depletion (Werner and Hall 1977; Keast 1978; Crowder and Cooper 1979, 1982; Werner et al. 1983; Engel 1985).

The growth rate of bluegills is obviously dependent on numerous factors. Biotic and abiotic factors combine to produce conditions which are favorable to or detrimental to rapid growth of bluegills (Beard 1982).

In many lakes environmental conditions exist which favor development of dense bluegill populations which show slow growth (Gerking 1962; Cooper et al. 1971). The variables studied included benthic invertebrate biomass from discrete lake zones, zooplankton density and size distribution, macrophyte density in the littoral zone as an index and throughout the lake as percent surface area coverage, water chemistry and nutrient content, chlorophyll concentration, secchi disk transparency and lake morphology.

The objective of this study was to observe several environmental factors in a group of lakes, exposed to similar climatic fluctuations, to produce a model which might be used to predict bluegill growth performance in a typical fishery (Laarman and Schneider 1972, Ryder 1974; Gailbraith 1975, Schneider 1975a, 1975b,1978; Mills et al. 1978; Mills and Schiavone 1982). Thirty lakes, in southern Michigan, were chosen for study and bluegills in these lakes exhibited a continuum of growth rates (Schneider 1981b). The continuum of growth rates allowed for comparisons among the lakes of relative importance of the limnological variables being investigated. The comparative approach is a powerful tool for field biologists in that it provides for a "natural experiment" (Diamond 1978). It allows biologists to observe systems in nature for which the variable of interest differs, thus avoiding the need
for manipulation by the investigator. This approach is appropriate in field ecology in that it avoids time consuming, manipulation of variables in the field and it can provide the significant amount of data necessary for ecological generalization. One drawback to this method is that it allows for, in this case, only one sampling effort on each of the sampling sites. The lack of multiple sampling can miss important ecological shifts such as changes in zooplankton distribution, benthic emmergence, or chemical changes. The purpose of this study was to test the following hypotheses:

1. The biomass of benthic invertebrate food items influences the growth rate of bluegills.
2. The density and size distribution of zooplankton influences the growth rate of bluegills.
3. The density of aquatic macrophytes influences the growth rate of bluegills.
4. Algal productivity of the lake influences the growth rate of bluegills.
5. The lake nutrient status influences the growth rate of bluegills.
6. Lake morphology influences the growth rate of bluegills.

## METHODS

Thirty lakes for which the MDNR had bluegill growth data were chosen by the following criteria:

1. Bluegills were the predominant (> $50 \%$ by weight) of all centrarchid species.
2. The lake had a history of consistently fast- or slow-growing bluegill populations based on at least two MDNR surveys over the past forty years.
3. The lake had a fish community free from extensive manipulation by fishery managers.
4. The lake received average fishing pressure (reported by district managers).
5. The lake had a relatively small surface area (< 180 hectares).

An effort was made to include equal numbers of lakes with slow and fast growth (Table 1).

Benthos was sampled during the winter (January-February) of 1988. A stratified systematic design was used to minimize sampling error and to insure that most habitat types were sampled. Strata were chosen to represent major zones of the lakes. The first stratum included the littoral

Table 1. Locations, growth index (deviation in inches from the Michigan average) and classification ("good" = G, "poor" = P) of the study lakes.

| Lake | County | Location | Growth index | Classification |
| :---: | :---: | :---: | :---: | :---: |
| Algonquin | Barry | T2N,R9W,Sec.1-2 | -1.25 | P |
| Baptist | Newago | T11N,R11W,Sec.23-24 | -1.6 | P |
| Bass | Kent | T10N,R1E,Sec. 9 | -1.0 | P |
| Bear | Hillsdale | T7S,R3W,Sec.8,17 | 0.06 | G |
| Big Brower | Kent | T9N,R10W, Sec. 34 | -0.9 | P |
| Big Pine Island | Kent | T8N,R9W,Sec.3,10 | -0.95 | P |
| Big Seven | Oakland | T5N,R7E,Sec.19,30 | -1.6 | P |
| Big Silver | Washtenaw | T15S,R4E,Sec. 3 | 0.6 | G |
| Blueberry | Livingston | - | 0.5 | G |
| Carter | Barry | T3N,R8W,Sec. 6 | -0.9 | P |
| Cassidy | Washtenaw | T1S,R3E,Sec. 33 | 0.0 | G |
| Crispell | Jackson | T4S,R1W,Sec. 20 | -0.05 | P |
| Crooked | Washtenaw | T1S,R3E,Sec. 5 | -0.2 | G |
| Dead | Washtenaw | T1S,R6E,Sec. 6 | 0.3 | G |
| Dickinson | Oakland | T5N,R7E,Sec. 29 | -1.4 | P |
| Eagle | Allegan | T1N,R14W,Sec. 35 | -0.5 | P |
| Gilead | Branch | T8S,R7W,Sec. 7 | 0.9 | G |
| Halfmoon | Washtenaw | T1S,R4E,Sec. 6 | 0.9 | G |
| Hall | Barry | T2N,R8W,Sec. 2 | -1.2 | P |
| Long1 | Kent | T10N,R11W, Sec. 31 | -1.0 | P |
| Long2 | St. Joseph | T6S,R12W,Sec. 7 | -0.6 | P |
| Loon 1 | Oakland | T3N,R9E,Sec. 11 | 0.6 | G |
| Loon2 | Oscoda | T25N,rE,Sec. 36 | 1.8 | G |
| Muskellunge | Montcalm | T11N,R9W,Sec. 26 | -0.9 | P |
| Sand1 | Lenawee | T5S,R3E,Sec. 12 | 0.5 | G |
| Sand2 | Newago | T11N,R13W,Sec. 19 | -0.4 | P |
| Strawberry | Washtenaw | T1N,R5E,Sec. 27 | 1.7 | G |
| Sugarloaf | Washtenaw | T1S,R3E,Sec. 31 | -0.6 | G |
| Townline | Montcalm | T12N,R7W,Sec. 6 | -0.5 | P |
| Turk | Montcalm | T10N,R8W,Sec. 10 | -1.25 | P |

zone ( $0-3 \mathrm{~m}$ ). The second stratum included the sublittoral zone (3-9m). The third stratum was to represent the profundal zone (> 9 m ). To increase sampling efficiency, strata were sampled unevenly. The littoral strata were allotted ten samples due to habitat heterogeneity characteristic of the littoral zone of lakes. The sublittoral and profundal strata were allotted six samples each due to the more homogeneous habitat types found in deeper waters. Sampling locations were chosen by randomly picking a starting point on bathymetric maps, then spacing samples evenly around the lake. In the profundal stratum samples were evenly spaced in a line through the center of the strata.

Benthic samples were obtained using a $15 \times 15 \mathrm{~cm}$ Ekman dredge. Stations were located using landmarks on maps which could be identified from the lake. Holes for the dredge were made with a chain saw or by augering by hand.

The dredge was lowered slowly to the bottom, with care taken not to disturb the substrate. The dredge was then triggered and retrieved to the surface where samples were placed in plastic bags, labeled, sealed and returned to the lab for ellutriation.

An ellutriator was used to rinse the samples through a $500 \mu \mathrm{~m}$ mesh net and into a collecting jar. These samples were preserved in 10\%
formalin. Samples were sorted by sugar flotation and hand sorting. The sugar flotation method was abandoned due to inefficient separation of benthic organisms from plant material. Hand picking was done under a binocular dissecting scope at 7X magnification. Material in jars were subsampled when the grab sample was large or animal density was high.

Taxonomic identification was done to a level which allowed determination of the general size of the animal (Appendix 1). Insects were identified to genera except for chironomids and ceratopogonids which were identified to the family level. Dry-weight biomass estimates were made using length-weight regressions developed for this study and from the literature (Table 2). The invertebrates, excluding worms, were measured using a computer-aided video analysis system (JAVA 1988). The animals were placed under a video camera where there lengths were digitized and sent directly to computer files. Subsamples of thirty animals were measured when a particular taxon was extremely numerous. Taxa commonly subsampled for measurement included chironomids, amphipods, caenid mayflies and molluscs. Invertebrates were removed as they were encountered for use in the development of the length-weight regressions. Animals saved for regression estimates were remeasured, dried 24 hours at 105 oC and weighed on a Cahn model 4700 digital
electrobalance.

The length-weight regressions were computed using the formula:

$$
\text { In weight }=a+\ln \text { length (b) }
$$

where $a=$ the intercept and $b=$ the slope of the regression between the natural log transformed length and dry weight of the animals (Table 2). BASIC programs were written to convert length to weight and to produce biomass estimates for each stratum and for the entire lake. To facilitate analysis for this project, invertebrates were categorized as either "food items" or "large food items" (Table 2). Food items included all benthic invertebrates found in the study except for molluscs and worms. Large food items included those invertebrates which were known to reach a large size (odonates, Hexagenia sp., Sialis sp., ect.) and chironomids over 10 mm long. The biomass estimates were expressed as mg dry weight per square meter.

Zooplankton was sampled in summer (July - August) of 1988.
Vertical tows were made at cardinal compass points around the deepest point of the lake. A $159 \mu \mathrm{~m}$-mesh plankton net, fitted with a one-pint mason jar, was lowered to a depth at which the oxygen was below 0.5 $\mathrm{mg} / \mathrm{l}$ or to the bottom. The net was retrieved vertically, slowly to prevent back pressure, to the surface where the sides were washed to collect all
organisms. Samples were preserved in $10 \%$ fomalin in the field. In the laboratory, samples were stained with Eosin $Y$ to aid identification and counting. Samples were rinsed of formalin and concentrated to a volume of 400 ml . Subsamples of 5 or 10 ml were counted in a gridded Petrie dish under 40X magnification. Zooplankton taxa were identified to major groups (copepods, Daphnia sp. and other cladocera). Counts from subsamples were converted to number of plankters per cubic meter. Densities from the four samples taken were averaged to give lake-wide density estimates.

Zooplankton size distribution was determined by measuring random subsamples of 50 zooplankters from each sample ( 200 per lake). The subsamples were measured and recorded using JAVA (JAVA 1988). Daphnia sp . were measured from the anterior of the head to the base of the terminal spine along the line bisecting the eye spot. Other Cladocera were measured from head to posterior end along a line bisecting the eye spot. Copepods were measured from anterior of the head to end of the abdomen. Copepodites were measured similarly and recorded as copepods.

A BASIC program was written to produce size-frequency histograms, summary statistics (mean, minimum, maximum, and standard deviation) and quartile plots for each group and for the groups pooled. The sizes exceeded by $10 \%$ and $90 \%$ of the population, and the mean size of the

Table 2. Benthic invertebrates in food item and large food item categories, and the regression coefficients used to estimate dry weight (mg) from lengths (mm). All large food items were considered as food items as well.

| Taxa | Category | $N$ | r 2 | Size <br> range <br> (mm) | Intercept | Slope |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Planaria a | food | 23 |  |  | 0.359 | -- |
| Hirudinea | food | 36 | 0.823 | 3-44 | 0.048 | 1.759 |
| Hydracarina | food | 42 | 0.475 | 0.5-3 | 0.007 | 1.752 |
| Isopod | lg. food | 23 | 0.735 | 6-13 | 0.003 | 2.963 |
| Amphipod | food | 98 | 0.672 | 2-15 | 0.008 | 2.112 |
| Diptera |  |  |  |  |  |  |
| Chaoborus ${ }^{\text {b }}$ | food |  |  |  | 0.003 | 1.721 |
| sm. Chironomidae c | food |  |  |  | 0.0003 | 3.07 |
| lg. Chironomidae ${ }^{\text {c }}$ | lg. food |  |  |  | 0.0003 | 3.07 |
| Ceratopogonidae b | food |  |  |  | 0.007 | 1.473 |
| Trichoptera |  |  |  |  |  |  |
| Hydroptilidae | food | 50 | 0.756 | 1-4 | 0.008 | 2.027 |
| Polycetropodidae | lg. food | 42 | 0.803 | 3-14 | 0.005 | 2.225 |
| Leptoceridae | food | 37 | 0.750 | 2-11 | 0.006 | 2.271 |
| (Nectopsyche) ${ }^{\text {d }}$ | lg. food |  |  |  | 0.006 | 2.271 |
| Helicopsychidae d | food |  |  |  | 0.006 | 2.271 |
| Molannidae d | lg. food |  |  |  | 0.006 | 2.271 |
| Phryganeidae a | lg. food | 2 |  |  | 40.97 | -- |
| Ephemeroptera |  |  |  |  |  |  |
| Heptageniidae | food | 60 | 0.805 | 2-12 | 0.006 | 2.340 |
| Leptophlebiidae ${ }^{\text {e }}$ | food |  |  |  | 0.006 | 2.340 |
| Oligoneuridae e | food |  |  |  | 0.006 | 2.340 |
| Ephemerellidae e | lg. food |  |  |  | 0.006 | 2.340 |
| Baetidae e | food |  |  |  | 0.006 | 2.340 |
| Ephemeridae | lg. food | 31 | 0.942 | 3-25 | 0.002 | 2.715 |
| Caenidae b | food |  |  |  | 0.029 | 1.837 |
| Odonata |  |  |  |  |  |  |
| Zygoptera |  |  |  |  |  |  |
| Coenagrionidae | lg. food | 59 | 0.891 | 3-14 | 0.002 | 2.567 |
| Anisoptera |  |  |  |  |  |  |
| Corduliidae fi | lg. food | 43 | 0.755 | 2-28 | 0.008 | 2.498 |
| Libellulidae f | Ig. food |  |  |  | 0.008 | 2.498 |
| Macromiidae f | lg. food |  |  |  | 0.001 | 2.498 |
| Gomphidae | Ig. food | 7 | 0.945 | 4-23 | 0.007 | 2.064 |
| Coleoptera (larvae) | food | 20 | 0.650 | 3-7 | 0.002 | 2.593 |
| Hemiptera |  |  |  |  |  |  |
| Naucoridae a | lg. food | 5 |  | 13.4 | - |  |
| Megaloptera |  |  |  |  |  |  |
| Sialis | lg. food | 28 | 0.935 | 8-23 | 0.002 | 2.827 |

Table 2 continued.
$a=$ Too few animals were found to develop a regression so mean weights were used.
$b=$ Regression developed from data in Hall et al. (1970).
$\mathrm{c}=$ Regression from Wiley (1981).
$d=$ Families combined due to similar body forms.
$e=$ Families combined due to similar body forms.
$f=$ Families combined due to similar body forms.
population, were used to represent the size distribution of zooplankton. In this manuscript the $10 \%$ exceedence size is termed large zooplankton, the $90 \%$ exceedence size is termed small zooplankton.

Macrophyte densities were measured using two methods. The littoral macrophyte density index involved circling the lake in a boat and assigning a macrophyte density rank. The range was $1-5$ with $1=$ no plants and $5=$ complete cover. The percent coverage was determined by mapping the bottom with a Si-Tex depth sounder and chart recorder adjusted to distinguish plants from the bottom. Six evenly spaced transects were run on each lake. The percent of each transect containing plants was determined and the results of the transects were averaged to give a whole lake estimate. Some lakes had emergent plants so dense in the littoral zone that it was impossible to run transects. For these lakes, the percent coverage was estimated from maps assuming that plants covered $100 \%$ of the area under 4 m deep.

Water quality sampling was done in summer (July-August) of 1988. A temperature profile was constucted from measurements at 1-meter intervals using a thermister. An approximation of an oxygen profile was constructed from water samples taken at the; surface, mid epilimnion, thermocline, mid hypolimnion and bottom. Dissolved oxygen content was
measured using the Winkler method. Dissolved oxygen was ranked as follows for analysis; $1=$ oxygen to bottom $>2 \mathrm{ppm}, 2=$ oxygen to mid hypolimnion > 2 ppm, 3 = oxygen only to thermocline > 2 ppm . Alkalinity was measured at surface and mid hypolimnion. Dissolved nutrients were sampled from four depths; surface, mid epilimnion, thermocline and mid hypolimnion. Nutrients analyzed included; total phosphorus (total P), nitrates $\left(\mathrm{NO}_{3}\right)$, ammonia ( $\mathrm{NH}_{4}$ ) and orthophosphate ( $\mathrm{PO}_{4}$ ). Water samples for nutrient analysis were stored in plastic bottles in a cool, dark cooler in the field. Upon return to the lab, samples were filtered through $0.45 \mu \mathrm{~m}$ membrane filters. Nutrient analysis was done using a Technicon Autoanalyzer II at the University of Michigan Great Lakes Research Division (Davis and Simmons 1979). Secchi disk transparencies and water samples for chlorophyll analysis were taken at the deepest part of the lakes. chlorophyll samples were obtained from the filters used to filter epilimnetic nutrient samples. Filters were dissolved in 10 ml of acetone in amber glass vials and kept frozen until analysis (Davis and Simmons 1979).

Data on growth of bluegills was provided by MDNR. The data were obtained from routine fisheries surveys, during surveys for other projects and in some cases specifically for this project. Some of the lakes had been
surveyed as early as the 1940's. Fish samples were obtained by a variety of methods; trap nets, fyke nets, gill nets, seines and electroshocking. Due to the variety of gear used and the lack of consistent sampling efforts population size and fish community composition were difficult to estimate.

A growth index developed by MDNR (Schneider 1981b) was used to summarize growth characteristics of fish. The growth index is a measure of deviation in mean size of a particular bluegill population from the state's mean bluegill size. Growth index is calculated by comparing mean total length at each age for a fish population, measured at one of four times of the year, to the state mean total length at each age for that species. The lakes were classified as "good" or "poor" based on this growth index. "Good" lakes had bluegills greater than eight inches and the growth index was greater than -1.0 inch. "Poor" lakes had no bluegills greater than eight inches and the growth index was below the state average.

Statistical analysis was done using SYSTAT software on an IBM personal computer. Exploratory analysis included one - way analysis of variance (ANOVA) and Pearson correlation matrices. One - way ANOVAs were done on each variable measured. Lakes were classified as "good" ( $\mathrm{n}=$ 13) or "poor" ( $\mathrm{n}=17$ ) using bluegill growth as the classification factor.

The Pearson correlation matrix of all variables was used to explore relationships between variables and bluegill growth as well as among variables themselves $(\mathrm{N}=30$, alpha $=0.05)$.

Multiple linear regression (MLR) was used in the final analysis to produce a model which would use several limnological variables as independent variables to explain variance in bluegill growth rates. Stepwise MLR (alpha for inclusion $=0.15$ ) was used to aid in development of the model.

RESULTS

There were no significant correlations between any nutrients measured and bluegill growth performance (Table 3). Ammonia ranged from $10 \mu \mathrm{~g} / \mathrm{l}$ to $510 \mu \mathrm{~g} / \mathrm{l}$, with a mean concentration of $148 \mu \mathrm{~g} / \mathrm{l}$. Nitrates ranged from nondetectable levels to $1.31 \mathrm{mg} / \mathrm{l}$, with a mean concentration of $0.167 \mathrm{mg} / \mathrm{l}$. Orthophosphate ranged from $2 \mu \mathrm{~g} / \mathrm{l}$ to 18 $\mu \mathrm{g} / \mathrm{l}$, with a mean concentration of $6 \mu \mathrm{~g} / \mathrm{l}$. Total phosphorus ranged from $11 \mu \mathrm{~g} / \mathrm{l}$ to $27 \mu \mathrm{~g} / \mathrm{l}$, with a mean concentration of $17 \mu \mathrm{~g}$.

There were no significant relationships between dissolved oxygen rank or alkalinity (Table 4) and bluegill growth. Dissolved oxygen varied among the lakes; shallow, unstratified lakes had oxygen to the bottom while deeper lakes with highly organic substrates occassionally had oxygen depletion in the hypolimnion. The lakes were primarily hardwater lakes; $\mathrm{CaCO}_{3}$ concentrations ranged from $62 \mathrm{mg} / \mathrm{l}$ to $227 \mathrm{mg} / \mathrm{l}$, with a mean of $147.7 \mathrm{mg} / \mathrm{l}$.

There were also no significant correlations between bluegill growth and secchi disk depth or chlorophyll a. concentration (Table 4). Secchi transparencies ranged from 1.3 m to 4.6 m with a mean of about 3 m .

Table 3. Mean nutrient concentrations for each study lake. Lakes are listed in rank by bluegill growth rate with the slowest growth first.

| Lake | $\begin{aligned} & \mathrm{NO}_{3} \\ & (\mathrm{mg} / \mathrm{l}) \end{aligned}$ | $\begin{aligned} & \mathrm{NH}_{3} \\ & (\mu \mathrm{~g} / \mathrm{l}) \end{aligned}$ | $\begin{aligned} & \mathrm{PO} 4 \\ & (\mu \mathrm{~g} / \mathrm{l}) \end{aligned}$ | Total P ( $\mu \mathrm{g} / \mathrm{l}$ ) |
| :---: | :---: | :---: | :---: | :---: |
| Baptist | 0.01 | 10.7 | 5.6 | 13.7 |
| Big Seven | 0.73 | 77.6 | 5.0 | 13.7 |
| Dickinson | 0.47 | 57.0 | 4.1 | 11.3 |
| Algonquin | 0.02 | 19.0 | 3.2 | 24.5 |
| Turk | 0.06 | 325.6 | 3.5 | 16.8 |
| Hall | 0.04 | 39.1 | 4.1 | 27.3 |
| Bass | 0.02 | 51.2 | 5.3 | 14.7 |
| Long 1 | 0.02 | 51.5 | 3.4 | 13.6 |
| Big Pine ls. | 0.00 | 19.3 | 5.3 | 15.6 |
| Big Brower | 0.00 | 163.7 | 5.4 | 12.7 |
| Carter | 0.03 | 452.1 | 17.6 | 22.3 |
| Muskellunge | 1.15 | 115.0 | 4.7 | 16.2 |
| Long2 | 0.01 | 265.3 | 5.8 | 19.1 |
| Sugarloaf | 0.06 | 479.7 | 5.0 | 13.2 |
| Eagle | 0.01 | 51.4 | 6.4 | 21.5 |
| Townline | 0.02 | 180.6 | 6.4 | 21.4 |
| Sand2 | 0.00 | 15.2 | 7.7 | 21.5 |
| Crooked | 0.02 | 153.7 | 2.0 | 13.4 |
| Crispell | 0.01 | 25.5 | 4.5 | -- |
| Cassidy | -- | -- | -- | 19.4 |
| Bear | 0.03 | 325.6 | 2.3 | 14.8 |
| Dead | 0.02 | 224.2 | 1.6 | 14.6 |
| Blueberry | 0.04 | 25.3 | 3.4 | 21.7 |
| Sand1 | 0.06 | 129.2 | 13.5 | 21.5 |
| Big Silver | 0.06 | 46.3 | 4.2 | 14.5 |
| Loon ${ }^{1}$ | 0.47 | 133.9 | 8.6 | 12.2 |
| Gilead | 0.05 | 164.8 | 5.2 | 15.8 |
| Halfmoon | 1.31 | 157.6 | 5.2 | 15.0 |
| Strawberry | 0.07 | 510.3 | 16.6 | 14.7 |
| Loon2 | 0.06 | 37.8 | 4.2 | 12.8 |

Chlorophyll a concentrations ranged from nondetectable to $0.034 \mathrm{mg} / \mathrm{l}$, with a mean concentration of $0.005 \mathrm{mg} / \mathrm{l}$. Chlorophyll a concentrations are lower than would be expected for the types of lakes in this study. The low values may be due to chlorophyll degradation during 60 days storage in the freezer before analysis, however, there was still a significant relationship between chlorophyll $a$ and secchi disk depth ( $r=-0.480$, alpha $=0.05$, indicating the inverse relationship between secchi disk depth and algal density).

The correlation between the two macrophyte density measurements was also strong and significant ( $r=.750$, alpha $=0.05$ ) (Table 5). Macrophyte ranks ranged from $1=$ very few plants to $5=$ complete coverage of the littoral zone. The percent surface area coverage ranged from $8.5 \%$ to $100 \%$, with a mean of $48.8 \%$.

There was a significant negative correlation between the macrophyte density and bluegill growth performance (Figure 1). This correlation was significant using either measurement of macrophyte abundance (littoral rank: $r=-0.443$, alpha $=0.05$ ), percent surface area coverage: $r=-0.431$, alpha $=0.05)$. The relationship between macrophyte density and growth is not linear, (Figure 1) and fish growth was highly


Figure 1. The relationship between plant percent coverage (natural log) and growth index of bluegills for each study lake.
variable for plant densities of $30-100 \%$.

There was considerable variation in the size distribution of the zooplankton in the lakes studied (Table 6). Small zooplankton (size exceeded by $90 \%$ of the population) size ranged from 0.26 mm to 0.72 mm , with a mean of 0.43 mm . The average size of zooplankton ranged from 0.49 mm to 1.15 mm , with a mean of 0.76 mm . Large zooplankton (size exceeded by $10 \%$ of the population) size ranged from 0.73 mm to 1.86 mm in the lakes with a mean of 1.15 mm .

There was a significant positive correlation between the average size of large zooplankton and bluegill growth performance (Figure 2, $\mathrm{r}=$ 0.622, alpha $=0.05$ ).

Zooplankton densities ranged from $5900 / \mathrm{m}^{3}$ to $212,000 / \mathrm{m}^{3}$ in the lakes, with a mean density of $51,566 / \mathrm{m}^{3}$. There was a significant negative relationship between zooplankton density and bluegill growth performance $(r=-0.361$, alpha $=0.05)$.

Surprisingly, there was only one significant correlation between bluegill growth and any benthic biomass variable (Appendix 7). This correlation was for large food items from the profundal stratum ( $r=$ 0.444 , alpha $=0.05$ ). There was high variability among lakes' benthic

Table 4. Secchi disk transparency, chlorophylll a, dissolved oxygen rank and average alkalinity for the study lakes. Lakes are in rank order as in Table 3.

| Lake | Secchi depth <br> (meters) | Chlorophyll a <br> $(\mathrm{mg} / \mathrm{l})$ | Dissolved <br> oxygen <br> rank | Alkalinity <br> $(\mathrm{mg} \mathrm{CaCO} / \mathrm{l})$ |
| :---: | :---: | :---: | :---: | :---: |


|  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- |
| Baptist | 2.7 | 0.002 | 2 | 62 |
| Big Seven | 3.5 | 0.002 | 3 | 170 |
| Dickinson | 3.5 | 0.000 | 2 | 208 |
| Algonquin | 1.9 | 0.007 | 2 | 174 |
| Turk | 1.2 | 0.011 | 3 | 142 |
| Hall | 1.8 | 0.004 | 1 | 148 |
| Bass | 3.2 | 0.001 | 1 | 125 |
| Long1 | 3.2 | 0.002 | 1 | 106 |
| Big Pine Island | 2.4 | 0.004 | 2 | 144 |
| Big Brower | 1.8 | 0.010 | 3 | 142 |
| Carter | 1.8 | 0.010 | 3 | 221 |
| Muskellunge | 1.6 | 0.003 | 1 | 158 |
| Long2 | 2.6 | 0.002 | 3 | 113 |
| Sugarloaf | 2.0 | 0.015 | 1 | 186 |
| Eagle | 4.5 | 0.002 | 3 | 149 |
| Townline | 2.9 | 0.003 | 3 | 153 |
| Sand2 | 1.8 | 0.005 | 1 | 73 |
| Crooked | 2.9 | 0.000 | 1 | 103 |
| Crispell | 2.3 | 0.002 | 2 | 153 |
| Cassidy | 2.4 | 0.002 | 1 | 124 |
| Bear | 3.2 | 0.002 | 3 | 147 |
| Dead | 3.0 | 0.003 | 3 | 114 |
| Blueberry | 2.0 | 0.034 | 3 | 105 |
| Sand1 | 3.7 | 0.001 | 3 | 149 |
| Big Silver | 2.6 | 0.000 | 2 | 152 |
| Loon1 | 3.5 | 0.001 | 3 | 201 |
| Gilead | 2.7 | 0.001 | 3 | 161 |
| Halfmoon | 2.6 | 0.006 | 3 | 189 |
| Strawberry | 1.4 | 0.004 | 3 | 227 |
| Loon2 | 3.7 | 0.001 | 1 | 137 |
|  |  |  |  |  |

Table 5. Macrophyte densities expressed as littoral density index and as percent surface area coverage. Lakes are in rank as in Table 3.

| Lake | Macrophyte littoral density index | Macrophyte percent cover |
| :---: | :---: | :---: |
| Baptist | 4.5 | 21.9 |
| Big Seven | 5 | 89.9 |
| Dickinson | 3 | 33.6 |
| Algonquin | 5 | 70.3 |
| Turk | 4 | 50.8 |
| Hall | 5 | 100 |
| Bass | 3.5 | 56.8 |
| Long 1 | 5 | 76.6 |
| Big Pine Island | 4 | 44.4 |
| Big Brower | 1 | 35.2 |
| Carter | 5 | 82.2 |
| Muskellunge | 5 | 53.0 |
| Long2 | 5 | 52.3 |
| Sugarloaf | 4 | 98.3 |
| Eagle | 4.5 | 35.4 |
| Townline | 4 | 58.5 |
| Sand2 | 4 | 52.9 |
| Crooked | 4 | 67.8 |
| Crispell | 4 | 88.1 |
| Cassidy | 3 | 60.0 |
| Bear | 3 | 22.1 |
| Dead | 5 | 40.7 |
| Blueberry | 5 | 82.8 |
| Sand ${ }^{1}$ | 2 | 12.0 |
| Big Silver | 2 | 9.1 |
| Loon1 | 2 | 15.8 |
| Gilead | 3 | 20.4 |
| Halfmoon | 1.5 | 11.2 |
| Strawberry | 2 | 13.9 |
| Loon2 | 1 | 8.5 |

Table 6. Zooplankton sizes for three categories (all zooplankters, large zooplankters and small zooplankters) and total densities (no./m3). Large and small zooplankton represent the size exceeded by $10 \%$ and $90 \%$ of the population, respectively. Lakes are in rank as in Table 3.

| Lake | Mean <br> size <br> (mm) | Large <br> size <br> (mm) | Small <br> size <br> (mm) | $\begin{gathered} \text { Density } \\ \text { X103 } \\ \text { (no./m3) } \end{gathered}$ |
| :---: | :---: | :---: | :---: | :---: |
| Baptist | 0.65 | 0.88 | 0.30 | 57.8 |
| Big Seven | 0.84 | 1.22 | 0.48 | 77.7 |
| Dickinson | 0.83 | 1.22 | 0.50 | 11.1 |
| Algonquin | 0.77 | 1.15 | 0.41 | 54.5 |
| Turk | 0.60 | 0.81 | 0.38 | 74.6 |
| Hall | 0.49 | 0.76 | 0.28 | 85.5 |
| Bass | 0.49 | 0.73 | 0.26 | 51.7 |
| Long1 | 0.63 | 0.85 | 0.43 | 44.4 |
| Big Pine Island | 0.65 | 0.87 | 0.38 | 59.9 |
| Big Brower | 0.53 | 0.77 | 0.29 | 107.8 |
| Carter | 0.81 | 1.15 | 0.43 | 34.2 |
| Muskellunge | 0.79 | 1.20 | 0.53 | 73.6 |
| Long2 | 0.70 | 0.92 | 0.43 | 51.7 |
| Sugarloaf | 0.54 | 0.80 | 0.28 | 211.7 |
| Eagle | 0.95 | 1.72 | 0.49 | 15.4 |
| Townline | 0.66 | 0.86 | 0.43 | 45.3 |
| Sand2 | 0.50 | 0.74 | 0.27 | 129.6 |
| Crooked | 1.00 | 1.37 | 0.72 | 49.9 |
| Crispell | 0.72 | 0.96 | 0.42 | 47.9 |
| Cassidy | 0.77 | 0.99 | 0.52 | 19.0 |
| Bear | 0.84 | 1.85 | 0.27 | 24.9 |
| Dead | 0.86 | 1.33 | 0.49 | 16.9 |
| Blueberry | 0.78 | 1.15 | 0.41 | 5.9 |
| Sand1 | 0.90 | 1.29 | 0.68 | 28.3 |
| Big Silver | 0.98 | 1.59 | 0.40 | 9.6 |
| Loon 1 | 0.86 | 1.54 | 0.45 | 22.3 |
| Gilead | 0.74 | 1.29 | 0.41 | 48.4 |
| Halfmoon | 0.77 | 1.18 | 0.49 | 20.2 |
| Strawberry | 0.93 | 1.46 | 0.52 | 52.8 |
| Loon² | 1.15 | 1.86 | 0.52 | 14.4 |



Figure 2. The relationship between large-size zooplankton and growth index of bluegills for each study lake.
populations (Tables $7-8$ ) as might be expected, but there were no trends noted such as high benthic biomass in dense vegetation. The littoral stratum consistently had the highest biomass of benthos ( $703 \mathrm{mg} / \mathrm{m}^{2}$ ) while the profundal stratum consistently had the lowest biomass (190 $\mathrm{mg} / \mathrm{m}^{2}$ ). The food-item category contributed the bulk of benthic biomass. Since benthic sampling and the other sampling was done at different times of the year, one must use caution in relating benthos to other variables, and this may be the reason for the lack of association between variables.

The morphological measurements (strata sizes as percent of the total area, total area and maximum depth) varied among lakes (Table 9). Total lake area ranged from 10.1-178.1 ha, with a mean size of 60 ha. The percent of area within each specific stratum was variable; some lakes were entirely littoral zone while others had large sublittoral and profundal areas. The size of each stratum was a function of the basin morphology and maximum depth (which ranged from 3.4-25m). There were no significant correlations between the morphological variables and bluegill growth performance.

In addition to the relationships between each variable measured and bluegill growth, the relationships among the variables were examined
through the use of a Pearson correlation matrix (Appendices 2-7).
Algal standing crops correlated negatively with size of zooplankton and maximum depth. Algal density was positively correlated with abundance of zooplankton, macrophyte cover and littoral strata size (Appendices 3 4).

Macrophyte densities were positively correlated with the size of the littoral zone ( $r=0.683$, alpha $=0.05$ ) and negatively correlated with the size of the profundal zone $(r=-0.646$, alpha $=0.05)$ and maximum depth ( $\mathrm{r}=-0.676$, alpha $=0.05$ ). Another significant positive correlation was found between macrophytes and total phosphorus concentration.

There were several significant correlations between zooplankton and other variables (Appendices 4-7). Generally algal content, size, morphology and macrophyte density of the lake determined size distribution of zooplankton. Zooplankton mean and large size classes were negatively correlated with both macrophyte density and the littoral strata size and positively correlated with the algal content, the size of profundal strata and the maximum depth. Zooplankton densities were negatively correlated with zooplankton size (mean size: $r=-0.631$, large zooplankton size $r=-0.595$, small zooplankton size $r=-0.447$, alpha $=0.05$ ).

Table 7. Benthic invertebrate biomass for food items from the study lakes. Lakes are in rank as in Table 3. (--) indicates no profundal stratum.
Benthic invertebrate biomass ( $\mathrm{mg} / \mathrm{m}^{2}$ ).

| Lake | Littoral | Sublittoral | Profundal | Whole lake |
| :---: | :---: | :---: | :---: | :---: |
| Baptist | 607.1 | 188.4 | 409.0 | 376.7 |
| Big Seven | 109.8 | 96.9 | 18.3 | 102.3 |
| Dickinson | 616.8 | 699.7 | 213.1 | 418.7 |
| Algonquin | 765.3 | 62.4 | 12.9 | 481.1 |
| Turk | 804.1 | 95.8 | -- | 780.4 |
| Hall | 372.4 | 164.7 | -- | 303.5 |
| Bass | 461.8 | 258.3 | -- | 389.7 |
| Long1 | 299.2 | 215.3 | -- | 263.7 |
| Big Pine Island | 768.5 | 603.9 | 1188.3 | 664.1 |
| Big Brower | 456.4 | 359.5 | -- | 429.5 |
| Carter | 2300.2 | 153.9 | -- | 1657.6 |
| Muskellunge | 1099.0 | 392.9 | 185.1 | 589.9 |
| Long2 | 1610.3 | 92.6 | 167.9 | 586.6 |
| Sugarloaf | 1364.9 | 102.3 | -- | 1362.7 |
| Eagle | 1928.9 | 442.4 | 557.6 | 920.3 |
| Townline | 297.1 | 455.3 | 582.3 | 371.4 |
| Sand2 | 129.2 | 64.6 | -- | 117.3 |
| Crooked | 607.1 | 329.4 | -- | 543.6 |
| Crispell | 241.1 | 148.5 | -- | 194.8 |
| Cassidy | 550.0 | 81.8 | -- | 527.4 |
| Bear | 1103.3 | 1022.6 | 204.5 | 845.0 |
| Dead | 622.2 | 306.8 | -- | 526.4 |
| Blueberry | 360.6 | 173.3 | -- | 292.8 |
| Sand1 | 883.7 | 418.7 | 727.6 | 722.3 |
| Big Silver | 226.0 | 216.4 | 494.1 | 268.0 |
| Loon ${ }^{1}$ | 298.2 | 89.3 | 119.5 | 195.9 |
| Gilead | 1202.3 | 394.0 | 265.9 | 711.5 |
| Halfmoon | 318.6 | 43.1 | 20.5 | 130.2 |
| Strawberry | 141.0 | 39.8 | 25.8 | 59.2 |
| Loon² | 554.3 | 297.1 | 517.7 | 405.8 |

Table 8. Benthic invertebrate biomass for large food items from the study lakes. Lakes are in rank as in Table 3. (--) indicates no profundal stratum. Benthic invertebrate biomass ( $\mathrm{mg} / \mathrm{m}^{2}$ ).

| Lake | Littoral | Sublittoral | Profundal | Whole lake |
| :---: | :---: | :---: | :---: | :---: |
| Baptist | 115.2 | 76.4 | 0.0 | 63.5 |
| Big Seven | 5.4 | 0.0 | 0.0 | 3.2 |
| Dickinson | 345.5 | 15.1 | 0.0 | 103.3 |
| Algonquin | 104.4 | 0.0 | 0.0 | 64.6 |
| Turk | 334.8 | 0.0 | -- | 323.0 |
| Hall | 94.8 | 0.0 | -- | 63.5 |
| Bass | 4.3 | 16.1 | -- | 8.6 |
| Long 1 | 47.4 | 31.2 | -- | 40.9 |
| Big Pine Island | 159.3 | 1.1 | 0.0 | 65.7 |
| Big Brower | 239.0 | 21.6 | -- | 179.8 |
| Carter | 1080.7 | 22.6 | -- | 773.9 |
| Muskellunge | 350.9 | 43.1 | 1.5 | 139.9 |
| Long2 | 72.1 | 9.7 | 0.0 | 28.0 |
| Sugarloaf | 271.2 | 0.0 | -- | 271.2 |
| Eagle | 808.4 | 5.4 | 0.0 | 265.9 |
| Townline | 26.9 | 9.7 | 0.0 | 18.3 |
| Sand2 | 65.7 | 0.0 | -- | 52.7 |
| Crooked | 164.7 | 109.8 | -- | 152.8 |
| Crispell | 35.5 | 0.0 | -- | 22.6 |
| Cassidy | 162.5 | 7.5 | -- | 155.0 |
| Bear | 253.0 | 0.0 | 0.0 | 110.9 |
| Dead | 144.2 | 11.8 | -- | 105.5 |
| Blueberry | 155.0 | 10.8 | -- | 117.3 |
| Sand 1 | 523.1 | 20.5 | 14.0 | 315.4 |
| Big Silver | 76.4 | 0.0 | 0.0 | 35.5 |
| Loon 1 | 33.4 | 14.0 | 0.0 | 18.3 |
| Gilead | 431.6 | 1.1 | 0.0 | 187.3 |
| Halfmoon | 42.0 | 5.4 | 0.0 | 16.1 |
| Strawberry | 49.5 | 6.5 | 0.0 | 14.0 |
| Loon² | 274.5 | 93.6 | 17.2 | 152.8 |

Table 9. Depth and area (as percent of the total) of the littoral ( $0-3 \mathrm{~m}$ ), sublittoral ( $3-9 \mathrm{~m}$ ) and profundal ( $>9 \mathrm{~m}$ ) zones, as well as whole lake area and maximum depth for the study lakes. Lakes are in rank as in Table 3.

| Lake | Littoral (\%) | Sublittoral (\%) | Profundal (\%) | Whole lake (ha.) | Maximum depth (m) |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Baptist | 29.9 | 37.7 | 32.4 | 34.8 | 19.8 |
| Big Seven | 74.6 | 19.6 | 5.8 | 68.8 | 16.1 |
| Dickinson | 29.3 | 17.8 | 52.9 | 17.8 | 20.7 |
| Algonquin | 62.0 | 28.2 | 9.8 | 97.1 | 13.7 |
| Turk | 51.0 | 49.0 | 0.0 | 64.3 | 6.1 |
| Hall | 75.1 | 24.9 | 0.0 | 17.0 | 3.7 |
| Bass | 66.0 | 34.0 | 0.0 | 74.5 | 6.2 |
| Long1 | 58.3 | 41.7 | 0.0 | 19.4 | 8.2 |
| Big Pine Island | 40.6 | 48.8 | 10.6 | 90.2 | 13.7 |
| Big Brower | 72.5 | 27.5 | 0.0 | 34.4 | 8.2 |
| Carter | 71.0 | 29.0 | 0.0 | 28.3 | 7.6 |
| Muskellunge | 32.8 | 57.9 | 9.3 | 54.2 | 11.3 |
| Long2 | 32.8 | 50.2 | 17.0 | 85.4 | 12.5 |
| Sugarloaf | 98.3 | 1.7 | 0.0 | 72.8 | 5.5 |
| Eagle | 32.6 | 36.6 | 30.8 | 91.1 | 18.0 |
| Townline | 52.7 | 41.7 | 5.6 | 100.0 | 14.9 |
| Sand2 | 81.0 | 19.0 | 0.0 | 23.5 | 4.6 |
| Crooked | 79.1 | 20.9 | 0.0 | 45.7 | 6.1 |
| Crispell | 64.8 | 35.2 | 0.0 | 33.2 | 7.6 |
| Cassidy | 71.0 | 29.0 | 0.0 | 17.4 | 3.4 |
| Bear | 44.1 | 34.7 | 21.2 | 47.3 | 15.2 |
| Dead | 71.3 | 28.7 | 0.0 | 23.1 | 9.8 |
| Blueberry | 74.2 | 25.8 | 0.0 | 10.1 | 6.7 |
| Sand1 | 65.2 | 23.7 | 11.1 | 178.1 | 16.2 |
| Big Silver | 45.9 | 33.7 | 20.4 | 82.6 | 14.3 |
| Loon 1 | 46.2 | 20.5 | 33.3 | 98.3 | 22.3 |
| Gilead | 43.3 | 32.4 | 24.3 | 52.6 | 14.9 |
| Halfmoon | 34.6 | 26.8 | 38.6 | 95.5 | 25.0 |
| Strawberry | 24.7 | 33.5 | 41.8 | 105.2 | 15.8 |
| Loon2 | 36.5 | 55.0 | 8.5 | 36.4 | 15.2 |

Correlations with other variables showed relationships opposite those shown for zooplankton size. Zooplankton densities were positively correlated with algal density and macrophyte density. They were negatively correlated with profundal stratum size and maximum depth.

Relationships between benthic invertebrate biomass and other variables measured are difficult to distinguish in this study since they were measured at different times of the year (Appendices 5-7). One unexpected negative correlation was between invertebrate biomass and macrophyte density. This result is opposite of expectations from a review of the literature (Hayne and Ball 1956; Pardue 1973; Crowder and Cooper 1979, 1982; Gilinsky 1984; Wiley et al. 1984). These results may be due to benthic invertebrates distributing differently when plants die back in the fall and recolonize macrophyte beds again in spring (Gilinsky 1984). Sampling benthos in winter and plants in summer would not account for such invertebrate migrations and the relationship between invertebrates and plants could be biased. A second significant correlation was a positive relationship between size of zooplankton and biomass of the large-sized benthic invertebrates.

There were several significant correlations between lake morphology and variables affecting bluegill growth (as noted above) but no
direct correlations with growth (Appendices 2-7).
One-way analysis of variance (ANOVA) was used to examine differences observed among the variables between "good" and "poor" lakes. Bluegill growth performance was the classification factor used to separate the two lake types by criteria described above (Table 10). Only two variables were significantly different between the two lake types (macrophytes and zooplankton) and these were examined more closely using regression analysis.

Stepwise regression using all variables (alpha for inclusion $=0.15$ ) was also used to aid development of the final multiple linear regression (MLR) model. As in the previous analysis, stepwise techniques identified plant cover and large-sized zooplankton as variables which should be included in the model for bluegill growth.

In an effort to produce a better model, each of the other variables was forced into the model. Several variables improved the fit of the model but were difficult to explain biologically or introduced too much multicollinearity. The final equation was:

$$
Y=1.362-0.685(X 1)+1.005(X 2)+0.083(X 3)
$$

where: $Y=$ bluegill growth index (deviation in inches from state avgerage).
$\mathrm{X} 1=$ natural log of percent surface area coverage by macrophytes. $\mathrm{X} 2=$ size exceeded by $10 \%$ of the zooplankton population (mm). X3 = biomass of the large size benthic invertebrates from the profundal stratum.
$\left(r^{2}=0.589\right)$.
Analysis of Variance

| Source | Sum of Squares | df | Mean Square | F-ratio | P |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |
| regression | 14.544 | 3 | 4.848 | $12.290<0.0005$ |  |
| residual | 10.256 | 26 | 0.394 |  |  |

The assumptions of homogeneity of variance and normality were tested and met. There were no problems with multicollinearity among variables included in the model. Macrophytes contribute most to the variance explained by the model (75\%), large-size zooplankton $10 \%$ and large profundal benthos contribute $15 \%$ despite there being only three data points with non-zero values.

## DISCUSSION

The regression model developed here is consistent with bluegill biology as it is currently understood. Macrophytes contribute greatly to the variance explained by the model, and they play a key role in structuring bluegill population density (Beard 1982; Lemly and Demmick 1982; Engel 1985) and growth (Crowder and Cooper 1979, 1982). The contribution of macrophytes is, presumably, through their role in providing refuge from predation for small bluegills (Mittelbach 1981; Savino and Stein 1982, 1989; Werner et al. 1983; Engel 1985). When small fish engage in this behavior, and successfully avoid predation, their populations grow to a point where they may overexploit their food resources (Grice 1957; Parker 1958; Gerking 1962).

Zooplankton are an easily obtained food resource in lakes. They are distributed into fairly discrete zones with larger zooplankton found predominantly in open water areas while smaller zooplankton concentrate in littoral areas (Brooks and Dodson 1965; Pennak 1978). Among the study lakes, large zooplankton were common in lakes with large open water areas and small zooplankters were common to lakes consisting predominantly of littoral habitat. Bluegills forage optimally by

Table 10. Results of One-way ANOVAs for each variable measured and its relation to bluegill growth. The classification factor is bluegill growth performance (good or poor) alpha $=0.05$, d.f. $=1$.

Variable $N$| Relation |
| :---: |
| to growth |$\quad$ Mean square $\quad F . \quad \mathrm{p}$

| $\overline{\text { nitrates }}$ | 30 | NS | 0.004 | 0.032 | 0.860 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| ammonia | 30 | NS | 44001.863 | 2.161 | 0.153 |
| orthophosphate | 30 | NS | 0.833 | 0.057 | 0.813 |
| total phosphorus | 30 | NS | 21.966 | 1.282 | 0.267 |
| dissolved oxygen |  |  |  |  |  |
| rank | 31 | NS | 0.979 | 1.223 | 0.278 |
| alkalinity | 31 | NS | 581.048 | 0.378 | 0.544 |
| secchi disk |  |  |  |  |  |
| transparency | 31 | NS | 25.236 | 3.643 | 0.066 |
| chlorophylll a | 31 | NS | 0.0 | 0.119 | 0.732 |
| macrophyte density |  |  |  |  |  |
| index | 31 | - | 11.715 | 8.481 | 0.007 |
| percent coverage | 31 | - | 5664.980 | 7.984 | 0.008 |
| zooplankton |  |  |  |  |  |
| mean size | 31 | + | 0.332 | 18.122 | <0.0005 |
| large size | 31 | + | 1.585 | 26.066 | <0.0005 |
| small size | 31 | + | 0.067 | 5.901 | 0.022 |
| density | 31 | NS | $0.86 \mathrm{e}+10$ | 3.587 | 0.068 |
| benthic invertebrate biomass |  |  |  |  |  |
| stratum 1 | 31 | NS | 422.998 | 0.161 | 0.691 |
| stratum 2 | 31 | NS | 129.693 | 0.194 | 0.592 |
| stratum 3 | 31 | NS | 141.152 | 0.194 | 0.663 |
| whole lake |  | NS | 418.603 | 0.367 | 0.550 |
| large benthic invertebrate biomass |  |  |  |  |  |
| stratum 1 | 31 | NS | 242.683 | 0.478 | 0.495 |
| stratum 2 | 31 | NS | 2.346 | 0.347 | 0.561 |
| stratum 3 | 31 | NS | 0.304 | 2.406 | 0.132 |
| whole lake | 31 | NS | 31.777 | 0.155 | 0.697 |
| size stratum 1 | 31 | NS | 105.582 | 0.242 | 0.627 |
| stratum 2 | 31 | NS | 140.959 | 0.790 | 0.382 |
| stratum 3 | 31 | NS | 490.529 | 2.204 | 0.148 |
| whole lake | 31 | NS | 14404.375 | 1.683 | 0.205 |
| max. depth 31 | 31 | NS | 735.823 | 0.490 | 0.171 |

selecting large zooplankton when zooplankton density is high (Werner and Hall 1974; Mittelbach 1981; Werner et al. 1983). When zooplankton density is low or large zooplankters are rare, bluegills feed on items as they are encountered or switch food items. This pattern has been observed in all bluegills when predators are absent but only in large (> 100 mm ) bluegills when predators are present. Small bluegills in the presence of predators confine themselves to the cover provided by macrophytes (Werner and Hall 1976; Werner et al. 1983; Osenburg et al. 1988), a behavior which may limit their growth by reducing foraging opportunities.

Large immature insects (primarily Chironomus) in the profundal zone of lakes provide a valuable food resource for bluegills when the insects emerge. It is unlikely that these insects are utilized extensively in their profundal habitat due to anoxic conditions which occur during large portions of the year; some bluegills, however, are taken at depths $>10 \mathrm{~m}$ in the winter (J. C. Schneider, personal communication). Insects emerging from the profundal zone may provide a valuable, though seasonally distributed, food resource (Wiley, personal communication). The seasonal distribution of profundal insects emerging is similar to the seasonal distribution, shown by occurrence in gut samples, of terrestrial insects in the diet (Laarman and Schneider 1972; Beard 1982; Engel 1985).

Anecdotally, this is also consistent with high success rates of anglers fishing with dry flies and surface baits.

Macrophyte density, zooplankton abundance and size, and profundal benthic biomass were significantly related to bluegill growth performance. These variables were, similarly, related to other limnological variables sampled in this study. Although there were not strong correlations between depth or the size of the lake zones and bluegill growth I believe that these factors are important in the structuring bluegill populations because of their influence on distribution and abundance of macrophytes, zooplankton and benthic invertebrates.

Lake morphology was correlated to the density of macrophytes (Appendix 2). Macrophyte distribution is largely, though not entirely, limited by the depth of light penetration (Wetzel 1985). Large, deep (>10 $\mathrm{m})$ lakes with small littoral areas and extensive profundal zones had few macrophytes while shallow, unstratified lakes had dense macrophyte cover. Macrophyte distribution is a key component in the ecology of freshwater lakes (Wetzel 1985). Macrophyte distribution influences zooplankton species diversity (Brooks and Dodson 1965; Pennak 1978), nutrient concentrations in water (Wetzel 1985), algal densities and distribution (Wetzel 1985) and benthic invertebrate species diversity,
abundance and distribution (Gilinsky 1984). Lakes included in this study, as most lakes in the region, are of glacial origin whose sediments are rich in organic matter after 10,000 years of existence. They commonly have macrophytes ringing the shoreline to the depth at which light penetration becomes limiting, indicating a rich supply of nutrients in the sediments.

Zooplankton size distribution and abundance were also related to lake morphology (Appendix 4). The size of zooplankton was positively correlated with size of the profundal stratum and depth and were negatively correlated with size of the littoral strata. Abundance of zooplankton was positively correlated with size of the littoral stratum and negatively to size of the profundal stratum and depth. Littoral zooplankton are often small and commonly occur in large numbers (Brooks and Dodson 1965; Pennak 1978), but the small size class did not correlate significantly with the morphologic variables.

Benthic invertebrate species diversity and abundance can be affected by lake morphology. Many benthic animals have strict habitat requirements which can only be met in certain areas of a lake. For example, many odonates cling to vegetation, some mayflies (Hexagenia sp.) require firm sediments for burrowing and other insects (Chironomidae) can live virtually anywhere (Merrit and Cummins 1984). The invertebrates
correlated with the growth of bluegills, in this study, were profundal animals not commonly found in shallow regions of lakes. Their distribution is undoubtedly limited by the amount of profundal habitat available.

The results of this study identify several variables which influence growth of bluegills. One factor, not studied here, which may be as important as all others combined is density of the bluegill population and structure of the entire fish population. Schneider (1981a) identified "good" lakes as having 50-78\% fish biomass comprised of bluegills while "poor" lakes had $>78 \%$ of the fish community comprised of bluegills. He also determined a predator biomass at $20 \%$ of the total fish biomass as being correlated with good bluegill growth.

Mechanisms determining year-class strength of bluegills have been studied extensively. It appears that most eggs laid will produce larvae and that year-class strength may be determined during the first four days of life (Beard 1982; Engel 1985). Lakes with extensive littoral zones and dense macrophytes are likely to produce large year-classes. Food for larval bluegills (small zooplankton) may be abundant in the littoral habitat and there is protection from predation in the macrophytes. Lakes with small littoral habitats and few macrophytes are likely to produce
small year-classes due to high rates of predation on small bluegills. It is unlikely that food for small bluegills in these lakes would be much less abundant because they can feed on the limited number of littoral zooplankton or on immature members of pelagic zooplankton.

The processes by which large populations of bluegills can limit their own food resources have been identified. Bluegills have been shown to reduce benthic invertebrate size, species abundance and morphology (Hayne and Ball 1956; Gerking 1962; Werner and Hall 1974; Crowder and Cooper 1979, 1982; Gilinsky 1984; Butler 1988; Pierce 1988). In cases where zooplanktivores are size selective in their feeding habits, they can alter size and species composition of the zooplankton community (Brooks and Dodson 1965; Gailbraith 1975; Hall et al. 1976; Mills et al. 1978; Zaret 1980; Mills and Schiavone 1982). Thus, large populations of bluegills feeding in the limnetic region of lakes could cause species changes in the zooplankton community towards smaller species and thus limit this food resource. This effect would be most pronounced in small lakes with high densities of planktivores.

One goal of fishery managers is to provide an ample supply of harvestable fish for anglers. To accomplish this goal for bluegills the manager must consider many factors. Sufficient cover must be available
for some young fish to survive but not so much that most survive. Habitat for important food types, as mentioned above, must be available. Finally, piscivorous fish and forage fish populations must be balanced so the piscivores can control bluegill population abundance.

Plant management can be an effective tool for fishery managers. Using chemicals, fertilizers and mechanical control, they can keep macrophyte density low which may promote good growth of bluegills. Low plant density should provide habitat for large zooplankton species and keep bluegill recruitment low through higher rates of predation.

## LITERATURE CITED

Beard, T. D. 1982. Population dynamics of young-of-the-year bluegill. Wisconsin Dept. Nat. Res., Technical Report No. 127., Madison, Wi., USA

Brooks, J. L. and S. I. Dodson. 1965. Predation, body size and composition of plankton. Science 150:28-35.

Butler, M. J. 1988. Community responses to variable predation: field studies with sunfish and freshwater macroinvertebrates. Ecological Monographs 59:311-328.

Cooper, E. L., C. C. Wagner and G. E. Krantz. 1971. Bluegills dominate production in a mixed population of fish. Ecology 52:280-290.

Crowder, L. B. and W. E. Cooper. 1979. Structural complexity and fish-prey interactions in ponds: a point of view. Pages 2-10 in D. L. Johnson and R. A. Stein, eds. Response of fish to habitat structure in standing water. North Central Div., Am. Fish. Soc., Special Pub. No.6.
$\qquad$ 1982. Behavioral interactions between fish predators and their prey: Effects of plant density. Anim. Behav. 37:311-321.

Davis, C.O. and M. A. Simmons. 1979. Water chemistry and phytoplankton field and laboratory procedures. Great Lakes Research Division Special Report No. 70., Univ. of Mich., Ann Arbor, Mi. USA.,

Diamond, J. M. 1978. Niche shifts and the rediscovery of interspecific competition. Am. Sci. 66:322-331.

Engel, S. 1985. Aquatic community interactions of submerged macrophytes. Wisconsin Dept. Nat.Res., Technical Report No. 156, Madison, Wi., USA.

Gailbraith, M. G. Jr. 1975. The use of large Daphnia as indices of fishing quality for rainbow trout in small lakes. Verh. Internat. Verein. Limnol. 19:2485-2492.

Gerking, S. D. 1962. Production and food utilization in a population of bluegill sunfish. Ecological Monograph 32:31-78.

Gilinsky, E. 1984. The role of fish predation and spatial heterogeneity in determining the benthic community structure. Ecology 65:455-468.

Grice, F. 1957. Effect of removal of panfish and trashfish by fyke nets upon fish populations of some Massachusetts' ponds. Trans. Am. Fish. Soc. 87:108-115.

Hall, D. J., E. C. Cooper, E. E. Werner. 1970. An experimental approach to the production dynamics and structure of freshwater animal communities. Limnol. Oceanogr. 15:839-928.

Hall, D. J., S. T. Threlkeld, C. W. Burns, and P. H. Crowley. 1976. The Size Effeciency Hypothesis (S.E.H.) and the size structure of zooplankton communities. Ann. Rev. Ecol. Syst. 7:177-208.

Hayne, D. W., and R. C. Ball. 1956. Benthic productivity as influenced by fish predation. Limnol. and Oceanogr. 1:162-175.

JAVA. 1988. Jandel Video Analysis Software. Jandel Scientific, Corte Madera, CA., USA.

Keast, A. 1978. Feeding interelations between age groups of pumpkinseeds (Lepomis gibbosus) and comparisons with bluegill sunfish (Lepomis macrochirus). J. Fish. Res. Board Can. 35:12-27.

Latta, W. C. and J. W. Merna. 1977. Some factors influencing size of the year class of bluegills (Lepomis macrochirus) in ponds. Mich. Acad. IX:483-502

Laarman, P. W. and J. C. Schneider. 1972. The food and feeding habits of the bluegill and yellow perch in lakes with good and poor fishing. Mich. Dept. Nat. Res., Fish. Res. Rep. no. 279.

Lemly, D. A. and J. F. Dimmick. 1982. Growth of young-of-the-year and yearling centrarchids in relation to zooplankton in the littoral zone of lakes. Copea 1982(2):305-321.

Merrit, R. W. and K. W. Cummins. 1984. An introduction to the aquatic insects of North America. Kendall/Hunt Pub. Co. Dubuque, IA., U.S.A. pp. 722.

Mills, E. L. and A. Schiavone Jr. 1982. Evaluation of fish communities through the assessment of zooplankton populations and measures of lake productivity. N. Am. J. Fish. Mgmt. 2:14-27.

Mills, E. L., D. M. Green and A. Schiavone Jr. 1978. Use of zooplankton size to assess the community structure of fish populations in freshwater lakes. N. Am. J. Fish. Mgmt. 7:369-378.

Mittelbach, G. G. 1981. Foraging efficiency and body size: a study of optimal diet and habitat use by bluegills. Ecology 62:1370-1386.

Osenburg, C. W., E. E. Werner, G. G. Mittelbach and D. J. Hall. 1988. Growth patterns in bluegill (Lepomis macrochirus) and pumpkinseed (L. gibbosis) sunfish: environmental variation and the importance of ontogenetic niche shifts. Can. J. Fish. Aqat. Sci. 45:17-26.

Pardue, G. B. 1973. Production responce of the bluegill sunfish (Lepomis macrochirus Rafinesque) to added attachment surface for fish food organisms. Trans. Am. Fish. Soc. 102:622-626.

Parker, R. A. 1958. Some effects of thinning on a population of fishes. Ecology 39:304-317.

Pennak, R. W. 1978. Freshwater invertebrates of the United States, 2nd ed. John Wiley and Sons, New York, NY, USA.

Pierce, C. L. 1988. Predator avoidance, microhabitat shift, and risk sensitive foraging in larval dragonflies. Oecologia 77:81-90.

Ryder, R. A., S. R. Kerr, K. H. Loftus and H. A. Regier. 1974. The morphoedaphic index, a fish yield estimator - review and evaluation. J. Fish. Res. Board Can. 31:663-688.

Savino, J. F., and R. A. Stein. 1979. Predator-prey interaction between largemouth bass and bluegills as influenced by simulated submersed vegetation. Trans. Am. Fish. Soc. 111:255-266.
1982. Habitat structural complexity and the interaction between bluegills and their prey. Ecology 63:1802-1813.
$\qquad$ 1989. Behavioral interactions between fish predators and their prey: effects of plant density. Anim. Behav. 37:311-321.

Schneider, J. C. 1975a. Typology and fisheries potential of Michigan lakes. Mich. Acad. VIII:59-84.

1975b. Fisheries classification of Michigan lakes. Mich. Dep. Nat. Res., Fish. Res. Rep. no. 1822.
$\qquad$ 1978. Predicting the standing crop of fish in Michigan lakes. Mich. Dep. Nat. Res, Fish. Res. Rep. no. 1860.
$\qquad$ 1981a. Fish communities in warmwater lakes. Mich. Dep. Nat. Res., Fish. Res. Rep. no. 1890.
___ 1981b. Growth Index for Michigan fish. In Manual of Fisheries Survey Methods. Michigan Dept. Nat. Res. Lansing Mi., USA

Weiner, J. G., and W. R. Hanneman. 1982. Growth and condition of bluegills in Wisconsin lakes: effects of population density and pH . Trans. Am. Fish. Soc. 111:761-767.

Werner, E. E. 1979. Species packing and niche complementarity in three sunfishes. Am. Nat. 111:553-578.

Werner, E. E. and D. J. Hall. 1974. Optimal foraging and the size selection of prey by the bluegill sunfish (Lepomis macrochirus). Ecology 55:10421052.
$\qquad$ 1976. Niche shifts in sunfishes:experimental evidence and significance. Science 191:404-406
$\qquad$ 1977. Competition and habitat shift in two sunfishes (Centrarchidae). Ecology 58:869-876.
$\qquad$ 1979. Foraging efficiency and habitat switching in competing sunfishes. Ecology 60(2):256-264.

Werner, E. E., D. J. Hall, D. R. Laughlin, D. J. Wagner, L. T. Wilsman and F. C. Funk. 1977. Habitat partitioning in a freshwater fish community. J. Fish. Res. Board Can. 34:360-370.

Werner, E. E., and G. G. Mittelbach. 1981. Optimal foraging: field tests of diet choice and habitat switching. Am. Zool. 21:813-829.

Werner, E. E., J. F. Gilliam, D. J. Hall, and G. G. Mittelbach. 1983. An experimental test of the effects of predation risk on habitat use in fish. Ecology 64:1540-1548.

Werner, E. E., G. G. Mittelbach, D. J. Hall and J. F. Gilliam. 1983. Experimental tests of optimal habitat use in fish: the role of relatave habitat profitability. Ecology 64:1525-1539.

Wetzel, R. G. 1983. Limnology. W. B. Saunders Company, Philidelphia, PA., USA.

Wiley, M. J. 1981. Chironomids of Michigan trout streams. Mich. Acad. vol. XIII

Wiley, M. J., R. W. Gordon, S. W. Waite and T. Powles. 1984. The relationship between aquatic macrophytes and sport fish production in Illinois ponds: a simple model. No. Am. J. Fish. Mgmt. 4:111-119

Zaret, T. M. 1980. Predation in freshwater communities. Yale University Press, New Haven, USA.

Appendix 1. Benthic invertebrates found in the study lakes.

| Class | Order | Family | Genus |
| :---: | :---: | :---: | :---: |
| Turbellaria | Tricladida | Planariidae |  |
| Annelida | Oligochaeta |  |  |
|  | Hirudinea |  |  |
| Crustacea | Isopoda |  |  |
|  | Amphipoda | Taltridae | Hyalella |
|  |  | Gammaridae | Gammarus |
| Arachnoidea Insecta | Hydracarina Ephemeroptera |  |  |
|  |  | Baetidae | Baetis |
|  |  |  | Calibaetis |
|  |  | Ephemeridae | Ephemera |
|  |  |  | Hexagenia |
|  |  | Heptageniidae | Stenacron |
|  |  | Oligonueridae | Isonychia |
|  |  | Caenidae | Caenis |
|  |  | Ephemerellidae | Eurylophella |
|  | Odonata | Coenagrionidae | Argia |
|  |  |  | Enallagma |
|  |  |  | Nehalennia |
|  |  | Corduliidae | Cordulia |
|  |  |  | Epitheca |
|  |  |  | Neurocordulia |
|  |  |  | Stomatochlora |
|  |  | Gomphidae | Arigomphus |
|  |  |  | Gomphus |
|  |  |  | Stylurus |
|  |  | Libellulidae | Celethimis |
|  |  |  | Erythemis |
|  |  |  | Ladona |
|  |  |  | Leucorrhinia |
|  |  |  | Libellula |
|  |  |  | Pachydiplax |
|  |  | Macromiidae | Didymops |
|  |  |  | Macromia |
|  | Hemiptera | Naucoridae | Pelocoris |
|  | Megaloptera | Sialidae | Sialis |

Polycentropodidae
Neuroclipsis Nyctiophylax Polycentropus
Hydroptilidae Hydroptila
Orthotrichia
Oxyethira
Helicopsychidae
Helicopysche
Leptoceridae Ceraclea
Leptocerus
Nectopsyche
Oecetis
Triaenodes
Molanidae Molanna
Phryganeidae Fabria
Agriypnia
Pyralidae
Lepidoptera
Coleoptera (larvae)

Diptera
Dytiscidae Haliplidae Elmidae Stenelmis Ceratopogonidae Chaoboridae

Chaoborus
Chironomidae
Tabanidae
Chrysops

Appendix 2. Correlations between macrophyte density and other variables. ( $\mathrm{N}=30$, alpha=.05). $\mathrm{NS}=$ not significant.

| Variable | Littoral index | \% Coverage |
| :---: | :---: | :---: |
| nitrate | NS | NS |
| ammonia | NS | NS |
| orthophosphate | NS | NS |
| total phosphorus | 0.449 | 0.394 |
| dissolved oxygen | NS | NS |
| alkalinity | NS | NS |
| macrophyte density index | -- | 0.750 |
| \% coverage | 0.750 | - - |
| size |  |  |
| stratum 1 | NS | 0.683 |
| stratum 2 | NS | NS |
| stratum 3 | -0.401 | -0.646 |
| whole lake | NS | -0.360 |
| maximum depth | -0.428 | -0.676 |

Appendix 3. Correlations between algal standing crop and other variables. ( $\mathrm{N}=30$, alpha=.05). $\mathrm{NS}=$ not significant.

| Variable | Secchi disk | chlorophyll a |
| :---: | :---: | :---: |
| secchi diskdepth | ---- | -0.480 |
| chlorophyll a | -0.480 | ---- |
| nitrate | NS | NS |
| ammonia | -0.372 | NS |
| orthophosphate | NS | NS |
| total phosphorus | NS | NS |
| dissolved oxygen | NS | NS |
| alkalinity | NS | NS |
| macrophyte density index | NS | NS |
| \% s.a. coverage | NS | 0.436 |
| size |  |  |
| stratum 1 | NS | 0.409 |
| stratum 2 | NS | NS |
| stratum 3 | NS | NS |
| whole lake | NS | NS |
| maximum depth | 0.501 | -0.394 |
| growth index | NS | NS |

Appendix 4. Correlations between zooplankton and other variables. ( $\mathrm{N}=30$, alpha= .05). NS $=$ not significant.


Appendix 5. Correlations between food items and other variables. ( $N=30$, alpha $=.05$ ). NS $=$ not significant.

| Variable | Littoral | Sublittoral | Profundal | Whole lake |
| :---: | :---: | :---: | :---: | :---: |
| secchi disk depth | NS | 0.402 | 0.381 | NS |
| chlorophyll a | NS | NS | NS | NS |
| nitrate | NS | NS | NS | NS |
| ammonia | 0.401 | NS | NS | 0.521 |
| orthophosphate | NS | NS | NS | NS |
| total phosphorus | NS | NS | NS | NS |
| dissolved oxygen | NS | NS | NS | NS |
| alkalinity macrophyte | NS | NS | NS | NS |
| index | NS | NS | NS | NS |
| \% s.a. coverage | NS | NS | NS | -0.417 |
| size |  |  |  |  |
| stratum 1 | NS | NS | NS | NS |
| stratum 2 | NS | NS | NS | NS |
| stratum 3 | NS | NS | NS | NS |
| whole lake | NS | NS | NS | NS |
| maximum depth | NS | NS | NS | NS |
| zooplankton |  |  |  |  |
| large size | NS | NS | NS | NS |
| small size | NS | NS | NS | NS |
| density | NS | NS | NS | NS |
| growth index | NS | NS | NS | NS |

Appendix 6. Correlations between large food items and other variables. ( $\mathrm{N}=30$, alpha= .05). $\mathrm{NS}=$ not significant.

| Variable | Littoral | Sublittoral | Profundal | Whole lake |
| :---: | :---: | :---: | :---: | :---: |
| secchi disk depth chlorophyll a | NS | NS | NS | NS |
|  | NS | NS | NS | NS |
|  | NS | NS | NS | NS |
| ammonia | NS | NS | NS | 0.455 |
| orthophosphate | 0.483 | NS | NS | 0.472 |
| total phosphorus | NS | NS | NS | NS |
| dissolved oxygen | NS | -0.416 | NS | NS |
| macrophytes |  |  |  |  |
| index | NS | NS | NS | -0.443 |
| \% s.a. coverage | NS | NS | NS | -0.362 |
| size |  |  |  |  |
| stratum 1 | NS | NS | NS | NS |
| stratum 2 | NS | NS | NS | NS |
| stratum 3 | NS | NS | NS | NS |
| whole lake | NS | NS | NS | NS |
| maximum depth zooplankton | NS | NS | NS | NS |
| mean size | NS | NS | NS | 0.389 |
| large size | NS | NS | 0.367 | NS |
| small size | NS | 0.437 | 0.412 | NS |
| density | NS | NS | NS | NS |
| growth index | NS | NS | 0.444 | NS |

Appendix 7. Correlation between growth index and other variables. ( $\mathrm{N}=30$, alpha $=.05$ ). NS $=$ not significant.

| Variable Growth | performance |
| :---: | :---: |
| secchi disk depth | NS |
| chlorophyll a | NS |
| nitrate | NS |
| ammonia | NS |
| orthophosphate | NS |
| total phosphorus | NS |
| dissolved oxygen | NS |
| alkalinity | NS |
| macrophytes |  |
| index | -0.443 |
| \% s.a. coverage | -0.431 |
| zooplankton |  |
| mean size | 0.582 |
| large size | 0.622 |
| small size | NS |
| density | -0.361 |
| benthic invertebrate biomass |  |
| stratum 1 | NS |
| stratum 2 | NS |
| stratum 3 | NS |
| whole lake | NS |
| large benthic invertebrate. biomass |  |
| stratum 1 | NS |
| stratum 2 | NS |
| stratum 3 | 0.444 |
| whole lake | NS |
| size |  |
| stratum 1 | NS |
| stratum 2 | NS |
| stratum 3 | NS |
| whole lake | NS |
| maximum depth | NS |

