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Trophodynamics of the Fish *Valenciennellus tripunctulatus*. III. Energetics, Resources and Feeding Strategy

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ABSTRACT: A model was constructed which relates energy in diet to metabolic requirements for the mesopelagic zooplanktivore fish *Valenciennellus tripunctulatus*. Linear and Von Bertalanffy growth models were used to explore the energetic consequences of a number of feasible age/growth strategies. Maximum efficiencies were obtained with linear growth to about 30 mm in 1 y. Efficiencies declined rapidly for slower growth strategies. It was hypothesized that an 'annual' life history strategy was optimum based on energetic considerations. Copepod prey in the 1-2 mm size class were an order of magnitude more abundant and over 4 times higher in caloric content than larger size classes. *V. tripunctulatus*, however, is highly efficient in obtaining calories from larger prey size classes (2-4 mm) which contain a high caloric content per individual. The resource field places severe constraints on predator growth and at 38 mm no surplus energy is estimated available from daily ration over and above metabolic requirements. *V. tripunctulatus* should be sensitive to 'nearfield' competitors as well as 'remote' predators grazing zooplankton populations in different space/time.

INTRODUCTION

Ecologists have recently begun to explore organism-environment interactions in terms of genetic systems (populations *sensu* Mayr, 1963) which delineate 'adaptive strategies' that allow these systems to persist. These strategies are envisioned as being adaptively directed or 'optimized' through natural selection (e.g. Levins, 1968; Morse, 1971; Baird, 1974). The strategic concept has been applied to feeding and a sizable literature on models of feeding strategies now exists (e.g. MacArthur and Pianka, 1966; Emlen, 1966, 1968; Slobodkin, 1968; Holling, 1968; Schoener, 1971). In this paper we consider the role of energetics in the trophodynamics of the small mesopelagic fish *Valenciennellus tripunctulatus* and explore the energetic consequences of patterns of resource availability in relation to the more general concepts of life history and feeding strategies in mesopelagic zooplanktivores.

ENERGY BUDGETS

We have attempted to construct an energy budget for *Valenciennellus tripunctulatus* based on known prop-

erties of the fish and its diet. Direct metabolic measurements on deep sea fishes are rare (Torres et al., 1979) and only two parameters in the equation that follows were directly measured. While the metabolic cost functions used in the model are consistent with our present understanding of the processes involved and serve as first approximations only, they are necessarily based on extrapolations from existing data in the literature. Nonetheless the energy budget model allows us to develop biologically reasonable scenarios about critical ecological processes that are not well known and about which information is exceedingly difficult to obtain.

An equation developed by investigators of fish trophodynamics was used to express the dynamics of energetics and metabolism (e.g. Warren and Davis, 1967; Weatherly, 1972). The equation, expressed in terms of calories, states that:

$$Q_c = Q_g + Q_w + Q_d + Q_s + Q_a$$

or

$$Q_c = Q_g + Q_s + Q_r$$

where:

Q_c = energy of ingested ration

Q_g = increase in potential energy through growth

$$Q_s = \begin{cases} Q_w = \text{energy lost as feces, urine and} \\ \text{secretions through skin and gills} \\ Q_d = \text{energy cost of digestion,} \\ \text{assimilation and storage of food} \\ \text{(often referred to as SDA, or} \\ \text{specific dynamic action)} \end{cases}$$

$$Q_r = \begin{cases} Q_s = \text{energy of metabolism of unfed} \\ \text{resting fish, i.e. standard or} \\ \text{basal metabolism} \\ Q_a = \text{energy of activity in excess of} \\ \text{standard metabolism, e.g.} \\ \text{swimming} \end{cases}$$

Assumptions and approximations for each term of the equation were as follows:

Q_c : Estimated from approximations of total food ingested over a given time period multiplied by its caloric value. Since copepods constituted 95 % or more of the diet of *Valenciennellus tripunctulatus* over all sizes of fish examined, the caloric content of the ingested ration was based on a copepod diet. As determined by Comita and Schindler (1963) for the genus *Calanus*, 5914 cal g⁻¹ ash-free dry weight (AFDW), adjusted for 5 % salt error in our prey weights (i.e. 5914 × 0.95 = 5617 cal g⁻¹), was used for caloric conversion of prey biomass.

Q_g : Increase in potential energy through growth of an individual (5617 cal g⁻¹ DW). Weights were based on direct measurements of the length/dry weight relationship in *Valenciennellus tripunctulatus* (Hopkins and Baird, 1981).

Q_w : A figure of 0.22 Q_c was used to account for losses to urine and feces, a conservative estimate based on a wide spectrum of teleost fishes (Brett and Groves,

1979). Losses from leakage and skin secretions were assumed to be minimal.

Q_d : Estimates of specific dynamic action or 'entropic tax' are as high as 0.46 Q_d (Warren and Davis, 1967) and considerable variation has been reported in teleosts (Brett and Groves, 1979). Recent estimates on the bleak are about 0.15 Q_c (Muir and Niimi, 1972; Ware, 1975), and we have chosen the latter as representing the most efficient estimate reported in the literature.

$Q_s + Q_a$: Neither standard metabolism or active metabolism have been precisely determined for mid-water fishes. Based on estimates of 'routine' or 'normal' activity (*sensu* Childress, 1975), a value of approximately 0.115 $\mu\text{l O}_2 \text{ mg}^{-1} \text{ wet wt h}^{-1}$ was calculated for the mesopelagic hatchetfish *Argyropelecus sladeni* by Childress and Nygaard (1973), and that value was used here for respiration. *A. sladeni* exhibits limited vertical migration in the upper mesopelagic environment (minimum depth of occurrence: 100 m; see Baird, 1971; Childress and Nygaard, 1973), and is thought to be phylogenetically the least remote from *Valenciennellus tripunctulatus* of the fish they examined. Respiration was assumed constant per unit weight over all sizes considered. Weight specific respiration rates are known to vary in fishes (e.g. Paloheimo and Dickie, 1966; Weatherley, 1972) with higher rates per unit weight often observed in smaller individuals. The size classes considered here are quite small in range (15-30 mm), however, and per unit weight differences in respiration rate were assumed not to greatly effect our first order approximations (i. e. other sources of variability are likely to exceed any size effect). For *V. tripunctulatus*, AFDW is 0.87 DW, and, using Childress

Table 1. *Valenciennellus tripunctulatus*. Regressions on fish standard length (X; mm) of various diet characteristics (Y). Regressions 1 and 2 were non-significant (N. S.); 5 was significant only to F = 0.05; all other regressions were significant to at least F < 0.01. Stomach and diet data are from fishes taken during period of most active feeding. Y values require the conversion indicated in the transformation column*

Dependent variable	Regression equation	Transformation	Corr. Coef. r	Range of Y	No. fish in sample	Comments
1. Avg. no. prey items per stomach	N.S.	(Y+0.5) ^{1/2}	0.08	0-24	195	Empty stomachs included in average
2. Max. no. prey items per stomach	N.S.	none	0.27	9-24	13	Based on stomachs with most prey in each fish size class
3. Fish biomass (mg AFDW)	Y = -0.116 + 0.061X	Log ₁₀ Y	0.95	6.1-31.8	6	
4. Avg. prey biomass in stomachs (mg DW)	Y = 0.637 + 0.013X	(Y+0.5) ^{1/2}	0.22	0.0-1.87	195	See comment for Equation 1
5. Max. prey biomass in stomachs (mg DW)	Y = -1.379 + 0.103X	none	0.86	0.57-1.87	13	Based on stomachs with most biomass in each fish size class

* Log₁₀ conversion was necessary for Regression 3 to normalize skewed data. The (Y+0.5)^{1/2} transformation was used in Regressions 1 and 4 because of the large number of zero values and the resulting skewness

and Nygaard's (1973) wet weight (WW) to DW ratios, our AFDW regression (Table 1), and integrating over the desired time interval (e.g. days, years), total respiration is obtained. This total actually represents mean respiration over a given time interval (i.e. it averages respiration rates which vary over shorter time intervals such as a diel period). To convert respiration to energy cost or calories, values of 5007, 4686, and 4500 cal l⁻¹ O₂ were assumed for the combustion of carbohydrate, lipid, and protein (Giese, 1962). The ratio of these components in the copepod genus *Calanus* is reported to be 1.0:5.4:33.6 (Raymont, 1963). By the assumption of those proportions and their oxy-caloric equivalency, a representative value for copepods was estimated to be 4538 cal l⁻¹ O₂.

We have used the distributional information, diet composition data, and minimum/maximum estimates of daily ration developed in Hopkins and Baird (1981) as input to the energetic and growth models used here. Regression Equations 1-5 were used to determine fish weight (AFDW), prey numbers, and biomass of food consumed in the daily ration for any given fish length. The daily ration-fish length regression can then be integrated over the various time periods and growth rates being considered, to calculate total food consumed during that period. For example, if growth is linear, the total prey biomass ingested during growth from 10 to 30 mm during a time period of 11 months is estimated at 133 and 255 mg DW, respectively, for average and maximum ration. This is the method used to calculate caloric intake for the various periods and growth rates considered in Table 2.

No direct means were available for the determination of actual age/growth relationships, and such data are non-existent for mesopelagic species in low latitude oligotrophic environments. Several feasible growth strategies were examined in relation to

ingested energy. An arbitrarily determined period of 30 d was allowed for larval development and commencement of metamorphosis at approximately 10 mm SL. This was based on developmental times for certain clupeid species having pelagic larvae (e.g. Houde and Palko, 1970; Houde and Swanson, 1975). Energy consumed during the pre-metamorphic period was not considered in the analysis.

The maximum length of an individual observed in our collections was 35 mm SL, with the great majority of the population ranging from 15-30 mm SL (Hopkins and Baird, 1981; their Fig 3). Similar size ranges have been reported for other North Atlantic populations (Krueger, 1972; Badcock and Merrett, 1976). Energy budgets were calculated for a maximum fish length of 38 or 40 mm SL depending on the growth model used.

Two basic models were used to calculate potential rates of growth for various time periods. One is a linear model in which,

$$l_t = l_{t_0} + t/t_m (l_{t_m} - l_{t_0})$$

where: l_t = length at time t ; l_{t_0} = length at t_0 (i.e. 10 mm SL); l_{t_m} = length at maximum assumed age t_m ; t_0 = 0.08 y (approximately 30 d); time of larval development; t_m = assumed maximum age. The other is a curvilinear or Von Bertalanffy (1938) model, which expresses growth rate as decreasing with increasing size. In this model, l_t is expressed as follows:

$$l_t = l_{\infty} (1 - e^{-R(t-t_0)})$$

where: l_{∞} = a theoretical maximum length (i.e. arbitrarily set at 38 mm SL); R = growth term or the rate at which growth curve reaches the asymptotic length.

Energy budgets were then calculated based on maximum and minimum daily ration for various assumed rates of linear growth (Table 2). Various time periods (t_m) from 5 to 23 months corresponding to maximum ages of 0.5 to 2 y, were used to determine linear rates of

Table 2. *Valenciennellus tripunctulatus*. Caloric budget assuming various size ranges, rations and periods of growth. Linear growth assumed

Daily ration	Maximum length (l max, mm)	Time at l max (y)	Energy budget (calories)				Balance		Conversion efficiency Q _g /Q _c (%)
			Q _c	Q _g	Q _s	Q _r	cal.	% Q _c	
Maximum	30	0.5	652	245	261	184	- 38	(-) 6	(-)38
	30	1.0	1435	245	575	405	210	15	17
	30	1.5	2220	245	888	625	462	21	11
	30	2.0	3006	245	1202	847	712	24	8
Minimum	30	0.5	341	245	136	184	-224	(-)66	(-)72
	30	1.0	750	245	300	405	-200	(-)27	(-)33
	30	1.5	1161	245	464	625	-173	(-)15	(-)21
	30	2.0	1571	245	629	847	-150	(-)10	(-)16
Maximum	35	1.0	1868	510	747	664	- 53	(-) 3	(-)27
	35	1.5	2891	510	1156	1036	189	6	18
Minimum	35	2.0	1889	510	755	1391	-767	(-)41	(-)27
Maximum	40	2.0	4894	1046	1958	2428	-538	(-)11	(-)21

growth to maximum sizes (l_{∞}) of 30, 35, and 40 mm. The Von Bertalanffy model was used to assess the energetic consequences of a longer life cycle in which the growth rate is reduced as age increases. The actual values used in the Von Bertalanffy model produced a rate of growth which is less than that reported for several mesopelagic myctophid species (Odate, 1966; Halliday, 1970; Smoker and Pearcy, 1970) and represents very slow growth compared to known rates in a number of epipelagic marine species. The assumed Von Bertalanffy values were $R = 0.83$, $t_0 = 0.08$ y and $l_{\infty} = 38$ mm, which result in estimates of a mean length of about 20 mm for age class one, 30 mm for age two, and 35 mm for age class three fish.

Graphs of the various modeled growth strategies are depicted in Figure 1. The energy budgets and conversion efficiencies calculated from these growth curves appear in Tables 2 and 3. Considering first the ration estimate, the caloric equivalents for most crustaceans

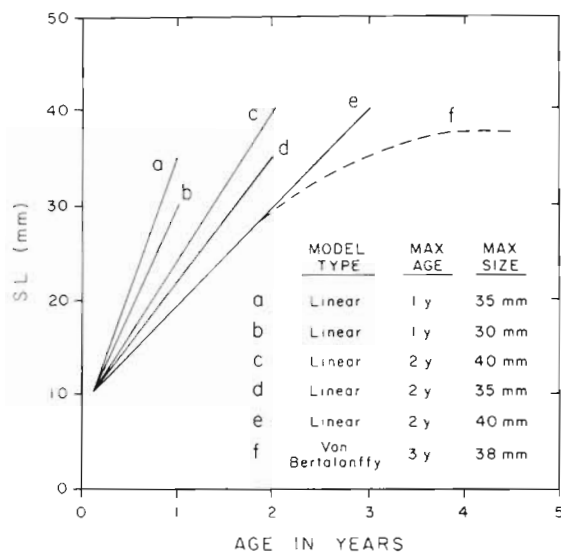


Fig. 1. *Valenciennellus tripunctulatus*. Projected growth curves based on maximum estimated daily ration (see text)

Table 3. *Valenciennellus tripunctulatus*. Conversion efficiency in percent of caloric intake at maximum ration to caloric equivalent of fish weight for various lengths and growth rates. Linear growth rates are assumed for Years 1 and 2; decline in growth rate (Von Bertalanffy effect) assumed for third year (see Fig. 1). In parentheses: efficiency at balance (% $Q_g = 0$) obtained by subtracting our excess in energy balance equation (i.e. Q_c considered = $Q_c - \text{surplus}$)

Maximum length	Year 1	Year 2	Year 3
			(Von Bertalanffy growth)
30	17 (23)	8 (12)	< 5
35	27 (30)	13 (17)	6

fall within narrow limits (5200–5900 cal g^{-1} AFDW; Cummins and Wuycheck, 1971), and a major error in caloric conversion is unlikely. The minimum ration as calculated here is considered an underestimate. That is given some support by the data in Table 2 (e.g. growth to 30 mm appears possible only after 18 mo to 2 y). The population is apparently not feeding in synchrony and in any given daylight sample empty stomachs may occur, lowering the value for minimum ration. The actual energy intake is predicted to fall between average and maximum as calculated here.

Considering the right side of the energy equation, it is apparent that growth (Q_g) and respiration ($Q_s + Q_a$) are the major factors contributing to energy costs. The other items are a function of ration intake and were conservatively estimated by assuming low values reported in the literature for teleost fishes. Q_g is a function of biomass, determinations of which are relatively accurate. Gonad weight was estimated to be 8–10 % of body weight in mature individuals and was included in biomass estimates. Gonadal material, however, is generally higher in energy content than somatic tissue, and an underestimation of caloric conversion at larger fish sizes may result.

Respiratory metabolism ($Q_s + Q_a$) was the most difficult term to estimate as little metabolic or behavioral data exist. Consequently, a higher uncertainty must be associated with respiratory cost factors. Recent measurements of resting metabolism in several mesopelagic myctophids range from 0.059 to 0.09 (μl O_2 $mgWW^{-1} h^{-1}$) (Torres et al., 1979) and are comparable to the 0.115 used here. Our estimate represents 'low routine' activity for 24 h (Childress and Nygaard, 1973; Childress, 1975). Unexpectedly large metabolic demands from feeding, escape or reproductive behavior (including, for example, increases in metabolism from egg production) might result in underestimation, while periods of very low activity would tend to correct in the other direction.

ENERGETICS AND GROWTH

A number of important ecological consequences follow directly from a consideration of the energy budgets presented in Table 2. It is, for instance, unlikely that a size of 30 mm SL could be reached by *Valenciennellus tripunctulatus* in 6 mo at either maximum or minimum daily ration. Significant positive energy balances are obtained at maximum daily ration and growth to 30 mm in 1 y. If the true daily ration lies between our mean and maximum, then growth to 27–30 mm in 1 y can be achieved.

The minimum ration yields negative energy budgets under the age-growth models depicted. Only

strategies involving long age (2+ y), slow growth and small adult size can theoretically produce viable populations of *Valenciennellus tripunctulatus* at ingestion rates approaching the estimated minimum ration. Daily ration determines the upper limit to any viable age-growth strategy which in this case theoretically limits growth to about 27–32 mm in the first year.

The energetic implications of various growth strategies in terms of conversion efficiencies (i.e. tissue accumulation efficiency *sensu* Kozlovsky, 1968) which compare caloric intake (at maximum daily ration) to caloric equivalent of fish weight as growth are apparent in Table 3. Calculations were made on the basis of ingested energy and energy required to exactly balance the energy budget. It can be seen that as the time to reach a given size increases, conversion efficiency decreases. At ages beyond 2 y, efficiencies fall below 10 %. Conversely growth to 35 mm in 1 y represents an high conversion efficiency (27 %) which may exceed the metabolic capabilities of vertebrate physiology. Maximum conversion efficiencies of about 25 % have been reported for young fishes under experimental conditions (e.g. Brett et al., 1969, for salmonids) and the estimated 17–23 % efficiencies predicted for growth to 30 mm in 1 y cannot be excluded on physiological grounds. Growth rates of 35 mm SL in 2 y also yield efficiencies in excess of 13 %. Slower growth options (30–35 mm 3 y⁻¹) of the Von Bertalanffy type yield lower efficiencies (i.e. less than 10 %), representing levels of utilization which would theoretically put individuals at a competitive disadvantage.

ENERGETICS, FOOD AVAILABILITY, AND RESOURCE UTILIZATION

A number of fundamental relationships in the trophodynamics of *Valenciennellus tripunctulatus* can be depicted in general form from a consideration of energy budgets, estimates of the caloric value of various sized prey items, prey density in the environment, and prey utilization by size class. The information is

Table 4. Size, weight and caloric value as a function of size for the copepod genus *Pleuromamma*. Caloric equivalency = 5600 cal g⁻¹ DW; DW = dry weight

Size (mm)	mg DW	Calories
1.0	0.007	0.039
1.5	0.025	0.140
2.0	0.044	0.246
2.5	0.068	0.381
3.0	0.110	0.618
3.5	0.185	1.037
4.0	0.294	1.647
4.5	0.457	2.559
5.0	0.710	3.976

summarized in Tables 4–7. Table 4 lists the estimated caloric equivalency of a size-series of copepods of the genus *Pleuromamma* using a caloric conversion value for the genus *Calanus*. From those data, it is possible to obtain a first approximation of the caloric content represented in each prey size class in the environment (Table 5). For purposes of estimate, the caloric equivalency of an individual of length one half the size class range (e.g. 1.5 mm for size class 1–2 mm) was assigned to every individual in that size class. Prey density was taken from data presented by Baird and Hopkins (1981). Information on prey utilization and encounter rates were obtained from the same source and estimates of the caloric contribution of each prey size class to the diet and to total prey available in the expected search volume were made. Data on caloric intake and % daily prey production removed are listed in Table 6. Daily energy budgets for various adult fish sizes based on the maximum daily ration were calculated (Table 7) assuming a zero daily growth rate (Q_g set at 0). No assumptions were made as to utilization of surplus calories over metabolic costs in this case.

The energetic consequences of diel vertical migration of prey in various size classes are readily apparent (Table 5). The reduction in available energy at night is dramatic and appears to affect all prey size classes. If perception distances remain constant (Tables 6 and 7) and grazing efficiency were set at 100 %, available energy at night would be insufficient to support an

Table 5. Estimated caloric content of copepod prey per m³ of water at depths of 300–350 m proportioned by prey size classes. Caloric equivalency based on members of the genus *Pleuromamma* and a mean prey size of 1/2 the size range of an individual prey size class. D: day; N: night. Prey availability data from Table 1 of Baird and Hopkins (1981)

Prey size class (mm)	Estimated calories item ⁻¹	Individual items m ⁻³		Total calories m ⁻³		N/D (%)
		D	N	D	N	
0.5–0.9	0.006	31.92	28.73	0.19	0.17	90
1.0–1.9	0.140	24.44	16.23	3.42	2.27	66
2.0–2.9	0.381	1.99	1.17	0.76	0.45	59
3.0–3.9	1.036	0.70	0.30	0.72	0.31	43
4.0–4.9	2.559	0.10	0.04	0.26	0.10	38

Table 6. *Valenciennellus tripunctulatus*. Estimated caloric intake of copepods at mean and maximum daily ration as a function of prey size class for individual of 26 mmSL (< 1 mm category not considered)

Parameters	Prey size class (mm)			
	1.0–1.9	2.0–2.9	3.0–3.9	4.0–4.9
Minimum no. eaten d ⁻¹	3.51	2.52	1.80	0.63
Maximum no. eaten d ⁻¹	8.00	5.74	4.10	1.43
No. encountered d ⁻¹	29.08	5.17	4.01	1.18
Minimum caloric intake d ⁻¹	0.49	0.96	1.87	1.61
Maximum caloric intake d ⁻¹	1.12	2.19	4.25	3.66
Caloric value of prey encountered (day)	4.07	1.97	4.16	3.02
Caloric value of prey encountered (night)	2.68	1.16	1.83	1.15
% calories encountered in daytime removed at minimum intake	12	49	45	53
% calories encountered in daytime removed at maximum intake	28	111	102	121
No. items to produce caloric equivalent of largest size class	18.3	6.7	2.5	1.0

individual of 27 mm SL at maximum ration. Further, if, because of decreased light intensity, perception distances were reduced at night by about 1/2 those used for calculations in Table 6, energy intake could be reduced by about 2/3 that of daytime values if energy costs for search remain constant.

Considering resource utilization, *Valenciennellus tripunctulatus* appears highly efficient in obtaining energy from the 2–5 mm prey size classes (Table 6). Efficiency falls off, however, for smaller (< 1.9 mm) and larger (> 4 mm) prey. It is also apparent that relatively large increases in density of prey less than 1 mm in length would be needed to support larger predators while modest increases in larger prey size classes could have profound effects on growth rate and maximum attainable size.

As rations increase with fish length, so do metabolic demands (Table 7). However, both the resource field

Table 7. *Valenciennellus tripunctulatus*. Daily energy budget as a function of adult size at maximum daily ration. Growth arbitrarily set at zero (i.e. no assumptions made as to how the surplus is used metabolically). Values expressed in calories day⁻¹

Fish size (mmSL)	Q _c	– Q _w	– Q _d	– (Q _s +Q _a)	= Surplus
27	8.72	– 1.92	– 1.31	– 2.36	= 3.13
30	12.08	– 2.66	– 1.81	– 3.59	= 4.02
35	15.38	– 3.38	– 2.31	– 7.27	= 2.42

and predator morphological-physiological characteristics interact to limit the extent to which increased energy can be obtained by simply taking larger prey. Under the present assumptions, there exists an optimum in the vicinity of 30 mm SL in which daily surplus over metabolic requirements reaches a maximum (Table 7). Theoretically, at a size of 37–38 mm no surplus would accrue, and no increase in size or other

metabolic activity would be possible; to exceed these limits, major changes in perception distance, search volume, or utilization patterns of large prey items must occur. Conversion to prey larger than 4 mm, however, is likely to involve severe metabolic penalties in terms of search and pursuit, since energetically rich prey (e.g. euphausiids, amphipods) are relatively uncommon and more mobile than copepod prey.

EFFICIENCY, OPTIMIZATION, AND COMPETITION

The trophodynamics of *Valenciennellus tripunctulatus* may relate to the nature and degree of 'optimization' in adaptive repertoires and to competitive interactions in depauperate mesopelagic environments. A number of characteristics indicative of high efficiency and diet optimization are evident. Feeding is restricted to that time of day when potential available energy intake per unit time is highest (Table 6). Feeding efficiencies are maximum over those prey size-classes in which the product of calories per individual copepod and total calories available is highest (i.e., 2–4 mm). Larger fish appear increasingly efficient at grazing larger prey size-classes. Smaller prey (less than 1 mm) are very inefficiently grazed by *V. tripunctulatus* and the energetic profit per item cannot be high.

Adaptations in both predator (Baird and Hopkins, 1981) and prey dictate compromises to the degree to which actual diets approach a theoretical optimum based on simple economic considerations. Specialization in *Valenciennellus tripunctulatus* apparently includes mechanisms for the efficient utilization of the copepod fraction of the available resource field. It should be noted that copepods constitute much smaller percentages of available prey at sizes greater than 4 mm. Not only are large prey items comparatively rare

but they also represent groups which are taxonomically and adaptively remote from copepods in terms of behavior, life history, and morphological characteristics. Intraspecific competitive interactions should in turn increase with fish size as more rare, larger prey items are required to meet daily energy requirements. The larger the fish, the greater the impact on the resource field by reason of increased pressure on calorically important large prey, a broader spectrum of size classes in the diet and higher absolute energy requirements.

A mechanism to reduce competition among *Valenciennellus tripunctulatus* size classes would be vertical segregation by size. This has been documented (e.g. Krueger and Bond, 1972; Clarke, 1974), and evidence published by Badcock (1977) suggests that it may be quite marked. Low density and noncontagious distributions among individuals, particularly as size increases, would also effectively reduce competitive grazing.

Concerning inter-specific competition, *Valenciennellus tripunctulatus* should be sensitive to both co-occurring 'nearfield' as well as 'remote' zooplanktivore predators. Most copepod prey populations occupy broad depth ranges (e.g. Roe, 1974) and many migrate vertically. They are impacted by a large number of mesopelagic fish and shrimp predators grazing over various depths and times. Higher trophic levels, in fact, appear co-exploitably organized wherein many predator species in the mesopelagic zone are competitively 'linked' to a common resource field and considerable diet overlap is observed (Foxton and Roe, 1974; Merrett and Roe, 1974; Donaldson, 1975; Walters, 1976; Hopkins and Baird, 1977; Clarke, 1978; Heffernan and Hopkins, in press). The premium on ecological efficiency, however, imposes a high degree of trophic organization in which mechanisms for partitioning resources can be expected, from theoretical considerations at least, to be well developed. Knowledge of the interaction of body size, energetics, and abundance of predator and prey exhibiting various patterns of distribution in space/time, then, is essential to the understanding of the organization of oceanic ecosystems. It should be emphasized that the present model has concentrated on an intermediate predator and its resource field, the zooplankton. Not considered here, because of inadequate information is the effect of higher level predation (Hayward and McGowan, 1979) on the population dynamics of this species.

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