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Growth Rate and Biological Production of Yellow Eel *Anguilla anguilla* and Production of Silver Eel and the Number of Glass Eel to Fulfill the Danish EMP

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Keywords: otolith; ageing; natural mortality; linear growth; growth rate; silver eel production; biological production of yellow eel; von Bertalanffy; EMP; stocking



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Article

Food, Growth and Biological Production of European eel *Anguilla anguilla* in a Small Stream in Jutland, Denmark

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Abstract: Yellow eels were sampled by electrofishing in 1979, 1980 and 1981 in Vester Vedsted Baeck, Denmark, stem 6 km, catchment 24 km², with outlet to the North Sea. Yellow eels were aged by burning the otoliths. The eels were not sexed, and they varied from 6.5 to 44 cm in length. The ages varied from 0+ to 10+ year. The annual growth rate Δ varied from 3.4 cm for 0+ eel to 2.2 cm for 10+ eel with a mean of 3.1 cm. Body mass wet wt. was recalculated to Kcal energy, with an annual growth rate Δ of 5.33 Kcal. In contrast to body mass, the annual growth rate Δ in Kcal increased with age. Von Bertalanffy growth trajectory (cm) of length-at-age was calculated and $L_{\infty} = 118.4$ cm. Annual natural mortality M was estimated, and M was significantly depending on body mass, i.e., high M and low body mass vs. low M at high body mass. The biological production was calculated to 13.5 g wet weight m⁻² year⁻¹. Total 780 eel stomachs were analyzed, of which 287 (37%) were empty. The weight of food content (2.1% of eel body mass) increased proportional with eel body mass. Chironomid larvae, Ephemeroptera nymph, Simulium larvae and *Gammarus pulex* were the dominating food taxa, followed by Trichoptera larvae. Size of prey were independent on length of eel.

Keywords: otolith; ageing; growth rate; von Bertalanffy; L_{∞} , natural mortality; biological production; feeding

1. Introduction

In Danish streams, the European eel and brown trout (*Salmo trutta*) populations constituted about 10% and 75%, respectively, of the total fish populations (Larsen 1955).

Since then, extensive studies were carried out on the biology of brown trout in fresh and salt water, and the brown trout is today very important in recreational fishing in fresh and salt water.

In the mid-1950s, fishing for eels in fresh and salt water was of great economic importance in the recreational and commercial fishery. In fresh water, yellow eels were caught in large numbers in streams and lakes in fykes and bottom nets, and a large number of fixed eel catching facilities throughout the rivers fished out-migrating silver eels. In salt water, silver eels were fished in coastal areas, many of them produced in Danish freshwater, and high number of silver eel migrated from the Baltic areas and passed through the Danish straits and the commercial fishery.

In contrast to brown trout, there are surprisingly few population studies about the Danish streams as production areas for eels, e.g., Larsen 1972, Rasmussen 1977, Rasmussen 1983, Rasmussen & Therkildsen 1979, Bisgaard & Pedersen 1991 and Rasmussen & Pedersen 2023.

The European eel population declined very much since the early 1960s, and the current recruitment of glass eel to North Europe is today a few percentage of the 1960–1979 reference level (ICES 2023). The causes of this decline are probably multiple and complex. These may include a combination of oceanic factors (Castonguay et al. 1994, Friedland et al. 2007. And continental factors such as reduction of grow-up habitats in rivers, fishing in fresh and saltwater, obstructions to up- and downstream migration for eel at all life stages, mortality in water turbines, pollution, diseases,

parasites (e.g., *Anguillicola crassus*) and bird predation (ICES 2005, Pedersen & Rasmussen 2013, Pedersen & Rasmussen 2018, Pedersen et al. 2024). The reduction in glass eel recruitment may have directly affected the density of local eel populations, and thus indirectly (through intraspecific and maybe interspecific competition) the growth dynamics and sex ratios of eels, but very little is known about this.

To aid recovery and conservation of the eel stock, the European Council adopted a framework regulation (EU 2007), requiring each EU Member State to issue, a country national eel management plan (EMP). The goal of this EMP is to ensure spawning escapement of at least 40% of the silver eel production, relative to the best estimate of escapement before start of anthropogenic influence. In Danish freshwater (rivers 15,000 ha and lakes 45,000 ha), the pristine annual silver eel production was estimated to 750 tons in rivers and 360 tons in lakes, totally 1,110 tons. Therefore, 40 % corresponds to 440 tons of silver eel escapement annually of pristine condition in freshwater (Pedersen & Rasmussen 2013). With the present implemented regulations in Denmark (reduced fishing effort, closed fishing periods, increased minimum size of yellow eel), the target of the 40% silver eel production without stocking and fishing in rivers, was estimated to be achieved in year 2080. To reach the goal earlier, eel stocking may be a useful method. To know how much to stock we need to establish the relationship between number of glass eel or on-grown eels and the number of produced silver eel (Rasmussen & Pedersen 2023, Pedersen et al. 2024).

Bisgaard & Pedersen (1991) compared mortality and growth of wild and on-grown stocked eels in a small river with reference to tagging techniques. Aged wild eel showed close agreement with annual length increment in tagged and recaptured wild-tagged eel.

Rasmussen & Pedersen (2023) used information from the 1970s about silver eel and yellow eel populations in two rivers in Danish streams, and calculated that at that time, before the decline of eel populations, the annual production of silver eel was about 50 kg ha⁻¹. In one study, the calculations (growth and survival) were based on sampled and otolith aged silver eels. In the other study the calculations (yellow eel density, growth and survival) were based on electrofished caught yellow eel, that were otolith aged and used to estimate silver eel production. In both studies, the data enabled to estimate growth rates and von Bertalanffy length growth trajectories. In the last study, feeding habits and feeding rates were also analyzed. These data and results possibly represent the pristine status of eel population in Danish freshwater before the decline in glass eel recruitment.

Measurement of biological production in fish species is an important aspect of population dynamics (e.g., Ivlev 1966, Chapman 1967). Production is the total elaboration of fish tissue (g wet weight or energy) during any time interval Δt , and the production and rates integrate changes in number and body mass. Part of the standing biomass might be able to migrate away from the production site, in this case as silver eel. Purely logically and mathematically, the migrating biomass of silver eel will always be less the total biological production. Calculation of silver eel production in the present study was not possible, because most of the yellow eels were less than about 35 cm, i.e., less than silver ageing of eel, maybe except for a small unknown number of youngest male silver eel, that might have left the population (see 4.1).

The objective of the present paper is to 1) examining growth and annual length increment (cm) and Kcal of yellow eel; 2) from yellow eel density and eel body mass calculate annual mortality M of yellow eel, and 3) calculate the biological production (g wet weight) of yellow eel. Finally, 4) feeding habits of yellow eel was examined.

2. Material and Methods

2.1. Location

Vester Vedsted Bæck (location 55.07768, 8.67902) has a catchment area of 24 km², main stem of 6 km estimated downstream areas of wetlands, mean annual water flow estimated to 293 L sec⁻¹, and the bæck goes out into the Wadden Sea, North Sea. Water temperature was measured (20 observations) with a min-max thermometer in all three years, and temperature was calculated to water temperature $^{\circ}\text{C} = 2.52(\pm 2.89) + 12.12(\pm 2.72) \cdot \exp(-0.09(\pm 0.04) \cdot (t - 6.5(\pm 0.2)))$, $R^2 = 0.78$, where t is day. Mean water temperature from January to December was calculated to 9.94 $^{\circ}\text{C}$ with a max

temperature in July to about 14.7 °C, with some observed values up to 18 °C. The baeck is heavily regulated, but there are no obstacles in the stem beside a sluice at the outlet to the sea. The baeck widths vary from 2.5 to 3.6 m, the bottom is dominated by sand, and four species of waterweeds can be seen in the baeck. Fascines supports the riverbank, which provide good hiding opportunities for eel. Besides eel, sticklebacks (*Gasterosteus aculeatus* and *Pungitius pungitius*) and plaice (*Platichthys flesus*) are common.

2.2. Sampling, Ageing, Growth and Food Analysis

In 1979, 1980 and 1981 electrofishing (Rasmussen 1983) were performed every sixth week from March to November. Four stations each of 50 m were electrofished (Bohlin et al. 1989), and the total electrofished area was 546 m². Glass eel from the North Sea into the baeck was fully recruited mid-June.

All eels caught were measured (0.5 cm), and five eel of each length group were weighed (0.01 g) at the laboratory, and 553 otoliths were aged (0+ to 10+ year) by burning the otoliths (Christensen 1964, Moriarty & Steinmetz 1979).

The mean length 7.19 (cm) and body mass 0.29 (g) of glass eel was calculated from a sample of 194 glass eel in another river (Pedersen et al. 2023).

Annual growth rate (Δ cm) from glass eel to yellow eel age stage of individual yellow eel, was calculated, assuming linear growth of yellow eel, as: $\Delta = (\text{length}_{\text{yellow eel}} - 7.19 \text{ cm}) / \text{age}_{\text{yellow eel}}$ (Lester et al. 2004, Rasmussen & Pedersen 2023). This was also done for body mass calculated to Kcal assuming that one glass eel has an energy content of 0.3496 Kcal.

Body mass (g wet weight) was calculated as $w = 0.00077 * \text{length}^{3.23}$, and body mass (Kcal) was calculated as $w_{\text{kcal}} = 0.00021 * \text{length}^{3.76}$ (Rasmussen 1983). Annual instantaneous growth rate G was calculated as $\ln(w_1/w_0)$, where w_1 is the body mass one year after w_0 . Mean body mass (MBM) was calculated as $\text{MBM} = w * (\exp(G) - 1) / G$.

Natural mortality M was calculated as $M = \ln(n_1/n_0)$, and mean number (MN) was calculated as $\text{MN} = n_1 * (1 - \exp(M)) / -M$, where n_1 is the number of eel one year after n_0 .

Natural mortality M as function of body mass BM was calculated as $M = a * BM^b$.

Biological production (g wet weight) was calculated as $\text{Production} = \text{MN} * \text{MBM} * G$.

Von Bertalanffy growth trajectory (cm) was calculated from Δ (cm) vs. age.

In this paper data for density, length, body mass and age class from all stations and years were pooled to month October, i.e., mean of data from three years. It has smaller significance for the calculation of growth, mortality and biological production, because these responses were calculated from age class to age class. Seasonal variations in density and some other results can be found in Rasmussen 1983.

Stomach ($N = 780$) sampled from all three years were distributed to months, the first in March and the last in October. Stomach content was weighed (0.01 g) and determined to main taxa. Number of Occurrence (NoC) was calculated as number of eel with the specific food topic divided with the number of eel with food in the stomachs. Some of the prey taxa were length (0.1 cm) measured.

2.3 Data Treatment

The results were calculated and tested using Excel ver. 5.0 and Real Statistics

Resource Pack version 6.7 and <http://statpages.org/nonlin.html>. Level of significance for statistical tests was 0.05.

3. Results

3.1. Number, Growth and Biological Production

Figure 1 shows length distribution of yellow eel in October as a mean for all three years. The population was dominated by eel sizes up to about 20 cm, and rather few eel above.

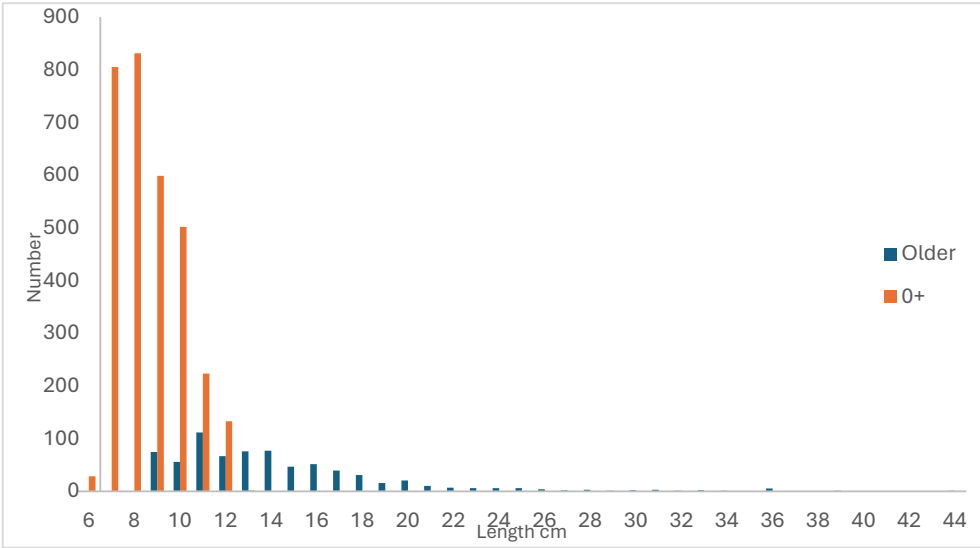


Figure 1. Number of yellow eel versus length cm.

The total population number of yellow eel in October (Table 1), ages 0+ to 10+, and excluding the permanent annual input of glass eel that was reduced to 0+ four months later, amounts to about 550 yellow eel pr. 100 m², or about 5-6 yellow eel pr. 1 m².

The glass eel started after recruitment in June with a size of about 7.19 cm and grow the following four months up to a size of 9.13 cm as 0+ in October. Over the next 10 year from age 0+ to age 10+ they grow up to a mean size of about 35.2 cm (Table 1), few single eels were found up to 48.5 cm.

The annual natural mortality M (Table 1) started from 3.63 (glass eel to 0+ in four months recalculated to annual mortality) and final mortality 0.13 from age 9+ to 10+.

The total annual biological production (Table 1) was calculated to about 1,347 g 100 m⁻² from glass eel to age 10+.

Table 1. Number (0+ to 10+), length cm and annual growth rate Δ cm of yellow eel from glass eel to age 10+ (column 2 to 4). Body mass g, annual instantaneous growth rate G and mean body mass g (column 5 to 7). Natural mortality M, mean number and mean biomass g 100² (column 8 to 10). Annual biological production g (column 11).

	No. 100	Length cm	Δ cm	Body mass	G	Mass	M	No.	Biomass g	Production g
Glass eel	563.13	7.19		0.29						
					1.21	0.56	3.63	488.30	275.67	333.95
0+	420.41	9.13	1.94	0.97	1.49	2.25	1.49	192.17	432.40	645.38
1+	66.74	14.49	5.36	4.33	0.49	5.59	0.49	44.63	249.51	122.27
2+	28.04	16.86	2.37	7.07	0.44	8.89	0.44	21.82	194.05	85.81
3+	16.59	19.34	2.47	11.00	0.25	12.53	0.25	11.04	138.34	35.21
4+	6.89	20.92	1.59	14.19	0.33	16.78	0.33	5.00	83.81	27.27
5+	3.49	23.14	2.22	19.65	0.38	23.88	0.38	3.18	76.03	28.74
6+	2.90	26.01	2.87	28.68						

					0.42	35.61	0.42	2.58	91.82	38.39
7+	2.28	29.61	3.59	43.57	0.14	46.87	0.14	1.93	90.34	13.04
8+	1.61	30.96	1.35	50.33	0.28	58.14	0.28	0.90	52.26	14.73
9+	0.44	33.79	2.82	66.73	0.13	71.24	0.13	0.28	19.73	2.55
10+	0.16	35.17	1.38	75.94						
Total	549.55								1703.95	1347.34

3.2. Length vs. Age

Figure 2 shows the relationship between length and age from 450 observations of length-at-age. The mean annual growth rate Δ was about 2.4 cm.

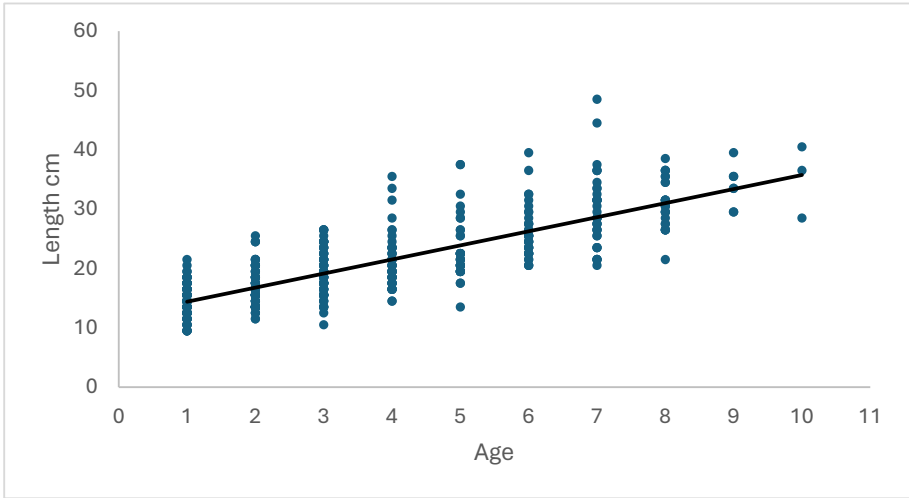


Figure 2. Relationship between (N= 450) lengths (cm) of yellow eel at age 1+ to 10+. The length-at-age was fitted to $L_t = 12.02(\pm 0.36) + 2.374(\pm 0.082) \cdot \text{Age}$, $R^2 = 0.65$.

The relationship between Δ cm and age from 450 observations is shown in Figure 3. The mean annual growth rate Δ was about 3.1 cm.

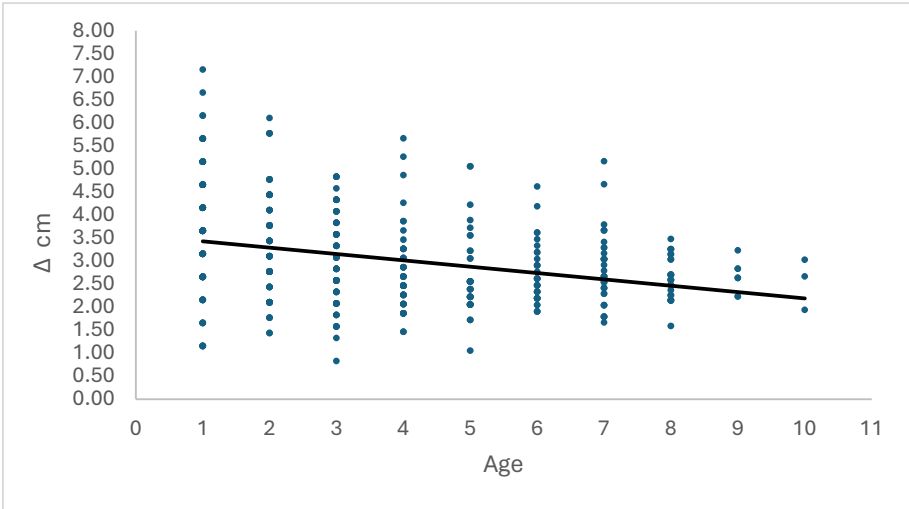


Figure 3. Annual growth Δ cm of yellow eel (N = 450) at age 1+ to 10+ from ages of otoliths. The Δ was fitted to $\Delta = 3.565(\pm 0.17) - 0.1378(\pm 0.0397) * \text{Age}$, $R^2 = 0.09$. Mean Δ cm was calculated to 3.1 cm.

The relationship between Δ Kcal and age from 450 observations is shown in Figure 4. The mean annual growth Δ was about 5.33 Kcal. Twenty observations were statistical outliers with values of Δ from 17.4 to 57.2 and lengths from 31.5 to 48.5 cm.

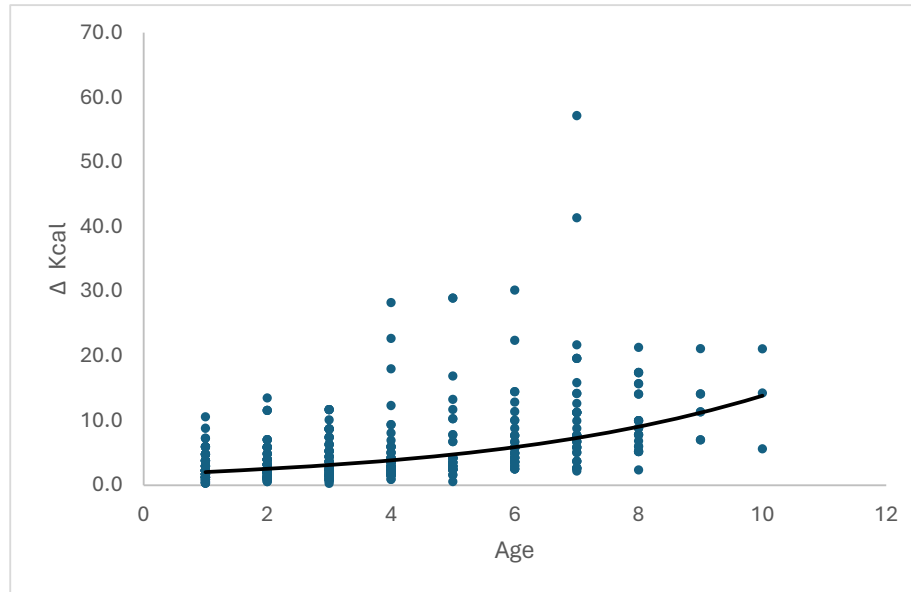


Figure 4. Annual growth Δ Kcal of yellow eel (N = 450) at age 1+ to 10+. The Δ Kcal was fitted to $\Delta = 2.37(\pm 0.25) * \exp(0.1918(\pm 0.0162) * \text{Age})$, $R^2 = 0.24$. Mean Δ Kcal was calculated to 5.33 Kcal. The observations had a skewness of 3.5671, and 20 observations are outliers with a higher relative energy content compared to the other observations.

The Bertalanffy growth trajectory length (cm) of yellow eel is shown in Figure 5 using the Δ values in Figure 3.

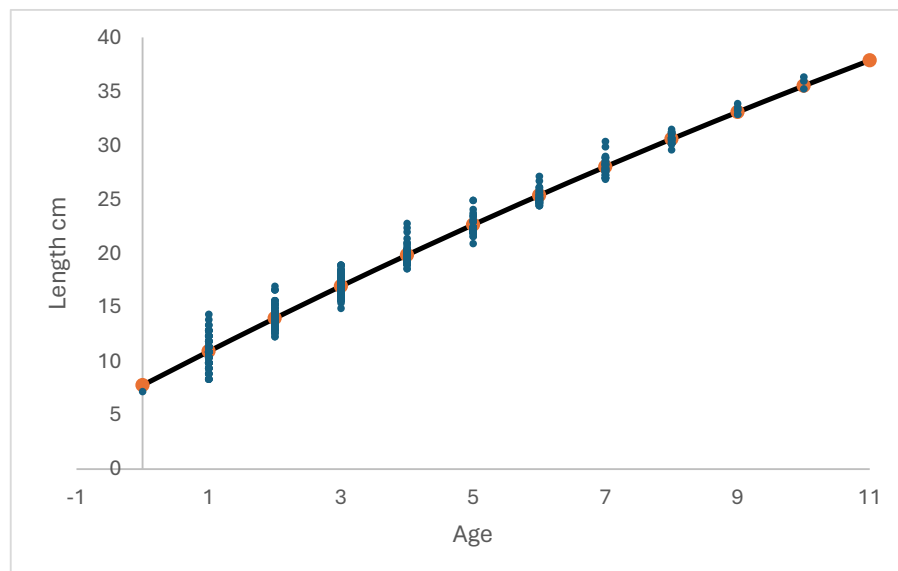


Figure 5. von Bertalanffy growth trajectory (cm) of yellow eel (N = 450) using the Δ values in Figure 2. $L_t = 118.4(\pm 21.4) * [1 - \exp(-0.0280(\pm 0.0289) * (t_t + 2.35(\pm 0.12)))]$, $R^2 = 0.98$.

3.3. Natural Mortality M vs. Body Mass

Figure 6 shows the natural mortality as a function of body mass. M vs body mass BM was calculated from 11 observations of M given in Table 1. 0.5*Ursin was calculated from $M = 0.5 \cdot BM^{-1/3}$. M calculated from the observations of M and BM fits very well with 0.5*Ursin.

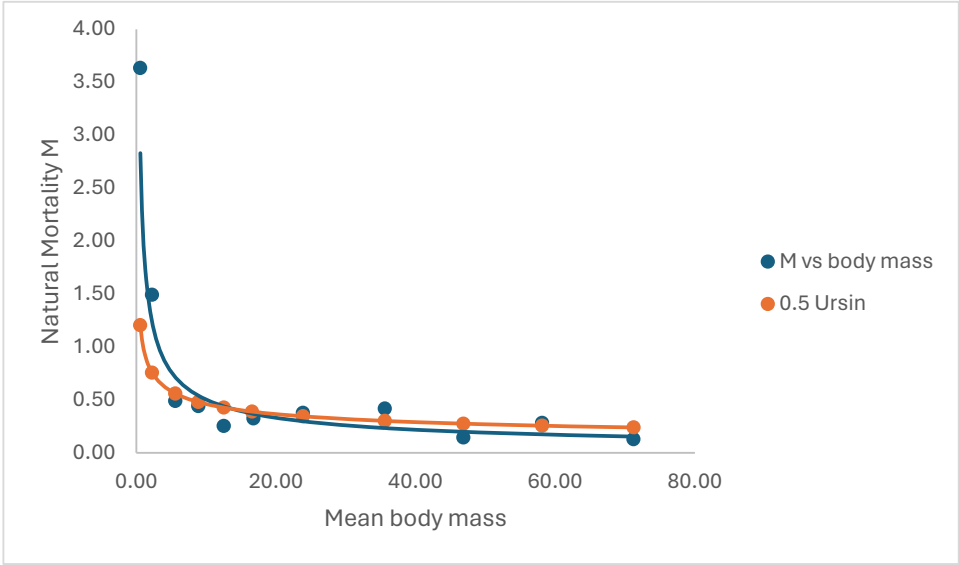


Figure 6. Annual mortality M vs. Body Mass. $M = 2.002(\pm 1.731) \cdot BM^{-0.6006(\pm 0.1864)}$, $R^2 = 0.86$. $M_{0.5 \cdot Ursin} = 0.5 \cdot BM^{-1/3}$.

3.4. Feeding Results

An overview of the food items is shown in Table 2. The stomach contents can be regarded as composed of five most important food groups. Isopoda (*Asellus aquaticus*), Amphipoda (*Gammarus pulex*), Ephemeroptera nymph, Trichoptera larvae, Chironomidae larvae and Simuliidae larvae.

Table 2. Number of taxa, No. of Occurrence and number of taxa pr. Eel with food. Taxa with smaller items are marked with x.

		25	21	23	22	16	6	3	3	April –
		March	April	May	June	July	Aug	Sept	Oct	October
	Number of eel	57	63	103	107	112	62	140	136	723
	Empty %	93	24	37	25	43	35	24	37	26
Asellus	% No.		4.2	10.8	7.5	-	-	4.7	7.0	5.3
	Occurence									
	No. Pr fish		2.0	1.6	2.0			1.4	2.5	1.9
Gammarus	% No.		4.2	52.3	38.8	45.3	20.0	43.4	39.5	39.5
	Occurence									
	No. Pr fish		2.0	4.0	4.0	4.0	3.8	3.7	3.8	3.8
Ephemeroptera	% No.	x	14.6	20.0	45.0	42.2	45.0	83.0	58.1	48.9
	Occurence									
	No. Pr fish		2.3	1.4	7.2	5.5	4.2	5.1	3.1	4.7
Tricoptera	% No.		2.1	15.4	6.3	17.2	10.0	0.9	3.5	7.2
	Occurence									
	No. Pr fish		1.0	3.3	1.8	1.9	1.3	1.0	1.0	2.1
Diptera	% No.		14.6	44.6	65.0	28.1	35.0	18.9	18.6	31.9
	Occurence									
	No. Pr fish		2.4	7.2	3.2	1.9	1.7	1.6	5.4	3.6

Diptera Simuliidae	% No. Occurrence	X	91.7	-	1.3	7.8	30.0	14.2	-	15.7
	No. Pr fish		112.0		5.0	1.4	8.9	6.3		66.8
Other Diptera		X		x	x	x			x	
Lumbricidae			x	x	x	x	x	x		
Dytiscidae larvae & adult				x	x					
Corixa					x					
G. aculeatus egg					x	x				
G. aculeatus								x		
Hirudinea					x	x				
Gastropoda					x					
Hemiptera								x		
Lepidoptera l									x	

In addition to the five important taxa are Lumbricidae, Dytiscidae larvae and adults, Hemiptera (*Corixa*), Hirudinea and Gastropoda, and eggs (June and July) from *Gasterosteus aculeatus*, in one single eel 29 cm in August, two *Gasterosteus aculeatus* was observed, but all the last taxa are in smaller amounts.

Chironomidae larvae, *Gammarus pulex* and Ephemeroptera nymphs are the dominating taxa. Simuliidae larvae was found from March to September, and especially in April they were important and in high number pr. Eel.

The mean number of empty stomachs is 26% (April – October) and the relationship between empty and stomachs with food is given in Figure 7. The distribution of empty stomachs vs length is different compared to stomachs vs length with food (Kruskal-Wallis test, $p < 0.05$).

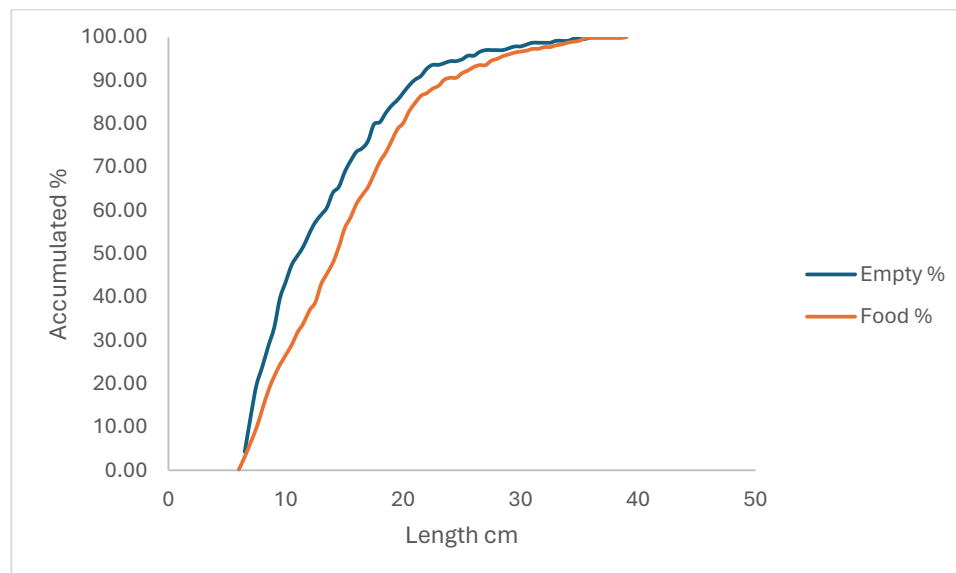


Figure 7. Relationship between number of empty stomachs and stomachs with food.

The relationship (April to October) between stomach content (g) and body mass (g) of eel is given in Figure 8, and stomach content = $0.0214(\pm 0.0031) \cdot \text{body mass}^{0.9397(\pm 0.0876)}$, $R^2 = 0.62$, which mean that the mean stomach content is about 2.1% of body mass, and that stomach content is more or less proportional to body mass.

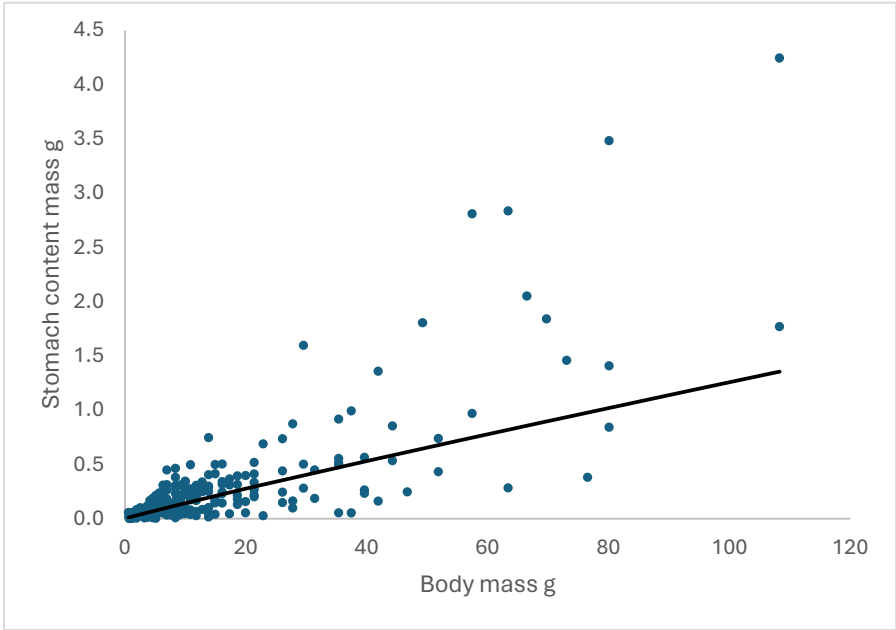


Figure 8. Relationship between stomach content (g) vs body mass (g).

Table 3 shows NoC predation on Simuliidae and Chironomidae larvae, Ephemeroptera nymphs, Trichoptera l and Gammarus vs four eel length groups. Ephemeroptera and Chironomidae larvae are preferred of smaller eel and at decreasing amount at decreasing length, and Trichoptera larvae are preferred at increasing eel length.

Table 3. NoC and number of prey pr. Eel for the five most important food taxa.

	Simuliidae l				Chironomidae l			
	NoC	% NoC	%	No. Prey pr eel	NoC	% NoC	%	No. Prey pr eel
Eel length cm								
6.5 – 13.1	19	3.8	23.8	3.5	99	20.0	61.9	3.5
13.1 – 19.7	31	6.3	38.8	15.7	39	7.9	24.4	4.6
19.7 – 26.3	14	2.8	17.5	59.5	16	3.2	10.0	3.4
26.3 – 39.5	16	3.2	20.0	241.2	6	1.2	3.8	1.5
	80	16.2			160	32.3		
	Ephemeroptera n				Gammarus			
	NoC	% NoC	%	No. Prey pr eel	NoC	% NoC	%	No. Prey pr eel
Eel length cm								
6.5 – 13.1	103	20.8	43.1	3.1	48	9.7	24.1	1.6
13.1 – 19.7	96	19.4	40.2	5.7	99	20.0	49.7	3.4
19.7 – 26.3	30	6.1	12.6	6.8	37	7.5	18.6	5.5
26.3 – 39.5	10	2.0	4.2	4.8	15	3.0	7.5	9.5
	239	48.3			199	40.2		
	Trichoptera l							
	NoC	% NoC	%	No. Prey pr eel				
Eel length cm								
6.5 – 13.1	4	0.81	10.5	1.3				
13.1 – 19.7	11	2.22	28.9	2.3				

19.7 – 26.3	13	2.63	34.2	1.5
26.3 – 39.5	10	2.02	26.3	2.7
	38	7.7		

The size (cm) of Simuliidae and Chironomidae larvae and Ephemeroptera nymph and Gammarus vs. eel length is shown in Figure 9. The mean size of prey is independent of eel length.

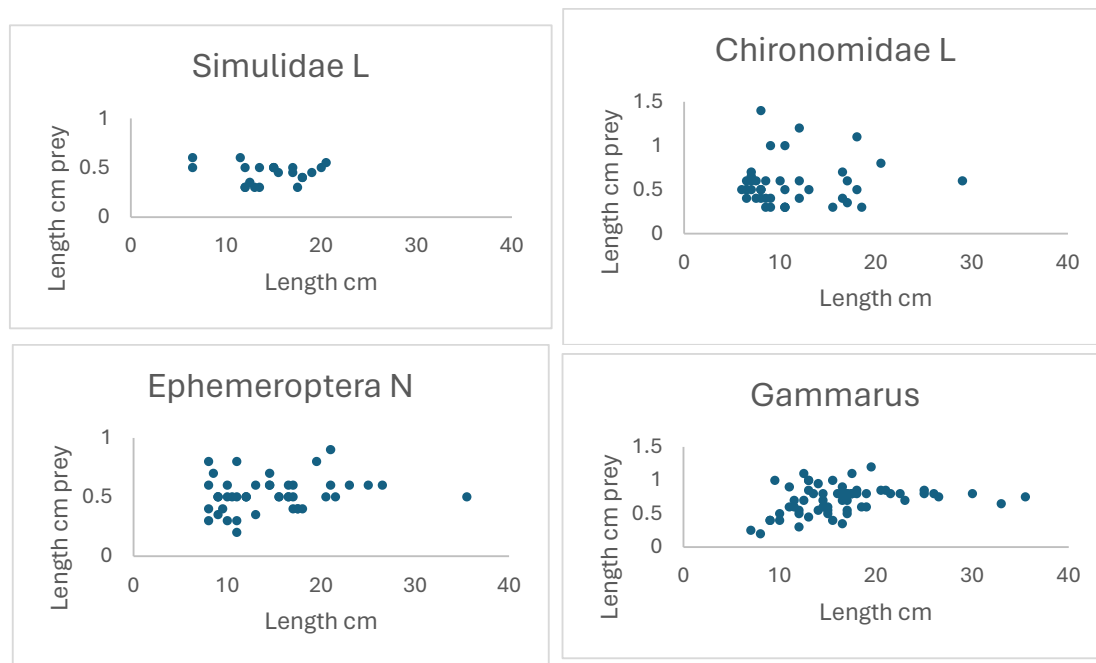


Figure 9. Size (cm) of four prey taxa vs length (cm) of eel.

4. Discussion

4.1. Linear Growth and Von Bertalanffy Trajectory of Yellow Eel

Bioenergetics model developed for fishes indicates that somatic growth rate scales allometrically with fish body mass: $dw/dt = a \cdot w^m - b \cdot w^n$, where w is somatic body mass, $a \cdot w^m$ is the rate of energy acquisition, and $b \cdot w^n$ reflects energy losses owing to respiration (e.g., standard metabolism, SDA and activity) (Ursin 1967, Lester 2004).

Setting $m = 2/3$ and $n = 1$, integrate dw/dt , we get the von Bertalanffy growth trajectory: $w_t = w_\infty [(1 - \exp(-k \cdot (t - t_0)))]^3$.

References in the literature e.g., Lester et al. (2004 and references herein) suggests, that m and n have similar values, that both lie close to $2/3$, suggesting the following equation holds: $dw/dt = (a - b) w^{2/3}$.

Given that body mass increases with the cube of length ($w = al^3$), and substituting w with l , the potential growth rate for length (cm yr⁻¹) is $dl/dt = h_0$, where $h_0 = (a - b)/3a^{1/3}$. Integrating dl/dt , thus length is a simple linear function of time, $l_t = k + h_0 \cdot t - l$, where the constant k is set to 7.19 cm (i.e., length of glass eel). $(a - b)$ is the net growth, and with increasing individual food intake the growth rate will increase, but looked over increasing ages the annual growth rate of individuals is constant.

Linear growth of immature fish is debatable (Lester et al. 2004). For example, Rasmussen 1952 sampled and aged yellow eel caught in Lake Esrum, Denmark. The otoliths were age determined due to thin slip, and the individual growth rate showed constant annual linear growth. Dahl 1967 stocked glass eel in ponds in Denmark and aged the otoliths due to thin slip, and individual eel showed constant annual linear growth. Sinha & Jones (1967a) aged yellow eel (2 – 12 years) in different rivers in UK by grinding the otoliths, and individuals all showed linear growth. Frier & Rasmussen (2020)

aged and calculated annual growth rate of sea trout (*Salmo trutta*) in marine water. The growth rate from smolt up to first spawning in autumn was linear and rather fast. The sea trout loose up to about 40% of body mass when spawning, and they have to build body mass up before they can restart growing in length (Frier & Rasmussen 2020). Hereafter they grow in linear length and body mass up to next spawning the following autumn. The growth period is thus shorter than the year before, so following a cohort of sea trout through several spawning, the growth curve follow a van Bertalanffy growth trajectory. However, the individual growth in length through each year is always linear, and can vary from individual to individual.

The linear growth of fish species assumes, that (a - b) does not change, and that all surplus energy is allocated to somatic growth, and does not account for the change in somatic growth, that occurs when fishes become sexually mature and some energy is allocated to reproduction (Day & Taylor 1997). Yellow eels are immature during the entire growth period up to the silver eel stage, and individual yellow eel will therefore always grow with a linear growth trajectory. This does not mean that all individual yellow eel within an age class grow with the same Δ cm, see Figure 2, and Figure 2 in Rasmussen & Pedersen (2023), where the variance of the age individual observations decrease at increasing age. Probably it is a combination of genetics and varying feeding rate $a \cdot w^m$ of each single eel.

We propose that rather early at age, some yellow eel in the cohort distribute to fast growth rate and probably early silvering, while the lower growing yellow eel silver later at increasing higher age. The yellow eel that mature later make it possible to calculate a von Bertalanffy trajectory.

Rasmussen & Pedersen (2023) assumed linear growth of yellow eel in River Brede. Male (4 – 25 year) and female (7 – 24 year) silver eel with known length and age at silvering enabled to calculate length-at-age through a cohort of yellow eel from low age at silvering to high age at silvering. For both sex length-at-age decreased considerable going from low age at silvering to high age at silvering, and male yellow eel grow significant slower than female silver eel. For both sex and for a given yellow eel age, there was rather high variation in calculated yellow eel size. This means that through a cohort starting from glass eel to silvering, some yellow eel grow rather fast, and reach size of silvering in a few number of year, the decreasing growth rate at increasing age postpones the age at silvering. This means that even if we assume individual linear growth, for a given age there will be fast growing yellow eel vs slow growing yellow eel. It also means that the given potential growth rate vary for length-at-age (cm year⁻¹) or $h_0 = (a - b)/3a^{1/3}$. One can imagine that the fastest growing yellow eels simply eat relatively more than slow growing yellow eels, i.e., values of 'a' are greater for the fast-growing eels compared to the slow growing eels.

Figure 2 shows the traditional mean length-at-age relationship, where annual Δ is the slope of the regression and calculated to 2.4 cm. This is a rather slow growth rate. Figure 3 shows the annual Δ cm calculated from otoliths and is calculated to mean 3.1 cm. Δ is decreasing as function of age.

The difference between the two values of Δ is probably explained by the two different calculation data.

More interesting is the declining value of Δ in Figure 3 compared to increasing values of Δ in Figure 4. Most probably, it is explained by the relative increasing energy content in yellow eel at increasing age. This means that at increasing length the length/energy content is higher for a given length compared to length/weight relation in wet weight at the same length. The calculated Δ in Figure 3 was calculated in wet weight, so the corresponding increasing length-at-age in energy means decreasing Δ . In Figure 4 is calculated Δ Kcal as function of age and Δ is increasing. The observations are skewed, and 20 values of Δ Kcal are outliers, with lengths cm varying from 31.5 to 48.5 cm. They were all yellow eel, and we suggest that they have so high relative energy content, that they prepare for early silvering and out migrating.

Using the calculated Δ cm in Figure 3, which are decreasing and therefore follow the prerequisites (i.e., decreasing Δ at increasing age) for the von Bertalanffy growth trajectory which is shown in Figure 5. The slow decreasing Δ at increasing age means that L_∞ , here calculated to 118.4 (± 21.4) cm is rather high, but not unlikely, because eel can grow up to 133 cm (Fishbase.org). The 95% confidence limit of L_∞ is rather high and in general, the von Bertalanffy equation should be used with

precaution (Day & Taylor 1997, Lester 2004), especially with data that does not distinguish between male and female. Rasmussen & Therkildsen (1979) calculated in a Danish river L_{∞} for unsexed eel to 59.8 (± 5.01) cm. Rasmussen & Pedersen (2023) calculated in a Danish river L_{∞} for males to 49.7 (± 0.56) cm and females to 74.4 (± 1.52) cm. Bisgaard & Pedersen (1991) calculated in a Danish river L_{∞} to 90.3 cm in a population of unsexed eel.

4.2. Mortality M vs. Body Mass

Annual natural mortality M of fish species, including eel, is probably a function of e.g., sex, body mass, water temperature, predators and stock density (e.g., Pedersen et al. 2023, Bevacqua et al. 2012, Ursin 1967, Vøllestad & Jonsson 1988, Berg & Jørgensen 1994, De Leo & Gatto 1996 and Lorenzen 1996). This means that M is decreasing with increasing body mass and male versus female eel, and increasing with water temperature, and M is increasing with increasing density of yellow eel. Ursin (1967) proposed as a “rule of thumb” that $M = M^{-1/3}$, but he overlooked that M should be scaled including other predictors other than only body mass, so M is better described as $a \cdot M^{-1/3}$ (Pedersen et al. 2023, Bevacqua et al. 2011). Pedersen & Rasmussen (2013) propose that $M = 0.5 \cdot M^{-1/3}$. The two calculations in Figure 6 show reasonable agreement between M from observations compared to M calculated by $0.5 \cdot M^{-1/3}$, and this model of M has also been used in Pedersen & Rasmussen (2013) and Rasmussen & Pedersen (2023). Down to about body size of 10 g, there is a slight increase of M , but below body size of 10g the mortality increase rather fast.

4.3. Biological Production

The biological productions of yellow eel in this study was calculated to 13.5 g m⁻² year⁻¹. Rasmussen & Pedersen 2023 calculated biological production in two Danish rivers to 11.0 and 6.6 g m⁻² year⁻¹, respectively. Lobon-Cervia et al. 1995 calculated biological production along the course of River Esva in North Spain, mostly dominated by males, aged up to 4+ (i.e., five growth seasons) and up to 40 cm length. Numbers of yellow eel were estimated from electrofishing, and the eels were aged by clearing the otoliths. The biological production depended on distance to the estuary, and values up to 35.3 g m⁻² year⁻¹, but with large variations, was recorded. Biological production in running waters depends on number of recruitment of glass eel, mortality and growth rate, so production can be expected to vary quite much. However, there are so few yellow eel studies to compare. Lobon-Cervia and Rasmussen (2024) showed that the biological production of brown trout in rivers in England, Denmark and Spain had a limit up to 40-45 g m⁻² year⁻¹ primarily as a function of recruitment of fry in spring. Yellow eel and brown trout more or less eat the same taxa in rivers, so the high production of yellow eel is natural.

4.4. Feeding

The food taxa in the baeck in this study was dominated by five important taxa, other taxa were in small number. This is in contrast with River Køge-Lelling (Rasmussen & Therkildsen 1979), where a much higher number of different taxa constituted the diet of eel. In the baeck in this study, only one single stickleback was found, whereas in the other river several six fish species was found in the food. In this study *Gammarus pulex* was an important taxa, but only sporadic in the other study. We propose this is because of the small size and physical bad habitat condition of the baeck in this study, whereas in the other study the river was larger and with better habitat condition.

The number of empty yellow eel stomachs in both studies was surprisingly high, and one wonder why in contrast to stream brown trout (*Salmo trutta*), where more or less all fish always have food items in their stomachs. Feeding of eel depends on water temperature (Dörner & Berg 2015), and they eat from water temperatures 5-6 to 25-30 °C. The water temperature in 25 March in this study was estimated about 6.1 °C, so the eels had just started eating and this might explains the high percentage of empty stomachs. The water temperature on October 3rd was estimated to about 9.7 °C and this might explains the lower number of empty stomachs. In Figure 7 is shown, that the proportion of empty stomachs is more pronounced in smaller eel. It could be because of agonistic

behavior between eels because of the high density of eel up to about 20 cm length and because of the rather high eel density about 5 – 6 eel pr. one m².

Sinha & Jones (1967b) looked at the stomach content of a high number of eel in English rivers. The food consisted mainly of Trichoptera and Diptera larvae, Plecoptera, and Ephemeroptera nymphs, through at times, Gastropoda, Annelida, Crustacean and fish were common in the stomachs, when eaten the last the fish mostly taken were elvers and bigger eels. Frost (1946) analyzed the food of eel in the Windermere area, including two running water, total 74 eel. The food items were Limnephilids larvae, Plecoptera and Ephemeroptera nymphs, Chironomids and other Diptera larvae, *Gammarus pulex* and Mollusca.

In all the mentioned few studies available, there was no identification of river food items, so we don't know what the eel prefer from the food items, but the conclusion is probably that the yellow eel eat what is available in the river, but eat fish in low number (Dörner & Berg 2015).

The size of the food items in Figure 9 did not vary as function of eel length. Collecting eels for food analysis only gives a snapshot of the food on the day in question. One could explain the results in this study by the fact, that there is a constant grazing of new, recruited small individuals of food animals, and that these smaller items are quickly eaten, especially by the smallest eels under, for example, 10 cm in length, which then have to eat the slightly larger food items. The size of *Gammarus pulex* shows an increase in eaten Gammarus all the way down to 2 mm in length, then up to a constant length of Gammarus at a size of eel of approx. 12 cm length. *Gammarus pulex* constantly produces new individuals depending on the water temperature. This is in contrast to the aquatic insects, which have a more constant number of generations. This might explain Figure 9. However, this requires extensive research.

Data Availability: The data used can be obtained from the corresponding author.

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Conflicts of Interest: The authors certify that they have no affiliations with or involvement in any organization or entity with any financial interest (such as honoraria; educational grants; participation in speakers' bureaus; membership, employment, consultancies, stock ownership, or other equity interest; and expert testimony or patent-licensing arrangements), or non-financial interest (such as personal or professional relationships, affiliations, knowledge, or beliefs) in the subject matter or materials discussed in this manuscript.

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