**Daphnia** body size and population dynamics under predation by invertebrate and fish predators in Lago Maggiore: an approach based on contribution analysis

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**ABSTRACT**

Predation on Daphnia is size-specific: while zooplanktivorous fish select large, ovigerous females, carrying larger clutches, predation by invertebrates, particularly Cladocera, is generally regarded as acting mainly on young, small, non-ovigerous females. The two types of predators therefore produce different effects on the population of their prey: fish predation results in decreases in Daphnia fecundity, proportion of adults, and maximum body size, while predation by the invertebrate predators leads to decreases in the number and proportion of juvenile Daphnia. We investigated the effect of predation by three predators (one vertebrate, two invertebrates) on Daphnia. The study combines contribution analysis of Daphnia birth rate dynamics, with a body-size-oriented analysis of both predator and prey populations. Contribution analysis showed that during April-May, when Bythotrephes densities were low, changes in Daphnia birth rate were due to both changes in fecundity (ConF) and in the proportion of adults (ConA), with ConA being much larger (by absolute value) than ConF: such a pattern is expected under fish predation. Whereas during mid May to June, when Bythotrephes densities were high, changes in Daphnia birth rate were again mainly due to both ConF and ConA, but now the difference between these contributions was less pronounced. Although Bythotrephes predation produced similar effects on Daphnia population dynamics as fish predation, affecting both fecundity and the proportion of adults, it can be distinguished in two ways from fish predation. Firstly, the ConF-ConA pattern is more uniform under invertebrate predation than under fish predation and secondly, the mean body size of Daphnia increased continuously whereas the mean size of ovigerous females remained approximately constant except for the last 3 weeks of June, indicating a high mortality of juveniles during the whole period and an increased mortality of young females with eggs during the last 3 weeks. We conclude that both zooplanktivorous fish and invertebrate predators affect population dynamics of Daphnia in Lago Maggiore. However, the invertebrate predator Bythotrephes has the largest impact on Daphnia dynamics in June, both because of its high densities and its relatively large size at that time. Predation pressure exerted by Bythotrephes is high enough to reduce the Daphnia densities at the end of June to low values.

Key words: Bythotrephes, Leptodora, Coregonus, size-selective predation

**1. INTRODUCTION**

Predation on *Daphnia* is size-specific, while zooplanktivorous fish positively select large, ovigerous females carrying larger clutches, predation by invertebrate predators generally acts on small, young, non-ovigerous females (Zaret 1980). In the pelagic zone of Lago Maggiore, a deep subalpine lake in Northern Italy and the second largest lake in Italy (surface area = 212.2 km², $Z_{\text{mm}} = 370$ m) two cladoceran invertebrate predators (i.e. *Bythotrephes longimanus* and *Leptodora kindtii*) and zooplanktivorous fish (coregonid spp.) play a dominant role as feeders on zooplankton (de Bernardi et al. 1987; Manca et al. 2000; Volta 2000).

*Bythotrephes longimanus* (Cladocera: Onychopoda) is a native invertebrate predator in the open water of Lago Maggiore. Before (pre-1960) and during (1960-1978) cultural eutrophication *B. longimanus* densities were low (25 ind m⁻³) and the species was only present in late summer and early autumn (Manca et al. 1992; Manca & Ruggiu 1998). Following re-oligotrophication a 10-fold increase in densities from the 1980s to the early 2000s was accompanied by a 3-month earlier onset of population growth and a nearly 6-month increase in the duration of its occurrence in the water column (Manca et al. 2007).

*Bythotrephes* is a cruising predator (Gerritsen & Strickler 1977) that swims continuously throughout the water column and strikes at zooplankton prey as they pass close by. It detects prey by mechanoreceptors located on the first antennules, mandible, labrum, maxillary process and thoracic limbs (Martin & Cash-Clarck 1995) or visually by its large medial compound eye. Unlike *Leptodora*, *Bythotrephes* does not have a feeding basket for prey capture. Instead it grasps a prey item with its long feeding appendages and shreds its prey, ingesting only soft parts (Schulz & Yurista 1999). Because it is not restricted by the capacity of its feeding basket, large *Bythotrephes* are probably able to feed successfully on larger prey items.
There is an increasing amount of field observations from N. America, where the exotic *Bythotrephes* has invaded lakes, showing that this predator often has a serious impact on the microcrustacean zooplankton communities. These results show further that in general cladocerans are more vulnerable to *Bythotrephes* predation than copepods and that smaller and intermediate sized cladocerans are more vulnerable than large cladocerans (Yan et al. 2001, 2002). Average species richness was 30% higher in the reference as in the invaded lakes (Boudreau & Yan 2003). Total zooplankton biomass was significantly lower too in the invaded lakes, mainly because of lower abundances of all common epilimnetic cladoceran species. In a more recent but similar field study in North American lakes, Strecker et al. (2006) showed that *Bythotrephes* significantly reduced cladoceran species richness, diversity and abundance. The larger species *Daphnia longiremis* and *Daphnia mendotae* also had lower abundances in invaded lakes as compared to reference lakes, suggesting that small body size alone does not provide refuge from *Bythotrephes* predation. This is corroborated by the results of Barbiero & Tuchman (2004) who showed that in lakes Michigan, Huron and Erie, *Daphnia reticulata* and *Daphnia pulicaria* declined dramatically after the introduction of *Bythotrephes*.

Since *Bythotrephes*, unlike *Leptodora*, does not have a feeding basket for catching its prey, prey size cannot be estimated from the dimensions of the feeding basket as was done previously for *Leptodora* (Manca & Comoli 1995; Branstrator 1998) and, therefore, vulnerability of prey species and prey size classes has to be inferred from experiments. *Bythotrephes* is difficult to keep under laboratory conditions, therefore published information about selective feeding is scarce, to our best knowledge only two experimental studies exist which report contrasting results. Schulz & Yurista (1999) concluded that *Bythotrephes* is selecting medium sized (1.4-1.6 mm) daphnids (one experiment) and/or daphnids larger than 2.0 mm (another experiment), whereas Vanderploeg et al. (1993) concluded on basis of their predation experiment that mainly cladocerans in the size range of ca. 0.5-0.9 mm were selected as food items.

*Leptodora* is a cruising predator, and employs strike tactics to capture prey. The prey location space for *Leptodora* is limited to a small forward-directed area and direct contact with the prey is required before an attack is initiated (Browman et al. 1989). *Leptodora* tears the cladocerans apart, the wider the ventral carapace gape of the prey the easier for *Leptodora* to tear it apart (Browman et al. 1989). An important limiting factor for prey capture success is the size of the feeding basket, a structure formed by thoracic appendages enabling *Leptodora* to catch and handle prey for ingestion (Manca & Comoli 1995; Branstrator 1998).

*Leptodora* feeds mainly on small-bodied cladocerans and avoids copepodite copepods (e.g. Lunte & Luecke 1990; Herzig 1994). Although relatively large, *Diaphanosoma* and *Daphnia* spp. are eaten too, from a newborn size (ca 0.5 mm) until a maximum size of ca. 1.0 mm (Browman et al. 1989; Herzig 1995). Only very few *Leptodora* consume *Daphnia* >1.0 mm and these *Leptodora* were always >11.0 mm (Branstrator & Lehman 1991). Copepod nauplii and rotifers are eaten by small juvenile *Leptodora* (i.e. <5 mm), but contribute in terms of biomass generally little to the *Leptodora* diet as a whole (Arndt et al. 1993).

The invertebrate (*Bythotrephes*, *Leptodora*) and vertebrate (coregonid spp.) types of predators produce different effects on the populations of their *Daphnia* prey: fish predation results in decreased *Daphnia* fecundity, decreased proportion of adults, and a reduction of adult body size, while predation by the invertebrate predators leads to a decrease in the proportion of young *Daphnia* and, therefore, in an increase in the proportion of adults (Gliwicz & Pijanowska 1989; Polishchuk 1995).

Here we present the results of a field study of predation by two types of predators (one vertebrate, two invertebrates) on *Daphnia*, which was aimed at investigating the predator’s, rather than the prey’s, size-specific response. The study combines contribution analysis of *Daphnia* birth rate dynamics (Polnishchuk 1995), with a body-size-oriented analysis of both predator and prey populations.

2. MATERIAL AND METHODS

2.1. Sampling of zooplankton

Zooplankton samples were collected in May-June 2001 at weekly intervals at a station corresponding to the maximum lake depth. They represented the content of at least 1000 L of lake water filtered through a 126 μm nylon net mounted on a Clarke-Bumpus plankton sampler, which was towed along a sinusoidal trajectory within the upper 50 m, the water layer where cladocerans are distributed. *Daphnia* and *Bythotrephes* size estimates were the result of measurements of body length (from the apex of the head to the base of the carapace spine) of 200 individual *Daphnia* per sample and all *Bythotrephes* individuals found in each sample.

2.2. Contribution analysis and population dynamics

Contribution analysis (Caswell 1989; Polishchuk 1995, 1999; Polishchuk & Vijverberg 2005) was used to investigate the relative importance of different environmental effects to changes in *Daphnia* birth rate. The idea behind the analysis is to assess the role of major environmental factors such as food, predation, and temperature in terms of corresponding population parameters.

In *Daphnia*, temperature mainly acts on the developmental time of eggs, while food conditions affect fecundity (number of eggs or embryos per adult female where being adult is normally determined on the basis of body
size). Fish predation, being directed against larger adult females carrying larger clutches, influences both fecundity and the proportion of adults. Predation by invertebrates is thought to be mainly directed against juveniles, and therefore is related to the increase in the proportion of adults. Also, if invertebrate predators take off some of the smaller adults with few or no eggs, they will affect fecundity too, though probably to a lesser extent than fish. We expect therefore that fish, as compared with invertebrate predators, will produce a more pronounced and potentially less uniform (with more room for variation) effect on fecundity and the proportion of adults of their prey.

The per capita birth rate \( b \) of *Daphnia* is calculated according to equation:

\[
b = V \ln(1 + FA)
\]  
(1)

where: \( V \) is egg developmental rate (day\(^{-1}\)), the reciprocal of egg developmental time, obtained from mean water temperature of the sampled water layer; \( F \) (\( F = E/N_{ad} \)) is fecundity, the ratio of eggs numbers to adult females numbers, irrespective of whether those females carry eggs or not; \( A \) (\( A = N_{ad}/N \)) is the proportion of adult females to the total number of individuals in the population. The product \( FA \) is equal to \( E/N \); therefore, by substituting \( FA \) in equation (1) we obtain:

\[
b = V \ln(1 + E/N)
\]  
(2)

which represents the Edmondson and Paloheimo equations for birth rate estimation (Edmondson 1968; Paloheimo 1974). The rate of population increase \( r \) was estimated by:

\[
r = (\ln N_{t_2} - \ln N_{t_1}) / (t_2 - t_1)
\]  
(3)

where \( N_{t_1} \) and \( N_{t_2} \) are the number of individuals in the population at times \( t_1 \) and \( t_2 \).

Death rates \( d \) are estimated by:

\[
d = b - r
\]  
(4)

Starting from equation (1) contributions of \( F \), \( A \) and \( V \) (\( ConF \), \( ConA \), \( ConV \)) to changes in \( b \) can be expressed as partial derivatives with respect to the corresponding parameters times changes in those parameters:

\[
db/dt = \left[ VA/(1+FA) \right] (dF/dt) + \left[ VF/(1+FA) \right] (dA/dt) + \left[ \ln(1+FA) \right] (dV/dt)
\]

where: \( ConF = \left[ VA/(1+FA) \right] (dF/dt) \), \( ConA = \left[ VF/(1+FA) \right] (dA/dt) \), and \( ConV = \left[ \ln(1+FA) \right] (dV/dt) \).

However, the sum of the contributions determined in this way is not exactly equal to \( db/dt \) (Polishchuk 1995). Hence, one of us (D.A.V.) proposed taking integral of the contributions over time, with the result that the sum of contributions is precisely equal to the change in \( b \) over sampling interval, \( \Delta b \). Such integrals can be calculated numerically. This method is used in the present study. Provided that changes in temperature are not large, \( ConV \) is relatively small, and changes in *Daphnia* birth rate can be analysed in terms of \( ConF \) (food conditions) or both \( ConA \) and \( ConF \) (predation), with the \( ConA-ConF \) pattern being less uniform (i.e. the difference between \( ConA \) and \( ConF \) being more pronounced) under fish predation than under invertebrate predation.

3. RESULTS

*Daphnia* population density increased during April and May (Fig. 1); the subsequent decline corresponded to an increase in *Bythotrephes* abundance. *Leptodora*, although appearing in late May, started its numerical
increase in June, when Daphnia numbers were already low.

During April to mid May the per capita birth rate \((b)\) of Daphnia was approximately 0.05 day\(^{-1}\) and showed only little variation over time. At the end of May it reached its lowest value (0.02 day\(^{-1}\)) after which it steadily increased to values of 0.08-0.13 day\(^{-1}\) during the second half of June (Fig. 2). The per capita death rate \((d)\) showed much more variation in time. Part of it was due to sampling errors, because population densities were estimated on one station only. To reduce variation we calculated the moving average of the death rate over time (Fig. 2). We can distinguish two main phases in the seasonal death rate \((d)\) pattern of the Daphnia population. The first phase, in April to mid May when the death rates slowly increased but were still very low, was concomitant with low Bythotrephes and Leptodora densities and a sharp increase of Daphnia densities during early May. The second phase started during the last week of May with a strong increase in the death rate, which reached high values in early June and remained high during that whole month. This phase was concomitant with sharp increasing Bythotrephes densities (until mid June) and steady increasing Leptodora densities from mid June onwards. This high death rate was associated with a sharp decrease in Daphnia densities (Fig. 1).

The body size of the Daphnia ovigerous females decreased during April and early May, then remained more or less constant till mid June, and increased in size in the second half of June. The mean clutch size started high in April-mid May, then decreased during the second half of May and remained low during early June and then from mid June onwards started to increase again until the end of June (Fig. 3). The mean size at maturity (MSAM) varied from 1.6 mm in early April to 1.85 mm in the second half of June.

Not only was Bythotrephes becoming more abundant in May-June, but also the maximum body size it attained was increasing, reaching its maximum value at the end of June (Fig. 4). Daphnia mean size decreased somewhat during April, but then from early May onwards increased steadily during May and June; largest body sizes were reached during the second half of June. From early May onwards increase in body size of Daphnia and Bythotrephes occurred approximately with the same rate.

Contribution analysis showed that during April-mid May, when Bythotrephes densities were low, changes in Daphnia birth rate were due to both ConF and ConA, with ConA being much larger (by absolute value) than ConF (Fig. 5a). During mid May to June changes in Daphnia birth rate were again mainly due to both ConF and ConA, but now the difference between ConA and ConF was less pronounced (Fig. 5b). During late June, when Leptodora population density increased and Daphnia densities fell to low values, changes in birth rate were mainly related to ConA (Fig. 5c).

4. DISCUSSION

Predation on Daphnia by fish or invertebrates is size-selective; while fish prey on larger ovigerous females carrying large clutches, invertebrates are generally thought to remove only small, immature females, though removal of some of the smaller adults with few eggs is probably also possible. The two types of predation produce different patterns of the population dynamics of the prey, which can be identified by an analysis of Daphnia birth rate dynamics and body size. When fish predation prevails, changes in Daphnia birth rate are determined by both changes in the proportion of adults \((\text{ConA})\) and in fecundity \((\text{ConF})\), with ConA being much larger (by absolute value) than ConF becau-
se fish typically produce a stronger effect on the size structure than on fecundity of their prey. When invertebrate predation prevails, changes in *Daphnia* birth rate are also determined by ConA and ConF, but now the ConA-ConF pattern is expected to be more uniform. This is because the effects produced by invertebrates are apparently weaker than those by fish, and weak effects are perceived as less variable.

During April-mid May, contribution analysis revealed a ConA-ConF pattern that is expected under fish predation. Moreover, the selective removal of larger ovigerous females expected on the basis of the contributions, was accompanied by a decrease in *Daphnia* body size (maximum and mean body size, as well as the mean size of ovigerous females) and a reduced mean clutch size observed in the lake. This pattern typically occurs in Lago Maggiore during April-mid May, when *Daphnia* is the preferred prey of coregonid fish (Giussani 1974; Volta 2000).

During mid May-June, contribution analysis revealed that changes in *Daphnia* birth rate also resulted from both ConA and ConF, but the difference between these contributions was less pronounced than that observed in the previous period. This is what we expect under invertebrate predation. In addition, not only did *Daphnia* mean body size increase but also the mean size of the ovigerous females remained almost constant, which can be explained by the prevailing removal of small immature individuals. In the second half of June the mean size of ovigerous females started to increase, which indicates that also small ovigerous females were removed. This shows that invertebrate predators, most probably

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**Fig. 3.** Seasonal changes in size of *Daphnia* ovigerous females and in the clutch size (mean ± 1.96 SE).

**Fig. 4.** Seasonal changes in *Daphnia* mean (± 1.96 SE) body size, along with *Bythotrephes* maximum body size.
Bythotrephes, which was in terms of biomass the most abundant one, were the dominant predators on Daphnia. Changes in Daphnia body size and the relative contributions of population traits to changes in birth rate can be explained by viewing Bythotrephes as a predator able to select not only young, but also some of the small sized ovigerous Daphnia, in parallel with the increase in predator body size. Although Bythotrephes predation seems to produce similar effects on Daphnia population dynamics as fish predation, affecting both fecundity and the proportion of adults, it can be distinguished in two ways from fish predation. Firstly, by an increase in Daphnia mean body size, whereas the mean size of ovigerous females remains approximately constant (mid May-mid June) or increased in size (mid June-end of June). Secondly, because the pattern of contributions to Daphnia birth rate dynamics observed under the implied Bythotrephes predation is more uniform than that observed under implied fish predation.

During late June, contribution analysis revealed that changes in Daphnia birth rate were determined by ConA, a pattern in that respect somewhat similar to that observed in April (see Fig. 5). It can be hardly attributed to fish pressures however, because at that time the dominant predator in the lake was Leptodora rather than fish. On the other hand, it can be hardly indicative of invertebrate pressures because for invertebrates we expect a more uniform ConA-ConF pattern, a sort of that shown in Fig. 5b. Apparently, contribution results do not reflect the pressures by Leptodora at that time, because the density of Daphnia was extremely low in late June (Fig. 1) and Leptodora might switch to another food source.

We conclude that both predation by zooplanktivorous fish and by invertebrate predators are affecting population dynamics of Daphnia in Lago Maggiore. Among the invertebrate predators Bythotrephes has the largest impact on Daphnia dynamics, both because its relatively high population biomass and its relatively large body size. In the second half of June its size is large enough to predate also on small ovigerous Daphnia, and its predation pressure is high enough to reduce the Daphnia densities to low values.

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