

# Carbon transfer in herbivore- and microbial loop-dominated pelagic food webs in the southern Barents Sea during spring and summer

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**ABSTRACT:** We compared carbon budgets between a herbivore-dominated and a microbial loop-dominated food web and examined the implications of food web structure for fish production. We used the southern Barents Sea as a case study and inverse modelling as an analysis method. In spring, when the system was dominated by the herbivorous web, the diet of protozoa consisted of similar amounts of bacteria and phytoplankton. Copepods showed no clear preference for protozoa. Cod *Gadus morhua*, a predatory fish preying on copepods and on copepod-feeding capelin *Mallotus villosus* in spring, moderately depended on the microbial loop in spring, as only 20 to 60% of its food passed through the microbial loop. In summer, when the food web was dominated by the microbial loop, protozoa ingested 4 times more bacteria than phytoplankton and protozoa formed 80 to 90% of the copepod diet. Because of this strong link between the microbial loop and copepods (the young cod's main prey item) young cod (<3 yr) depended more on the microbial loop than on any other food web compartment, as >60% of its food passed through the microbial loop in summer. Adult cod (≤3 yr) relied far less on the microbial loop than young cod as it preyed on strictly herbivorous krill in summer. Food web efficiency for fish production was comparable between seasons ( $\sim 5 \times 10^{-4}$ ) and 2 times higher in summer ( $5 \times 10^{-2}$ ) than in spring for copepod production.

**KEY WORDS:** Food web · Microbial loop · Protozoa · Copepods · *Gadus morhua*

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## INTRODUCTION

The structure of most pelagic food webs can be thought of as a hybrid between the traditional food web, dominated by herbivory, and the microbial loop (Legendre & Rassoulzadegan 1995). Systems dominated by the herbivorous web, hereafter called 'herbivorous food webs', mainly occur in nutrient-rich environments, rely on phytoplankton production and are characterised by large phytoplankton cells with a tendency to aggregate and, thus, suffer high sedimentary losses. In contrast, microbial loop-dominated food webs, hereafter termed 'microbial food webs', are typically associated with low nutrient status and are fuelled by picoplanktonic primary production with considerable recycling by bacteria. This recycling may be enhanced by production and consumption of dis-

solved organic carbon (DOC). DOC may be released by phytoplankton exudation (Fasham et al. 1999), by sloppy feeding from zooplankton (Jumars et al. 1989) and by viral lysis of phytoplankton and bacterial cells (Fuhrman 2000). This DOC is taken up by bacteria and may be transferred via protozoa to zooplankton and higher trophic levels in successive grazing steps (Azam et al. 1983).

Because of its role in the global carbon cycle, the structure and function of pelagic food webs has been the main object of study in large-scale field studies carried out in recent years. These studies include the Joint Global Ocean Flux Study (JGOFS) (e.g. Smith et al. 2000, Steinberg et al. 2001), the ROAVERRS Program (Research on Ocean-Atmosphere Variability and Ecosystem Response in the Ross Sea) (Arrigo et al. 1998) and the Palmer Long-Term Ecological Research

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(PAL-LTER) Program (Ross et al. 1996). Apart from abiotic quantities (e.g. nutrient levels), sampling campaigns typically include measurements of primary and bacterial (secondary) production and of standing stocks of the main phytoplankton species, bacteria and mesozooplankton. Because of the focus on the microbial components of the pelagic food web, mesozooplankton are often the highest trophic level considered. Relationships between (bacterial and/or primary) production and consumption (by [proto]zooplankton) tend to be estimated by grazing experiments (as in Vargas et al. 2007) or by numerical modelling (Fasham et al. 1999).

Whereas the microbial components of the pelagic food web play a crucial role in biogeochemical cycles (Sabine et al. 2004), they also provide the necessary resources for higher trophic levels such as fish, marine mammals and humans. The differences between phytoplankton food webs and microbial food webs in terms of the efficiency of elemental cycling may thus lead to differences in terms of production rates of higher trophic levels. This concern is being included in emerging end-to-end approaches that amalgamate the food web's different trophic levels next to physical drivers to investigate the effect of environmental perturbations on marine ecosystems (e.g. Cury et al. 2008, Pedersen et al. 2008). Although relationships between lower and higher trophic levels have been established for some time, and summarised in the top down versus bottom up paradigm, it is unclear how the structure of the planktonic community (microbial versus herbivorous) determines the production rate of top predators in natural ecosystems. We addressed this important issue by studying how food web structure (microbial versus herbivorous) determines carbon transfer from primary producers to predatory fish in a natural ecosystem.

Food web flows in a highly productive ecosystem (the ice-free southern Barents Sea) were estimated during the spring bloom, when the food web is predominantly herbivorous, and during summer, when a microbial food web structure prevails (Wassmann et al. 2006). Food web flows were estimated with inverse models (Klepper & Vandekamer 1987, Vezina & Platt 1988, Soetaert & Van Oevelen 2009) that use an extensive data set gathered from the literature. The inverse modelling approach is comparable with other techniques that use mass balance to estimate elemental budgets such as Ecopath (Pauly et al. 2000), yet it differs in the following ways. While Ecopath to a large extent depends on diet compositions defined *a priori*, the inverse method only requires food web topology and estimates the quantitative importance of food web flows upon model solution. Also, parameters do not have to be defined by a unique value, but instead a

range of physiologically realistic values can be assigned. This uncertainty is then used to estimate uncertainty associated with the food web flow solutions. The inverse food web models set up here include bacteria, protozoa, 3 types of phytoplankton, meso- and macrozooplankton and the main fish species. Fish production is a food web output. Based on estimated food web flows, we estimated the (direct plus indirect) dependency of fish on lower trophic levels (Szyrmer & Ulanowicz 1987) and the efficiency (sensu Rand & Stewart 1998) of fish production in both food web structures. Differences in food web efficiency or in dependencies between both food web structures are discussed based on differences in estimated carbon budgets. The southern Barents Sea was chosen as a model ecosystem because (1) its distinct seasonality allows studying both food web types in one ecosystem, (2) its fish population sustains one of the world's largest fisheries (Bogstad et al. 2000), and (3) it contains characteristics that are present in many polar systems such as the seasonal migration of species induced by spatial heterogeneity (Carmack & Wassmann 2006). The choice for the Barents Sea is also a practical one, as data on lower trophic levels (Wassmann 2002) and the most important fish stocks (ICES 2008) are abundant, which facilitates the quantitative reconstruction of food web flows.

## MATERIALS AND METHODS

**Study region and conceptual food web.** The southern part of the Barents Sea is characterised by permanently ice-free waters and inflow of water from the Atlantic Ocean. The largest data set available in the literature that covers both spring and summer was for May 1998 (spring) and July 1999 (summer). Microbial and zooplankton compartments are a reflection of the species groups found during the Arktisk Lys og Varme (ALV) sampling campaign, as described in a special issue dedicated to Barents Sea C-flux (Wassmann 2002). Fish compartmentalisation was inferred from Bogstad et al. (2000). Representative food web compartments for the area are DOC, detritus, bacteria, heterotrophic flagellates, heterotrophic ciliates, phytoplankton (pico- and nanoplankton, diatoms and *Phaeocystis* sp.), mesozooplankton (copepods), macrozooplankton (krill and chaetognaths), cod *Gadus morhua* and herring *Clupea harengus*. The spring food web also contains capelin *Mallotus villosus*, but as capelin migrate out of the southern Barents Sea before summer, they are excluded from the summer food web. Because the diet of cod changes drastically during their life cycle (Mehl 1989), adult cod ( $\geq 3$  yr) and young cod ( $< 3$  yr) were considered as 2 different populations.

We acknowledge that models focusing on the fish community often make use of more than 2 age classes (e.g. Hjermann et al. 2004). The number of age classes in our model, which describes a complete food web and not only the fish community, is a trade-off between realism and complexity. Additionally, the use of 2 age classes is not unrealistic and has been successfully applied elsewhere (Hjermann et al. 2007, Durant et al. 2008).

The food web topology within the microbial community was set as follows. Phytoplankton fix dissolved inorganic carbon (DIC), an external food web model input, and transform this carbon into a particulate and dissolved form. Bacteria take up DOC and are consumed by protozoa. Bacterial cells may lyse and thereby release DOC that can be re-used for >90% (Nagata 2000, Ogawa et al. 2001, Davis & Benner 2007). Only a small percentage enters the DOC pool, which is to a large extent refractory. Exact data on the size of the DOC pool were not available for the years considered. Yet the DOC pool is not used in any of the constraints imposed on the inverse model (see below) and, therefore, the size of this pool does not influence the outcomes of the model. Heterotrophic flagellates are preyed upon by ciliates, and both protozoan groups eat all 3 phytoplankton groups and can be consumed by copepods (Calbet & Saiz 2005). Krill in the southern Barents Sea mainly consists of the herbivorous species, *Thysanoessa inermis* (Falk-Petersen et al. 2000), and therefore also fed exclusively on the phytoplankton groups in the model. Chaetognaths eat copepods (Tonnesson & Tiselius 2005). The food web topology within the fish community and their food was set as in Bogstad et al. (2000): capelin and herring eat krill, chaetognaths and copepods, while young cod feed on krill, chaetognaths, copepods, capelin and herring. Adult cod is the top predator and can eat all zooplankton and fish, including young cod. Respiration and excretion of all populations was introduced by flows to DIC and DOC, respectively. Sedimentation of phytoplankton and detritus was considered a loss from the food web, which is important to consider since vertical export can be high in the Barents Sea ecosystem (Wassmann et al. 2006). Detritus is produced as faeces by all populations except bacteria and protozoa and can be transformed to DOC. Trophic levels higher than cod were not explicitly included, but instead, chaetognaths, krill and fish compartments were equipped with an export flow to allow for predation by whales, seals, birds and humans. Additionally, zooplankton and fish were provided with an additional export flow to allow for net population growth. The possibility of advection of copepod biomass with water from the Atlantic Ocean into the Barents Sea was incorporated as well, as deemed necessary in contemporary literature on the

Barents Sea carbon budget (Wassmann et al. 2006). The resulting food web, including all components and flows is shown in Fig. S1 (Supplement 1, [www.int-res.com/articles/suppl/m398p093\\_app.pdf](http://www.int-res.com/articles/suppl/m398p093_app.pdf)).

The described food web topology does not contain any information regarding the quantitative importance of the flows and without additional data each flow can theoretically range from zero to infinity. In the next section we describe the data and constraints used to restrict these flows to ranges that are biologically realistic and consistent with the characteristics of the Barents Sea.

**Data and constraints for setup of the linear inverse models.** Two types of data are typically used to setup a linear inverse model (LIM): standing stocks and rate measurements. As a typical spring month we chose May 1998 and as a summer month July 1999 was chosen, mostly because a rich data set exists for these months. The year 1999 was a warm year (4 to 6°C in summer and an annual primary production of 100 to 125 g C m<sup>-2</sup>), while 1998 was a cooler year (2 to 3°C in spring and an annual primary production of 50 to 75 g C m<sup>-2</sup>) (Wassmann et al. 2006). The late 1990s represented the late recovery phase for capelin, a relatively low cod stock and a high herring stock (ICES 2005, 2008). Standing stocks of all microbial compartments, phytoplankton and zooplankton were found in Wassmann (2002). Sampling locations from which data were used were not ice-covered during May 1998 and July 1999 and are listed and georeferenced (latitude/longitude) in Table 1 of Arashkevich et al. (2002) as those locations with ‘%Ice cover’ = 0. A map of the region can also be found in the supporting information (Fig. S2, Supplement 1). Standing stocks of fish were found in ICES reports (ICES 2005, 2008). Table 1 provides a complete overview of the data used and references. In the absence of year-specific data, we assumed equal distribution among Barents and Norwegian sea subregions for standing stock estimation of cod, (see footnote j to Table 1). Spatial patterns of the cod stock for 1998 to 1999, which are also available for earlier years (Huse et al. 2004), may refine these cod stock estimates. In cases where multiple data points for the standing stocks of one model compartment were available, the median value was used. Measured processes included production rates of faecal pellets by zooplankton, sedimentation rates of detritus and phytoplankton, bacterial production, primary production and ingestion rates by protozoa.

We acknowledge that the use of inverted microscopes may yield somewhat conservative estimates for the standing stocks of picoplankton and that higher concentrations may have occurred. However, with no information on the quantitative importance of such

Table 1. Data and constraints for inverse model construction of the southern Barents Sea food web. NPE: net production efficiency; AE: assimilation efficiency; GPE: gross production efficiency; na: not available; BW: body weight; protozoa: heterotrophic nanoflagellates and ciliates; *P. pouchetii*: *Phaeocystis pouchetii*; NPPP: net particulate primary production, i.e. corrected for excretion and respiration losses; NPP: net primary production, i.e. corrected for respiration losses; GPP: gross primary production; POC: particulate organic carbon. 'h' denotes the fraction of nanoflagellates that is heterotrophic. In May, this was unknown and assigned a value of 0, 0.5 and 1 in HF0, HF50 and HF100, respectively. In July, this value was 0.3 (Verity et al. 2002). Actual standing stocks per m<sup>2</sup> were calculated from depth profiles given in the corresponding papers. Single values indicate that this value is included as fixed values and 2 values give the range that is imposed on the model. Conversion of wet weight to carbon for invertebrates was done as: 1 g wet weight = 1 × 0.10 g dry weight = 1 × 0.10 × 0.5 g C (Hendriks 1999). Data categorized under spring and summer were used for both seasons

Population	Characteristic	Unit	Spring	Summer	Source
<b>Phytoplankton</b>					
Data					
Picophytoplankton	Standing stock	g C m <sup>-2</sup>	0.241	0.045	Rat'kova & Wassmann (2002)
Nanophytoplankton <sup>a</sup>	Standing stock	g C m <sup>-2</sup>	(1 h) × standing stock of nanoflagellates		Rat'kova & Wassmann (2002)
Nanoflagellates	Standing stock	g C m <sup>-2</sup>	5.14	5.8	Rat'kova & Wassmann (2002)
Diatoms	Standing stock	g C m <sup>-2</sup>	3.3	1.50	Rat'kova & Wassmann (2002)
Diatoms	Sedimentation rate	g C m <sup>-2</sup> d <sup>-1</sup>	0.05–0.15	na	Olli et al. (2002)
<i>P. pouchetii</i>	Standing stock	g C m <sup>-2</sup>	4.35	1.25	Rat'kova & Wassmann (2002)
<i>P. pouchetii</i>	Sedimentation rate	g C m <sup>-2</sup> d <sup>-1</sup>	0.6–0.8	na	Olli et al. (2002)
Total phytoplankton	Sedimentation rate	g C m <sup>-2</sup> d <sup>-1</sup>	0.5–1.8	0.04–1.3	Olli et al. (2002)
Total phytoplankton	NPPP rate	g C m <sup>-2</sup> d <sup>-1</sup>	1–1.5	0.2–0.8	Matrai et al. (2007)
Constraints					
All phytoplankton	Excretion rate	Fraction of NPP	0.05–0.6	0.05–0.6	Vezina & Platt (1988) and Matrai et al. (2007)
All phytoplankton	Respiration rate	Fraction of GPP	0.05–0.3		Vezina & Platt (1988)
All phytoplankton	Standing stock-specific GPP	d <sup>-1</sup>	0.5–1.5		MacIntyre et al. (2002)
<b>Microbial loop</b>					
Data					
Ciliates	Standing stock	g C m <sup>-2</sup>	0.04–0.05	0.12–0.55	Rat'kova & Wassmann (2002)
Heterotrophic bacteria	Standing stock	g C m <sup>-2</sup>	1.8	2.20	Howard-Jones et al. (2002)
Heterotrophic bacteria	Production/biomass	d <sup>-1</sup>	na	1.1–3.9	Howard-Jones et al. (2002)
Heterotrophic nanoflagellates	Standing stock	g C m <sup>-2</sup>	h × standing stock of nanoflagellates		Rat'kova & Wassmann (2002)
POC	Standing stock	g C m <sup>-2</sup>	26.0	13.6	Olli et al. (2002)
POC	Sedimentation rate	g C m <sup>-2</sup> d <sup>-1</sup>	0.8–1.7	0.2–0.4	Olli et al. (2002)
Protozoa	Ingestion rate	Fraction of NPPP removed daily	na	0.7–1	Verity et al. (2002)
Constraints					
Detritus	Dissolution rate	d <sup>-1</sup>	<0.02		Donali et al. (1999)
Protozoa	Biomass-specific respiration rate	d <sup>-1</sup>	>0.08		Vezina & Platt (1988)
Protozoa	GPE		0.1–0.6		Vezina & Platt (1988)
Protozoa	Uptake rate	Proportion of BW d <sup>-1</sup>	<7		Vezina & Platt (1988)
Protozoa	Excretion	Fraction of respiration rate	0.33–1		Vezina & Platt (1988)
Heterotrophic bacteria	GPE		0.01–0.6		del Giorgio & Cole (2000)
Heterotrophic bacteria	Sedimentation rate	Fraction of bacterial production rate	<2%		Donali et al. (1999)
Heterotrophic bacteria	Viral mortality of bacteria	Fraction of bacterial production rate	10–40%		Fuhrman (2000)
<b>Zooplankton</b>					
Data					
Chaetognaths	Standing stock	g C m <sup>-2</sup>	0.6965	0.0897	Arashkevich et al. (2002)
Copepods	Standing stock	g C m <sup>-2</sup>	1.79	1.50	Arashkevich et al. (2002)
Copepods <sup>b</sup>	Biomass advection with Atlantic current	g C m <sup>-2</sup> d <sup>-1</sup>	<0.23		Edvardsen et al. (2003)
Total zooplankton	Fecal pellet production rate	g C m <sup>-2</sup> d <sup>-1</sup>	>0.11	>0.18	Wexels Riser et al. (2002)
Krill	Standing stock	g C m <sup>-2</sup>	0.003	0.00188	Arashkevich et al. (2002)
Constraints					
Copepods	Respiration rate	d <sup>-1</sup>	0.015–0.038		Drits et al. (1993)
Copepods	Ingestion rate	d <sup>-1</sup>	0.008–0.14		Tande & Bamstedt (1985)
Copepods	AE		0.5–0.9		Besiktepe & Dam (2002)

Table 1 (continued)

Population	Characteristic	Unit	Spring	Summer	Source
Copepods	GPE		<0.4		Vezina & Platt (1988)
Krill	Ingestion rate	d <sup>-1</sup>	<0.4		Vezina & Platt (1988)
Krill	Respiration rate	d <sup>-1</sup>	>0.03		Vezina & Platt (1988)
Krill	GPE		<0.4		Vezina & Platt (1988)
Krill	Production/biomass	d <sup>-1</sup>	>0.0058		Siegel (2000)
Chaetognaths	Ingestion rate	d <sup>-1</sup>	0.1–0.3		Tonnesson & Tiselius (2005)
Chaetognaths	NPE		0.15–0.35		Welch et al. (1996)
Chaetognaths	AE		0.7–0.9		Reeve & Cosper (1975)
Chaetognaths	Respiration rate	d <sup>-1</sup>	0.0072–0.0252		Welch et al. (1996)
All zooplankton <sup>c</sup>	AE		0.5–0.9		Vezina & Platt (1988)
All zooplankton <sup>c</sup>	Excretion	Fraction of respiration rate	0.33–1		Vezina & Platt (1988)
<b>Fish</b>					
<b>Data</b>					
Adult cod	Ratio of ingestion x/y of prey item x over prey item y		7–9 (krill/herring) 2–4 (krill/young cod)		Bogstad et al. (2000)
Adult cod <sup>d</sup>	Maximum net growth rate during summer and spring	d <sup>-1</sup>	0.0005–0.0015		Pogson & Fevolden (1998)
Adult cod <sup>e,f,g</sup>	Standing stock	g C m <sup>-2</sup>	0.053	0.033	ICES (2008)
Capelin <sup>f,g,h</sup>	Standing stock	g C m <sup>-2</sup>	0.38	0	ICES (2008)
Herring <sup>f,g,i</sup>	Standing stock	g C m <sup>-2</sup>	0.055	0.132	ICES (2005)
Young cod <sup>l, f, g</sup>	Standing stock	g C m <sup>-2</sup>	0.006	0.003	ICES (2008)
Young cod	Contribution of herring in diet		<contribution of copepods, <contribution of capelin, <contribution of krill		Dalpadado & Bogstad (2004) and references therein
<b>Constraints</b>					
All fish <sup>a</sup>	AE		0.7–0.9		Hendriks (1999)
Adult and young cod	Ingestion rate	d <sup>-1</sup>	0.017–0.054		I. Y. Ponomarenko & N. A. Yaragina (unpubl. data)
Adult and young cod <sup>k</sup>	Excretion rate	Fraction of assimilation rate	0.045–0.135		Holdway & Beamish (1984)
Adult cod	GPE		0.2–0.5		References in Hansson et al. (1996), e.g. Daan (1975)
Capelin	Ingestion rate	d <sup>-1</sup>	0.013–0.022		Ajiad & Pushchaeva (1992)
Capelin <sup>l</sup>	Respiration rate	d <sup>-1</sup>	0.01–0.04		Karamushko & Christiansen (2002)
Herring	Ingestion rate	d <sup>-1</sup>	0.01–0.1		Megrey et al. (2007)
Herring <sup>m,n</sup>	Maintenance respiration rate	d <sup>-1</sup>	0.019–0.024		Megrey et al. (2007)
Herring <sup>m,n</sup>	Growth respiration rate	d <sup>-1</sup>	0.0875–0.263		Rudstam et al. (1988)
Herring	AE		0.76–0.92		Arrhenius (1998)
Herring	Excretion rate	Fraction of ingestion rate	0.05–0.15		Klumpp & Vonwesternhagen (1986)
Young cod	GPE		0.3–0.5		Daan (1975)

<sup>a</sup>The sum of these 2 groups forms the small autotrophs

<sup>b</sup>Assuming  $7 \times 10^6$  t C advection to occur during 1 mo

<sup>c</sup>Only used when species-specific data not available

<sup>d</sup>Calculated from  $0.1–0.3 \text{ yr}^{-1}$  assuming growth occurs mainly during 6 mo

<sup>e</sup>Total stock for fishing areas I, IIa and IIb = 0.84 million t (spring 1998) and 1.1 million t (summer 1999); stock in area I = total stock/3; standing stock concentration in southern Barents Sea = stock in area I  $\times$  (surface area southern Barents Sea)<sup>-1</sup>. In summer, half of the adult cod stock follows capelin migrating North, thus lowering the concentration by 50%

<sup>f</sup>Surface area of southern Barents Sea = 0.7 million km<sup>2</sup>

<sup>g</sup>1 g wet weight =  $1 \times 0.33$  g dry weight =  $1 \times 0.33 \times 0.4$  g C; valid for fish (Sakshaug et al. 1994)

<sup>h</sup>2 million t (spring 1998) and 0 t (summer 1999) in southern Barents Sea

<sup>i</sup>0.29 million t (spring 1998) and 0.7 million t (summer 1999)

<sup>j</sup>Spring 1998: total stock for fishing areas I, IIa and IIb = 6620 age 1 fish (10 g each) + 1254 age 2 fish (47 g each) = 92 645 t; summer 1999: total stock for fishing areas I, IIa and IIb = 3004 age 1 fish (12 g each) + 1062 age 2 fish (55 g each) = 94 458 t; standing stock concentration in southern Barents calculated as for adult cod

<sup>k</sup>Data for North Sea cod

<sup>l</sup>Calculated from  $0.05$  to  $0.15 \text{ ml O}_2 \text{ g}^{-1} \text{ h}^{-1}$

<sup>m</sup>Energy density ratios used are between 0.8 and 1.2 as in Megrey et al. (2007)

<sup>n</sup>Range taken as reported value  $\pm$  50%



deviations, we relied on the original data rather than applying some arbitrary scaling factor. Also the used value for krill is particularly low ( $0.003 \text{ g C m}^{-2}$ ) compared with stock assessments for earlier years (Dalpadado & Skjoldal 1996). Differences in sampling gear (Multiple Opening/Closing Net and Environmental Sampling System [MOCNESS] versus WP2 nets) have been suggested as a possible cause (Arashkevich et al. 2002), yet Gjosaeter et al. (2000) found only small differences (factor of 1.4 to 2) between these 2 sampling methods and only at 3 out of 9 sampling locations. This original krill biomass value was therefore maintained.

A large number of constraints on the food web flows were included in the inverse model (see Table 1). Briefly, these constraints reflect limits on the physiology and biological functioning of marine organisms and include constraints on biomass-specific respiration, ingestion and production rates, excretion processes, and assimilation and/or production efficiencies.

No information was available on the fraction of flagellates that was heterotrophic in spring. Thus, the standing stocks of heterotrophic flagellates and pico- and nanoplankton (includes autotrophic flagellates) were unknown. Therefore, we ran 3 scenarios: HF0, HF50 and HF100, in which we assumed 0, 50 and 100% of the flagellates to be heterotrophic, respectively (or 100, 50 and 0% of the flagellates to be autotrophic, respectively).

**Setup and solution of the LIMs.** A LIM uses the mass balances of the food web compartments (the food web topology) to estimate  $n$  flows of matter  $\mathbf{f}$  between compartments of a food web (Klepper & Vandekamer 1987, Vezina & Platt 1988, Soetaert & Van Oevelen 2009). Mathematically, a LIM is expressed as a set of  $q$  linear equality equations:

$$\mathbf{A}_{q,n} \mathbf{f}_n = \mathbf{b}_q \quad (1)$$

and a set of  $q_c$  linear inequality equations, i.e. constraints:

$$\mathbf{G}_{q_c,n} \mathbf{f}_n \geq \mathbf{h}_{q_c} \quad (2)$$

Each element  $f_i$  in vector  $\mathbf{f}_n$  represents a food web flow ( $\text{g C m}^{-2} \text{ d}^{-1}$ ). The equality equations contain the mass balances over the different compartments and the measurements (Table 1), which are all linear functions of the flows. Each row in  $\mathbf{A}$  and  $\mathbf{b}$  is a mass balance or data point expressed as a linear combination of the food web flows. Numerical data enter  $\mathbf{b}$ , which are the rates of change of each compartment or the measured value in case of field data (Table 1). The inequality equation is used to place upper and/or lower bounds on single flows or combinations of flows (Table 1). The absolute values of the bounds are in  $\mathbf{h}$ . The inequality coefficients, quantifying how much a flow contributes to the inequality, are in  $\mathbf{G}$ . LIMs are

solved assuming steady state of all compartments, an assumption that has very little effect on the derived food web flows as changes in standing stocks in most cases are much smaller than the food web flows, even in highly dynamic systems (Vezina & Pahlow 2003).

Inverse food web models typically have less equality equations than unknown flows, i.e.  $q < n$  (Vezina & Platt 1988), making the problem mathematically underdetermined. Consequently, the  $n$  elements of the vector  $\mathbf{f}_n$  are quantifiable within certain ranges only. The models were solved in the R environment for statistical computing (R Development Core Team 2009) using specific R-packages. These ranges are derived using the R function 'lp' available in the R package 'limSolve' (Soetaert et al. 2008). From these lp-derived uncertainty ranges,  $N$  food web realisations can be sampled using a Markov Chain Monte Carlo procedure included in the function 'xsample' (Van den Meersche et al. 2009) of the R package limSolve. It is important to note that each of the  $N$  realisations corresponds to a set of  $n$  values (1 value per food web flow,  $\text{g C m}^{-2} \text{ d}^{-1}$ ), which obeys mass balance within the food web, as well as data and constraint equations (Eqs. 1 & 2). Hence, each food web realisation is equally likely. In this exercise,  $N$  was set at 1000, a value which proved to be large enough so that the  $N$  food web realisations covered the solution ranges as derived by lp. For each of the 4 food webs, we analysed each of these  $N$  food web realisations. Four LIMs were constructed and solved: HF0, HF50 and HF100 for spring (the herbivorous food web) and one for summer (the microbial food web). Spring and summer LIMs had 91 flows, 15 compartments, 5 externals and 107 inequalities. The spring LIMs had 15 equalities that expressed mass balance over all compartments. The 10 flows associated with capelin were set to zero in the summer LIM, which resulted in the summer LIM having 25 (i.e. 10 + 15) equalities. As an example, we show how the data in Table 1 were used to define the equalities (and inequalities) for the LIM for spring HF0 (Supplement 2, [www.int-res.com/articles/suppl/m398p093\\_app.pdf](http://www.int-res.com/articles/suppl/m398p093_app.pdf)). All constraints used to solve these LIMs are given in Table 1. The data in all 3 inverse models for spring were consistent and the models could be solved to recover food web flow values. For the summer food web model, this was not the case. The data dictated a DOC consumption that could not be met by DOC production under the assumed steady state conditions for DOC, indicating an exceptionally dynamic DOC pool. The smallest depletion rate of the DOC stock that resulted in a model solution was  $1.5 \text{ g C m}^{-2} \text{ d}^{-1}$ . Assuming such a depletion rate continued throughout July (31 d), the pool of labile DOC should have been  $\geq 46.5 \text{ g C m}^{-2}$ . For the southern Barents Sea, no data are available for the years considered. In the northern Barents Sea, the DOC concentration in the surface water

(<200 m) was about  $90 \mu\text{mol l}^{-1}$  in summer 2004 (Gasparovic et al. 2007), i.e.  $216 \text{ g C m}^{-2}$ . If this stock size is representative for the system under consideration, the labile DOC would have been about 22% of the total DOC. Although such data are not available for the southern Barents Sea, this percentage is still within the range derived for the Chukchi Sea, another pan-Arctic shelf sea (Davis & Benner 2007). Food web flow ranges (i.e. lp ranges) can be found in the SM (Table S1, Supplement 1).

**Analysis of LIM solutions.** The LIM solutions were analysed in 2 steps. Firstly, carbon flows of the herbivorous food webs (HF0, HF50 and HF100) and of the microbial food web were synthesised and compared. More precisely, the fate of gross primary production (GPP), consumption of bacteria by protozoa, carbon recycling and consumption of protozoa by copepods were quantified for all 4 food webs. Carbon recycling by the whole food web was calculated using Finn's cycling index (FCI), which is the fraction of the sum of flows that is devoted to cycling, and is equal to the amount of carbon that is recycled divided by the sum of all flows in the food web (Allesina & Ulanowicz 2004). The R package 'NetIndices' (Soetaert & Kones 2006) was used to calculate FCI (see Kones et al. 2009 for details).

Secondly, the effect of food web structure on cod grazing and production was analysed by calculating the direct and indirect dependency of cod on the other food web compartments and the efficiency of cod production for the 4 food webs. The dependency of a given food web compartment  $a$  on another compartment  $b$ , is the fraction of the diet of  $a$  that has passed through  $b$  (Szyrmer & Ulanowicz 1987). It is often termed the extended diet of  $a$  and was calculated using the R package 'NetIndices'. The food web efficiency (FWE) for production of cod was calculated as (Rand & Stewart 1998):

$$\text{FWE} = \frac{\sum_{i=1}^n \text{COD} \rightarrow i}{\text{NPP} + \text{ATL}} \quad (3)$$

where  $\text{COD} \rightarrow i$  are flows leaving the cod compartment that represent growth and export to higher trophic levels including humans; NPP is the total net primary production of all 3 phytoplankton groups (i.e. fixed carbon minus phytoplankton respiration); ATL is the advection of copepods into the Barents Sea by the Atlantic current. All terms in Eq. (3) have  $\text{g C m}^{-2} \text{ d}^{-1}$  as a unit. Calculations were performed separately for adult cod and for young cod and for the whole fish community as well. For comparison with literature data, where copepods are often the highest trophic level considered, FWE was also calculated for copepod production, now

using  $\sum_{i=1}^n \text{COP} \rightarrow i$  instead of  $\sum_{i=1}^n \text{COD} \rightarrow i$  in Eq. (3),

where COP stands for copepods.

Per food web structure (spring, HF0, HF50 and HF100), each of the previously mentioned quantities was calculated for all  $N (= 1000)$  food web realisations. We report the 90% CIs of these quantities.

## RESULTS

### Food web flows

From the DIC taken up by the 3 phytoplankton groups (GPP), about 20% was respired regardless of season resulting in a net primary production (NPP, sum of particulate and dissolved) of about  $3 \text{ g C m}^{-2} \text{ d}^{-1}$  in spring and  $2 \text{ g C m}^{-2} \text{ d}^{-1}$  in summer (Fig. 1). Group-specific NPP was highly variable and differences between phytoplankton groups were not pronounced. The only exception was the pico- and nanoplankton group in HF100, which provided between 7 and 20% of the NPP in spring and between 30 and 100% of the NPP in summer. Between 20 and 40% of the total GPP

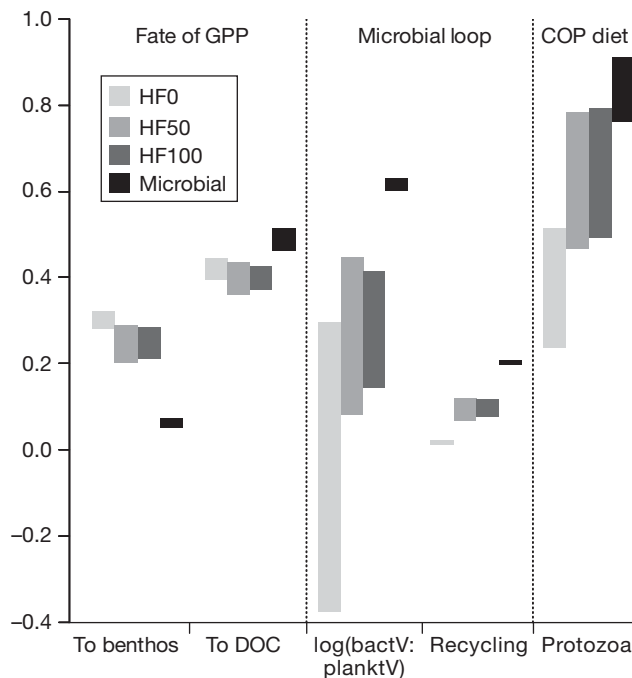


Fig. 1. Differences between the 3 herbivorous food webs (grey: HF0, HF50 and HF100, from left to right, respectively) and the microbial food web (black). Left section—fate of gross primary production (GPP): proportions sinking to the benthos (To benthos) and excreted as dissolved organic carbon (To DOC). Middle section—microbial loop: the logarithm of the ratio of bacterivory over planktivory by protozoa ( $\log[\text{bactV} : \text{planktV}]$ ; unitless), and the proportion of flows resulting from recycling (Recycling) as quantified by Finn's cycling index. Right section—diet of copepods: proportion of protozoa in copepod diet (Protozoa). Bars represent 90% CI and were calculated by analysing all  $N$  realised solutions of the food web model

was lost by sinking in spring, while in summer this was only 5 to 10% (Fig 1). The percentage of GPP released as DOC was about 40% in spring and 50% in summer. In spring, release of DOC by phytoplankton represented between 64 and 88% of all flows to the DOC compartment; in summer this was 41%. The remaining flows to DOC came from excretion by protozoa.

In the food web models, all DOC was taken up by bacteria, which were subsequently grazed by protozoa. This bacterivory appeared at least as important as the phytoplankton ingestion by protozoa because the pooled carbon flows from bacteria to protozoa (heterotrophic nanoflagellates and ciliates) were in general 1.2 (spring) to 4 times (summer) higher than pooled carbon flows from the 3 phytoplankton groups to protozoa (Fig 1). For HF0, the protozoan diet was highly uncertain. Carbon recycling, as quantified by the FCI, was between 2 and 15% in spring, and reached 20% in summer.

During summer, the majority of the copepod diet (80 to 90%) consisted of protozoa while estimates for spring are less decisive and depend on the scenario chosen (20 to 80% heterotrophs in diet) (Fig 1).

Within the fish community, differences in food web flows between spring and summer could be largely attributed to the absence of capelin in the southern Barents Sea in summer. Adult cod that did not follow the capelin migrating north in summer (50% of the spring stock, see Material and methods), experienced a diet shift from capelin (spring) to krill (summer) (Table 2). The diet contribution of young cod and herring in adult cod was marginal. The diet of young cod always consisted of about 90% copepods (Table 2).

### Dependency of cod on other food web compartments

In general, young cod were most dependent on copepods, their main food item (Fig. 2). In summer, the extended diet of young cod was dominated by ciliates, heterotrophic nanoflagellates, DOC and bacteria (hereafter termed 'the microbial loop'), together with copepods. In spring, when the food web was herbivorous, the dependency of young cod was more equally distributed over other food web compartments. Each of the phytoplankton groups alone was of moderate importance for young cod with no apparent minima or maxima and, apart from pico- and nanoplankton, no between-month differences. The dependencies of adult cod were much more evenly distributed over all other food web compartments than for young cod, and seasonal differences were far less pronounced (Fig. 3). Adult cod were 3 times less dependent on the microbial loop than were young cod. In contrast, adult cod were 6 times more dependent on macrozooplankton (chaetognaths and krill) and 3 times more dependent

Table 2. Diet of young and adult cod (three scenarios; % of food intake  $\pm$  1 SD)

	Krill	Capelin	Copepods	Herring
<b>Young cod</b>				
Spring HF0	4 $\pm$ 3	6 $\pm$ 4	89 $\pm$ 6	1 $\pm$ 1
Spring HF50	3 $\pm$ 2	5 $\pm$ 4	90 $\pm$ 7	2 $\pm$ 1
Spring HF100	4 $\pm$ 3	5 $\pm$ 4	90 $\pm$ 6	1 $\pm$ 1
Summer	2 $\pm$ 1		97 $\pm$ 2	1 $\pm$ 1
<b>Adult cod</b>				
Spring HF0	30 $\pm$ 6	54 $\pm$ 9	12 $\pm$ 3	4 $\pm$ 1
Spring HF50	31 $\pm$ 7	53 $\pm$ 11	12 $\pm$ 4	4 $\pm$ 1
Spring HF100	30 $\pm$ 6	55 $\pm$ 10	11 $\pm$ 3	4 $\pm$ 1
Summer	64 $\pm$ 2		28 $\pm$ 2	8 $\pm$ 1

on planktivorous fish (capelin) than were young cod. Similar to our findings for young cod, differences between both months in dependencies on phytoplankton groups were not apparent for adult cod.

### Food web efficiency

The FWE based on young cod production was between  $2 \times 10^{-5}$  and  $5 \times 10^{-5}$  for spring and almost an order of magnitude higher in summer (Fig. 4). When

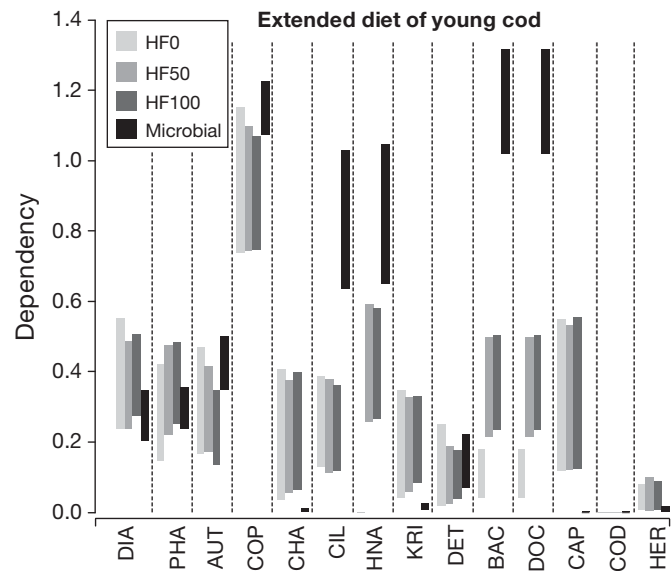


Fig. 2. Proportions of the different food web compartments in the extended diet ('dependency', unitless) of young cod for 3 scenarios for spring (grey: HF0, HF50 and HF100) and for summer (black). Bars represent 90% CI and were calculated by analysing all N realisations per food web structure. DIA = diatoms, PHA = *Phaeocystis pouchetii*, AUT = autotrophic nanophytoplankton, COP = copepods, CHA = chaetognaths, CIL = ciliates, HNA = heterotrophic nanoflagellates, KRI = krill, DET = detritus, BAC = bacteria, DOC = dissolved organic carbon, CAP = capelin, COD = adult cod, HER = herring



based on adult cod production, FWE was about 2 times higher in spring than in summer. When based on production of all fish (cod, capelin and herring), the food web was equally efficient in both seasons. The FWE based on with copepod production was 1.5 times higher in summer than in spring.

## DISCUSSION

### DOC production

The estimates for DOC release by phytoplankton (in summer, up to 50% of GPP) are among the highest reported (Nagata 2000), but are consistent with most of the observations in other Arctic systems such as the Eastern North Water Polynya where Tremblay et al. (2006) found DOC release to be 30% (spring) and 50% (summer) of the net particulate production. Similar values were also found for Antarctic oceans (Anderson & Rivkin 2001). Modelling studies revealed relatively high DOC release by phytoplankton in temperate waters as well, e.g. 35% of the GPP in the northeastern Atlantic Ocean during spring (Fasham et al. 1999) and 65% of the photosynthetic carbon products in estuary enclosures during a simulated bloom experiment (Van den Meersche et al. 2004). DOC release in the ice-covered Arctic is probably lower than in the ice-free regions discussed here, as suggested by data from the Chukchi Sea (Mathis et al.

2007) and from the West Antarctic Peninsula and the Ross Sea (Ducklow et al. 2006).

Processes resulting in phytoplankton DOC release include incomplete digestion by grazers (Jumars et al. 1989), cell lysis by viruses and exudation (Anderson & Williams 1998). Inferring the importance of viral activity is not straightforward as quantitative information is scarce. A number of studies in the polar freshwater environment exist (Anesio et al. 2007, Sawstrom et al. 2007), but information on the ice-free waters of the Barents Sea was not available. In the Chukchi Sea, another panarctic shelf sea, Hodges et al. (2005) showed that high bacterial and viral abundances coincided with algal blooms during summer. Hodges et al. (2005) also found that the bacterial community tends to converge towards less diverse, more specialist-dominated assemblages in summer, a process they attributed to viral activity. Clearly, the exact cause for high DOC release by phytoplankton certainly deserves more attention in future experimental work.

The importance of DOC excreted by heterotrophs, relative to the prime DOC source, i.e. release by phytoplankton, is still under debate. Anderson & Williams (1998) considered the heterotrophic production to be negligible compared with DOC production by phytoplankton for the English Channel, while others authors (Jumars et al. 1989, Strom et al. 1997) support the opposite view. Our results indicate that in spring, DOC re-

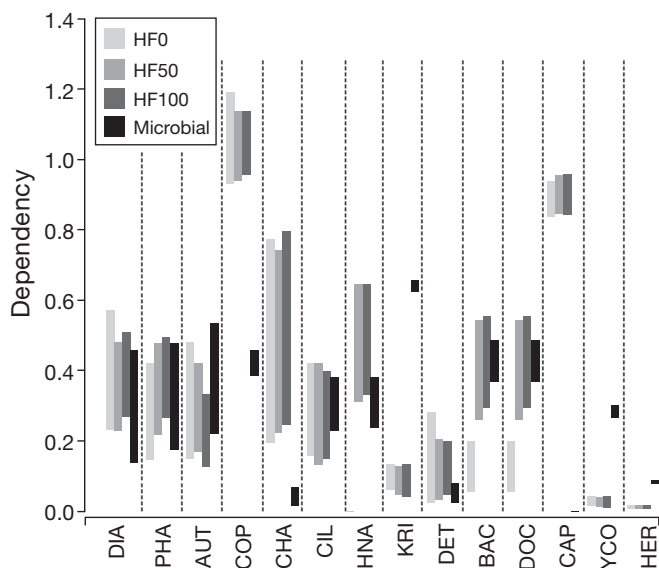


Fig. 3. Proportions of the different food web compartments in the extended diet ('dependency', unitless) of adult cod for 3 scenarios for spring (grey: HF0, HF50 and HF100) and for summer (black). Bars represent 90% CI and were calculated by analysing all N realisations per food web structure

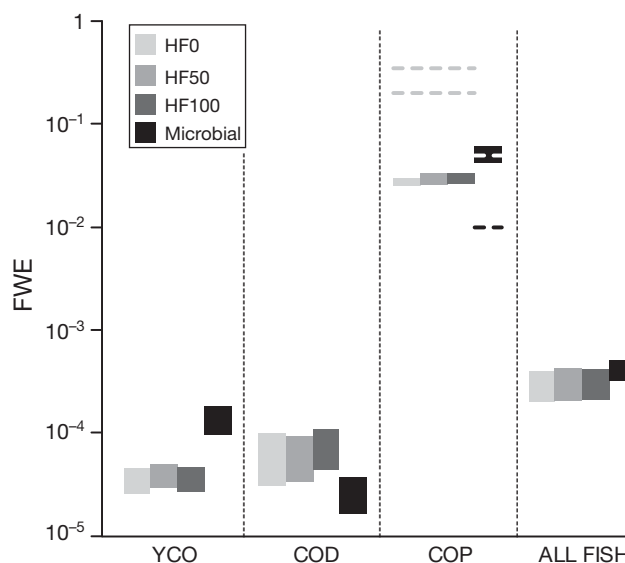


Fig. 4. Food web efficiency (FWE, expressed as fraction) calculated by dividing the production of young cod (YCO), adult cod (COD), copepods (COP) and all fish (ALL FISH) by the sum of net primary production and input of copepod biomass by the Atlantic current. Dashed line: FWEs for copepod production found by Berglund et al. (2007). Bars represent 90% CI and were calculated by analysing all N realised solutions of the food web model

lease by phytoplankton is the major source of DOC (>50%). In summer, the opposite is true as 59% of flows to DOC come from protozoa and not from phytoplankton. These findings are in line with other modelling exercises that show that DOC excretion by heterotrophs varies with season and is most important in summer when food webs are microbial (Fasham et al. 1999).

### The microbial loop

The DOC produced by all processes was consumed by bacteria because it is the only sink of DOC. Such a tight control of the DOC stock by bacteria is realistic in marine ecosystems (Vargas et al. 2007) and in polar systems in particular. Evidence from experimental studies in the Bering Strait region and across to the Canadian Basin indicates that microbial production is primarily controlled by dissolved organic matter (DOM) availability rather than by physical forcing such as by temperature (Rich et al. 1997, Kirchman et al. 2005). Likewise, the distribution of heterotrophic bacterial activity in the Kara Sea (Meon & Amon 2004) and Barents Sea (Thingstad et al. 2008) was controlled by the availability of DOC.

Bacterial production, essentially a reflection of DOC production because of the strong coupling discussed in the previous paragraph, was of comparable importance as primary production to fulfill carbon requirements of protozoa. In summer, protozoa fed 4 times more heavily on bacteria than on phytoplankton. This trend is amenable to other Arctic and Antarctic ecosystems. Becquevort et al. (2000) found that in the Indian sector of the Southern Ocean, protozoa principally ingested bacteria (87 to 99%) in both early spring and late summer. In summer, the diet of protozoa almost completely consisted of bacteria. Simek & Straskrabova (1992) found bacterivory by protozoa in a reservoir in Southern Bohemia to be negligible during spring, but protozoa consumed all bacterial production in summer. The same was found for the Canadian Arctic and McMurdo Sound in Antarctica (Anderson & Rivkin 2001).

An additional sink of DOC, not included in our model, might be photolysis of DOC. Photochemical mineralisation rates of terrestrial DOC have been found to exceed biological rates, although not for freshwater DOC (Obenosterer & Benner 2004). In the marine environment, photolysis is mostly considered as a transformation of aged refractory DOC to labile DOC (or directly to CO<sub>2</sub>) (Mopper et al. 1991). Although we do not claim that photolysis of marine labile DOC is unimportant, this process is currently too poorly constrained for reliable incorporation in food web models (Kieber 2000). Additionally, the loss of

labile DOC by photolysis will to some extent be compensated by the creation of new labile DOC through photolysis of refractory DOC.

The dominance of bacteria over phytoplankton as a food source for protozoa, combined with the high protozoan excretion of DOC, which is again readily taken up by bacteria, enhanced recycling of carbon in summer as compared with spring. This recycling was quantified as the percentage of all flows that is generated by recycling and is referred to as the Finn's cycling index (FCI). FCI was between 2 and 15% in spring and covered the range reported for open oceanic systems (Heymans & Baird 2000). However, recycling in summer was nearly 20%, i.e. no longer in the range expected for open oceanic systems but rather representative of estuarine systems (Heymans & Baird 2000) rather than open oceanic systems. The role of protozoa in this carbon recycling is crucial, as can be seen from the very low FCI for HF0 (Fig. 1), i.e. the spring scenario where all nanoflagellates were assumed to be autotrophic. Although the HF0 scenario might be judged as unrealistic, given recent findings on mixotrophy (Zubkov & Tarran 2008), it appears to be a useful exercise that increases our insight into the role microbes play in marine systems.

### Feeding on microbial carbon by copepods

Intensive feeding of copepods on protozoa is found in different marine ecosystems (Tamigneaux et al. 1997, Mayzaud et al. 2002, Calbet & Saiz 2005) although the intensity of this feeding process differs between systems. For example, the grazing of copepods on protozoa was only half of the grazing of copepods on phytoplankton in all seasons and for different regions in the Greenland Sea (Moller et al. 2006). Carmack & Wassmann (2006), as well as Levinsen et al. (2000) argued that feeding by copepods on protozoa especially occurs at low phytoplankton concentrations. In a recent cross-ecosystem analysis, Calbet & Saiz (2005) suggested a cut-off value for phytoplankton biomass (50 µg C l<sup>-1</sup>), below which ciliates contribute at least as much to the copepod diet as do phytoplankton. As Calbet & Saiz (2005) only considered ciliates as protozoa, this cut-off value may be higher when other protozoa such as heterotrophic nanoflagellates are included (as in the present study). The phytoplankton concentration in southern Barents Sea is about 50 µg C l<sup>-1</sup> in summer and 100 µg C l<sup>-1</sup> in spring in the upper 90 m of the water column (Rat'kova & Wassmann 2002). The fact that copepods feed on a mixture of autotrophs (20 to 80%) and protozoa in spring, while up to 90% of their diet consists of protozoa in summer, does agree with the cross-ecosystem trends found by

Calbet & Saiz (2005). This seasonal diet shift typically coincides with a shift from large copepod species in spring to smaller species in summer that are perfectly suited for grazing on protozoa (Moller et al. 2006). This shift was experimentally confirmed by Arashkevich et al. (2002) for the southern Barents Sea.

### **Copepods: the link between the microbial loop and higher trophic levels**

The diets we derived for adult and young cod, which corresponded well with independent stomach content data (Orlova et al. 2005, Link et al. 2009), have implications for the ecological role of copepods. Copepods served as the main food item for young cod in both seasons. In summer, copepod production was closely linked to protozoa production, as shown by the proportion of protozoa in the copepod diet (Fig. 1). In turn, protozoa in summer relied heavily on bacteria that controlled the stock of DOC. Because of their interdependency, not only copepods, but also protozoa, bacteria and DOC were important for young cod in summer (Fig. 2). The dependencies of young cod indicated that >60% of their diet passed through the microbial loop. As the fraction of heterotrophs in the copepod diet was lower in spring than in summer (Fig. 2), the dependency of young cod on the microbial loop was 2 to 3 times lower in spring than in summer. Instead, the dependencies on phytoplankton, protozoa, bacteria and DOC were comparable in spring. Still, the dependency on copepods was comparable for both food web structures indicating that copepods were always crucial for converting microbial carbon to forms consumable by cod. The unique coupling of the microbial domain with young cod through one important link (copepods), resulted in an interesting 'hourglass'-like food web structure for young cod. The same hourglass-like structure was found for adult cod, albeit only in spring when the adult cod's favourite food was capelin, a copepod feeder. This resulted in similar dependencies for young and adult cod in spring. In summer, the migration of capelin forced the adult cod population to feed on herbivorous krill, i.e. exclusively relying on phytoplankton and not on protozoa. This creates an uncoupling of adult cod from the microbial loop in summer, as reflected by the lower dependency for adult cod on the microbial loop than for young cod (Figs. 2 & 3).

Copepod biomass advection ( $0.03 \text{ g C m}^{-2} \text{ d}^{-1}$  in both seasons) was 1.25 to 5 and 2.5 to 5 times lower than locally produced copepod biomass in spring and in summer, respectively. These differences could be expected from current measurements between Bear Island and the northern coast of Norway. Currents

were about 2 times higher in May 1998 than in July 1999 (Ingvaldsen et al. 2004), i.e. the months for which data were gathered (Table 1), and were in general high ( $\approx 2.5$  Sverdrups [Sv]). One could thus expect that in years with lower net inflow rates, local phenomena become increasingly important, which would further strengthen the relationships between the microbial and fish communities established here.

### **Food web efficiency**

The lower FWE for adult cod production in summer than in spring reflects the lower standing stock of cod during summer in the southern Barents Sea. Because of the lower resource requirements of this reduced cod stock, less carbon is transferred to cod in summer than in spring. Instead, carbon is transferred to the other fish compartments. As such, the food web cannot be said to be less efficient in summer, as can be seen from the FWE for production of all fish (Fig. 4). In contrast with adult cod, the reduction of the young cod stock in summer did not cause the FWE for young cod production to be lower in summer than in spring. Apparently, the higher efficiency of the carbon transfer from phytoplankton to copepods in summer (Fig. 4), the main food for young cod in both seasons, compensated for the effect of a reduced young cod stock in summer.

The FWEs for fish production found here (Fig. 4) suggest a lower efficiency for fish production than the transfer efficiencies (TEs) assumed by Jennings et al. (2008) in their recent effort to estimate global fish production from primary production data. Using  $\text{TE} = \text{FWE}^{(\text{TL} - 1)^{-1}}$  as an approximation, our results indicate TEs of 0.05 to 0.13 for cod (with trophic level [TL] = 4.3 to 5.6), and of 0.01 to 0.11 for young cod (TL = 3.3 to 5), while Jennings et al. (2008) use a fixed TE of 0.125. Sensitivity analyses carried out by Jennings et al. (2008) demonstrated that the use of the TEs found here would lower their estimates of fish production by more than a factor of 2.

The FWE based on copepod production in summer was among the highest FWEs observed in an experimental microbial food web established by Berglund et al. (2007) (Fig. 4). However, the FWE for copepods in spring was 10 times lower than what Berglund et al. (2007) reported for an experimental herbivorous food web. For this apparent discrepancy between experimental data and our results, 2 explanations are offered. First, the use of mesocosms by Berglund et al. (2007) with 40 cm depth may have resulted in an underestimation of phytoplankton sedimentation, a loss term for the pelagic food web, and thus an overestimation of FWE. Second, the most abundant species in those mesocosms were of intermediate size (cryptophytes

and prasinophytes <10 µm) and were smaller than the phytoplankton that dominated the spring food webs discussed here (>20 µm) (Rat'kova & Wassmann 2002). Phytoplankton in the enclosures described by Berglund et al. (2007) are thus intrinsically less prone to sedimentation than in the Barents Sea food webs described here. A recalculation of the FWE for copepods using the net primary production minus sedimentation losses (i.e. using  $NPP + ATL - SED$  as a denominator in Eq. (3) with  $SED$  representing the sedimentation losses) reveals that FWEs are comparable across the 4 food webs discussed in the present study (Fig. S3, Supplement 1). This indicates that the carbon remaining in the water column is processed as efficiently in the summer (microbial) food web than in the spring (herbivorous) food webs.

A contemporary view is that microbial food webs have more trophic levels and, thus, higher overall metabolic requirements (Straile 1997) when transferring carbon from the primary producers up to the top predators. However, in this paper we show that a higher number of trophic levels do not necessarily result in lower FWEs for microbial food webs than for herbivorous food webs. Life forms that dominate in the microbial food web, i.e. bacteria and protozoa, essentially consume loss products from other food web compartments (e.g. excretion of DOC by protozoa and subsequent uptake by bacteria). As such, the fraction of the carbon fixed by phytoplankton that reaches higher trophic levels would be higher than in herbivorous food webs, as suggested by Vargas et al. (2007) and Calbet & Saiz (2005).

As our estimates of DOC production, bacterivory by protozoa and consumption of protozoa by copepods were shown to reflect what is observed for many other food webs, our results may well extend beyond this particular case for the Barents Sea. However, the heterogeneity of this system (Wassmann et al. 2006) would make extrapolation of our results to other regions or periods within the Barents Sea speculative. For example, for periods with a collapsing capelin stock (Dalpadado & Bogstad 2004) in years with a strong link between benthic production and adult cod feeding, the trophic position of cod would likewise change and, thus, so would its dependencies. It would be interesting to determine how such spatiotemporal variability changes the conclusions drawn here, and we encourage future studies to examine such issues using the inverse modelling framework or other quantitative tools.

## CONCLUSIONS

In spring (the herbivorous food web), release of DOC by phytoplankton dominated the carbon flows to the DOC compartment where it was consumed by bacte-

ria. Bacteria were of comparable importance as phytoplankton as food for protozoa. The diet of copepods was a mixture of protozoa (20 to 80%) and phytoplankton, which resulted in moderate dependency of young and adult cod on the microbial loop (DOC–bacteria–protozoa).

In summer (the microbial food web), protozoa excretion was more important than DOC release by phytoplankton. Bacteria consuming this DOC were 4 times as important as a food source for protozoa as phytoplankton. Protozoa in turn formed 80 to 90% of the copepod diet. Because of the strong relationships between the key players of the microbial loop (DOC, bacteria and protozoa) and copepods, the dependency of young cod on the microbial loop was high in summer. Adult cod were far less dependent on the microbial loop than young cod as they relied on strictly herbivorous krill in summer.

The efficiency of the food web for fish compared well between seasons and for copepod production; FWE was 2 times higher in summer than in spring. For the summer case, FWE for copepod production agreed well with available data; for spring, our estimates were an order of magnitude lower than literature estimates from shallow enclosures with relatively small phytoplankton species. Our estimates on DOC production, bacterivory by protozoa and consumption of protozoa by copepods agreed well with what has been observed for many other food webs (both polar and nonpolar), which suggests our results to be amenable to other parts of the world's oceans.

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